

Department of Energy

Washington, DC 20585

January 30, 1991

Mr. Robert Browning U.S. Nuclear Regulatory Commission Washington, D.C. 20555

Dear Mr. Browning:

Please find enclosed a copy of the Trip Report for the Department of Energy, Office of Environmental Restoration and Waste Management, November 1990, fact-finding mission to Japan. Any further questions may be directed to me at (301) 353-7954.

Sincerely,

Donald H. Alexander, Ph.D. Program Manager International Technology Exchange Program Support Division Office of Technology Development

412

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Enclosure

PDR



FOREIGN TRAVEL REPORT

ENVIRONMENTAL RESTORATION & WASTE MANAGEMENT TECHNOLOGY EXCHANGE DISCUSSIONS IN JAPAN

NOVEMBER 3 - 14, 1990



CONTRIBUTORS:

LEO P. DUFFY, DONALD H. ALEXANDER, ELLEN LIVINGSTON-BEHAN, SATYENDRA (JOHN) MATHUR, DONALD T. OAKLEY, W. MELINDA DOWNING, WILLIAM C. SCHUTTE CONTENTS

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Technologies discussed with Kobe Steel and Bibliography



TRIP REPORT SUMMARY

1. TRAVELER'S NAMES:

Leo P. Duffy Director Office of Environmental Restoration and Waste Management U.S. Department of Energy

Donald H. Alexander Program Manager International Technology Exchange U.S. Department of Energy

Ellen A. Livingston-Behan Environmental Regulatory Specialist Office of Environmental Restoration and Waste Management U.S. Department of Energy

Satyendra P. (John) Mathur Program Manager TRU Waste and WIPP R&D Programs, U.S. U.S. Department of Energy

Donald T. Oakley Senior Advisor to the Department of Energy's Office of Environmental Restoration and Waste Management Los Alamos National Laboratory

W. Melinda Downing
 Program Review Coordinator
 Office of Environmental Restoration and
 Waste Management
 U.S. Department of Energy

William C. Schutte Group Manager Technical Integration Idaho National Engineering Laboratory





2. ITINERARY:

November 3 -Travel to Minneapolis

November 4 -Leave USA for Tokyo, Japan

November 5 - Tokyo, DOE Tokyo Staff, Atomic Energy Bureau/Science and Technology Agency, Embassy, JAERI and JGC Corporation.

November 6 - Mito, National Research Institute for Pollution and

November 7- Mito, Power Reactor and Nuclear Fuel Development Corporation, Tokai Works and Oarai Works.

November 8 - Mito, Japan Atomic Energy Research Institute

November 9 -Takasaki, Takasaki Radiation Chemistry Research Establishment Tokyo, Power Reactor and Nuclear Fuel Development Corporation

November 10-Kyoto, Kyoto University

November 13-Tajimi, Power Reactor and Nuclear Fuel Development Corporation, Chubu Works.

November 14-Osaka, Ministry of International Trade and Industry/Agency for Industrial Science and Technology.

November 15-Depart Osaka, Japan for U.S.

3. TRIP COST: Estimated cost per traveler of \$4 378.00

4. ABSTRACT: The Department of Energy Of ce of Environmental Restoration and Waste Management (DOE/EM) participated in a series of fact finding meetings and facility tours in Japan. Meetings with Power Reactor and Nuclear Fuel Development Corporation (PNC) were conducted in accordance with the bilateral agreement between the United States Department of Energy and Power Reactor and Nuclear Fuel Development Corporation (PNC) In the opening session with Japan's Science and Technology Agency the EM delegation, led by Mr. Leo P. Duffy, jointly underscored the continuing U.S. commitment that the technologies to be discussed would be limited to peaceful uses in the areas of environmental restoration and waste management. The delegation visited government, quasi-government and private organizations during the November 3-14, 1990, fact finding mission including the Science and Technology Agency (Tokyo), the Ministry of Trade and Industry (Tokyo, Mito, Osaka), the Power reactor and Nuclear Fuel Development Corporation (Tokyo, Tokai, Oarai, Chubu), the Japan Atomic Energy Research Institute (Tokai, Tokyo), Kyoto University, JGC Corporation (Oarai), and Kobe Steel. The delegation concludes that expanded collaboration in several technical areas described in this report appear to be of potential mutual benefit. Several areas of potential technical collaboration appear to have considerable merit in addition to successful on-going interactions related to vitrification, decommissioning, and TRU. This report provides meeting summaries and selected information on a wide range of technologies by organization.



5. PURPOSE OF TRIP: This trip was one in a series of fact-finding visits to Japan on environmental restoration and waste management. The following four pre-trip objectives were successfully met.

OBJECTIVE 1: IDENTIFY SPECIFIC TECHNOLOGIES OF MUTUAL INTEREST

The facility and site visits were designed to encourage useful dialogue for identifying:

- 1) transferrable technologies,
- 2) technologies under development, and
- 3) areas for cooperative technology development.

The delegation identified key technologies that have potential for the highest payoff for use in EM site characterization and restoration efforts. The delegation was provided with a substantial amount of technical information through presentations and published materials with broad applicability to the EM mission. A number of technology breakthroughs with potential for significant impact for EM site characterization and environmental restoration were presented to the delegation.

OBJECTIVE II: DETERMINE IF AN ANNEX TO THE CURRENT DOE-PNC AGREEMENT IS REQUIRED

The current agreement appears to broadly cover DOE-EM/PNC exchanges. However, the agreement may need to be modified if new initiatives are pursued with PNC. The agreement will be revisited after joint workshops are conducted in early 1991 to determine if joint technical collaboration with organizations other than PNC will be pursued. Collaboration and possible agreements with JAERI, MITI and Kyoto University needs to be pursued since they are developing technologies or conducting research in areas relevant to DOE-EM.

OBJECTIVE III: IDENTIFY TECHNICAL AREAS OR SPECIFIC TECHNOLOGIES THAT SHOULD BE INCORPORATED IN "DOE-EM/PNC RECORD OF MEETING"

DOE and PNC agreed that several workshops would be arranged over the coming months to focus on areas of joint technical interest in preparation for the Bilateral Coordination Meeting in the spring of 1991. DOE appointed Dr. Donald Alexander as the DOE-EM coordinator. PNC appointed Mr. Takao Yagi as PNC-EM coordinator.



The workshops will provide reports recommending areas of technical collaboration to the DOE-EM/PNC Coordinating Committee in the Spring of 1991. The U.S. delegation identified several technical areas for continued collaboration in the November 9th meeting with PNC including decontamination and decommissioning, vitrification and TRU handling, treatment and disposal.

Areas of particular interest to the delegation which will be considered for future collaboration with PNC include:

- the use of the plasma-arc saw for dismantling reactor vessels and related contaminated structures;
- 2. smelting of slightly contaminated ferrous metals for recycling;
- 3. methods for removal of contaminated concrete;
- 4. methods for reusing slightly contaminated concrete;
- 5. methods of waste reduction and minimization;
- 6. partitioning and transmutation;
- 7. robotics;

Areas of particular interest to the delegation which will be considered for future collaboration with MITI, JAERI and Kyoto University include:

- 1. methods for the removal of uranium from seawater;
- applications of fiber-optics and lasers for in situ analysis of groundwater;
- 3. methods for the removal of organics such as trichloroethylene;
- 4. simulation modeling of groundwater contaminant migration;
- 5. research on actinide chemistry;
- 6. methods of underground characterization;
- 7. inorganic microencapsulated adsorbents;
- 8. optical microsensors for gases; and,



9. gold-metal oxide catalysts for sensor applications.

OBJECTIVE IV: INITIATE MECHANISM FOR COLLABORATION WITH OTHER JAPANESE ORGANIZATIONS

Based on the findings of the delegation, as outlined in Objective III, mechanisms for collaborating with MITI, JAERI, and Kyoto University should be pursued. A bilateral agreement between the USNRC and JAERI for exchanges in waste management currently exists and could serve as a model for DOE/EM-JAERI and DOE/EM-MITI. Amending the existing USNRC/JAERI agreement does not appear to be a viable option. Agreements with MITI and JAERI would be initiated by developing a broad scope of work, seeking State Department concurrence and then arranging meetings with MITI and JAERI representatives in Washington to discuss the U.S. proposal. Upon general agreement between parties to collaborate, formal agreements would be written, approval and signing of which would be coordinated through the DOE Office of International Affairs, IE-12.

Collaboration with Japanese Universities should be pursued through the Ministry of Education, Science and Culture or through laboratory to laboratory agreements.

One option being explored is to expand the existing DOE/JAERI Agreement on Decommissioning Nuclear Facilities (term 7-2-87 to 7-2-92) to include environmental restoration and waste management activities.





DETAILED TRIP REPORT

INTRODUCTION:

As the next step in the development of a cooperative initiative between the U.S. and Japan for technology development in the area of environmental restoration and waste management, Leo P. Duffy led a U.S. delegation to Japan on November 3-14, 1990.

The trip was closely coordinated with and supported by the Power Reactor and Nuclear Fuel Development Corporation (PNC). The PNC is a Japanese quasi-governmental agency responsible for developing fuel cycle technologies (including technologies related to reactor design, uranium mining and enrichment, spent fuel reprocessing, and radioactive waste disposal).

The Department of Energy (DOE) currently has an agreement (finalized in 1986) with the PNC for the cooperative development of technology and techniques for radioactive waste management. Objectives of the delegations trip to Japan focused on (1) identifying with the PNC specific technologies either in existence or under development that are applicable to EM efforts and that should be considered for exchange; (2) determining whether an annex to the existing DOE-PNC Agreement is required to support technology exchange and cooperative development efforts; and (3) identifying (in cooperation with PNC) appropriate future initiatives with other Japanese organizations having technologies that appear applicable to DOE's clean-up and waste-management activities.

Meetings and on site visits during the trip focused on specific Japanese research and development (R&D) initiatives of particular interest to the Office of Environmental Restoration and Waste Management (EM), including robotics, plasma-arc saw, recycling of slightly contaminated metals and concrete, methods of waste reduction and minimization, partitioning and transmutation, fiber optics, filtration, sensors, research on actinide chemistry, inorganic micro-encapsulated adsorbents, gold-metal oxide catalysts, trichloroethylene extraction and uranium extraction from groundwater.

GENERAL OBSERVATIONS:

The delegations general impressions of the activities and facilities reviewed during the trip are as follows:

- o technology development initiatives were generally at the same level of development for similar DOE initiatives;
- as is the case in many U.S. research facilities, many Japanese research facilities are greatly under-utilized. Collaboration will reduce underutilization for both countries and eliminate costs of erecting new redundant facilities;



- o the facilities were clean and well managed:
- As is the case in the U.S., the number of scientists that have specialized in Environmental Restoration and Waste Management research is insufficient to support future government needs for technology development. Collaboration will help reduce the manpower shortage for both nations.
- o <u>The Japanese are developing foreign-scientist enclaves at major research centers to</u> <u>attract foreign post doctoral candidates.</u> This approach is being discussed with <u>Frank Parker</u>. National Academy of Sciences, for possible implementation in U.S. <u>Universities and at U.S. National Laboratories</u>.
- o <u>Collaborative efforts with PNC in decomissioning. TRU</u>, waste reduction and minimization, and vitrification continue to be mutally beneficial.
- o New initiatives with MITI, JAERI, and MESC should be pursued.
- o <u>U.S. mixed waste management capabilities should be shared with responsible</u> Japanese organizations.

MONDAY, November 5

Abstract of Activities for November 5, 1990:

The first day of the trip was devoted to meetings with the directors of Japan's Science and Technology Agency (STA), the Japan Atomic Energy Research Institute (JAERI), and the Power Reactor and Nuclear Fuel Development Corporation (PNC).

The organizations have the following responsibilities:

- o The STA formulates policies for nuclear research and development, and establishes and enforces technical and safety standards applicable to nuclear material management and disposal.
- The JAERI is a quasi-governmental research organization responsible for implementing national nuclear energy programs.
- o The PNC is a quasi-governmental research organization responsible for developing fuel cycle technologies (including technologies related to reactor design, uranium ining and enrichment, spent fuel reprocessing, and radioactive waste disposal). The Department of Energy (DOE) currently has an agreement (finalized in 1986) with



the PNC for the cooperation of technology development for radioactive waste management.

Principal spokesmen for each organization during meetings with the U.S. delegation were Mr. Hiroto Ishida, Deputy Director-General for the STA's Atomic Energy Bureau; Mr. Toyojira Fuketa, Vice President of JAERI; and Mr. Takao Ishiwatari, President of PNC.

During the meetings with these and other representatives of each organization, Mr. Duffy discussed the following topics:

- Actions being taken by the DOE under the "Ten Point Plan" established by Secretary of Energy James Watkins to improve the DOE's performance and accountability in protecting the environment and public health and safety.
- o The development and implementation of DOE's program for environmental restoration, waste management, and related research and development initiatives under the "Environmental Restoration and Waste Management Five-Year Plan."
- o Objectives of the U.S. delegation's trip to Japan, with emphasis on:
- identifying with the PNC specific technologies either in existence or under development that are applicable to efforts of the Office of Environmental Restoration and Waste Management and that should be considered for exchange (specific Japanese research and development initiatives of preliminary interest to the Office of Environmental Restoration and Waste Management were noted to include robotics, fiber optics, partitioning and transmutation, filtration and sensors);
- 2) determining whether an annex to the existing DOE-PNC Agreement is required to support technology exchange and cooperative development efforts; and
- identifying (in cooperation with the PNC) appropriate future initiatives with other Japanese organizations having technologies that appear applicable to DOE's cleanup and waste-management activities.

The spokesmen for each organization expressed interest in working to identify areas of cooperative effort. However, they emphasized the need (under existing laws and treaties) to limit such cooperation to activities that are not directly related to U.S. nuclear defense activities.

Later that day, the STA responded to press inquiries on the DOE-EM visit. The STA reviewed the mission of the U.S. fact finding team, reviewed technologies of particular interest to the DOE, noted its intent to cooperate positively in the DOE's fact-finding mission and the identification of cooperative technology initiatives, described the potential



development of a government-to-government agreement by 1992, and emphasized the need to focus any such initiatives on the peaceful uses of nuclear energy.

Meeting at U.S. Embassy, Tokyo

Address:

EMBASSY OF THE UNITED STATES OF AMERICA 1-10-5 AKASAKA MINATO-KU TOKYO 107 FROM THE U.S.A. APO SAN FRANCISCO, CA 96503 PHONE: (03) 224-5066

(ADDRESS FOR MAIL FROM U.S.) U.S. DEPARTMENT OF ENERGY U.S. EMBASSY - TOKYO APO SAN FRANSISCO, CA 96503

Participants:

William T. Breer Minister

Richard S. Kanter Assistant Commercial Attache

Detailed Meeting Notes:

The Minister and his staff provided the delegation with an overview of local protocols, information on the Ascension of the Emperor, and background information on the role of the U.S. Embassy in international technology exchanges.

Meeting with US/DOE Japan Staff, Tokyo

Address:





UNITED STATES DEPARTMENT OF ENERGY AMERICAN EMBASSY 10-5 AKASAKA 1-CHOME MINATO-KU, TOKYO 107 PHONE (03) 224-5475

Participants:

Milton Eaton Senior DOE Representative

Toshiaki Okubo Senior Energy Affairs Specialist

Mayumi Kainuma Energy Affairs Specialist

Detailed Meeting Notes:

The DOE Representative and Staff are responsible for all in country coordination of DOE related activities. The staff can provide on-site secretarial and other support services. The logistics for the delegation including the arrangement of lodging, travel accomodations and coordination of the itinerary and coordinating agenda topics with Japanese organizations prior to the trip were all arranged by the DOE Embassy Staff. The delegation highly recommends that all future EM exchanges with Japan be coordinated with the Senior DOE Representative.

Meeting with Science and Technology Agency, Tokyo

Address:

SCIENCE AND TECHNOLOGY AGENCY ATOMIC ENERGY BUREAU 2-2-1 KASUMIGASEKI CHIYODA-KU, TOKYC PHONE (03) 581-527

Role ci STA in Nuclear Energy Development:



- The AEB formulates policies for nuclear energy R&D and supervises activities at JAERI, PNC, the National Institute of Radiological Sciences, and the Institute of Physical and Chemical Research (RIKEN). AEB also provides administrative support to the Atomic Energy Commission (AEC).
- The Nuclear Safety Bureau (NSB) determines technological standards and enforces safety regulations concerning new types of nuclear reactors, research reactors, nuclear fuel processing and reprocessing facilities, radioactive waste and transportation of nuclear fuel substances. NSB also provides administrative support to the Nuclear Safety Commission (NSC).

STA is also conducting the OMEGA Project, exchanging information of advanced waste treatment technology. The project involves two technological fields - partitioning and transmutation of TRU elements. A meeting of the OMEGA Project was convened in Tokyo during the first week of this visit.

Participants:

Hiroto Ishida Deputy Director-General Atomic Energy Bureau

Yukio Sato Director, Power Reactor Development Division Atomic Energy Bureau

Tomoyuki Murakami Research and International Affairs Division Atomic Energy Bureau

Yukihide Hayashi Director, Research & International Affairs Division Atomic Energy Bureau

Shizuo Hoshiba Director, Nuclear Fuel Cycle Back-End Office

Agenda:

*Meeting with Mr. Ishisda, Deputy-Director General, Atomic Energy Bureau.

Detailed Meeting Notes:

Mr. Ishida chaired the meeting with STA. The DOE delegation provided STA with a copy of the presentation by Mr. Duffy entitled "The Program of the United States Department of Energy on Environmental Restoration and Waste Management" (See References). Mr. Duffy underscored the Department's commitment to involve all affected parties in the restoration of contaminated sites. He noted by way of example that Indian Nations, with lands affected by environmental contamination resulting from DOE activities, are participants in the restoration process.

Environmental restoration is being undertaken by many technologically advanced nations. However, the resource requirements are staggering and the demands on scientific and engineering personnel will increase substantially. Therefore, the U.S. delegation suggested that the joint international sharing of technologies should be pursued to reduce the overall resource requirements and accelerate clean-up.

Mr. Duffy assured STA that EM's role is limited to environmental restoration and waste management. Once a facility is turned over to EM, EM has the responsibility for the management of site clean-up. Mr. Duffy also stated that Secretary Watkins created the EM organization as an entity with clear separation from military activities.

STA responded very positively to continued technology exchanges with DOE-EM. STA underscored that joint activities must be restricted to peaceful uses and that the waste generated by military activities which are recovered with jointly developed technologies must not be used for military activities again.

**Mr. Duffy underscored the U.S. commitment to peaceful uses. He stated that many of the environmental problems being dealt with by EM were derived from non-nuclear wastes such as solvents, PCBs, and heavy metals.

Meeting with Japan Atomic Energy Research Institute, Tokyo

Address:

JAPAN ATOMIC ENERGY RESEARCH INSTITUTE FUKOKU SEIMEI BLDG. 2-2-2 UCHISAIWAI-CHO CHIYODA-KU, TOKYO 100, JAPAN PHONE 03-592-2101





Role of JAERI:

JAERI is a quasi-governmental research organization which implements national long-term programs in nuclear energy, including joint projects and international cooperative efforts.

Research Activities:

- R&D of nuclear energy, nuclear safety, high temperature gas-cooled reactors, nuclear fusion, radiation applications, nuclear powered ships, basic research, decommissioning of nuclear reactors;
- Design, construction and operation of reactors;
- Education and training of researchers and engineers in the field of nuclear energy; and
- Dissemination of information obtained through R&D activities.

Participants:

Kazuo Sato Executive Director

T. Tsujino Deputy Director Office of Planning

Masashi Iizumi Director, Office of Planning

Keisuke Kaieda Senior Staff, Office of Int'l Affairs

Hideki Omichi Senior Staff, Office of Planning

Bibliography of Literature Received:

- "Development of Technologies on Decommissioning of Nuclear Fuel Cycle Technologies", Japan Atomic Energy Research Institute. 5 pages.
- "JPDR Decommissioning Program", written by T. Hoshi from the 9th TAG Meeting on October 8-12, 1990 at the Japan Atomic Energy Institute. 10 pages.



"New JRR-3, Compiled by the Research Reactor Operation at the Tokai Research Establishment", Japan Atomic Energy Research Institute. 6 pages.

"Progress Report on Safety Research of High-Level Waste Management for the Period April 1988 to March 1989", Edited by Haruto Nakamura and Susumu Muraoka, Department of Environmental Safety Research, Tokai Research Establishment, Japan Atomic Energy Research Institute, 2 pages.

"Reactor Decommissioning Technology Development and Actual Dismantling of JPDR," compiled by the Tokai Research Establishment, Japan Atomic Energy Research Institute. 9 pages.

"Safety Studies on Glass Waste Form", written by S. Muraoka at Japan Atomic Energy Research Institute. 10 pages.

"Summary of WASTEF Facility", from Japan Atomic Energy Research Institute. 10 pages.

"Volatilization of Cesium from Nuclear Waste in a Canister", Hiroshi Kamizono, Shizuo Kikkawa, Shingo Tashiro and Haruto Nakamura. at Japan Atomic Energy Research Institute. Department of Environmental Safety Research, 6 pages.

Agenda:

*Meeting with Executive Vice President and Directors of JAERI.

Detailed Meeting Notes:

The DOE delegation provided JAERI with a copy of the presentation by Mr. Duffy entitled "The Program of the United States Department of Energy on Environmental Restoration and Waste Management" (See References). Mr. Duffy stated that there are an enormous number of sites in the United States that need to be restored. He emphasized the delegation's interest in any technologies which may expedite the clean-up of these sites including sensors, fiberoptics, remote measuring techniques for groundwater, robotics, biotechnology, filter technology and waste forms.

Mr. Kazuo Sato stated that the JAERI representatives were favorably impressed with Mr. Duffy's presentation and looked forward to future collaboration with DOE-EM with the caveat that all technology exchanged be used exclusively for peaceful uses. He stated that JAERI would be very interested in reviewing any DOE-EM proposals for collaboration.



Address:

POWER REACTOR & NUCLEAR FUEL DEVELOPMENT CORPORATION 9-13, 1-CHOME, AKASAKA MINATO-KU, TOKYO, 107 JAPAN

Role of PNC:

Plays a central role in developing fuel cycle technologies, and fast breeder reactors. PNC has developed technologies for prospecting for uranium deposits, refinement and conversion, centrifugal uranium enrichment, spent fuel reprocessing and radioactive waste disposal and is preparing to cooperate with industry for demonstration and utilization of these technologies.

PNC operates the following facilities:

Tokai Works, Oarai Engineering Center, Fugen Nuclear Power Station (ATR prototype reactor), Monju Construction Office (FBR prototype reactor), Tsuruga Office, Chubu Works, and Ningyo Toge Works.

PNC has a plan to establish "Storage Engineering Center" in Horonobe, Hokkaido, to store vitrified HLW and to study technology for geological disposal in deep underground formations.

Participants:

Takao Ishiwatari President

Yoshikazu Hashimoto Executive Director

Masao Yamamoto Deputy Senior Director

Saburo Kikuchi Secretary to the President

Kiyoshi Kikuchi Ing. Geologue, Department des Ressources Nucleaires

Tadashi Mano General Manager, Conditioning Research Program Radioactive Waste Management Project



Takashi Yoshikawa Manager, Int'l Cooperation Office, Int'l Division

Reiko Nunome Int'l Cooperation Office, Int'l Division

Akira Wadamoto Engineer, Conditioning Research Prog. Radioactive Waste Management Project

Bibliography of Literature Received:

"FBR Development in PNC for Commercialization", PNC, 8 pages.

"Technical Draft for Comments RD&D Program on Low-Level TRU Bearing Waste Management Technologies", PNC, 43 pages.

Agenda:

*Meeting with PNC President.

Detailed Meeting Notes:

Dr. Takao Ishiwatari welcomed the delegation and stated that PNC was pleased with the long standing exchange program with the Department of Energy. He stated that future technology exchanges should be preceded by a statement of clear objectives, should be open to interested parties, and he underscored the need to limit collaboration to areas of peaceful uses of nuclear technology.

Mr. Duffy stated that the U.S. delegation appreciated the sensitivities. He stated that DOE-EM has nothing to do with weapons technologies. The technologies being developed by DOE-EM will have applications for the clean-up of contamination from industry, agriculture, as well as nuclear activities.

The two delegations agreed to meet on Friday November 9, 1990, to discuss the tours by the U.S. delegation and to prepare the Record of Meeting.



TUESDAY, November 6

Abstract of Activities for November 6, 1990:

A tour of STA's National Research Institute for Pollution (MITI, Tsukuba Science City) focused on the review of research on a water treatment system to promote biological treatment of hazardous chemicals; decomposition of CFC's through use of a thermal plasma reaction; measurements of organic pollutants in groundwater through fiber optics and the use of laser enhanced ionization methods for detecting trace amounts of inorganic pollutants; and remote sensing techniques using satellites for air pollution analysis. The laboratory appeared to be greatly under-utilized.

The delegation then traveled to Mito and visited the JGC Corporation's Oarai Center, where the following technology developments were reviewed: uranium removal from waste liquids; advanced cement solidification of hazardous wastes; on-site stabilization processes using fixing agents and cement or bentonite; automated waste-container inspection system; and incineration technologies for hazardous and radioactive waste.

Meeting with National Research Institute for Pollution & Resources (NRIPR):

Address:

NATIONAL RESEARCH INSTITUTE FOR POLLUTION & RESOURCES ONOGAWA 16-3 TSUKUBA SCIENCE CITY IBARAKI 305, JAPAN

Role of NRIPR:

NRIPR is 1 of 16 research institutes of the Ministry of International Trade and Industry (MITI); 9 are located within Ibaraki Prefecture and 7 are located elsewhere. The NRIPR has 9 departments conducting research in 4 overall areas: Resources, Industrial Safety, Energy, and Environmental Protection. The R&D budget for FY 1990 was 1,899 Million Yens (Approximately \$32.1 Million.) The total staff consists of 319 people, out of which 243 are technical officials.

The departments and the research activities are described on P.5 of the NRIPR brochure. (See Attachments) Major R&D activities of the NRIPR are:

- Energy: Development of new fuels as alternatives to oil, including oil shale and geothermal resources development.
- Coal: Gasification and liquefaction, and clean energy production.
- Fuel: Production of high quality natural gas, gas generation from bioma s in sludges, and extraction of useful material from sea.



- Combustion: Project on control of oxides of Sulfur and Nitrogen emissions, development of a system for high combustion efficiency, and heat pipes for clean energy.
- Resources: Materials processing, mining and geotechnology development.
- Materials: Extraction of rare earths, development of fine silicone particles, and deashing and desulfurization of coal.
- Mining: Recovery of manganese and cobalt from sea bottom, and research on geothermal technology
- Industrial Safety: Safety in coal mines and other industries (prevention of fires and explosions)
- Environmental Protection: environmental assessments, atmospheric environmental protection, and water pollution control. Examples of environmental assessments are: impacts of housing projects, land reclamation, discharges of pollutants to land and sea, including remote sensing of pollutants in land and sea. Examples of atmospheric R&D are: Decrease/eliminate atmospheric discharges of pollutants from industries, mr asurement and control of exhausts from automobiles, and prediction of pollutant movements. Examples of water pollution control are: control of pollutants from agriculture and industries, and wastewater treatment (biophysical and biological).

Participants:

Akira Takata Director

Osayuki Yokoyama Deputy Director

Akira Miyazaki Chief of Water Analysis Laboratory Water Pollution Control Department

Seiji Matsumoto Research Planning Office

Yasumasa Yamashita Director, International Cooperation Office

Masanao Hirai Director, Water Pollution Control Department



Shooichi Taguchi Interspheric Environments Lab.

Bibliography of Literature Received from NRIPR:

"Summary of National Research Institute for Pollution and Resources", NRIPR, 44 pages.

Agenda:

*Tour National Research Institute for Pollution and Resources (NRIPR)

*Attended Presentations at NRIPR on following topics:

- New Water Treatment System "Aqua Renaissance '90":

 Nitrogen Removal by an Activated Sludge Process with Cross-Flow Filtration
 Evaluation Technique for Organic Membrane Materials
- 2) Treatment of Industrial Types of Waste Containing Halogenated Organic Compounds
- Prediction of Groundwater Pollution

 Measurement of Pollutants
 Measurements of Pollutants in Groundwater
- Treatment and recovery of Biological Refractory Chemicals in Wastewater with Supercritical Fluid.
- 5) Biological Treatment of Hazardous Chemicals
- 6) Remote Sensing Technologies
- 7) Mechanisms of Environmental Pollution
- 8) Biomass Energy
- Waste Forms (Research on Solidification and Storage Techniques of High-Level Nuclear Wastes)
- 10) Research on High Performance Chemical Sensors
- Anti-Pollution Technology

 Research on Automation Techniques for Monitoring Pollution in Lakes



-Research on High-Performance Materials for Treatment of Hazardous Waste -Research on Optical Micro-Sensors for Gases

Detailed Meeting Notes:

Following the formal presentations, the DOE delegation went through a tour of the R&D facilities. The following is a brief description of the projects.

NEW WATER TREATMENT SYSTEM "Agua Renaissance '90"

By: Dr. Y. Urushigawa, Chief, Ecological Chemistry and Microbiology Laboratory, Water Pollution Control Department.

The goals of the project are: energy recovery, and increasing treatment efficiency to allow reuse of treated waste water. This project was initiated in 1985, as 6-year project, for meeting energy and water needs of Japan. The project is co-sponsored by 4 government organizations and about 20 industrial companies. The total budget of the project is about 12 Billion Yen (about \$1 Billion 1990 dollars). The project elements are: selection of microorganisms for activated waste treatment; development of efficient filter membrane which is resistant to deterioration by sewage and microorganisms; bioreactor for denitrification, and R&D for on production of microorganisms and organic material; R&D on high efficiency bioreactor for methane production; development of monitoring system; and design and pilot plant operation by the end of 1990.

The DOE group saw the pilot plant which consists of a 20 cubic meter reactor for treating 7 different types of simulated wastes. The filter membrane had microorganisms which survived on the nutrients in the sewage. The pilot plant has an H_2S control system, but did not have any need for CO₂ controls. The pilot plant consists of a vertical tank into which wastewater is fed from the bottom. Fermentation of the solids occurs in the tank and methane gas is collected from the top of the tank. Supernatant liquid is pumped off from the top of the tank; the liquid goes through membrane filtration and settling. The effluent from this system is ready for reuse.

CFC Decomposition By Thermal Plasma Reactor:

By: Dr. T. Wakabayashi, Senior Researcher, Organic Chemicals Laboratory, Fuel Department

Management of organic chemical, CFC in particular, is a part of a Global Environmental Program. The major effort of this program is to reduce environment emissic is that cause acid rain and global warming. CFCs are attributed to ozone layer depletion and the greenhouse effects.



The sources of CFC in Japan are: Production of CFCs which are used as solvents, propellants, refrigerants, and in semiconductor industries. The types of solutions are being worked: Development of degradable CFC, restrict emissions of CFC to the atmosphere, where possible reuse it, and "break up" CFC. CFCs used in the semiconductor industries cannot be reused because of the impurities in it.

The tour focused on thermal degradation of CFC. The NRIPR has built a pilot plant Thermal Plasma Reactor to treat 30 liters per hour of CFC. The reactor is heated by induction heating, developing maximum temperature of 10,000° C and up to 7,000° C in the reaction zone. Briefly, the process is: CFC is evaporated and mixed with water vapor. The mixture is thermally treated in the plasma reactor in the presence of Argon gas. Off-gases, consisting of HC!, HF, and water vapor, are treated in beds of KOH and CaO. Problems with the current system are: deposition of Carbon in the nozzles and efficiency of 70%. R&D is in progress to solve the carbon deposition problem and to improve the efficiency. Estimated cost of the process is 500 Yens (about \$4) per kg of CFC treated.

Remote Sensing Techniques for Air Pollution Analysis

By: Dr. S. Taguchi, Senior Researcher, Interspheric Environments Laboratory, Environmental Assessment Department

NRIPR is currently working on the Interspherometric Monitoring for Greenhouse Gases (IMG) project to remotely measure concentrations of gases in the atmosphere, that are associated with greenhouse effect. The gases under consideration are: water vapor, CO₂, oxygen, and chlorinated hydrocarbons. This project is sponsoged by MITI, with participation of many other organizations, such as STA. Sensors will be attached to a satellite ADEOS, which will be in synchronous orbit; the satellite is scheduled to be launched in February, 1996. Sensors are being developed by TOSHIBA. Gas concentrations will be measured as a function of the difference between transmitted and reflected lights.

Measurement of Pollutants in Groundwater

By: Dr. A. Miyazaki, Chief, Water Analysis Laboratory, Water Pollution Control Department

The NRIPR is developing remote sensors for characterization of contaminated groundwater. Rapid analytical methods will be used to monitor groundwater quality. The R&D consists of two parts: development of remote optical fibers for measuring organic chemical contaminants; and development of a laser ionization method for determining concentrations of inorganic pollutants. Sensors have been developed for measuring TCE in ppb range and chloroform in 200ppm range.

During the lunch with NRIPR, Dr Akira Takata, Director-General, said that Mr. Yasumasa Yamashita, Director, International Office will be the contact point.



Leo Duffy gave his presentation on EM's programs. Some of the points he discussed are briefly stated below:

- In the U.S., public hearings are held on power plant applications.
- In the environmental clean-up and restoration of ecology programs, DOE has problems with heavy metals, contamination of groundwater from buried wastes.
- We do not want to pass the problems to future generations.
- We are looking for cheaper and efficient extraction technologies to remove the contaminants.
- Dr. Donald Alexander is developing a 3-D model for contaminant transport in groundwater.
- We need to get more analytical capabilities to measure environmental releases in low concentrations. These measurements are to be made for demonstrating regulatory compliance and are expensive.
- The technologies DOE develops would be passed to industry. DOE would like to transfer some of these cleanup technologies to Eastern Europe.
- DOE has problems with chlorinated compounds. Dr. Donald Oakley said that groundwater contamination by TCE is at all DOE sites. The pump and treat method is not suitable. DOE is drilling horizontal wells and is vaporizing the volatile organic contaminants. Some U.S. sites are using stripping columns packed with activated carbon; this method is very expensive, including the cost of treating carbon. The first treatment system, utilizing horizontal drilling and vapor extraction, is being done at Savannah River Site. This system was alle to reduce concentrations to below detectable limits (less than 2000ppb) in 90 days. Vie are looking into screening systems (biological or otherwise) also.
- We have looked into the use of ultra-violet light and catalysts (Hydrogen peroxide) to break down hazardous chemicals, but the system is not very efficient and is costly.

The cheapest solution is in situ microorganisms with the appropriate nutrients.

Briefly described RDDTE Plan.

Q. Mr. Duffy asked a question regarding materials extraction technology development, the response was that no research on materials extraction was being carried out because Oak Ridge has 500,000# of mercury in soil.

Q. Mr. Duffy asked if there were any problems with heavy metals in sewage. A. Except Tokyo, there are no problems.

Meeting with JGC Corporation:

Address:

JGC CORPORATION ENGINEERS & CONSTRUCTORS OARAI RESEARCH & DEVELOPMENT CENTER NUCLEAR & ADVANCED TECHNOLOGY DIVISION 2205, NARITACHO OARAIMACHI, HIGASHIIBARAKI-GUN IBARAKI PREFECTURE, 311-13 JAPAN PHONE: 03-279-5441

Participants:

Takao Nakajima Executive Vice President General Manager

Hiroshi Kuribayashi Director, Senior Deputy General Manager

Yasuhiro Moriya Manager, Oarai Research & Development Center

Norimitsu Koshiba Section Manager, Sales, No. 1 Sales Department

> Hiroshi Yamashita Manager, U.S. Marketing Dept. U.S. Project Operations

Mamoru Shibuya Oarai Research & Development Center

Takuro Yagi Manager, No. 1 Team Technology Development Department

Stephen D. Goetsch Market Development Coordinator No. 1 Team, Technology Development Dept.



Bibliography of Literature Received:

Advanced Waste Management Technologies, JGC Corporation, 100 pages.

Agenda:

- 1) Overview of Technologies to be Discussed.
 - RASEP/Uranium Chelate Resin
 - Soil Stabilization
 - Drum Inspection
 - Advanced Cement Solidification
 - Others, Included
 - 3H Separation by Column and Laser
 - Incineration
 - Induction Melting
 - Wet Oxidization
 - Radioactive Gas Monitoring Research
- 2) View Drum Inspection System in Operation.
- 3) View Induction Melting System.
- 4) View ³H Separation Column.
- 5) Transit to Cold Pilot Plant Building.
- 6) View Wet Oxidization Pilot Plant.
- 7) View High Temperature Incinerator.
- 8) Pass Through Cold Building Pointing Out Other Pilot Plant Installations.

Detailed Meeting Notes:

Dr. Kuribyashi opened the meeting and made the following remarks:

-JGC is modifying or developing technologies for treatment of mixed waste, with specific emphasis on mixed waste monitoring.

-JGC has developed some technologies that could be applied to US through US companies. -During the visit of the US delegation, there will be some discussion of the technologies and a visit of the plant.



Mr. Moriya briefly described the facilities included on the tour:

Cold Test Facility, Radioisotope Test Buildings & Pilot Plant Test Building:

Initial waste management R&D, using non-radioactive material, is conducted in the Cold Test facility. This facility is equipped with chemical and physical analytical capabilities, as well as electron microscope, plasma emission spectrochemical analyzers, etc.

The radioisotope test facility is used for radioactive waste treatment R&D, as well as for use of radioisotopes for liquid wastes generated at the Center, and demonstration of compliance with environmental regulations. The Center is not permitted to discharge any liquid wastes containing radioisotopes (allowable radioactivity is less than 10⁻⁷mC/liter. JGC reuses all waste water.

High Temperature Demonstration Plant:

High Temperature Demonstration Plant. This plant directly converts mixtures of combustible and non-combustible radioactive wastes, including sludges, into stable granular form. This system could be used for mixed waste processing also. Waste material is shredded and fed into a cyclone furnace sludge incinerator/melter. High temperature, between 1400°C and 1500°C, is obtained by induction heating. The bottom ash and slag is removed in a molten state and cooled to form granular material. Off-gases are cooled and passed through ceramic and HEPA filters, and then discharged to the atmosphere.

Robotics:

Robot that moves on the floors as well as climbs over obstacles on walls was viewed and discussed.

Steve Goetsch made the following comments:

Soil Stabilization:

JGC submitted a proposal to INEL (August 1990) in response to PRDA for On-Site Stabilization Process (OSSP). The concept of the process is to stabilize by chemical and physical methods soils contaminated with heavy metals, mercury, uranium and TRU isotopes. The process uses a proprietary fixing agent (non-silicate compound + cementitious agent) to render the contaminates insoluble. The insoluble reaction products will be adsorbed into the soil. The advantages of this process are: low leach rates, and minor volume increases. This process has application to <u>Oak Ridge and Rocky Flats Plant</u>.

Uranium-Selective Chelate Resin Process:

This process utilizes a uranium selective chelate resin (UR-3100) for removing uranium from liquid wastes. This resin is highly selective for uranium, and the ion can be easily eluted



from the resin by using NaHCO₃ to give a concentrated solution of uranium. SN-3100 we been tested with a solution containing zinc, copper, and iron. The resin did not a solution or copper, and absorbed very small quantities of iron. Solutions containing uranium up to 71.1 ppm could be processed through this system. JGC proposed this system for Weldon Springs (did not receive any response; HydroPure got the contract); and has proposed the system for the Portsmouth Plant. Jerry Westerbeck from <u>Fernald</u> will be going to Japan to look at this system.

Advanced Cement Solidification Process:

The process consists of pretreatment and mixing with cement. This process has been used for immobilization of spent resins, boric acid wastes and incinerator fly ash and bottom ash. In the case of spent resins and boric acid waste, an additional step is introduced in the system - solid liquid separation or waste concentration. For Incinerator wastes, pretreatment consists of treatment with Ca(OH)₂ and NaOH, while for spent resins, cement and water are used. In case of boric acid wastes, pretreatment is provided by using calcium borate. The leachability of the immobilized waste form is very low. Tests with wastes containing cadmium or mercury showed that bentonite, instead of cement works better as an immobilization agent. This system has potential application to the mercury problems at Oak Ridge.

Automated Waste Container Management System:

The system consists of the following steps: Visual inspection unit - top, sides and bottom of waste drums are inspected by three ITV cameras; Surface Contamination and Dose Rate Measurement Unit - Smear samples are taken from four locations of the drums to determine surface contamination density. At the same time surface dose rates are measured by sensors built into the smear sampling unit; Radioactivity/Weight Measurement Unit - Nondestructive determination of radioactivity concentrations of each radionuclide in the drum are made by spiral scanning Ge semi-conductor sensor, a plastic scintillator (radioactivity concentration is determined from total dose - plastic scintillator , g scaling factors, and nuclides), and weights are determined by a simple machine; and Labeling Unit - Identification labels with a serial number and surface dose rates are automatically printed on the drums. TRU radioassay and free liquid detection systems are available, but have not been incorporated in the system. JGC submitted a proposal in response to INEL PRDA. This system could be potentially used for analysis gas in head space of drums to be disposed of at <u>WIPP</u>.

Incinerators:

Different types of incinerators at JGC were very briefly described: rotary kiln for hazardous waste, radwaste, irradiated carcasses, and medical waste.





WEDNESDAY, November 7

Abstract of Activities for November 7, 1990:

A visit to the PNC's Tokai Works focused on R&D activities related to the (1) glass vitrification of high-level waste; (2) processing of transuranic waste through a process of incineration, and acid digestion, with the production of metal ingots, ceramics and liquid residues that are discharged into the sea; and (3) treatment of solid and liquid low-level waste through incineration, compaction and filtration processes. Presentations were also provided on environmental monitoring programs at the Tokai Works, and, upon request, the facility's building that houses a high-level waste tank was toured.

Meeting with Power Reactor and Nuclear Fuel Development Corporation, Tokai Works:

Address:

TOKAI WORKS, PNC MURAMATSU, TOKAI-MURA 319-11 IBARAKI-KEN, JAPAN

Role of PNC:

Plays a central role in developing fuel cycle technologies and fast breeder reactor and ATR. PNC has developed technologies for prospecting for uranium deposits, refinement and conversion, centrifugal uranium enrichment, spent fuel reprocessing and radioactive waste disposal and is preparing to cooperate with industry for demonstration and utilization of these technologies.

Participants:

Tanehiko Yamanouchi Director

Y. Kishimoto Director Technology Development, Coordination Division

> Yoshiro Asakura Director Waste Plants Operation Division



Takao Tsuboya Director Waste Technology Development Division

> Ken-Ichi Matsumoto Deputy Director

Shin-Ichiro Torata Assistant Senior Engineer Waste Tech. Devel. Division

Hiroyuki Umeki Manager, Geological Isolation Technology Section Waste Technology Development Division

Bibliography of Literature Received:

Budget Information from PNC, Tokai. PNC, Tokai, 1 page.

"Nuclear Fuel Cycle, Tokai Reprocessing Plant", PNC- Tokai Works, 15 pages.

"Present Status of R&D Activities on HLLW and TRU Conditioning in Tokai Works", PNC Tokai Works, 26 pages.

"Some Aspects of Natural Analogue Studies for Assessment of Long Term Durability of Engineered Barrier Materials, CEC 4th Natural Analogue Working Group Meeting", Prepared by Y. Yusa, G. Kamei and T. Arai, PNC, 19 pages.

"Tokai Vitrification Facility", PNC, Tokai Works, 3 pages.

"Pu-Contaminated Waste Treatment Facility", PNC, Tokai Works. 3 pages.

Agenda:

- 1) Attended Presentations at PNC Tokai Works on following topics:
 - Present status of R&D Activities on HLLW and TRU Waste Conditioning in Tokai Works. (T. Tsuboya/PNC)
 - Present Status of Waste Treatment Facilities in Tokai Works (Y. Asakura/PNC)
 - Present Status of other activities on Nuclear Fuel cycle in Tokai Works (Y. Kishimoto/PNC)



- 2) Tour of Tokai Vitrification Facility (TVF).
- 3) Tour of Engineering Demonstration Facility III (EDF-III).
- 4) Tour of Engineering Testing Facility (ETF).
- 5) Tour of Chemical Processing Facility (CPF).
- Tour of Plutonium Contaminated Waste Treatment Facility (PWTF) and Plutonium Contaminated Waste Storage Facility (PWSF).

Detailed Meeting Notes:

Director Tanehiko Yamanouchi gave a general introduction and overview of Tokai operations.

Mr. Duffy thanked the Tokai Directors for inviting the DOE-EM delegation to the Tokai plant.

T. Tsuboya presented information describing the status of the R&D activities on HLLW and TRU waste. Information relating to the Organization of PNC including staff and budget is given on page 2 and 3 of the booklet for the Power Reactor and Nuclear Fuel Development Corporation, PNC. The 1989 PNC budget is 2,3x10¹¹ yen or 1.8x10⁶ dollars with a 1989 staff of 2800. Activities discussed were uranium enrichment, reprocessing, plutonium fuel, and waste management.

Uranium Enrichment:

Uranium Enrichment method consisted of gas centrifuge enrichment where UF⁶ gas containing U-234 and U-238 is centrifuged. The gas centrifuge has shown to be more efficient. Presently the Japan Nuclear Fuel Industries is constructing a plant that will have a capacity of 1,500 tons SWU per year. For more information see page 9 of the PNC booklet.

Reprocessing Technology:

Reprocessing technology to recover uranium and plutonium from spent fuel are being investigated at its Tokai Reprocessing Plant. This plant opened in 1977 and has reprocessed 392 tons of spent fuel, this includes 5.2 tons of MOX fuel from ATR Fugen. Note they had initial problems with the acid recovery evaporator and dissolvers. The short-term goal is to operate the plant on a 90-ton per year basis to recover plutonium for FBR Monju, which is scheduled to reach criticality in 1992. The flow diagram of the LWR Spent Fuel Reprocessing is given on page 11. \$2


Plutonium Fuel Production:

The production of plutonium fuel for new reactors such as the ATR and FBR is produced by combining the plutonium with uranium to form a mixed oxide (MOX) fuel. The MOX fuel is produced by mixing the nitrates of plutonium and uranium and converting to the oxide by using microwave-heating to decompose the mixed uranium and plutonium nitrate to a mixed uranium and plutonium oxide. This system is automated and went into operation at the Tokai Plutonium Conversion Development Facility with a daily co-conversion capacity of 10kg MOX. PNC has fabricated more than 100 tons of MOX fuel and more than 40,000 fuel rods have been successfully irradiated as of March 1989. At the Tokai Plutonium Fuel Production Facility they have the capacity of 5 tons of MOX fuel per year and another line is now under construction that will supply 40 tons of MOX fuel per year. Flow diagram of the process is on page 13.

Waste Management:

Waste Management activities are described on page 14 of the PNC booklet. Briefly, they consist of research and development in vitrification using the liquid-fed ceramic melter; processing of TRU waste by incineration, melting the ash with microwave, and melt metal waste by electro slag remelting; and conducting extensive research and development related to long-term management of high-level radioactive waste.

HLLW and TRU:

T. Tsuboya indicated that he gave an extensive presentation to Larry Harmon and thus gave a brief presentation of the R&D activities on HLLW and TRU. Title of the presentation is Present Status of R&D activities on HLLW and TRU Waste conditioning in Tokai Works. Material presented was: R&D in the areas of Vitrification by LFCM and HLLW; Nuclide separation from low level liquid waste and decomposition and nuclide separation from spent solvent for TRU waste: Demonstration phases of Pu-Contaminated Waste Treatment Facility, Bituminization Demonstration Facility and Solvent Waste Treatment Facility.

Vitrification:

In the area of Vitrification a flow diagram was included in the presentation; it is important to note that this process includes glass fiber addition. The delegation noted during the tour of the vitrification facility that the facility was nearly identical in design to the <u>Savannah River</u> <u>Defense Waste Processing Facility (DWPF)</u> but approximately 20-30% larger. The Tokai vitrification facility is 1 to 2 years behind the DWPF plant. Interior stainless steel walls were not finished in the same manner as the DWPF plant.



LL Liquid Waste Treatment:

In the area of Low Level Liquid Waste Treatment silver nitrate and sodium sulfite are added to the low level liquid waste. The silver precipitates the iodide ions present, the silver iodide is removed via ultra filtration. Sodium hydroxide and ferric nitrate are then added to the supernatant and plutonium, uranium and the fission products are precipitated and removed via ultra filtration. Ion exchange is used to remove the cesium and strontium ions. Sodium is removed by evaporation and the liquid is released to the sea. The <u>Spent Solvent</u> <u>Oxidative Decomposition Process</u> consists of oxidizing the solvent with hydrogen peroxide in the presence of a copper (II) oxide catalyst producing carbon dioxide, water, and calcium hydrogenphosphate. Lanthanum (III) is then added to precipitate the plutonium phosphate.

Waste Treatment Facilities (PWTF and LWTF):

Y. Asakura, presented "Present Status of Waste Treatment Facilities in Tokai Works". This presentation basically covered the Pu-Contaminated Waste Treatment Facility (PWTF) and the Low Level Waste Treatment Facility (LWTF). The PWTF uses conventional incineration for metals, HEPA filters and cellulose. The ash is melted and converted to ceramics and the metals are melted and converted to an ingot. The PVC chloroprene is shredded, followed by cyclone incineration or acid digestion; the ash is melted and converted to ceramics. A flow diagram is included in the presentation. In the WTF for solids the combustibles are incinerated by open incineration, the Cl containing materials are incinerated by closed incineration, HCL is collected, the ash is melted and converted to ceramics. The non-combustibles are cut, lecontaminated and melted into ingots or pressure compacted. The low level liquid waste undergoes ultrafiltration, absorption, and is solidified. The flow diagram for these processes is also included in Asakura's presentation.

Pu Monitoring in Surface Waters:

Y. Kishimoto reviewed the Present Status of Other Activities on Nuclear Fuel Cycle in Tokai Works. He identified the sampling and monitoring points off site, the sampling points for surface water and sediment, and the analysis and measurement methods they are using. Included in his presentation was the concentration of Pu-239 and 240 in sea sediments off shore of Tokai-mura.

The Plant Tour from 10:00 to 12:00 consisted of the Tokai Vitrification Facility, the Engineering Demonstration Facility-III, the Engineering Test Facility and the Chemical Processing Facility. From 1:30 to 2:30 pm the delegation was given a plant tour of the PWTF (Pu-contaminated Waste Treatment Facility) by Y. Asakura.





Meeting with PNC, Oarai Engineering Center:

Address:

POWER REACTOR & NUCLEAR FUEL DEVELOPMENT CORPORATION OARAI ENGINEERING CENTER 4002, NARITA OARAI, IBARAKI, JAPAN PHONE: (0292)67-4141

Role of PNC:

Plays a central role in developing fuel cycle technologies and fast breeder reactor and ATR. PNC has developed technologies for prospecting for uranium deposits, refinement and conversion, centrifugal uranium enrichment, spent fuel reprocessing and radioactive waste disposal and is preparing to cooperate with industry for demonstration and utilization of these technologies.

Basically, the mission of OARAI is to conduct research and development of key technologies associated with FBR and ATR power plants. This includes design studies for framing plant systems with safety and economic competitiveness, research and development on base technologies and innovative technologies, research and development using experiences obtained through construction and operation of Joyo, Monju, and Fugen, and research and development on fuel recycling (this includes research in the area of transmutation.)

Participants:

Mitsuru Kamei Deputy Director Technology Development Division

> Kyoichiro Suzuki Deputy Director Oarai Engineering Center

Hidehiko Miyao General Manager Waste Management Section

Hiroki Kanemaru Manager, Administration Div. PNC Oarai Engineering Center



Masao Shiotsuki Senior Research Engr., Waste Mgmt. Sec. Oarai Engineering Center

> Shigeyoshi Kawamura Waste Management Section Oarai Engineering Center

Masahiko Itoh Research & Devel. Coordination Section Technology Development Division

Bibliography of Literature Received:

"Development of a Heat Resistant and Angle Beam Type Electro-Magnetic Acoustic Transducer", Compiled by K. Ara, H. Rindo, K. Nakamoto, T. Doi, K. Morimoto, and T. Sakamoto, Oarai Engineering Center, PNC, 5 pages.

"Development of Decommissioning Technologies for Nuclear Fuel Cycle Facility in Waste Dismantling Facility", Oarai Engineering Center, PNC, 18 pages.

"Research and Development in Oarai Engineering Center", Oarai Engineering Center, PNC, 16 pages.

Agenda:

1) Arrive Oarai Engineering Center

2) Tour Waste Dismantling Facility (WDF).

Detailed Meeting Notes:

Waste Dismantling Facility:

H. Miyao at OARAI presented the activities of the Waste Dismantling Facility. The efforts in decontamination included dry ice blasting (ice-blasting), electo-polishing, and redox processes. In the area of dismantling they were using plasma cutting, robotics, and laser cutting. In the area of monitoring they were using radiation image display through remote measurement.



Decontamination and Decommissioning:

S. Kawamura, discussed the technologies that were needed to decontaminate and decommission a facility. These needed technologies are in the areas of monitoring, radiation control, decontamination, dismantling, remote handling, waste treatment, and system engineering.

For the tour of the facility they distributed ear phones to the DOE contingency. This was extremely helpful for the people standing at the back of the group. Technologies viewed on the tour consisted of ice-blasting, electropolishing, the alpha facility, laser cutting, monitoring, Hot Isostatic Pressure (HIP), robotics, fiber optics, and electromagnetic acoustic sensors.

THURSDAY, November 8

Abstract of Activities for November 8,1990:

A tour of JAERI Tokai Research Institute facilities included review of safety evaluations of long-term storage and disposal of high-level wastes (with research focusing on confinement ability and durability of materials), reactor decommissioning and actual dismantling activities, the use of robotics, and heat/radiation resistant fiberscopes.

Meeting with JAERI, Tokai-Mura

Address:

JAPAN ATOMIC ENERGY RESEARCH INSTITUTE TOKAI RESEARCH ESTABLISHMENT TOKAI-MURA, IBARAKI-KEN JAPAN PHONE: 0292-82-5410

Role of JAERI:

JAERI is a semi-governmental research organization which implements national long-term programs in nuclear energy, including joint projects and international cooperative efforts.



Research Activities:

- R&D of nuclear energy, nuclear safety, high temperature gas-cooled reactors, nuclear fusion, radiation applications, nuclear powered ships, basic research, decommissioning of nuclear reactors;
- Design construction and operation of reactors;
- Education and training of researchers and engineers in the field of nuclear energy; and
- Dissemination of information obtained through R&D activities.

Participants:

Kakuzo Tomii Director Department of JPDR

Satoshi Yanagihara Senior Engineer Decommissioning Tech. Lab.

Tsutao Hoshi General Manager Reactor Decommissioning Op. Div.

Eiji Shirai Deputy Director, Department of Research Reactor Operation

> Yoshiki Wadachi Deputy Director Department of Env. Safety Research

Susumu Muraoka Head, Engineered Barrier Materials Laboratory

Bibliography of Literature Received:

"Development of Technologies on Decommissioning of Nuclear Fuel Cycle Technologies," Japan Atomic Energy Research Institute. 5 pages.



- "JPDR Decommissioning Program", written by T. Hoshi from the 9th TAG Meeting on October 8-12, 1990 at the Japan Atomic Energy Institute. 10 pages.
- "New JRR-3, Compiled by the Research Reactor Operation at the Tokai Research Establishment", Japan Atomic Energy Research Institute. 6 pages.
- "Progress Report on Safety Research of High-Level Waste Management for the Period April 1988 to March 1989", Edited by Haruto Nakamura and Susumu Muraoka, Department of Environmental Safety Research, Tokai Research Establishment, Japan Atomic Energy Research Institute, 74 pages.
- "Reactor Decommissioning Technology Development and Actual Dismantling of JPDR," compiled by the Tokai Research Establishment, Japan Atomic Energy Research Institute. 9 pages.
- "Safety Studies on Glass Waste Form", written by S. Muraoka at Japan Atomic Energy Research Institute. 10 pages.
- "Summary of WASTEF Facility", from Japan Atomic Energy Research Institute. 10 pages.
- "Volatilization of Cesium from Nuclear Waste in a Canister", Hiroshi Kamizono, Shizuo Kikkawa, Shingo Tashiro and Haruto Nakamura. at Japan Atomic Energy Research Institute. Department of Environmental Safety Research, 6 papers.

Agenda:

- 1) Visit JAERI Tokai Research Facility.
- 2) Tour of JRR-3 (Japan Research Reactor No. 3).
- Tour of Japan Power Demonstration Reactor-BWR TYPE (JPDR).
 -R&D for decommissioning technology.
- Tour of Waste Safety Testing Facility (WASTEF).
 -Glass solidification technology for high-level radioactive wastes management.

Detailed Meeting Notes:

Decomn issioning of JPDR;

The delegation was given a short overview of the JPDR decommissioning program which is continuing through Jim Fiore and Bill Murphie (EM40). Mr. Hoshi is responsible for



decommissioning of the JPDR. The program started in 1981. The first phase completed in 1986 involved technology development for cutting and disassembly and waste management. Phase II involving actual decommissioning started in 1986. Equipment surn and fing the reactor vessel was removed by 1989. Reactor internals were removed by 1909 and the pressure vessel was projected for removal in 1990. Removal of the biological shield concrete is projected by 1992 and site restoration should be completed in 1993. Mr. Duffy asked where the materials were being stored. All materials are being stored on site at Tokai.

Concrete Disposal:

Two thousand tons of concrete are proposed to be stored in shallow land burial sites. The Japanese are exploring ways to reuse minimally contaminated concrete.

Dismantling:

All dismantling of the biological shield is remote. Dose levels for workers in the U.S. and Japan is 5Rem in special cases. Plasma arc cutting is conducted underwater to prevent release of fumes. Shaped explosives are used to minimize air contamination. A central vacuum system is used to remove dust from cutting and blasting operations. The arc cutter system has been adopted from the U.S. The U.S. typically uses a vacuum system at the cutter head in addition to the central vacuum system with appropriate in line filters.

Studies of Activated Metals:

Studies of steel are being conducted to determine the extent of corrosior, and to evaluate "cracking" and embrittlement, particularly in weldment materials.

FRIDAY, November 9

Abstract of Activities for November 9,1990:

The delegation visited the JAERI Radiation Chemistry Research Establishment in Takasaki where presentations focused on research initiatives related to the practical application of nuclear energy (e.g., irradiation of food products to prevent spoilage). Of particular interest were studies on uranium extraction from seawater. The development of a new accelerator lab will depend heavily on the ability to attract foreign scientists committed to live with their families at the site in a foreign-scientist community development.

The delegation then traveled to Tokyo for a final meeting with PNC. A record of meeting was signed, and the delegation agreed that Dr. Donald H. Alexander would be assigned the lead for working with the Japanese to formalize cooperative R&D initiatives. The PNC close-out discussion focused on R&D initiatives of particular interest to the U.S. including:





- Application of Japanese decontamination and decommissioning techniques.

- Waste vitrification.

- Fiber-optics.
- Extension of studies of uranium extraction from seawater to contaminated groundwater.
- Robotics.
- Development of a "profile system" for tracking developments in international wastemanagement R&D initiatives.
- Exchange of scientists and students.

Meeting with JAERI Takasaki Radiation Chemistry Research Establishment:

Address:

TAKASAKI RADIATION CHEMISTRY RESEARCH ESTABLISHMENT JAPAN ATOMIC ENERGY RESEARCH INSTITUTE 1233 WATANUKI-MACHI, TAKASAKI GUNMA, 370-12, JAPAN PHONE: 0273-46-1211

Role of JAERI:

JAERI is a quasi-governmental research organization which implements national long-term programs in nuclear energy, including joint projects and international cooperative efforts.

Research Activities:

- R&D of nuclear energy, nuclear safety, high temperature gas-cooled reactors, nuclear fusion, radiation applications, nuclear powered ships, basic research, decommissioning of nuclear reactors;
- Design construction and operation of reactors;
- Education and training of researchers and engineers in the field of nuclear energy; and
- Dissemination of information obtained through R&D activities.

Participants:

Sueo Machi Director General

Waichiro Kawakami Deputy Director Department of Development



Shoichi Sato Director, Department of Research

Isao Ishigaki General Manager Radiation Processing Devel. Laboratory Department of Development

Bibliography of Literature Received:

- *Amidoxime-Group-Containing Adsorbents for Metal Ions Synthesized by Radiation-Induced Grafting*, Written by J. Okamoto, T. Sugo, A. Katakai and H. Omichi. JAERI, Takasaki Radiation Chemistry Research Establishment, 11 pages.
- "A New Type of Amidoxime-Group-Containing Adsorbent for the Recovery of Uranium from Seawater", Written by H. Omichi, A. Katakai, T. Sugo and J. Okamoto. JAERI, Takasaki Radiation Chemistry Research Establishment. 6 pages.

Agenda:

1) Tour Takasaki Radiation Chemistry Research Establishment.

Detailed Meeting Notes:

Activities at RCRE are centered upon the use of Co-Go and electron beam irradiation facilities; principal activities are the 1) synthesis and modification of polymers for industrial and medical applications, 2) environmental protection, and 3) development of materials for nuclear/space applications.

Industrial Applications:

The most interesting recent development is the irradiation of a chelating membrane (5 micron thickness), which has been used to absorb uranium from sea water (2 gr U/Kg in 20 days). Uranium is desorbed from the membranes at ~pH 2. Dr. Alexander suggested that the use of polyethylene beads may be more efficient for certain applications. He noted that polyethylene beads have been used as tracers and have been pumped through groundwater systems. We discussed the potential use of micron-sized U-adsorbent beads to remediate uranium-and heavy element-contaminated ground water through vertical/horizontal wells. Adsorbant micro-spheres are being developed at MITI, Osaka. Other industrial applications included the development of new slow-release drugs, vulcanization of latex (which leads to less incineration residue and SO² emissions), and curing of various coating materials.





Environmental Applications:

RCRE has developed technologies for the treatment of flue gases, municipal wastewater and sewage sludge. The latter two applications, although effective, have also met resistance in Japan. RCRE is evaluating electron beam irradiation of flue gases (SO² and NO^x) in the presence of ammonia. The resulting ammonium nitrate/sulfate solids can then be removed by an electrostatic precipitator.

Nuclear/Space Applications:

Irradiation of materials that are used for nuclear and space applications is performed at RCRE. This includes wire, cable, and insulating materials.

Scientific Exchange:

EM programs should consider scientific exchange with JAERI-Takasaki. STA offers scholarships for studies at these facilities. Irradiation of chelating membranes for adsorption of actinides, such as uranium or adsorption of organics such as trichloroethylene could be critical to meeting EM objectives.

Meeting with PNC, Tokyo

Address:

POWER REACTOR & NUCLEAR FUEL DEVELOPMENT CORPORATION (PNC) 9-13, 1-CHOME, AKASAKA MINATO-KU, TOKYO, 107 JAPAN

Participants:

Takao Ishiwatari President

Yoshikazu Hashimoto Executive Director

Masao Yamamoto Deputy Senior Director

Saburo Kikuchi Secretary to the President

Kiyoshi Kikuchi Ing. Geologue, Department des Ressources Nucleaires



Tadashi Mano General Manager Conditioning Research Program Radioactive Waste Management Project

Takashi Yoshikawa Manager, International Cooperation Office International Division

Reiko Nunome International Cooperation Office International Division

Akira Wadamoto Engineer, Conditioning Research Program Radioactive Waste Management Project

Bibliography of Literature Received:

"FBR Development in PNC for Commercialization', PNC, 8 pages.

"Technical Draft for Comments RD&D Program on Low-Level TRU Bearing Waste Management Technologies", PNC, 43 pages.

Agenda:

*Meeting with PNC President.

Detailed Meeting Notes:

DOE and PNC agreed that several workshops would be arranged over the coming months to focus on areas of joint technical interest in preparation for the Bilateral Coordination Meeting in the spring of 1991. DOE appointed Dr. Donald Alexander as the DOE-EM coordinator. PNC appointed Mr. Takao Yagi as PNC-EM coordinator.

The workshops will provide reports recommending areas of technical collaboration to the DOE-EM/PNC Coordinating Committee in the Spring of 1991. The U.S. delegation identified several technical areas for continued collaboration in the November 9th meeting with PNC including decontamination and decommissioning, vitrification and TRU handling, treatment and disposal.

Areas of particular interest to the delegation which will be considered for future collaboration with PNC include:





- the use of the plasma-arc saw for dismantling reactor vessels and related contaminated structures;
- 2. smelting of slightly contaminated ferrous metals for recycling;
- 3. methods for removal of contaminated concrete;
- 4. methods for reusing slightly contaminated concrete;
- 5. methods of waste reduction and minimization;
- 6. partitioning and transmutation;
- 7. robotics;

Areas of particular interest to the delegation which will be considered for future collaboration with MITI, JAERI and Kyoto University include:

- 1. methods for the removal of uranium from seawater;
- 2. applications of fiber-optics and lasers for in situ analysis of groundwater;
- 3. methods for the removal of organics such as trichloroethylene;
- 4. simulation modeling of groundwater contaminant migration;
- 5. research on actinide chemistry;
- 6. methods of underground characterization;
- 7. inorganic microencapsulated adsorbents;
- 8. optical microsensors for gases; and,
- 9. gold-metal oxide catalysts for sensor applications.



RECORD OF MEETING BETWEEN THE UNITED STATES DEPARTMENT OF ENERGY AND THE POWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION OF JAPAN IN THE RADIOACTIVE WASTE MANAGEMENT NOVEMBER 9, 1990 TOKYO

In accordance with the terms of the Agreement between the United States (U.S.) Department of Energy (DOE) and the Power Reactor and Nuclear Fuel Development Corporation of Japan(PNC) in the Area of Radinactive Waste Management, representatives of the two organizations met in Tokyo on November 9, 1990.

A technical delegation led by Mr. Leo Duffy, Director, Office of Environmental Restoration and Waste Management (EM), Department of Energy (DOE), met with officers and staff of the Power Reactor and Nuclear Fuel Development Corporation of Japan(PNC), to discuss potential areas of possible technology exchange during 1991. The wrap-up meeting was convened after participants completed tours of PNC sites and facilities.

Mr. Ishiwa ari, President, PNC welcomed Mr. Duffy and the U.S. delegation provided a brief overview of the status of PNC waste management activities. He encouraged continued technology exchanges.

Mr. Duffy thanked PNC for their hospitality and efforts in making the arrangements to visit PNC sites and facilities. He provided a brief overview of the status of the DOE-EM program and underscored his continued desire to pursue exchanges of technology.

DOE and PNC agreed that several workshops would be arranged over the coming months to focus on areas of joint technical interest in preparation for the Bilateral Coordination Meeting in the spring of 1991. DOE appointed Dr. Donald Alexander as the DOE-EM coordinator. PNC appointed Mr. Takao Yagi as PNC-EM coordinator.

Dr. Alexander and Mr. Yagi will meet in Washington, D.C. in December 1990, to define workshops in common areas of interest.

9 Nov 1990

Dr. Donald Alexander, DOE

9. Nov. 1990_

Mr. Kiyoshi KIKUCHI, PNC

SATURDAY, November 10

Abstract of Activities for November 10, 1990:

A meeting was held with Professor Higashi of the Kyoto University, where the delegation discussed actions taken by the DOE under the "Ten-Point Plan" and the development and implementation of the DOE's program for environmental restoration, waste management and related R&D under the "Five-Year Plan". Professor Higashi expressed great interest in the status of the U.S. high level waste repository program and in the concept of a student exchange. The University Staff expressed interest in pursuing opportunities for joint R&D projects related to neptunium chemistry.

Meeting at Kyoto University:

Address:

KYOTO UNIVERSITY DEPARTMENT OF NUCLEAR ENGINEERING YOSHIDA, SAKYO-KU KYOTO 606, JAPAN TEL 075-753-5831

Participants:

Kunio Higashi Professor

Hirotake Moriyama Associate Professor

Kazukuni Shimoura Associate Professor

Tetsuji Yamaguchi Faculty of Engineering

M. I. Pratopo Faculty of Engineering

Ichizou Kokaji Chief of Reprocessing Section



Agenda:

- 1) Meeting with Professor Higashi, Kyoto University.
- 2) Discussed following topics:
 - Yucca Mountain Pioject Overview
 - High Level Repository Siting
 - Treatment of Slightly Contaminated Soils

Detailed Meeting Notes:

Site Characterization and Remediation

Professor Higashi introduced faculty members and students. Mr. Duffy gave an overall presentation on the EM program and emphasized technology needs. Mr. Duffy invited Dr. Alexander to give a brief overview of performance assessment and site characterization activities at Yucca Mountain. Following the Yucca Mountain presentation, the participants discussed the three issues raised by Dr. Higashi:

- A brief explanation on the present status of Yucca Mountain Project (Conceptual Design of the repository, and related matters).
- 2) Permanent disposal of HLW.
- 3) Treatment of soils which are alightly contaminated with metal oxides.

Neptunium Chemistry:

Dr. Alexander opened discussion on the Kyoto University research in neptunium chemistry. Several new compounds which raise the solubility limit of neptunium in groundwater were mentioned. This information could have a significant impact on transport calculations for neptunium in groundwaters. Copies of recent publications are being acquired.

MONDAY, November 12, 1990 Ascension Day



TUESDAY, November 13, 1990

Abstract of Activities for November 13, 1990:

The delegation visited the PNC Chubu Works where presentations were provided on R&D activities conducted at the Tono Uranium Mine to support the development of geologic disposal technologies. Activities reviewed included the performance of engineered barriers and corrosion testing; geochemical investigations of groundwater; and the development and validation of migration models of radionuclides. PNC representatives noted strong public opposition to the development of a geologic repository in the area.

Meeting with FNC, Chubu Works:

Address:

CHUBU WORKS POWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION OFFICE 959-31, SONODO, JYORINJI TOKISHI, GIFUKEN, JAPAN PHONE: 0572-54-1271

Role of PNC, Chubu Works:

PNC Chubu Works is developing technical capability to characterize sites for the permanent disposal of CHLW.

Participants:

Yozo Sugitsue Director

Toshio Tomishige Deputy Director

Toshihiro Seo Geologist, Waste Isolation Research Section



Kozo Sugihara Research Engineer Waste Isolation Research Section

Shougo Fujita Manager, General Affairs Section

Bibliography of Literature Received:

- "Field Tour Guide for Tono Mine Gallery (Tsukiyoshi Deposit)," PNC Chubu Works, 28 pages.
- "Natural Analogue Study of Tono Sandstone Type Uranium Deposit in Japan," Written by C. Sato, Y. Ochiai and S. Takeda. Waste Management and Raw Materials Division, PNC-Chubu Works, 11 pages.
- "Natural Analogue Study of Tono Sandstone Type Uranium Deposit in Japan," Written by T. Seo, Y. Ochiai, S. Takeda and N. Nakatsuka. PNC-Chubu 6 pages.

Agenda:

- 1) Tour PNC Chubu Works.
- 2) Tono Mine Gallery

Stop 1- Engineered Barrier Materials Field Tests.
Stop 2,3- Hydrogeochemistry of Groundwater.
Stop 4- Geochemistry of Natural U-Th Series Nuclides Uranium Mineralization.
Stop 5- Mine-by Experiments of Excavation Responses.
Stop 6- Shaft Excavation Effect Project Site.

3) General Discussions re: Chubu Works Activities.

Detailed Meeting Notes:

The delegation was welcomed by Chubu Works Director Yozo Sugitsue and Deputy Diatestor Toshio Tomishige. We were told that Tono is a well known area for ceramics. The PNC office at Tono (Chubu Works) was established about 25 years ago with the discovery of uranium. The deposits represent approximately 2/3 of the known Japanese reserves.



Exploration has since shifted overseas and has proven to be more economic, especially in Canada, Australia, Africa and China. Since 1986 Chubu Works has been devoted to R&D.

Chubu Works has 5 sections: 1) administrative; 2) overseas exploration; 3) geologic research; 4) Uranium deposit evaluation; and 5) physics and chemistry technology.

The ore at Tono is 135 meters deep with 7500 meters of developed underground galleries. The stratigraphy consists of sandstone, mudstone, and conglomerates over a granite basement. The ore is covered by marine mudstones and clays.

PNC is conducting a variety of site characterization studies at Tono:

- 0 natural analogue studies;
- isotopic disequilibria; 0
- excavation response/disturbance studies; 0
- corrosion tests (glass and stainless steel) 0
- development of a hydrogeologic model; and 0
- fault studies. 0

The fault at Tono is a reverse fault with a displacement of approximately 30 meters and with an age of about 10 million years.

PNC has been conducting regional hydrogeologic studies at Shomasama where they maintain an extensive core library with several 1000 meters of core. Other important HLW site characterization studies are ongoing in Horonobe tuff.

Engineered Barrier Experiment:

Two stations in the underground facility are devoted to engineered barrier experiments. The first involves the long term evaluation of bentonite backfill in a weathered granite block. The principal approach is similar to the U.S. approach of vertical emplacement of the waste packages in the floors of the drifts. However, since the Japanese site is expected to be saturated, bentonite blocks would be emplaced around the container. The objective of the first station is to examine the durability and behavior of the barriers under field conditions and develop methods for monitoring and performance evaluation. The second station is designed to examine a range of barrier materials including glass, copper, bentonite, stainless steel and bentonite blocks. Experiments include waste glass leaching (heating test), overpack



materials corrosion test (heating test), and monitoring the migration of groundwater contaminents.

Nuclide Migration Tests:

The nuclide migration test is designed to examine uranium migration (and uranium series nuclides) along the Tsukiyoshi fault which has displaced the ore body by 30 meters. Geologic, mineralogic, geochemical, migration and retardation studies are being pursued. Groundwater chemistry tests indicate mixing of surface and deep waters through tritium studies. Uranium phases include andersonite $(Na_2Ca(UO_2)(CO_3)_3*10H_2O)$ and zippeite $(K_4(UO_2)_6(SO_4)_3(OH)_{10}*H_2O)$.

Excavation Response Tests:

Sever I drifts have been set aside for excavation response testing. Extensive cross hole testing and radial in-wall monitors are used to monitor excavation response. Measurements to date indicate minimal response.

Exploratory Shaft Construction:

An exploratory shafe is being excavated to the testing horizon. A radial set of instrumented boreholes are being used to monitor groundwater drawn down. Performance evaluations, similar to those at the <u>Canadian URL</u> and planned at <u>Yucca Mountain</u> are underway.

WEDNESDAY, November 14

Abstract of Activities for November 14, 1990:

The delegation visited the Government Industrial Research Institute in Osaka. The visit included a review of research on solidification and storage techniques (the research focused on glass vitrification and appeared to have been discontinued); water treatment systems for industrial waste using membrane materials and microorganisms; and pollution detection technologies using high-performance chemical sensors and optical microsensors for gases.

Meeting with MITI/AIST:

Address:

MITI/AIST GOVERNMENT INDUSTRIAL RESEARCH INSTITUTE, OSAKA MIDORIGAOKA 1, IKEDA, OSAKA PHONE: (0727) 51-8351



Participants.

T. Komiyama Director General

Takako Takahashi

Masatake Haruta Head of Catalysis Section

Teruo Kodama Director, Research Planning Office

Makoto Kinoshita Glass and Ceramic Materials Department

Steven N. Crichton Visiting U.S. Scientist, Glass Science Section Glass and Ceramic Materials Department

> Kunishige Higashi Senior Researcher

Tetsuhiko Kobayashi Research Chemist

Yoshiko Nakahara Director, Material Chemistry Department

Bibliography of Literature Received:

"AIST Summary", AIST, 35 pages.

"Aqua Renaissance '90 Project", National Research and Development Program, MITI, 6 pages.

"Budget, Staff and Scale Information", MITI, 7 pages.

"Fine Structure of Nobel Gold Catalysts Prepared by Coprecipitation", Written by M. Haruta, H. Kageyama, N. Kamijo, T. Kobayashi, and F. Delannay. GIRI-Osaka, 10 pages.



- *40th ISE Meeting -Extended Abstracts*, Volume 1, International Society of Electrochemistry, 3 pages.
- "Glass and Ceramics for the Future", Glass and Ceramic Material Department, GIRI-Osaka. 27 pages.
- "Gold Catalysts Prepared by Coprecipitation for Low-Temperature Oxidation of Hydrogen and of Carbon Monoxide", Written by M. Haruta, N. Yamada, T. Kobayashi, and S. Iijima. GIRI-Osaka, 9 pages.
- "Gold Supporting Tin Oxide for Selective Co-Sensing", Written by T. Kobayashi, M. Haruta and H. Sano. GIRI-Osaka, 4 pages.
- "Methodology for Making R&D Programs of Chemical Sensors", Written by M. Haruta, K. Hiiro, H. Tanigawa, H. Takenaka, S. Yoshikawa and H. Sano. GIRI-Osaka, 28 pages.
- "New Technology Japan", Vol. 17, No. 2, May 1989, 2 pages.
- "Outline of Researches", Government Industrial Research Institute, AIST, MITI, 9 pages.
- "Preparation and Catalytic Properties of Gold Finely Dispersed on Beryllium Oxide", Written by M. Haruta, K. Saika, T. Kobatashi, S. Tsubota and Y. Nakahara. GIRI-Osaka, 4 pages.
- "Preparation of Highly Dispersed Gold on Titanium and Magnesium Oxide", Written by S. Tsubota, M. Haruta, T. Kobayashi, A. Ueda, Y. Nakahara. GIRI-Osaka, 9 pages.
- *Proceedings- 9th International Congress on Catalysis", M. Haruta, T. Kobayashi and F. Delannay, GIRI-Osaka, 6 pages.
- "Proceedings of the 3rd International Meeting on Chemical Sensors", Cosponsored by the Edison Sensor Technology Center, Resource for Biomedical Sensor Technology, Electronics Design Center and Case Western Reserve University. 5 pages.

"Research on HLW Management in GIRI-Osaka", GIRI-Osaka, 2 pages.

"Selective Co Sensor Using Ti-DOPED Fe₂O₃ with Coprecipitated Ultrafine Particles of Gold". Written by T. Kobayashi, M. Haruta, H. Sano and M. Nakane. GIRI-Osaka, 11 pages.



Agenda:

- 1) Visit AIST/MITI's Government Industrial Research Institute in Osaka.
- 2) Dr. T. Komiyama outlined the research activities of the institute.
- 3) Dr. M. Kinoshita discussed solidification and storage techniques of high-level nuclear wastes. He also presented the first part of a presentation on the New Water Treatment System (Aqua-Renaissance '90 project).
- Dr. Y. Nakahara gave the second portion of the new water treatment system (Aqua-Renaissance '90 project) detailing micro-organisms. Dr. Nakahara also discussed anti-pollution technology.
- Dr. Masatake Haruta made presentations on the development of new catalysis for sensor technology and micro-encapsulated spheres for the adsorption of contaminants.

Detailed Meeting Notes:

The MITI Government Industrial Institute in Osaka was establishes in 1918. The Institue is well known for its work on batteries, metal alloys for storage of hydrogen, and development and adaptation of new compounds for optical applications. They have manufactured some large optical lenses (the delegation saw some display) and are conducting optical fiber research.

Perhaps one of the most promising efforts for collaboration with EM involves the efforts of the Glass and Ceramics Materials Department and Materials Chemistry Department to develop a wide range of sensors, adsorbants, and HLW glass waste forms.

GLASS AND CERAMIC MATERIALS DEPARTMENT

Glass and Ceramic Materials Development, Optical Glasses

The institute is developing glasses with high refractive index and low dispersion such as LaK and LaF utilized for lenses in the camera industry. They have succeeded in casting high quality discs 2m in diameter for telescopes with low thermal expansion.

Nuclear Waste Form Glass

Glass is being evaluated as a waste form for high level wastes. Chemical and physical properties, including ionic diffusion, electrical conductivity of the melt, thermal conductivity, volatilization, crystallization and phase separation of waste form glasses have been evaluated for over a decade. They have established glass compositions and melting technology necessary for solidification of waste glass. Basic research continues on phase separation,



nucleation and crystallization, volatilization, diffusion, electrical conductivity, and mixed alkali effect. They have also compared synthetic glass leachability with natural glasses.

Ion Conducting Glass for Sensors

The Institute has been investigating Lithium-ion-conducting glasses for sensor applications. Li enhances conductivity at 0.3 Li20*0.3B203*0.4Li2SO4.

Porous Glass for Biochemical Catalysis and Bioreaction

Porous glasses have been successfully developed for biochemical catalysis as carriers of enzyymes. Enzymes thus immobolized by adsorption on the porous glasses maintain high activity for a long period and may be used repeatedly.

The EM application may be the delivery of porous enzyme bearing glass beads (inorganic microcapsules) to the underground bioreactor zone. (Alexander)

New Water Treatment System (Aqua-Renaissance '90 Project)

The Institute is developing a series of membranes for water purification. The project is coordinated with MITI-Mito. For details see materials from Tuesday November 6, 1990.

MATERIAL CHEMISTRY DEPARTMENT

Chemical and Bio Sensor Technology

The Institute sponsors state-of-the-art research in sensor technology as the references attest. Emphasis is placed on the development of catalysts with an emphasis on gold plated metal oxides. The Institute is developing sensors for chemical detection in gas and liquids and enzyme immuno sensors. They have worked on multiple layer sensors. They are also working jointly with Belgium to develop sensors to detect gases such as CO

Catalysts for Gas Detection

Preparation of well defined metal compound catalysts is attempted by using homodisperse fine particles as a building block and by applying modern thin film techniques. Catalysts are characterized using EXAFS, XPS, and IR. The work is focused on the catalytic behavior of ultrafine gold particle and their interaction with the support oxides. These materials have widespread application as catalysts for low temperature oxidation of CO and as sensors for flammable gases. This technology is applicable to <u>Hanford tank characterization</u>.

Inorganic Shell Microcapsules

The Institute is developing inorganic porous spheres and inorganic-shell microcapsules for adsorption of contaminants. The chemical and physical properties governing adsorption can





be selectively controlled by preparation conditions. These microcapsules could have wide application for restoration activities.

Glass Composite Membranes

Containment of microbes in the reactor and reuse of the filtered water by separating microbes and pollutants from waste water is being researched. Glass and glass-ceramic membranes are being evaluated. A range of membranes are being selected for each reactor type.

Meeting with Kobe Steel:

Address:

KOBE STEEL, LTD. TEKKO BLDG. 8.2, MARUNOUCHI 1-CHOME CHIYODA-KU, TOKYO, 100 JAPAN PHONE: TOKYO (03) 218-6733

Participants:

Yoshimasa Yama 2000 Chief Manager, Sales & Marketing Nuclear Engineering and Equipment Department

Detailed Meeting Notes:

Waste Management, Incineration, Ash Melting, and Crud-Slurry Solidification

Kobe Steel provides technological support to PNC. Although we did not have a formal meeting scheduled, we met with <u>Mr. Yamamoto</u> who provided the delegation with information on their capabilities. They are involved in the management of alpha contaminated wastes, incineration and ash melting for plutonium-contaminated combustible wastes, and crud-slurry solidification systems. They have a U.S. Patent (4330698 My 18, 1982) on their Microwave Melter.









LEO P. DUFFY DIRECTOR, OFFICE OF ENVIRONMENTAL RESTORATION AND WASTE MANAGEMENT U.S. DEPARTMENT OF ENERGY

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VISIT TO JAPAN NOVEMBER 1990



PNL/1PS0-11/90

THE PROGRAM OF THE UNITED STATES DEPARTMENT OF ENERGY ON ENVIRONMENTAL RESTORATION AND WASTE MANAGEMENT

INTRODUCTION

The Department of Energy (DOE) sponsors energy research and development in the United States and is ultimately responsible for the disposal of nuclear and hazardous waste from its operations as well as commercial spent reactor fuel. The DOE's role has evolved over the last forty years from one of almost total control over nuclear-related activities to a more limited one with additional regulatory oversite, primarily by the Nuclear Regulatory Commission (NRC) and the Environmental Protection Agency (EPA). The original Atomic Energy Research and operations function provided by the Energy Research and Development Administration (ERDA), which was late* changed by the Congress into the DOE. The DOE collaborates with the "EPA on matters dealing with the environmental restoration of DOE sites and the disposal of nuclear and hazardous waste.

In fulfilling its mission, the DOE is responsible for a wide complex of facilities in which radioactive and other hazardous materials are used. These facilities have produced wastes that have led to varying degrees of physical plant and environmental contamination. The DOE facilities (at Hanford, Washington; Idaho Falls, Idaho; Savannah River, South Carolina; and Oak Ridge, Tennessee; and others) have produced a variety of radioactive wastes. In addition to radioactive wastes, the DOE facilities produce hazardous chemical wastes such as heavy metals, organic solvents, and acids. These wastes and mixed wastes, containing both radioactive and hazardous chemical constituents, have only recently received much attention in the United States and have added a new element to the DOE waste management program. Although waste generation has been reduced in recent years and waste handling techniques have been improved, the need remains for continuing safe waste management practices, and for correcting inadequate past practices.

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Commercial uranium mining and milling operations, along with processing of radioactive materials in the early years of nuclear development, have left some sites and facilities contaminated. Reactors at the Savannah River, Hanford, and Idaho sites have produced high-level radioactive wastes that are stored locally in underground tanks. Operation of DOE facilities has resulted in the creation of burial grounds, storage facilities, underground tanks and pipes, surface impoundments, treatment facilities, and accumulation areas that have the potential for releasing radionuclides and hazardous chemicals into the environment. The primary contaminants at major DOE sites are summarized in Table 1.

The United States has, since the inception of its nuclear program, managed the bulk of its wastes in a manner considered safe and consistent with the standards and understanding of environmental protection needs at the time. These standards have become more stringent and the old practices were, in some cases, found to be inadequate. Some unsatisfactory disposal practices have occurred in the use of injection wells, drainage trenches, and shallow-land burial sites. Most wastes since these early days are still stored or disposed of safely, but improvements, such as high-level waste vitrification plants now under construction, can add to long-term safety. The high-level wastes resulting from DOE programs, along with a small amount of high-leve' wastes produced in the reprocessing of commercial spent fuel at West Valley, New York, are safely isolated for the short term. Active programs are under way for vitrifying and disposing of the high-level portion of the wastes which are stored at the Savannah River, Hanford, Idaho, and West Valley sites, and for disposing of the low-level portion of the wastes as a cement-based waste form in engineered vaults.

The environmental restoration work now under way typically involves low levels of contaminants in relatively large volumes of soil, water, or structures. Characterization of the environmental problems created by the early practices used for disposing of the low-level liquid and solid radioactive, transuranic, and hazardous wastes is just beginning. A true measure of contamination has yet to be established and the ongoing characterization of site contamination will continue for some time.

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TABLE 1. Primary contaminants at major DOE sites.





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The United States Government is committed to effective waste management and correcting past inadequacies at its facilities. In concert with the growing public demands for reducing environmental pollution and correcting environmental problems, the DOE has embarked on a major effort to restore its contaminated sites to satisfactory conditions and to improve the management of wastes currently being generated. DOE's three goals are to reduce risk to human health and the environment, to responsibly manage and minimize the overall cost of the necessary restoration activities, and to complete this cleanup work within 30 years.

DOE'S ORGANIZATION FOR ENVIRONMENTAL RESTORATION AND WASTE MANAGEMENT

To implement this effort, the DOE has established a new Office of Environmental Restoration and Waste Management to consolidate the Department's activities in this area. The new Office, outlined in Figure 2, integrates management, budgets, and technologies for the DOE-wide waste management and cleanup program. It contains three programmatic offices and two support offices.

The Office of Waste Operations has program responsibilities for waste management at all DOE sites. Waste management includes the treatment, storage, and disposal of several types of waste: high-level radioactive wastes; transuranic wastes, including the Waste Isolation Pilot Plant (WIPP); low-level radioactive wastes; chemically hazardous wastes; mixed wastes; and sanitary solid wastes. Waste minimization efforts are managed within this Office, as are corrective activities at waste management facilities.





FIGURE 2. Office of Environmental Restoration and Waste Management organization chart.

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The Office of Environmental Restoration has program responsibilities for cleanup of inactive hazardous and radioactive waste sites at all DOE facilities and some non-DOE sites for which DOE has responsibility (e.g., uranium mill tailings sites). Included are remedial actions and decontamination and decommissioning (D&D) along with two ongoing programs, the Uranium Mill Tailings Remedial Action Program (UMTRAP) and the Formerly Utilized Sites Remedial Action Program (FUSRAP). Remedial actions are concerned with all aspects of the assessment and cleanup of inactive but potential release sites. D&D is primarily concerned with the safe caretaking of surplus nuclear facilities until either they are decontaminated for reuse or are completely removed. Excluded from DOE responsibility are sites under the authority of the electric power marketing administrations, the Office of Naval Reactors, and the Office of Fossil Energy.

The Office of Technology Development has program responsibilities for providing new and more effective technologies to meet DOE's goal of regulatory compliance and cleanup. Activities included are research and development of new technologies; demonstration, testing, and evaluation of technologies developed elsewhere; transportation; and educational programs to provide trained staff to maintain the momentum of technology development.

The Office of Planning and Resource Management supports the program offices in budget preparation and accounting and has the responsibility for coordinating the annual update of the DOE Environmental Restoration and Waste Management Plan. The Office of Environmental Quality Assurance and Quality Control performs independent internal oversight to ensure compliance with environmental and safety laws and regulations and to enhance the technical validity and cost effectiveness of programs and projects.

The regulatory oversight of hazardous and radioactive waste management is shared by several organizations. Establishment of overall standards for environmental protection is performed by the EPA, which reports directly to the U.S. President. The EPA's area of responsibility covers non-nuclear as well as nuclear areas. The NRC regulates commercial nuclear activities, long-

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term storage and disposal of spent fuel and high-level waste, and (together with the Department of Transportation) transportation of radioactive materials and wastes.

SCOPE OF THE ENVIRONMENTAL RESTORATION AND WASTE MANAGEMENT PROGRAM

The activities of the DOE in the areas of environmental restoration and waste management have been divided into four technical areas for management purposes: (1) Corrective Activities, (2) Environmental Restoration, (3) Waste Operations, and (4) Technology Development. Categories (1) and (3), being similar in nature, are both managed by the DOE Office of Waste Operations.

CORRECTIVE ACTIVITIES

Corrective Activities are those activities necessary to bring active and standby DOE facilities into compliance with local, state, and federal regulations. Because corrective activities must be completed in a timely and effective manner to protect the public health and safety and the environment, these activities will generally be accomplished by the application of existing technologies rather than new technologies that would require significant time for development. Examples of corrective activities are the installation of modern equipment to monitor air and liquid waste streams and replacement of obsolete waste handling and storage equipment.

ENVIRONMENTAL RESTORATION

Past operations connected with DOE nuclear programs have resulted in contamination of a large number of sites and facilities with quantities of radioactive, hazardous, and mixed wastes. Environmental restoration is concerned with assessment and cleanup of such sites and facilities to meet prescribed standards derived from federal and state laws. A listing of typical environmental restoration activities is presented in Table 2.

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TABLE 2. Typical environmental restoration activities.

Uranium Mill Tailings Remedial Action Program (UMTRAP) Formerly Utilized Sites Remedial Action Program (FUSRAP) Remediate contaminated surface waters Remediate contaminated underground waters Remediate contaminated soils Remediate improper burial grounds D&D contaminated inactive facilities

Note: Contaminants may include:

- Organics (polychlorinated biphenyls (PCBs), trichloroethylene (TCE), volatile organic compounds (VOCs), pesticides, petroleum products)
- Heavy metals (mercury, "barium, beryllium, lead, etc.)
- Radionuclides (including plutonium, thorium, and uranium)
- Chemicals (nitrates, asbestos, acíds)

The DOE has two major restoration programs underway. Since 1974, the FUSRAP has been working to restore 30 inactive federal facilities contaminated with radioactive and/or hazardous materials. Nine sites have been completed. The UMTRAP activity has been underway since 1978. The program was established by a federal law to remediate uranium mill tailings resulting from uranium production conducted between the early 1950s and the early 1970s. The program is stabilizing 24 sites. Stabilization has been completed at 5 of these sites.

Contaminated soil cleanup represents a major problem. The soil columns at several of the sites are contaminated with radioactive and hazardous substances resulting from the use of ponds and drainage fields for disposal of

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process effluents. Characterization and cleanup of these sites is difficult because of the low concentrations of the contaminants, their variability, and the large volumes of soil involved. The Hanford site, for example, covers an area of 1,450 square kilometers. The near-surface disposal sites on the Hanford Reservation contain 1.1 million cubic meters of solid waste which were buried before segregation of alpha-contaminated waste was required. There are also 32,000 cubic meters of soil contaminated with transuranic radionuclides.

Soils are often excavated and disposed of in secure landfills or processed at the surface to separate the contaminants. A technique (in situ vitrification) to convert soils in place to a vitreous and crystalline mass using electrical current is being developed. The resulting mass significantly lowers the solubility of inorganic contaminants. Such a treatment will also destroy organic hazardous chemicals. Other prospective treatments for organic contaminants include vacuum extraction, chemical oxidation, temporary removal and soil treatment, and bioremediation.

Groundwater contamination arises from leaking tanks, buried waste, and from lagoons previously used for the disposal of liquid wastes. For example, at the Portsmouth uranium enrichment facility, the groundwater has been found to be contaminated with trichloroethylene and polychlorinated biphenyls. If the level of contamination is high and the available technical means are not adequate, interdiction wells can be used to temporarily retard migration. New interdiction techniques under consideration are soil freezing, vitrified barriers underneath the contamination zone, and various forms of grouting and chemical injections to retard migration. It must be recognized, however, that in the case of complex aquifers contaminated with dense, non-aqueous phase liquids, current restoration technologies are probably inadequate.

Decontamination and decommissioning (D&D) of obsolete facilities produces significant amounts of additional radioactive and hazardous waste. D&D needs to rely on techniques to minimize the generation of waste by means of better characterization and selective decontamination using materials that can be recycled. Certain metals such as stainless steel, aluminum, nickel, and lead might be recycled and used for containment and shielding materials in future

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construction. The use of robotic manipulators can improve safety by reducing the radiation dose to workers.

WASTE OPERATIONS

Waste Operations include the treatment, storage, and disposal of wastes generated as a result of ongoing operations at active facilities. This includes the programs to build vitrification facilities for high-level wastes at the Savannah River, Hanford, Idaho, and West Valley sites, and the WIPP repository for transuranic wastes. One of the major waste operations objectives is to effectively manage its processes and facilities in a safe and environmentally responsible manner. Current efforts in this area also include such tasks as establishing management controls over nonradioactive wastes, including segregation of waste types and minimizing waste produced in DOE facilities.

Mixed wastes, which contain both hazardous chemicals and radioactive materials, can be reduced by minimizing the use of hazardous materials and identifying non-hazardous substitutes. The mixed wastes that already exist can be treated to separate or destroy the hazardous components. A number of physical and chemical processes are available to treat the waste: electrochemical recovery of metals, leaching, washing, chromatographic separation, evaporation and distillation, and smelting. Reactive metals can be deactivated by controlled chemical reactions, and organic compounds can be destroyed by heat provided by sources such as plasma arc furnaces, glass melters, and even solar energy.

High-level radioactive waste treatment, although well-advanced, can be tailored to address specific problems. Chemical compositions can be adjusted to reduce the volume and the associated disposal cost. Stored calcined highlevel waste can be processed into a ceramic or glass to provide more stable forms for geologic disposal.



TECHNOLOGY DEVELOPMENT

Many waste management methods used in the past are no longer adequate. DOE is striving to transcend current environmental restoration and waste management practices and tools, replacing them with more effective and efficient techniques. Where needed capability is not presently available, the Office of Technology Development will seek to develop it. This can occur through applied research and development by its laboratories, through adaptation of technology from other fields, or through development in concert with industry, academic institutions, and the international scientific community. Current environmental restoration must be performed effectively the first time and preclude the need for additional restoration in the future. Waste management must prepare and treat residual materials to produce a minimum volume of physically and chemically stable waste forms. Facility operations must minimize waste generation by eliminating or recycling hazardous materials.

To achieve these improvements, the attention of the scientific and engineering communities is required. The long-term protection of human health and the environment must be assured, and public confidence and respect for the technical community are at stake. The aims of the environmental restoration and waste management programs are to ensure that unacceptable risk of exposure to contamination is eliminated and that there is no lasting adverse impact on the environment resulting from radioactive and hazardous wastes.

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ENVIRONMENTAL RESTORATION AND WASTE MANAGEMENT PLAN

The 1992-96 update of the DOE Environmental Restoration and Waste Management Five Year plan has been completed. The plan describes the current state of technology and identifies improvements needed to fulfill DOE's three goals of reducing risk to human health and the environment, decreasing the overall cost of its restoration activities, and completing its cleanup work within 30 years. The technology development part of the plan provides for:

- o development of new technologies and techniques for waste minimization;
- o development of technologies for improved waste treatment and storage;
- o site characterization for environmental restoration using in-situ monitoring, modeling to predict and prevent migration of contaminants, and improved methods for assessing environmental and human health risks;
- o development of large promising, but underdeveloped, new technologies such as robotics, biotechnology, and remote sensing to add new capability and reduce the cost and/or risks of remediation;
- o establishment of an outreach program through schools and universities for educating students in science and waste management; and
- o international technology exchanges to reduce or eliminate duplication of efforts and to assist the international waste management community in solving similar types of problems.

The technology development plan will continue to evolve over time because the field of waste management is dynamic. Technology development will focus on near-term research, development, and application, and it will not constitute basic research. DOE intends to support development of innovative concepts



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that are mature enough to begin demonstration within the present 5-year planning horizon. Emphasis will be placed on creativity and the program will have a strong international component.

INTEGRATED DEMONSTRATIONS AND PROGRAMS

Integrated demonstrations and programs will be used as a means to rapidly develop, demonstrate, and transfer needed technology to the environmental restoration and waste operations efforts. The integrated demonstration is the cost-effective mechanism that evaluates the performance of related technologies as part of a complete system in correcting waste management and environmental problems from "cradle-to-grave". An integrated program is similar, but is focused to solve a specific aspect of a problem.

The integrated demonstration involves three major facets: 1) the various steps to solving the problem (e.g., planning, site characterization, waste treatment, waste disposal, site monitoring), 2) development and application of innovative technology solutions, and 3) evaluations relative to performance goals and/or applicable regulatory requirements. Selecting and moving promising technologies from research and development through final evaluation is a continuous process. The transition of technologies into more advanced phases of development requires the establishment of technical and regulatory criteria for ascertaining if and when a developmental project should be advanced. Because development costs increase dramatically as a project progresses to more advanced phases, funds will not be available to support full development of all concepts. Therefore, for technical, regulatory, and cost-related reasons, the number of projects moving forward will be selectively decreased by a filtering process. Technology development will be conducted with programmatic integration at all stages.

The DOE sites themselves are important resources for technology development and will be used for the demonstration and evaluation of new technologies. The three main areas of focus and planned integrated demonstrations in these areas are:

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1) groundwater and soils cleanup

- · cleanup of organics in saturated soils and groundwater
- · cleanup of organics in unsaturated soils and groundwater
- · cleanup of plutonium/uranium from contaminated soils

2) waste retrieval and waste processing

- · remediation of underground storage tanks
- remediation of buried wastes
- · decontamination and decommissioning

3) waste minimization

- uranium manufacturing waste minimization
- non-nuclear waste minimization

WASTE MINIMIZATION

Throughout the DOE environmental restoration and waste management programs, waste minimization will be a key objective. Waste minimization includes volume reduction technology, such as supercompaction, and concentration. However, true waste minimization is the avoidance of the generation of radioactive, hazardous, and mixed wastes before treatment, storage, or disposal. Waste minimization can be attained by various measures, including administrative action, material substitution, recycling, and process changes. Development and demonstration c? new processes to avoid the generation of wastes containing radioactive and hazardous constituents will be conducted. Equipment used in waste processing will be designed to clean with nonhazardous substances and/or to yield a nonhazardous product.

While waste minimization will significantly reduce the amount of waste that must be managed, waste generation cannot be altogether eliminated. Generated wastes must be managed more effectively than what was done in the past, which will require new and better ways to treat, store, and dispose of wastes. The



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Technology Development Program will seek to develop and demonstrate technologies to provide permanent solutions for generated wastes.

CONCLUSION

In the past, the highest-quality technological efforts have been directed toward other missions than waste management. The Technology Development Program will serve as a catalyst for applying today's technology to unresolved cleanup and waste management problems in ways never before considered, as well as the means for development and demonstration of new and innovative environmental protection technologies. DOE will be providing hundreds of millions of dollars to develop technical expertise through a series of new partnerships between DOE's national laboratories, industry, and universities. Integrated demonstrations will use experts from all sources in a systems approach, including collaboration with other agencies and countries. International technology exchange will be a major activity to help provide a global approach to solving waste management problems.







The Program of the United States Department of Energy on Environmental Restoration and Waste Management

Leo P. Duffy

Director, Office of Environmental Restoration and Waste Management U.S. Department of Energy

> Visit to Japan November 1990

Presentation Outline

- DOE Waste Management Legacy
- DOE ER & WM Organization
- ER & WM Program Scope
- Technology Development Program

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DOE Waste Management Legacy

- Some past practices are below present standards
- Many facilities and sites are contaminated
- Environmental restoration needed
 - Uranium mill tailings
 - Contaminated soils
 - Surface and groundwater contamination
 - Decommission inactive facilities
- Hazardous chemical and mixed wastes included

DOE ER & WM Goals

- Reduce risk to human health and environment
- Decrease cost of restoration
- Complete cleanup in 30 years









DOE's Organization for Environmental **Restoration and Waste Management**



ER & WM Program Scope

- Corrective activities
- Environmental restoration
- Waste operations
- Technology development

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- Goal is to develop new technologies that are:
- safer, faster, cheaper, better
- while achieving and maintaining compliance

Objective of the Research, Development, Demonstration, Testing, and Evaluation Program

technology to Environmental Restoration and Waste Rapidly develop, demonstrate, and transfer needed Management Operations for:

- A. Groundwater and soils cleanup
- Waste retrieval and waste processing 3
- Waste minimization and waste avoidance Ċ

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Site Characterization Sensors Fiber Optics Remote Measuring Technologies for Groundwater Robotics

Tank Characterization Robotics Sensors Piber Optics

leanup Biotechnology Filter Technology Waste Forms

\$9010075.20



\$9010075.15







Integrated Demonstration

- Cost-effective mechanism to evaluate the performance of related technologies as part of a complete system
- Components include:
 - All steps to solving the problem (planning, site characterization, waste treatment, waste disposal, site monitoring, etc.)
 - Development and application of innovative technology solutions
 - Evaluations relative to regulatory requirements

The Three Dimensions of the Integrated Demo



*Including regulatory linkage













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Components of WO Integrated Demonstrations*

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*Like ER, WO integrated demonstrations will have inputs on requirements, include assessment of multiple options, and make recommendations 53010075.18

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WO Integrated Demonstrations



Planned Integrated Demonstrations

- Groundwater and soils cleanup
 - Cleanup of organics in saturated soils and groundwater
 - Cleanup of organics in unsaturated soils and groundwater
 - Cleanup of plutonium/uranium contaminated soils
- Waste retrieval and processing
 - Remediation of underground storage tanks
 - Remediation of buried wastes
 - Decontamination and decommissioning
- Waste minimization
 - Radioactive waste minimization
 - Non-radioactive waste minimization

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Waste Minimization

- Avoidance of waste generation
 - Administrative actions
 - Material substitutions
 - Recycling
 - Process changes
- Volume reduction
 - Supercompaction'
 - Waste concentration
 - Waste processing

Sources to Solve ER & WM Problems DOE Will Utilize Expertise from All

National laboratories

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- Industry
- Universities
- International cooperation

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International Technology Exchange

- Provides global approach to waste management
- Transfers innovative technologies
- Reduces cost and remediation time
- ITE mechanisms include:
 - Joint projects
 - Exchange of staff
 - Exchange of students
 - Workshops
 - Exchange of documents

Summary

- DOE is committed to 30-year goal to clean up sites
- DOE established Environmental Restoration and Waste Management Organization
- Technologies that are safer, faster, better, and cheaper will be developed
- · DOE will utilize expertise from all sources
 - International technology exchange will be a major activity





TECHNOLOGIES DISCUSSED AT JAPAN ATOMIC ENERGY RESEARCE INSTITUTE (JAERI)

- High Level Radioactive Waste
- Mineralogical Research
- Leaching and Volatilization of Radionuclides from Glass Waste
- Spectroscopic Method
- Fixation
- Long-Term Reaction Path Modelling
- Plutonium
- Adsorption of Neptunium
- Irradiation of Materials
- Cold Neutron Source
- Reactor Decommissioning Technology Development
- Dismantling
- Decommissioning
- Safety Studies on Glass Waste Form
- Waste Safety Testing Facility
- Volatilization of Cesium from Nuclear Waste Glass in a Canister

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R&D of DECOMMISSIONING TECHNIQUES



- High Decontamination Factor (DF)
- Reduction of Arising Secondary Westes
- Applicable to Complicated Form
- Cost Effectiveness

ICE-BLASTING

ELECTRO-POLISHING

REDOX PROCESS

DISMANTLING

- Applicable to Miscellaneous Items
- Remote Handling Superior to Contact Handling in Economics and Safety

MONTORINA

- Remote Measurement
- High Reliability
- High Efficiency



PLASMA CUTTING

ROBOTTICS

LASER CUTTING

RADIATION IMAGE DISPLAY WDF PROCESS FLOW



- A : Receiving
- B : Decontamination (Ice Blasting)
- C : Plasma Cutting
- D : Hacksaw Cutting
- E : Classification
- F : Compression
- G : Packaging

- H : Reseiving
- 1 : Decontaimation (Electro-Polishing)
- J Evaluation Glove Box
- K : Experiment hood
- L : Plasma Cutting Robot
- M : Press Cutting
- N : Packaging







lot Isostatic Processing with the

JPDR DECOMMISSIONING PROGRAM

The 9th TAG Meeting October 8 -12, 1990 T. Hoshi, JAERI

1) Dismantling of the RPV

Actual cutting of the RPV was started in the end of April and was completed successfully in the beginning of october, 1990. The underwater arc-saw cutting technique was applied to cut the body portion of the RPV where it was highly activated. The lower spherical portion was cut by the conventional technique, a gas cutting technique. Chronologies on the removal operations of the K2V are as follows:

oct. 1989	*	Apr.	1990	1	Installation of the water tank, water treatment system, and
					arc-saw system
APT.		Jul.	1990	1	RPV cutting by arc-saw cutting system
Jul.	-	Sep.	1990	1	Removal of arc-saw cutting system
Sep.		Oct.	1990	1	RPV cutting by gas cutting method.

Arc-saw cutting system and cutting plan of the RPV are shown in Figs 1 and 2, respectively.

Cutting Characteristics of Arc-saw System

The body portion of the RFV was segmented into 65 pieces less than 900 X 900 mm in size underwater. Cut pieces were, then, hanged up onto the service floor in the reactor containment building and were put into the containers. Shielded containers were used for the pieces around the reactor core region (Sections of 4 to 8 shown in Fig.2), but the pieces of the upper portions (Sections of 1 to 3) were put into the standard 1 m² containers.

Cutting speeds were 0.2 - 0.5 mm/sec at the flange portion of which thickness was 250 mm and 1 - 5 mm/sec at the body of about 80 mm in thickness. These results are within the results obtained both in the developing tests and in the mock-up tests as shown in Fig. 3 . However, a difficulty was experienced at the beginning of this work, i.e., frequently over current trip in the system. This over current trip was observed in the vertical cutting operation of the flange potion , especially, at the first cutting . This was due to inappropriate setting of an arc current and a cutting speed as well as the most thick portion of the RPV. It took about 3 days to cut one vertical line of 700 mm at the beginning, but it decreased soon to 5 - 8 lines (4,000 -7,000 mm) per day. Planned and actual schedule is shown in Table 1 and scenes of operations are shown in Photos 1 - 4.

Working Days and Manpower

Total number of days required to cutting by the arc-saw was 50 days and it is considerably less than the planned ones (60 days). It was concluded that this was due to increasing the skill of workers by progressing the cutting operations. On the other hand, about 8 months were required for such preparatory works as installations of cutting system, water tank and water treatment system as well as removals of these systems.

Manpower expenditures were of 1,700 man-days for arc-saw cutting and of about 6,200 man-days for the preparatory works.

Radiation Exposure

Total radiation exposure of workers in the work was about 9 manrem. Radiation exposure for cutting with the remote operated arcsaw cutting system was only 0.2 man-rem. but the rest was comefrom the preparatory works as shown in Table 2. This larger radiation exposure during the preparatory works was resulted from the installation of the water tank since the workers were obliged to access to the RPV during this work, where the radiation dose rate was of 20 - 80 mR/hr (Maximum dose rate at the surface of the RPV was about 7 R/hr at the core center level).

Radioactive Contamination

Insignificant contamination in the air was observed. A little contamination ($3 \times 10^{-5} \, \mu \text{Ci/cc}$) was measured in water, but it is easily reduced by the water treatment system with filters. Almost all of dross generated by cutting was also removed by the dross collecting pan which was installed in the lower portion of the RPV without difficulty.

Cutting of the Lower Spherical Portion of the RPV

The lower spherical portion of the RFV was hang up onto the service floor and was cut by using a conventional gas cutting technique. Radiation dose rate was about 10 mR/hr at the surface of the RFV.

2) Removal of Components in Turbine Building

Dismantling work of the components in the turbine building has been started in April, 1990. Components of the condensate and feed water system, the auxiliary system, the sea water circulation system, etc., were removed. Turbine and generator will be removed in coming quarter.

3) Future Plan

Installation of the diamond and coring concrete cutting system will be started in October and cutting of the biological shield concrete at the highly activated portion using this system will be carried out during December, 1990 and January, 1991. Water jet cutting will be followed from March to July, 1991.

Removal of the components in the turbine building has been continued until next year. Removal of components in the waste treatment building, the fuel storage building, etc., will be initiated from April, 1990.
Table 1 Arc-saw Cutting Schedule of the RPV







Item	Man-days	Man-rem
Preparatory Works		
Installation of arc-saw Installation of water tank	2,470 1,710	0.24 8.39
water treatment system	460	0.17
Removal of systems	1,600	0.36
Sus-total	- 6,240	9.16
Cutting operation	1,700	0.21

Table 2 Man-power and Radiation Exposure in Cutting the RPV by Arc-saw

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Fig. 1 Basic concept of arc saw cutting system

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Cutting Plan of The RPV by Arc-saw System

Fig.

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Fig. 3 Cutting Characteristics of Arc-saw System









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Photo 4 Cut piece of RPV being put into container







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JPDR DECOMMISSIONING PROGRAM

Japan Atomic Energy Reseach Institute

NATIONAL STRATEGY ON DECOMMISSIONING OF POWER REACTORS IN JAPAN

- · Dismontling to reuse the site
- Development of dismontling techniques to assure safety and reduce costs .

Improvement of current technologies and development of new technologies

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In JAERI, development of technologies and demonstration







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PURPOSE OF JPDR DECOMMISSIONING PROGRAM

- To obtain actual experience on dismantling of nuclear power plant
- 2. To develop and demonstrate dismantling techniques
- 3. To obtain data concerning;
 - 1) radiation (exposure of workers, airbonne activities, radiation level, etc.)
 - 2) waste (measurement and segregation, decontamination, etc.)
 - 3) system engineering (number of workers, working time, cost, etc.)

STANDARD DISMANTLING SEQUENCE OF COMMERCIAL POWER REACTORS IN JAPAN

(planned for 1,000 MWe plant by MITI)



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Reactor BWR Power 90 Mwt (Initial 45 Mwt) 12.5 Mwe

RPV 8 mH x 2mlD x 67 mm[†]

Biological Shield Concrete

1.5~3 mt Reinforcing Bar (29 mmOD, (150~200 mmPltch

Operation 1963, 10 Final Shutdown 1976, 3



CROSS-SECTION OF REACTOR CONTAINMENT BUILDING

5



OPERATIONAL HISTORIES OF JPDR

\uparrow	1963.8		Criticol
JPD	1963.10		Electrical power generation
Ř I I		(45 WMt operation in natural) convection of coolant
1	<u>1969.9</u>		Shutdown reactor to increase power
Ţ J	1959.10 ~ 1971.12		Modifications of reactor core and plant to 90 MWt
PDR	1972.2		Critical in new core
 11	1972.5		Electrical power generation
		. (power increase to approx. 60 MWt)
	<u>1976.3</u>		Final shutdown
	1982.12		Apply the decommissioning permit
	1986,7		Apply the revised decommissioning report
	1988,12		Start physical dismantlement



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JPDR DECOMMISSIONING SCHEDULE



1981~1986 : RED



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Radloactivity in RPV and Biological Shield Concrete

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RADIATION DOSERATE IN JPDR FACILITY



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DISMANILING ECHNIQUES



Object	Technique	Example of performance	
Pressure vessel	Arc-saw	Carbon steel 250mm (In water) 200mm (In air)	
Reactor internals	Plasma arc	Stainless steel 130mm (in water)	
Plping connected	Rotary disk knife	Stainless steel 12in, Sch 80	
to pressure vessel	Shaped explosive	Carbon steel 261n, Sch 80	
4	Dlamond sawing and coring	Cutting efficiency 2.5m ² /hr	
Biological shield	Abrasive-Water jet	Depth of cut 450~600mm	
	Controlled blasting	Blasting efficiency 10hr/m ³	

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Fig. 4 PLASMA ARC CUTTING SYSTEM APPLIED TO REACTOR INTERNALS







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Fig. 5 ARC-SAW CUTTING SYSTEM APPLIED TO PRESSURE VESSEL

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Fig. 4 SHAPED EXPLOSIVE CUTTING SYSTEM

1.





Top view



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Fig. # ROTARY DISK KNIFE CUTTING SYSTEM



Fig. 8 DIAMOND SAWING AND CORING CUTTING SYSTEM

Table I SPECIFICATION AND PERFORMANCE OF ABRASIVE-WATER JET CUTTING SYSTEM

Specification	Water pressure	196 MPa
	Water flow rate	50 i Amin
	Abrasive feed rate	3-7 kg/min
	Nozzle traverse rate	0-120 cm/min
Performance	Cutting time*	60 min
	Concrete dust generation*	33 kg

* Cutting time and concrete dust generation per one block cut off Block size: 40 cm (depth) x 40 cm (height) x 300 cm (length)



Fig. 6 ABRASIVE-WATER JET CUTTING SYSTEM

Kinds and Activity of Radioactive Waste generated from the JPDR Decommissioning 1 1

Kinds	of Radioactive Haste	Activity (Ci)	Weight (Ton)
Activated Components	Core Internals Control rods, Core shroud, etc.	4,050	20
	Pressure Vessel	40	110
	Biological Shield Concrete	10	1.350 .
Contaminated Components	Components	4.2	1,640
	Concrete	0.2	830
	Resin, etc.	0.5	130
Total		4,100	4,100

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Management System of Radioactive Solid Waste in Japan 20

RADIATION EXPOSURE FOR WORKERS

Dismantling Item	Man-Days	Man-rem
Equipments	8,700	14
RPV Internals	9,200	17
RPV	4,100	26
Blo. Shleld Concrete	4,500	8
Containment	9,000	- 11
Sub Total	35,500	66
Other Buildings	37,500	34
Total	73,000	100





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High quality irradiation conditions and high neutron flux conditions have been required in recent years for neutron irradiation and neutron beam experiments in the research reactors. To fulfil these requirements, it was decided that the Japan Research Reactor No.3(JRR-3) should be reconstructed to be upgraded, and the project of new JRR-3 is now progressing. The JE 1-3 was constructed in 1962, originally for general studies of nuclear reactors, irradiation of materials, production of radioisotopes, and research and development 6, nuclear reactors.

The new JRR-3 is equipped with various kinds of experimental facilities for irradiation and 'eam experiments including a cold neutron source so that the new reactor may be utilized as a multi-purpose research reactor which is at the highest level in the world, with the maximum thermal neutron flux of about 2×10^{14} n/cm² s, one order higher than that of the old reactor. The safety for the new JRR-3 is fully preserved with highly reliable control systems and cooling systems (including a facility to prevent the core from being unflooded).

The work of the construction for the new JRR-3 started in August, 1985 and Initial criticality of the new reactor was achieaved in March, 1990.

Isometric View of Reactor Pool Gate No. Top Shield CNS Low Temperature Channe Tube Irradiation Hok Heavy Water Control Rod Tank Primary Cor Circuit Pipe Bottom Shie Control Rod Drive Mechanism

(APPENDING)

	Low Enriched Uranium.		
Reactor Type	Light Water Cooled and Moderated.		
	Swimming Poor Type.		
Rated Power	20 M W		
Size of Core	Approx. 60cm dia. and 75cm high		
	(with Beryllium Reflector).		
	UAlx-Al Dispersed, MTR Plate Tope		
	20% Enriched Fuel.		
ruel	26 Standard Fuel Elements and		
	6 Follower Fuel Elements.		
	6 Control Rods, Box Type		
Control Rod	Absorber, followed with Follower		
	Fuel Element.		
	Swimming Pool Type		
Reactor Pool	4.5m dia.		
	8.5m deep		
Exectimental	9 Horizontal Beam Tubes,		
Facilities	17 Vertical Irradiation Foles, and		

Major Specifications of the New JRR-3







Experimental beam facilities

Nine beam tubes and five neutron guide tubes are installed in order to lead out neutrons for beam experiments.

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* Reach -

Good quality thermal neutrons can be obtained for the experiments in the new JRR-3 with these beam tubes arranged tangentially to the core because the γ rays and fast neutrons not preferable to the experiment are much reduced in such a tangentially arranged beam tubes.

The neutron guide tubes lead out neutrons in the distance so that a sufficient number of beam experimental apparatus can be provided to many experimenters and users.

Cold neutron source (CNS)

Cold neutron has very low energy with a wave length of the same order as the molecular structure of substance. So, it is able to research the structure of macro-molecule such as a high polymer with the cold neutron.



Bacas stall up to 60 m from Reserve

Schematic Diagram



CNS and Neutron Guide Tube

CNS converts thermal neutron into cold neutron with liquid H_2 at 20K. The cold neutron beam is led to the beam experimental apparatus in the beam hall through neutron guide tubes.

COOLING SYSTEMS

Cooling circuit systems are composed of a primary circuit, a secondary circuit and a heavy water circuit. Core heat is finally removed into atmosphere through the cooling tower.

The primary circuit has two independent siphon break valves to prevent the core from being unflooded.



Flow Diagram of Cooling Systems



FUEL

Plate-type Aluminium-Uranium alloy fuels which have higher power density, more coolability and higher performance than Oxide-Uranium fuels of light water reactors, are used in the new JRR-3 as in other research reactors. The new JRR-3 adopts two kinds of fuels, standard fuel element and follower fuel element following the control rod.

Summary of Fuel Specification

Item	Standard Fuel Element	Follower Fuel Element	
Size of Fuel	$76 \times 76 \times 1150$ m m	$64 \times 64 \times 880$ mm	
U-235 Enrichment	20%	20%	
U-235 Contents	300g	190g	
Size of Fuel Plate	$1.52^{\text{t}}\times71^{\text{t}}\times770^{\text{t}}\text{m}\text{m}$	$1.52^{\circ} \times 60^{*} \times 770^{1} \mathrm{mm}$	
Fuel Plate Number	20 Element	16 Element	
Fuel Meat Material	Dispersed UAIx+A1		
Cladding Material	Aluminium Alloy		

UTILITY SERVICES

For irradiation, vertical irradiation holes are arranged both in the core and in the heavy water tank.

For beam experiment, horizontal beam tubes are arranged in the heavy water tank.



For the purposes of reactor fuels and materials exposure tests, radioisotope production and activation analysis, these facilities are used to irradiate by neutron the samples inserted into the vertical irradiation holes.



Arrangement of Experimental Holes

Name	No	Application	Feature
Hydraulic rabbit irradiation facility (HR)	2	General irradiation Radioisotope production	The rabbit is conveyed and cooled by water. This facility is used to irradiate the relatively heavy and high heat generating samples.
Pneumatic rabbit irradiation facility (PN)	2	General irradiation Radioisotope production	The rabbit is conveyed and cooled by N, gas. This facility is used to irradiate the light and low heat generating samples.
Activation analysis irrediation facility (PN3)	1	Activation analysis of the short life radio nuclides	The radiation measurement is started immediately after the irradiation. This facility is used to analyze the short life radio nuclides.
Uniform irradiation facility (S1)	-1	Material irradiation Silicon irradiation	The sample is rotated and moved up and down during the irradiation. This facility is used to irradia'e the sample uniformly.
Rotating irradiation facility (DR)	1	Large material irradiation	The sample is rotated during the irradiation. This facility is used to irradiate the sample uniformly in the radial direction.
Capsule irradiation	in	Exposure test	This facility is used to irradiate for long period or control the sample temperature in response to the

Summary


Rear Building

Beam Experimental Facilities

History of JRR-3

Year	Major Events	Year	Major Events
1959	Beginning of JRR-3 construction	'76	
'60		'77	
'61	Reactor completion	'78	Integrated power 300,000 MWH achieved
'62	Reactor critical	'79	
'63		'80	
'64	Rated power 10,000 KW achieved	'81	
'65	Beginning of RI production	'82	The twentieth anniversary since the reactor
'66	Beginning of common use		critical
'67	Beginning of homemade fuel use	'83	Close of common use
'68		34	Finish of safety review for rew JRR-3
'69	Medical irradiation for a brain turmor		Beginning of the construction work for
'70			new JRR-3
'71	Sample irradiation of nuclear fuel in LHTL	1903	
172	Beginning of shift to UO, fuel core	ar m	Construction
'73			
'74	Integrated power 200,000 MWH achieved	St. 16	
175	Completion of shift to UO, fuel core	. 90 en : 645, mars	Completion of new JRR-3

JAERI-M 89-192

Progress Report on Safety Research of High-Level Waste Management for the Period April 1988 to March 1989

(Eds.) Haruto NAKAMURA and Susumu MURAOKA

Department of Environmental Safety Research Tokai Research Establishment Japan Atomic Energy Research Institute Tukai-mura, Naka-gun, Ibaraki-ken

(Received October 23, 1989)

Researches on high-level waste management at the High Level Waste Management Laboratory and the Waste Safety Testing Facility Operation Division of the Japan Atomic Energy Research Institute in the fiscal year of 1988 are reviewed.

The topics are following studies on the long-term chemical behaviors of long-lived nuclides in geosphere.

- Mineralogical researches on the alteration layer of glass exposed to water were carried out by laboratory experiments and investigation of natural glass. Leaching experiments of Pu and Np were also conducted.
- The spectroscopic methods are applied to study the long-term reaction path modeling of radionuclide fixation using natural materials.

Keywords: High-level Radioactive Waste, Glass, Mineralogical Research, Leaching, Plutonium, Neptunium, Spectroscopic Method, Long-term Reaction Path, Fixation

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Reactor Decommissioning Technology Development and Actual Dismantling of JPDR



Tokai Research Establishment Japan Atomic Energy Research Institute

Preparing for Reactor Decommissioning

Necessity of Reactor Decommissioning Technology Development

The useful lifetime of a nuclear power plant is estimated to be 30 to 40 years. Worldwide, a few plants have reached this age. The number of plants reaching this age will increase substantially in this decade. As a result, the technology for nuclear power plant ascommissioning must be developed and made available in the near future.

The Elk River Reactor in the United States is an unusual example of a power plant which was completely dismaniled after its duty life ended. However, there have been only a few reactors decommissioned worldwide. Reactor decommissioning technology therefore is not well established. It must be advanced through the development of necessary techniques and these techniques must be applied to actual dismantling.

The Japan Atomic Energy Research Institute, JAERI, has been developing techniques needed for dismantling the Japan Power Demonstration Reactor, JPDR. This is being done under a contract with the Science and Technology Agency, STA. This work, begun in 1981, has progressed to the actual dismantling of the JPDR using the techniques developed

Schedule of Reactor Decommissioning Technology Development and Actual Dismantling of JPDR

The JPDR decommissioning program consists of two major phases. The purpose of Phase 1 was to develop techniques necessary for dismantling JPDR. It was essentially completed in 1986. The purpose of Phase 2 is to dismantle the JPDR. This is

now underway using the techniques developed in Phase 1. The major objectives of this dismantling program are to develop and demonstrate dismantling techniques and to accumulate power reactor decommissioning experience.

* Schedule

Fiscal Year Phase	1981	1982	1983	1984	1985	
Development of Reactor Dismantling Techniques (Phase 1)		PRIMICAL CONSTRAINTS				
Actual Dismantling of JPDR	. o		a	10 - + 10 + 10 + 10 + 10 + 10 + 10 + 10		ſ
(mase 2)			**********			1



Pear to: Decommissioning Alternatives

Decommissioning a power reactor can be accomplished by any of the three following methods. Dismantlement is considered to be the most suitable decommissioning option in Japan because it effectively uses scarce land.





2







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Safety Studies on Glass Waste Form

S. Muraoka

In order to achieve the safe disposal of radioactive waste, it is necessary to promote the development of waste management technology as well as the safety assessment study. In JAERI, safety studies for contribution to establishment national safety criteria and safety assessment are being conducted as well as basic research to support development of waste management technology which is adaptable to environmental conditions in Japan.

Characterization of radioactive waste has been carried out for some items to evaluate the safety, according to process of transportation, storage and disposal.

In this document, some of our recent activities are described briefly of the JAERI's studies on the glass waste form for high level waste disposal.

(1) Volatilization of ¹³⁷Cs and ¹⁰⁶Ru from Borosilicate Glass Containing Actual High Level Waste [1]

As the safety evaluation test in relation to operation of storage facility, the volatilization of Cs and Ru from borosilicate glass containing actual HLW was examined in an almost closed canister.

The HLW glass used for the present study was borosilicate glass. The reagents for the glass additives and the simulated HLW which should be converted into 1300g of oxide glass were mixed simultaneously and placed in a vitrification apparatus with about one liter of a denitrated actual HLW solution. About 50g of various oxides were assumed to come from the actual HLW solution. This mixture was calcined at about 750°C, melted at 1200°C, for 2 hours in the vitrification apparatus. Half the molten glass was poured into an 8.1cm-i.d., 24.4cm-high stainless steel canister, kept at 600°C for 2 hours, and then cooled to room temperature at a cooling rate of less than 40°C/h. The furnace temperature was raised in steps from 25°C to 1000°C (Fig. 1). The temperature rise by the decay heat of the HLW was so small in the present study that it did not affect the temperature control of the glass. During the course of heating, part of the air in the upper space of the canister was collected in an evacuated sampling bottle with a volume of about 7 cm

Radioactivity from volatile elements trapped by both the sampling bottle and the sampling needle was measured by an intrinsic Ge solid state detector. Figure 2 shows the temperature dependence of volatility of 197Cs and 100Ru at a fixed time of 24 hours when both nuclides are at apparent saturation concentrations. The solid line in the figure represents the data obtained in our previous work [2] in which the activation energy of about 140 kJ/mol was obtained on simulated HrW glass



containing about 1.6 x 10¹⁰ Bq of ¹³⁴Cs. The present data shows

fairly good agreement with previous one. The volatility of TOBRU measured at 600°C and 800°C is about one fifth that of 13°Cs. Since Gray [3] has pointed out that the activation energy for various elements are almost the same as each other, the air contamination of 100Ru at a usual storage temperature of 400° C is also expected to be one fifth that of 137Cs; thus, the normalized concentration of 105Ru would be about 5 x 10⁻¹⁰ at 400°C. This extrapolated value may be conservative, since 400°C is below the softening point of the present glass and diffusion may not be a significant mechanism at this temperature. It should be also mentioned that the volatility of 106Ru at 1000° C could not be measured; it was under the detection limit of 5 x 10^{-2} Bq/cm³. This is probably an example that the backward step plays an important role; the stainless steel canister is markedly oxidized at around 1000°C and reacted with 106Ru in the air inside the canister. It is also probable that RuOd is not stable at high temperatures, and this is one of the causes that 100Ru disappeared at around 1000°C.

(2) Release of Neptunium from a ²³⁷Np doped Borosilicate Waste Glass (4)

The MCC 1 static leach tests were performed for a 237Np doped borosilicate waste glass at 90 °C with deionized water and silicate water leachants to spe-ulate the release mechanism of Np from waste glasses. The composition of the Np doped glass is shown (n Table 1. Teflon, vessels were used as leach containers and SA/V ratio was 0.1cm

Deionized water and silicate vater were used as leachants. At the desired leach durations, the Np concentrations in the leachates were measured by gamma counting with a high purity germanium detector.

The amounts of Np released from the glass are plotted as a function of time as shown in Fig. 3 in terms of the normalized elemental mass loss (NL) Np. Although the release behaviors as a function of time are appreciably different between the deignized water and silicate water, the (NL) $_{\rm Np}$ values of about $5g/m^2$ are similar for the two leachates after 91 day leaching. The release behavior of Np in this study are compared with those of other elements (Fig. 4). A linear relation between log(NL) and log(time) is observed for Na, B and Cs within the studied leaching duration. As time proceeds, NL's for Np and Sr approach to constant values.

According to our previous study, Na, B and Cs were found in the leachates but not found in the surface layer; they are released from the bulk glass by decomposition of the glass and diffuse through the surface layer without being trapped. Sr was detected both in the surface layer and in the leachates. Probably Np, representing a similar time dependent release behavior to that for Sr. is also present in the surface layers.

It is reasonable to assume that Np concentrations in leachates are controlled by the solubilities of Np solid phases formed in the surface layer. This assumption leads to that the solid phases formed in the surface layers must be primarily







identified in order to predict Np concentrations in glass leachates. Since none of present analytical techniques is applicable to wet surfaces, the Np species in the surface layers can not be identified directly. Then, an attempt was made to speculate it based on the predicted species in the bulk glass and that in leachates.

In bulk glasses either the tetravalent or the pentavalent species possibly exists. Np species in aqueous solutions can be estimated by pH and Eh of the solutions. However, such redox parameters have not been studied for solutions contained in surface layers. Instead, we use pH and Eh values of the leachates. The measured pH and Eh of the leachates in the present experiments are plotted in Fig. 5. These values change with time, but they are similar for different leach durations except 3 days. Referring available pH-Eh diagrams, the tetravalent and the pentavalent species are possibly present in comparable amounts in the present leachates, and the trivalent and the hexavalent species are probably absent. Since the tetravalent and/or pentavalent species are expected to exist in both the bulk glasses and leachates, the valence of Np in the surface layer is also likely to be tetravalent and/or pentavalent.

Considering the above estimation on the valence and OH as a predominant complexing anion present in the leachates, we take NpO_2XH_2O(am) and NpO_OH(am) as Np solid phases in the surface layer, and assume the following three types of solubility equilibrium; (1) Np_OH(am)=NpO_2+, NpO_2OH(aq) and NpO_2CO_3=

(2) $NpO_2 \times H_2O(am) = NpO_2^+$

(3) NpO2xH2O(am)=Np(OH)4(aq) Apparent steady state concentrations of Np from MCC-1 leach tests are plotted in Fig. 6. Solubilities of NpO2OH(am) and NpO2xH2O(am) calculated from equilibrium constants at 25°C are also shown in the same figure by dotted and hatched regions, respectively. These regions include the predicted solubilities for different ionic strengths of aqueous solutions. As seen in this figure, Np concentrations in the leachates are apparently limited by the solubilities of NpO2xH2O(am); the above mentioned equilibria (2) and (3) are expected. Np concentrations obtained from the leachates are distinguishably than lower solubilities of NpO2OH(am). If Np in the surface layers had been NpOgOH(am), the Np concentrations in the leachates should have been higher, approaching to the NpOgOH(am) solubilities. These facts imply that Np exists as the tetravalent solid phase NpO2xH2O(am) rather than the pentavalent solid phase NpO2OH(am) in the surface layers of leached waste glasses.

(3) Leaching Behavior of Simulated High Level Waste Glass in Groundwater (5), [6]

The purpose of the work is to examine the leaching behavior of simulated high level waste glass in actual groundwater in Japan. In-situ burial tests were carried out by immersing the sample in groundwater coming through schalstein type rock in southwestern Japan. The results were compared with the ones of laboratory test obtained using synthesized groundwater and deionized water.





Figure 7 shows the scanning electron microphotographs of the surface before and after leaching. In the case of actual groundwater (Fig. 7(b)) and synthesized groundwater (Fig. 7(c)), many grooves occur on the specimen along with the surrounding flat surface, indicating that some parts of simulated high level waste glass dissolve more easily than others. On the other hand, in the case of deionized water (Fig. 7(d)), such grooves are not clearly observed, which means that leaching is progressing more uniformly than in the case of groundwater.

We assume that the leaching behavior of the simulated high level waste glass is divided into two categories; one is leaching from the flat surface and the other is that from the grooves. The extent of leaching from the flat surface can a measured by SEM EDX. Here we define the C/C₀ values a he ratio of the concentration of Na on the flat surface contaction of the to the initial concentration of Na before the local specimen (C)

We pave the way for estimating r of normalized elemental mass loss of Na (NLNa) of lass leached in groundwater by measuring the C/C₀ value stauring the size and the number of grooves without leachates xaminations. For instance, in the case of the specimen leached in actual groundwater for one year and seven months, is 5 °C, value is about 0.86 which corresponds to NLNa of 6.5 °C, value is flat surface. On the other hand, by measuring is ize and the number of the grooves, we obtain NLNa of 3.9 x 10 ° g/cm² which corresponds to the amount of Na leached from the grooves. The sum of 6.5 x 10 ° g/cm² for the flat surface id 3.9 x 10 ° g/cm² for the grooves is 4.6 x 10 ° g/cm², result rg in a leach rate of about 8 x 10 ° g/cm² day.

(4) Accelerated Alpha Radiation Stability Test [7]

An accelerated alpha radiation stability test started in connection with characterization of returnable waste forms from overseas reprocessing facilities, and the test equivalent to 10,000 years aging of actual waste forms has been finished.

Curium-244 and plutonium-238 were added to a simulated waste substituting transuragium elements and 90% of rare earths (244Cm: 43.3 GBq/g-glass, 238Pu: 4.4 GBq/g-glass). The waste was molten with borosilicate glass in three platinum crucibles of 14 mm in diameter. Twenty four specimens were prepared by cutting the crucibles into pieces 5 mm thick, and each specimen was stored in a helium leak protective capsule.

Four or five specimens were taken out from the storage pits for each time equivalent test including zero time tests. The tests were performed with a mass spectrometer, a differential scanning calorimeter (DSC), Soxhlet type leaching apparatus and a microscope for measurement of helium remained in the matrix, stored energy in the matrix, leachability and fine-structure alteration, respectively.

It was found by measuring the amount of helium released from the specimens that 97-99% of helium remained in the matrix at the room temperature. The total amount of helium generated in the specimen was obtained from the amount of helium released from the specimen kept at 600°C for 15 min and that at room temperature, because at 600°C the helium was completely depleted from the glass specimen.

The test results are shown in Fig.8. Density of the specimens decreased slightly with the increase of time and the decrement of 1.1% was observed at 10,000 years equivalent. Regarding leachability based on the total weight losses, some fluctuating results were obtained in the initial stages of the test but subseque the the curve is flat to the 10,000 years equivalent. Microscopic observation also did not show any change in the microstructure. Those results seem to suggest that alpha radiation has no significant influence on the performance of glass forms to confine high level wastes.

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Table 1 Composition of *** Np-doped JAERI glass.

Component	Content (wt %)	Component	Content (w1%)
Additive SiO2 B2O3 Al2O3 COO NO2O ZnO Li2O	45.15 13.90 4.01 9.79 2.00	Wosie Te O2 Cs 20 Bo 0 Lo 203 Ce O2 Pr 6011 Nd 203	0.23 0.97 0.62 0.48 0.95 0.45 1.51
Woste Rb2O SrO Y2O3 ZrO2 MoO3 MnO2 Ag2O CdO SnO2 Sb2O3	0.12 0.34 0.19 2.64 1.73 0.03 0.02 0.02 0.024	EU203 Gd203 Se02 RU02 Fe203 NiO Cr203 RU RU RU Pd	0.000 0.000000
		237 NpO2	1.15

+ Component contains both additive and waste.



Arrows show the sampling positions.



1.5

103/T (K-1)

2.0

1.0

0.5







Fig. A the normalized elemental mass losses as a function of time for Ha, 8, Sr, Cs and Hp released from JAERI glasses in the MCC-1 static leach tests. Data on negtonium in Fig. 8 for delonized water were replotted. Data on Ha, 8, Sr and Cs were cited from Ref. (12).



FIE.5 Measured pH and Eh values of the plass leachates. Rumbers next to the points denote the leach durations. The "initial"s denote the values for leachants.







0

2 1

FIG. 7 Scanning electron microphotographs of the surface of the unleached specimen (a) and the surface of the specimens leached in actuel groundwater (b), in synthesized groundwater (c)



Fig. 8 Results of accelerated alpha radiation stability test for



Neptunium dissolution from closues



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Fig. 6 Neptunium concentrations in glass leachates obtained from the MCC-1-type static leach tests for durations of 56 days or longer, and expected solubilities of NpO₂-xH₂O(am) and NpO₂OH(am) based on thermodynamic data.

pH of leachates



Fig. 7 Scanning electron microphotographs of the surface of the unleached specimen (a) and the surface of the specimens leached in actual groundwater (b), in synthesized groundwater (c) and in detonized water (d) for 1 year.



Fig. 8 Results of accelerated wint: "adlation stability test for the "aste glass.





VOLATILIZATION OF CESIUN FROM NUCLEAR WASTE GLASS IN A CANISTER

HIROSHI KAMIZONO, SHIZUO KIKKAWA, SHINGO TASHIRO, and HARUTO NAKAMURA Japan Atomic Energy Research Institute Department of Environmental Safety Research Tokai-mura, Ibaraki-ken, 319-11 Japan

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Volatilization of ¹³⁴Cs from simulated high-level waste glass in a canister during several reheatings up to a maximum of 1000 °C was examined. The results showed that the temperature dependence of the amount of ¹³⁴Cs suspended in the air inside the canister could be divided into two categories. As the temperature was increased above 500 °C, the amount of ¹³⁴Cs suspended in the air inside the canister also increased. On the other hand, for temperatures <500 °C, the amount of ¹³⁴Cs suspended in the air inside the canister had an almost constant value after several reheatings up to a maximum of 1000 °C. In this case, the air contamination by cesium-bearing material inside the canister is considered to be significant even at waste storage temperatures <500 °C.





Q

Fig.

Kamizono et al. VOLATILIZATION OF CESIUM



Temperature dependency of the 134Cs concentration that represents the amount of 134Cs suspended in the air inside the canister (Auto) at a fixed time of I day. The data were collected during the first cycle of reheating up to 1000°C (+), the second cycle up to 900°C (=), the third cycle up to 900°C (x), the fourth cycle up to 700°C (*), and the tifth cycle up to 400°C (A), which correspond to Fig. 2. The error bars show the standard deviation of each plot. Note: Thus far, we have not yet examined the exact cesium speciation. We speculate, however, that cesium may exist as a vapor phase for temperalures of more than 500°C and fine particles containing cesium play an important role in air contamination for temperatures of <500°C. Therefore, it is not appropriate to plot in partial pressures of single chemical specie in the present paper. lustend, we prefer to plot in Auton which does not need the cesium speciation.



1.1

11. Komisono et al. / Air contamination by cestium



Fig. 3

115

Scanning electron microphotographs of the essiumbearing materials trapped by the air sampler apositional analyses were carrier out by WDX within the circles.





at 600°C

Fig. 4





1. WASTEF

The Waste Safety Testing Facility (WASTEF) constructed at the Tokai Research Establishment, Japan Atomic Energy Research Institute (JAERI), has been operated since November 1982 to research safety evaluation of the long-term storage and disposal of high-level radioactive wastes (HLW).

The present research includes tests on characteristics, confinement ability and durability of materials applied to artificial barriers and natural barriers against the waste release under storage and disposal conditions.

The tests will continue to the research with actual wastes after accumulating the data obtained with radioactive synthetic wastes.

The research has been performed in accordance with the national waste management program and the results will contribute to establish a safety system for HLW management in Japan.



2. Layout of WASTEF

Five concrete shielded cells (3 beta-gamma cells (No.1-3) and 2 alpha-gamma cells (No.4&5)) and a lead shielded cell are maintained and operated in the first floor.

Liquid waste tanks, ventilation and exhaust system, emergency power supply system and utilities services are installed in the basement floor.



Layou of WASTEF in the first floor

3. Specification of Hot Cells

Cell		Inside	Shielded wall thickness (m)	Maximun activity (Ci) (HLLW & typical nuclides)			Pu
		WxDxH (m)		HLLW	Cs+137	Sr+90	(gr)
Beta-gamma	No.1	7.5x3.0x4.5	1.05*	5x10 ⁴ (1x10 ⁶ in storage)	1×104	1×104	-
	No.2	5.0x3.0x5.0	1.05*	5×10 ⁴	1×10 ⁴	1×10 ⁴	-
	No.3	5.0x3.0x4.5	1.05*	5×10 ⁴	1×10 ⁴	1×10 ⁴	
Alpha-gamma	No . 4	5.0x3.0x4.2	1.05*	1×104	2×10 ³	2×10 ³	12
	No.5	5.0×2.75×4.2	1.1**	5×10 ²	5×10 ²	5×10 ¹	12
	Lead	2.5x1.4x2.0	0.15	2×10 ¹	2×10 ¹	1×10 ¹	1
	Lauran	A REAL PROPERTY AND A REAL	A same last top of which the last of the same last of the state of the same last of the sam	A	Contracted and the state of the state		A construction of the local distance of the

* Magnetite concrete ** Ordinary concrete



Interior of the operation area





4. Tests in WASTEF

Cell No.	Carrying-in	Preparation	Test
No . 1	Carrying-in of RI	Canning)	Storage test
No , 2	Carrying-in of actual HLLW	Vitrification Vitrified product	
	(5x10 ⁴ Ci max.)	(5 1 max.)	
No . 3	Carrying-in of RI	Preparation (Cutting,bore- holing,powdering, etc.)	Disposal test
No . 4			Measurement of characteristics
NO.5	Carrying-in of alpha-nuclides	Preparation of product with alpha-nuclides	Alpha-radiation stability test
Lead cell			Measurement of radioactivity
			microscope

Flow chart of tests in WASTEF




Test item and its Purpose in WASTEF

Test item	Pirpose	Measurement items		
Storage	Safety evaluation on long-term storage of vitrified package	•Temperature distribution •Volatility at high temperature •Cooling efficiency •Corrosion rate of packaging		
Disposal	Safety evaluation on geological disposal	 Immobility of nuclides in rocks Compatibility of glass and engineered barriers 		
Characterization	Accumulation of basic data on glass forms	 Density Heat generation Thermal conductivity Leachability Activity distribution 		
Alpha-radiation stability	Long-term durability evaluation of glass forms	 Stored energy Number of helium Change of structure 		
Vitrification	Safety evaluation on vitrification facility	 Integrity of applied materials Performance of off-gas system Material balance of process 		









Vitrification apparatus in No.2 cell





Gamma-scanning and storage test apparatus in No.1 cell



Test specimens preparation apparatus in No.3 cell



Characterization test apparatus in No.4 cell



Alpha-radiation stability test apparatus in No.5 cell

5. WASTEF Operation Schedule

Fiscal year Item	1981 1982	1983	1984	1985	1986	1987
Construction Test run						
Simulated waste tests on vitrification, storage, disposal, alpha-radiation stability and character- istics using Cs-137, Sr-90 etc.				\rightarrow		
Actual waste tests on vitrification, storage and disposal						

Organization Chart of Environmental Safety Rasearch at JAERI



TECHNOLOGIES DISCUSSED AT THE NATIONAL RESEARCH INSTITUTE FOR POLLUTION AND RESOURCES (NRIPR)

- Environmental Assessment Activities
- Water Pollution Control Activities
- Remote Sensing Techniques for Marine Pollution Analysis
- Study of CO2 Behavior in the Environment
- Treatment of Individual Types of Waste Containing Halogenated Organic Compounds
- Measurement of Pollutants in Groundwater
- Bio Treatment of Hazardous Chemicals in Waste Water

BIBLIOGRAPHY OF LITERATURE RECEIVED FROM NRIPR

"Summary of National Research Institute for Pollution and Resources", NRIPR, 44 pages.



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History

公害實重研究所は、石炭、石油等の化石燃料の合理的利用 及び燃焼技術に関する研究を行ってきた「燃料研究所 (大正 9年設立)」と地下資源の開発,選鉱、製錬及び鉱山保安技術 に関する研究を行ってきた「鉱業技術試験所(昭和24年鉱業 研究所等3所の統合により設立)」が昭和27年に合併した資源 技術試験所を前身としています。さらに昭和45年に公害部門 を拡充強化し、公害資源研究所として組織を改め、昭和63年 には産業構造の転換に対応して組織改革を行い現在に至って います。

現在、当所の組織は企画室、乾養部、9研究部(石炭部、 燃料資源部、燃焼技術部、材料資源部、資源開発部、産業保 安部、立地環境部、大気環境保全部、水環境保全部)のほか、 技術相談所、北海道石炭鉱山技術試験センター及び九州石炭 鉱山技術試験センターからなっています。 National Research Institute for Pollution and Resources. (NRIPR) was established in 1952 as the Resources Research Institute by the merger of two institutes the Fuel Research Institute and Mining and Safety Research Institute. The former had carried out studies on the utilization of fossil fuels and their combustion technology, and the latter, the exploitation and utilization of underground resources and mining safety.

Resources Research institute was reorganized into NRIPR in 1970 so as to meet the social demands at the time and expanded its research field to include industrial pollution control

NRIPR was further reorganized in 1988 to accelerate its research activities on future industry and the global environment. The present NRIPR is composed of nine Research Departments, Administration Department, Research Planning Office. Technology Advice Office and two Coal Mine Safety Research Centers in Hokkaido and Kyushu.





所長 工学博士 高 多 明 Director-General Akira Takata, Dr. Eng

地球上のすべての産業活動にとって,資源とエネルギーの確保はその基盤であり,豊かな人間生活はその上 に支えられています。この美しい惑星,地球の環境の中で、資源とエネルギーをいかに確保し,また有効に利 用するかは、公害資源研究所に与えられた責務であります。

科学技術における研究開発の急速な進歩には目を見張るものがあり、それに伴う産業の発展は人類に大き な恩恵をもたらしてきました。しかし、無限とも思われた天然資源やエネルギー、自然環境にも地球規模での 制約が認識されてきたいま、研究開発の面でもその調和を図るべきでありましょう。

そのためには、高度の技術と同時に広い学際的知識が必要とされます。当所は、研究開発に日夜努める工業 技術院の一研究所として、資源・エネルギーの開発から利用までの一貫した研究、さらにそれに伴う安全と環 境を確保する研究に、今後とも積極的に取り組んで参ります。

For all the industrial activities in the world, natural resources and energy are of essential importance, and a more affluent human life greatly depends on a sufficient supply of these resources. National Research Institute for Pollution and Resources (NRIPR) is responsible for the development of technology on how to secure the natural resources and energy, and how to utilize them effectively while preserving our environment of the beautiful planet, the earth.

Research and development in the field of science and technology have advanced at an amazing speed, and the developments of these progressive industries have brought about a great deal of benefit to mankind. During the development of technology it has been also recognized that the natural resources and energy are not infinite, and that the environment can be destroyed even to a global scale by industrial pollution. Therefore, future research and development should be carried out to harmonize human activities with the surrounding environment. Highly advanced technology and wide-ranging knowledge based on various fields of science are necessary for achievement of this desired harmony.

Agency of Industrial Science and Technology (AIST) dedicates itself to the welfare of people through the development of technology. NRIPR, as one of the members beionging to AIST, makes supreme efforts in carrying out studies on the safety of various industrial processes, on environmental protection of a global scale and on the exploitation and effective utilization of the natural resources and energy.

Role of the Institute

National Research Institute for Pollution and Resources is concerned with a wide range of research fields related to exploitation, processing and utilization of mineral resources and energy, mine and industrial safety and environmental protection. The Institute is affiliated with the Agency of Industrial Science and Technology under the Ministry of International Trade and Industry.

A great effort is being focussed on the following research subjects in each field at the institute.

1. Mineral Resources Development and Utilization

- Exploitation and development of marine mineral resources at offshore or at deep seabeds, such as manganese nodules, hydrothermal deposits and cobalt-rich manganese clusts.
- Advanced construction technology for underground space utilization.
- Production of new materials, such as functional silicune materials and ultrafine powder.
- Processing and refining technology for low quality ore and unexploited resources, especially rare metals.
- 2. Energy Development and Utilization
- Comprehensive utilization technology of oil-alternative fuel resources such as coal, natural gas, oil sand, oil shale

- and biomass, including organic materials technology.
- Advanced combustion technology utilizing various low grade fuels and energy saving technology.
- Geothermal energy exploitation and heat extraction technology
- 3 Environmental Protection
 - Comprehensive industrial pollution control technology for emission abatement, pollutants measurement and environmental assessment.
 - Pollution control and measurement technology of new chemical substances.
 - Advanced assessment technology for regional scales.
 - Global scale environmental studies on climatic change, acidic rain formation and transformation of chemicals in the troposphere.
- 4. Mine and Industrial Safety
 - Coal mine safety technology, such as gas and coal-dust explosions, mine fires and gas outbursts in support of the domestic coal mining industry.
 - · Safety assessment for utilization of underground space.
 - Safe demolition of old facilities using explosives.



Resources

大都分の褒要を商外に依存する我が国にとって, 鉱産物資 厳の長期安定供給の確保とその多様化,低質化に対応する有 効利用技術の確立は,我が国経済社会の発展と国民生活の向 上にとって不可欠の要件です。

そこためには、毎選鉱物質類など新しい質類の開発や低品 位、末利用資源の開発・利用、さらにはその高付加価値化が 強く望まれています。また産業の新たな展開を図るため、水 質額(工業用水類)の確保並びに地下空間利用技術の開発も 急務とされています。

当所では、マンガン団塊など商店鉱物資源の開発・利用, 花崗岩、蛇紋岩など国内資源の有効利用,レアメタルなどの 頼製業材化、炭素資源からの化学原料・新業材などの製造, 及びパイオテクノロジーと課分離を組合わせた水処理、再生 システムの研究並びに地下空間開発のための各種要素技術の 研究を推進し、これらの社会的な要請に応えています。 For Japan, as poorly endowed with mineral resources as it is, the sufficient supply of resources and the development of advanced technologies for utilization are of particular importance to support its vital economic activities and to provide for a more amiable life.

The Institute has conducted R & D on the techniques for mining and processing of deep sea mineral resources (manganese nodules, cobalt-rich crusts, polymetallic sulfides...), the advanced utilization of granite and serpentinite for industrial materials, the purification and preparation of rare metals into high-grade materials, the production of carbon materials from organic carbon resources, the reclamation of water resources with high-density bioreactor and organic membrane filter, and the underground space development and utilization



代表的なレアメタルの主要理蔵国とそのシェア Main Countries with Rare Metal Resources and these Shares



中国湖北省臺灣産レアメタル鉱石 傷一整液鏡 透過・オープンニコルを使用 (R. 金町石,Ho 魚閃石,Ep 緑れん石 ()



Land

オーストラリア産業砂鉱石 傷光勤焼筒、反射・クロスニコルを使用 (Mo:モナズ石.E、ユークセナイト, Cas:保石



CONTRACTOR AND AND INC.

南島島海域より採取したコパルトリッチ クラスト (Mn 12:0, Fel 29:4, N=0:82, Col:1:08, Col:0:0:55 x142:08



潔海底鉱物資源の開発

Development of Deep-Sea Mineral Resources

海洋は實際の宝庫といわれていますが、海底にもさまざまな鉱物資源が眠っています。経済的に有望なものとして、太平洋の深海底に砂利を敷きつめたように分布しているマンガン団塊、海底が拡大している箇所に湧き出したように分布している海底熱水性鉱床、梅山の頂上や斜面を覆うように分布しているカバルトリッチクラスト鉱床などが発見されています。

マンガン団塊については、世界の各国で採鉱技術の開発が 進められており、我が国においても工業技術院の大型プロジェ クトとして研究開発が進められています。当所では、このプ ロジェクトの基礎的な研究や、熱水性鉱床、コパルトリッチ クラスト鉱床の開発のための基礎的な研究を進めています。 Several mineral deposits deep at the bottom of ocean floors. Manganese nodules on the pelagic sea floor, polymetallic sulfide (PMS) around oceanic spreading centers, and cobalt rich manganese crusts of seamounts have been identified as potential economic resources.

Some groups around the world have been developing mining technology for manganese nodules. In Japan, AIST is promoting it as a national R & D program. The NRIPR is conducting fundamental research on nodules under this program and fundamental studies for PMS and Corrich manganese crusts as well.



海底鉱物資源の分布模式図 Distribution of Deep Sea Mineral Deposits



コパルトリッチクラスト酒度写真



マンガン団塊ボックスコア試料



熱水性鉱床試料

マンガン団塊の開発技術

マンガン団機探鉱実験システムは下図に示したように、面 底でマンガン団塊を採用する集鉱システム、海底から輸送す る場鉱システム、海中機器ハンドリングシステム、採鉱実験 船などから構成されています。当所では、このうち集低シス テム及び場鉱システムに関する基礎的な研究を進めています。 また、マンガン団塊の開発が海洋環境に及ぼす影響について も研究を進めています。

Manganese Nodule Mining Technology

The test mining system of manganese nodules consists of collecting, lifting and machinery handling systems, and a mining ship. The NRIPR is conducting fundamental research on the collecting and lifting systems. An environmental impact study is also being carried out.



マンガン団境採転システム Manganese Nodule Mining System

海底塊状鉱床の開発

マンガン団塊に次ぐ,第2,第3の菜商底鉱物資源として 注目されている商底熱水性鉱床やコパルトリッチクラスト鉱 床を開発するためには、その試存特性や物理的、工学的特性 を把握する必要があります。本研究では、商店の微小地形を 工学的に評価する方法の開発や、対象物の比重や弾性波伝播 速度などの物理的特性及び圧縮強さや被堀削性などの工学的 特性の把握を行っています。

Research on PMS and Co-Rich Manganese Crusts

An accurate assessment of the topographical characteristics and engineering properties of PMS and Co-rich manganese crusts are required for their development. An evaluation method of micro topography is being developed. Physical and engineer

探撮性の研究 Research on Collecting System

深海底の軟弱な堆積層の表面に分布している団塊を安全に、 効率良く採掘する集鉱装置を開発するためには、深海底堆積 層のきまぎまな性質を知り、集鉱装置との相互作用を把握す る必要があります。本研究では、実海域から採取した堆積層 の強き、支持力などの選定、模型実験による集鉱装置の芽動 の観測、集鉱装置の採掘機構の検討などを実施しています。



集鉱機模型曳航実験 Towing Test of Model Collector

撮鉱特性の研究 Research on Lifting System

マンガン団塊を海底から海面まで輸送する方法としては、 掲載管内に発生させた水流による水力掲載方式がとられます。 その方式として、ポンプリフト方式とエアリフト方式が提案 されています。これらの技術を確立するためには管内での固 体粒子の流れの状態を把握する必要があります。本研究では、 固液二相流、気液固三相流について、動揺管路での実験や遺 度変動実験などにより輸送特性の解明を行っています。



動摇傾斜管路爆鉱実験 Ore Lifting Test Using Inclined and Swaying Pipe



コバルトクラストー面せん断世録

岩盤開発と利用技術

Underground Space Development and Utilization



鉱物資源の開発

高い地圧を受けている地層あるいは軟弱な地層中に試存す る鉱物資源を開発するためには、従来とは異なる技術を確立 することが必要です。当所ではこの技術開発に関する研究を 現場実験、模型実験および数値解析などによって進めていま す。

また、南極などの極地における鉱物資源開発技術に関する ご売したっていうご。

Mineral Resources Development

It is necessary to establish a special method for development of mineral resources in highly stressed strata, or very soft strata. At our institute, studies on this problem have been carred out with in situ and laboratory model tests as well as numerical analyses.

Also, a study on a method for development of mineral resources in polar regions is now being research.



立方供試体に三方向から独立して載何する真三軸圧聴試験装置 True Triaxial Compressive Testing Apparatus

水嘯流利用技術

「木禰石を穿つ」という謎に示されるように、また、河川 の優食作用にみられるように、木の流れは時として大きな破 壊力を示します。これらの例は、長い時間を要する木の破壊 作用ですが、木の速度を通常より速めると、この作用を著し く急速に、一瞬のうちに生じさせることができます。

小さな孔から噴射させて人為的に高速化した水の流れ(水 噴流。あるいはウォータージェットと呼びます)の破壊力を 道具として利用しようとする技術が各方面で期待されていま す。当所では、この技術を創止での採掘作業や土木工事に利

地下の利用

近年,他下発輩所。石油地下備審基地の建設など、大規模 な地下の利用が見られるようになりました。しかし、快適な 生活環境を得るためには、今後、特に都市部の地下をもっと 有効け利用するべきであり、当時でも、若盤特性の調査方法。 地下空間の規則及び維持方法などに関する研究開発を変通し ています。

Underground Space Utilization

Recently, some construction of underground power plants and oil storage plants are taking place. However, only a few underground space utilization projects have been undertaken in Japan.

For a better human life, underground spece in urban areas has to be more affectively utilized.

The Institute is conducting research on a method for determining properties of underground, design of opening, and support system of opening



地下周辺における応力解析結果 Stress Distribution around an Underground Opening

Water let Cutting Technology

The prosive capabilities of water are well known. This erosive action can be concentrated and accelerated by creating localized coherent jets of high-velocity water. For over 20 years, water jetting has been used as a cutting and cleaning tool for industrial purposes.

The NRIPR is doing fundamental research on water jet cutting technology, particularly in abrasive jets, oscillating jets and air coated jets, to apply it to hydraulic mining and civil engineering.



鉱物資源の高度有効利用

Firetessing of Minerry Resources

花崗岩の資源化技術

花崗岩を集成する石英・長石、雲母という鉱物は、次のよ うな薫製な用途を終っています。石英はガラス対料として大 量に利用されるほか、高純産のものはハイキク用のシリコン 原料に、混石はガラス・セラミック原料として、雲母は電気 絶縁材料のほか、各種樹脂・215時の充填材としても使われ ています。

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本研究では我の国に豊富に産する花崗岩から石英,長石, 費用をそれぞれ分離して取りだし、様々な用途に向けて工業 原料化することを試みています、また整母の中には、リチウ ムやガリウム等の重要な元素もわずかながら含まれているた め、これらの元素の抽出についても検討を行っています。

Utilization of Granite for Industrial Materials

Quartz, feldspar and mica. If hich are also the main component minerals of granite, have the following important uses. Quartz is consumed in large quantities for making glass. Extra pure quartz is used for high tech products such as semiconductors and solal cells. Feldspar is indispensable for glass and ceramic industries. Mica is mixed with various plastics and rubbers to reinforce them in addition to being used as electric insulators. This study arms to recover those minerals from granite and process them for various industrial uses.

The extraction of rara elements such as lithium and gallium from mica is also investigated.



蛇紋岩の高付加価値化処理

国内に大量に賦存する蛇紋岩を監護処理でることにより、 比表面5~500m¹/gの繊維束状及び小粒集合体形状のアモル ファスシリカの生成方法を開発した。このシリカはこれまで にない素材適性を有しており、これを出発物質としてゼオラ イト、炭化珪素、水和珪酸カルシウム等数種のハイテク用珪 素系材料に合成するための研究を行っています。

また蛇紋岩の種類と酸処理条件に対応した生成物の形態制 御とその用途開発及びマグネシウム分の回収法を検討してい ます。

Advanced Utilization of Serpentinite

Amorphous silica with high purity and high surface area has been prepared from serpentimite by acid dissolution. The micro textures of the product are dependent on the original ores and acid treatment conditions.

中国レアメタルの分離精製技術

当所は、東北工業技術試験所と協力して、中国有色金属工 業総公司広州有色金属研究院との間で、国際産業技術研究事 業に基づく「中国レアメタルの分離精製に関する共同研究」 を実施しています。

- 具体的には、広衝栗木地区のニオブ、タンタル鉱石及び湖 計構踊地区のルチル発電処理レアメタル鉱石について、レア
- ・ 「ル元素の迅速定量技術」高援な鉱物相互分離技術ならび に高純度精製技術を開発しています。

To evaluate the potential of this amorphous silica for industrial uses, some siliceous materials such as high silica zeolites, calcium silicate hydrates and silicon carbide are being synthes, ized using the amorphous silica as a source of silica.



アモルファスシリカの微細組織

シリカの微細組織

SEM Photographs of Amorphous Silica with Different Texturer.

Separation and Refining of Raro Metal Ores from China

Our Institute, "Government Lidu/Itrial Research Institute. Tohoku and Guanzhou Research Institute of Nichtfervous Mictals have conducted this joint research under a five-year program. (FY 1988-1992).

Main research activities

- () Mineralogical study and chemical analysis
- Studies on various separation methods, gravity/maghetic/electrostatic separation and flotation for rare metal ores.

Resources

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鉱物資源の志行加度的材料化

レアメタルの精製業材化技術

エレクトロニクスなど先端技術産業にはレアメタルが不可 免となっています。これを含有する鉱石は低品位化が進み、 分離回収に多量のエネルギーを消費すると同時に、高先度化 への粉束がますます高くなるため、頼製工程は一段と複雑と なっています。このため低き有車資源からの有価金属の分離 回収と報製技術の開発を図るとともに、高純度化により金属 の潮在特性が発現され、付加価値の高い新機能材料への利用 が図られるような高純産業対化技術の研発の必要です。新業 材としてのレアメタルおよびレアメタル酸化物の敷粉体は、 単に微細であるばかりでなく、高純度であること、紅度分布 が狭いこと、均衡性があり、凝集性を示きないこと等が要求 され、単なる物理的な処理では対応が困難であり、精製段階 で均質なものを作る必要があります。水研究では、複雑な工 程を相し、しかも経済的に(三利な手段として、有機試薬と シマメタルの化学反応性を利用し、裏択的作業と認ら性の思 提 図ることにより高純度化と機能性の発現を密射に関連さ せてがらレデメタルの精製業材化技術の開発に取り組んでい ます。

Purification and Preparation of Rare Metals into High Grade Materials

Rare metals are indispensable to high-tech industry and the need of rare metal materials of high-requality is increasing more and more. Therefore, the initivation of purification and preparation process becomes very important. The important features of rare metals polyders into high gride materials are high purity, uniform particle size, homogeneity and high dispersibility in addition to having small particle size. The controlled preparation process through organometallic compound formation may realize those effectively instead of conventional physical methods. At this institute, the interaction cheracteristics and selectivity in chemical reaction between the urganic reagents and the formetals are utilized for high purification and powder synthesis, and the above mentioned highly controlled powders are prepared due to the ments of process simplification with a great reduction in posts.



レアメタルの相製業料10フロージート

Processing of Mineral Resources

超微粉碎技術

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本研究では、サブミクロン粒子を効率よく製造する新しい 超微粉砕機の開発を目的として、類斜軸ボールミルと円板粉 砕機について研究を進めています。類斜軸ボールミルは、円 筒状粉砕室の中心軸を回転軸に対して類斜させたもので、ボー ル及び砕料には回転運動と粉砕延円筒の軸方向の往復運動と が同時に作用するため通常のボールミルよりも粉砕能力が高 くなります。一方、円板粉砕機「写真参吹」「2、汚筒は粉砕 室に数枚が 3 央を穿った円板 1.4 206 つくを貫く一本のロッ ドを粉砕媒体として用い、機体を回転しと語」発生する鏖耗 現場を利用して砕料を粉砕するものです。

Rolling

各円板は各々独立して勧 この 詳細に対するパランスが 良く、接触点が増し、粉気効率が向上するものと期待されて います。評決ご幹の操作条件として、ボールミンキミッドミ 小等に比較、こ、同転速度が極めて大きいのが特徴でありま す。また、粉砕記つに張り 粉砕媒体はすべて「ルミナ製セ ラミックスで加工されているため、不純物の混入かなく、ク リーンな粉砕産物が得られます。

Ultrafine Grinding Process

At the institute, is new ultrafine grinding machine to produce submicron material with higher ultrafine grinding than other mills is under development. One example is called multiplics mill consisting of a cylindrical vessel charged with several centerperforated discs and a rod as the grinding bodies. The breakage of material is carried out mainly by the abrasive stresses caused through the rotation of the vessel. In this mill, each disc behaves independently, then the mill contents are distributed almost homogeniously and the number of contact points not only between grinding bodies but also between grinding bodies and the wall of the vessel increases so that desirable grinding efficiency is brought about. The operating condition of this mill for ultra fine grinding is characterized by an extremely higher rotating speed combared with the ball or rod mill case.



THERE IS NO . THE .









金鳳超微粒子の合成と機能化

RF プラズマやDC プラズマの超高盤を利用して磁性のある 金属・合立磁像粒子(0.1µm以下)を含成し、それを重ちに 有機板欄でくるんだのち酵素や薬剤を付けることによーニー 粒一粒が機能を持ったコロイド分散系を作ることができます。 当所ではこのように超微粒子が分散した状態で特異な機能が 発現する材料(ゼロ次元機能性材料)の割製のための研究を 進めています。現在はより強い磁石になりうるレアアース系 の合金・化合物超微粒子の合成技術とその超微粒子の表面機 能化および分散技術についての研究を始めています。



ニックル超微粒子の透過型電子顕微鏡写真 TEM Imilitie of Ni-UPPs



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炭業系資源の原料化・葉材化

炭素系質要に含まれている行加価値の高い化合物、あるい は炭素系質要を燃料化した際の副生成物を分離し、原材料化 することは炭漆系質要の経済的かつ酸合的利用を図る上から 重要な研究課題です。

すなわち石炭、石油、オイルシェール、バイオマス等種々 の焼料貨類について、それらのエネルギー類としての利用技 術開発と供行して、高付加価値化、すなわち原材料化のため の技術開発研究を行っています。

炭素系新材料

提其材料の多様な構造、物性は有機物から反応してはじめ に生成した固体(炭素前駆体)の構造によって大きく支配さ れます。当所では今後発展が期待できる気相及び固相炭素化 に重点をおいて研究しています。気相炭素化では熱ブラズマ 及び低級ブラズマ法などによってダイヤモンド硬質炭素など を析出させています。

また、固朽炭素化とし、は化学的手法によって特長ある高 分子を合成し、それを炭素化して機能性炭素材料を得る研究 を進めています。さらに、種合材料、多孔質炭素の製造につ いて基礎的研究を行っています。

Carbon Materials

An understanding of the structure of carbon prenursors is very important for obtaining characteristic materials in view of the relation to the mechanical electric and chemical properties of the end products. The study is primarily being directed to structural modifications of the precursors though gas phase and solid-phase processes which have a great potential to be developed. In the gas phase process of carbon deposition from chemical species activated by plaums OVD ato, an attempt is Preparation of Functional Colloids of Ultrafine Powder

Ultrafine powder technology offers a limitless potential in the near future. Fundamental studies on the preparation of functional colloids are being sarried out at this institute, in w⁻, h magnetic ultrafine particles (UFPs, less than 0.1 µm) are produced using ultrafigh temperature plasma reactors and their surfaces are coated with functional organic materials. Current ly, highly magnetized superparamagnetic colloids of vare earth metals or compounds are one of the target.



RF プラズマ装置 RF Plasma Reactor (for UFPs Processing)

Advanced Utilization of Carbonaceous Resources

Studies on the separation, functionalization and carbonization of useful components or by products from fuel-derived carbonaceouce resources are important both now and particularly in the future for the economical and total utilization of carbonaceous resources.

For the same reasons, studies should be carried out on hydrocarbon resources, petroleum, oil shale and biomass.

being made to modify the surface of substrates and produce high grade or hybrid carbon materials. Specific polymers synthesized by electrochemical polymerisation are carbonized during the solid phase process to prepare high performance carbon materials.



炭素の生成 Formation of Carbon 皮素が駆体が最終的にできる皮素材料の性質を支配する Structural modifications of precursors to produce

Stran and Bar



燃料速源の有機原材料化

石俊、石柚、オイルシェール、パイオマス等種々の燃料費 夢をエネルギー夢としてだけでなく、有機原材料の製造のた めに利用することは、これらの質要の配合的かつ経済的利用 を図る上から重要です。そのためには、右図の示すように、 飲業系燃料解算から有用成分を取り出すための分離プロセス、 分離して得られた成分を化学原材料または化学製品に変換す るための反応プロセスが必要です。現在、これらに関して効 家の高い新規技術を開発するために様々な研究を行っています。

Production of Chemicals from Fuel Resources

Fuel resources such as coal, petroleum, oil shale, biomass and so on can be utilized for the production of organic chemicals as well as for energy production. The production of useful chemicals from fuel resources is very important for the total and economical utilization of those resources. Separation r poesses for obtaining useful components or their mixturer from fuel resources and chemical reaction processes for curverting the components separated into chemical females or products are necessary for the production of chemicals from fuel resources, as shown in the diagram. At our research institute, various studies aimed at the development of efficient technologies and processes both for separation and chemical reaction are being carried out.



参料から原料、エネルギー同時主要システムの概念図 A Concept of CO, Free Utilization of Fuels

レーザー化学反応

レーザー光は、単一の波長と、そろった位相を持つ光線で、 かつ強力なパワーを持っています。本研究ではレードー主の 特徴を利用して、化学反応の制御、物質の起急純度にあらい は、クラスターの生成など、燃料質算を付加価値の大きい物 質へ転換する新技術の開発を目指しています。

Laser Chemistry

There have been many examples in which new technology came into being from a new scientific discovery and innovated an industry is just the case, laser light in coherent and monochromatic and gives very high photon density. We are working on projects to apply laser light with its special features to fuels to produce utirapure compounds or time clusters us new materials.



教科資源の有機原材料行の工程 Flow Diagram of Production of Organic Chemicals from Fuel Resources

炭酸ガスの原材料化

概頼からエネルギーを取り出した後の形態である抜酸ガス を利用して有用な化合物を合成できれば、有機資源の節約と 其酸ガス公害の軽減に役当ちます。当所では、遷移金属維体 等の敏媒を用いて装蔵ガスとオレフィン、アセチレン、アル コール、アミン署の小分子を反応させ、プラスチックや医農 薬原料となる基礎化成品の製造方法を研究しています。

Utilization of CO; as a Chemical Carbon Source

Fixations of CO, into organic materials are studied with transition metal complexes as the catalyst, which may serve to some extent for reducting the amount of CO, released into the atmosphere.



レーザー解射装置 Laser Irradiation System for Selective Photochemical Reaction





Resources

水総合再生利用システム

New Water Treatment System "Aqua Renaissance '90"

現在、我が国では、中長期的な木幣額の運迫化、上木、工 量用水の水源水質の悪化などが社会問題になりつつあります。 通聴産業省では、これらの問題に対応するため、近年、発展の著しいパイオテクノロジーと額分離技術を組み合わせた システムにより、都市下水や産業廃水を、その中に含まれる 有機汚満物をメタンガスに転化、回収しながら、工業用水な どに再利用できるように処理する技術の研究開発(アクアル オサンス '90)を昭和60年度より6年計画で実施しています。 本計画には、工業技術院の試験研究機関並びに協会と民間 企業で構成されている技術研究組合が参加しており、当所で は次に示す研究を担当しています。

As a means of solving the problems concerning the larger demand for water in future and the pollution of the source of water supply for the city and industrial water, a six-year plan entitled New Water Treatment System (Aqua Renaissance '90) was set up by MITI and research on this project was started as of 1985. The new system consists of a bioreactor having a high concentration of microorganisms and a membrane as the sepa rator of microorganisms from the bioreactor.

What is attractive about the new system is that it will be able to recover marsh gas from the organic material contained in the sewerage and industrial wastewater and will be able to reuse the treated water as industrial water

窒素除去の効率化

本研究では活性汚死法の固種分離過程に腹分離装置を組み 込むことによって、従来に比べはるかに高い微生物濃度を離 持すると同時に、汚死部留時間を長くして窒素除去に係わる 微生物(硝化酮、粉窒菌等、をリアクター内に保持し、窒素 除去を効率化することを目指しています。本リケッジ、で硒 化を行わせたとき、従来系を上回る窒素負荷で運転できるこ とが示されました。同時に、暖気槽内で脱窒が進行すること。 も認められました。本リアクターは一種式で BOD と窒素が除 去できる経済的な装置として期待されています。

Nitrogen Removal by an Activated Sludge Process with Cross-Flow Filtration.

The capability of ammonium oxidation of an activated sludge process with cross-flow filtration while retaining a higher concentration of activated sludge, and operating with very long sludge retention time is investigated. By this process, higher nitrogenous and organic loadings were attained, and some amount of oxidized nitrogen were denitrified. Thus, organic carbon removal, nitrification, and also denitrification may possibly all occur at the same time when using this process with a single reactor.



線分離リアクター

有機屢繁材の評価技術

本研究では有機または無機性の膜を用いて衛生物、有機物 等を分離し、リアクター内の微生物の保持、透過水の再生利 用を可能とするここを目標に、汚濁物質負荷に強く、微生物 により劣化せず、長期使用が可能な透過膜を適定しようとす るものです。当所では既存及び新規に開発される有機膜の検 討・評価を行っています。

: Evaluation Technique for Organic Membrane Materials

"o develop a means by which the microbes within the bioreactor can be maintained and the wastewater recycled for reuse, we have been conducting research related to high performance organic type membranes which exhibit excellent contaminant load capabilities, are resistant to microbial deterioration, and which can be used over long period of time due to their favorable durability characteristics.

Permeability test has been carried out on already existing organic membrane materials using wastewater containing anaerobic microorganisms.

Those materials, exhibiting superior performance, have been selected and the optimal operating conditions have been investigated.



後洋伴討餘装置

我が国はエネルギーの質量の約6割を商外の石油に依存していますが、石油質量は世界的に偏在しており、また埋蔵量も少なく、長期的な安定供給は保証されていません。そこで 我が国では環境問題にも配慮しつつ。将来のエネルギー問題 に対応できるような石油代替エネルギーの開発技術やエネル ギーの有効利用技術を確立しておく必要があります。

当所では、資源が豊富にあり安定して輸入可能な海外産石 酸のクリーンな利用技術としてその液化及びガス化技術、ス ラリー化技術、天然ギスやパイオマスからの酸体燃料製造技 術、劣質・重質相等の軽質化技術などの研究のほか、石油の 燃焼技術の改良による低公害化と熱エネルギーの有効利用に よる省エネルギー化の研究を行っています。さらに国内エネ ルギー質要である地熱エネルギーに着目して高温の地下岩盤 からの地熱エネルギーの回収技術、地熱抗井の効率的握制技 術等の研究を行い、エネルギー窒萎の多様化と有効利用を図っ ています。 About 60% of Japanese primary energy consists of petroleum which is all imported. Petroleum resources are irregularly distributed in the world and cannot cover the worlds enormous demand for oil consumption in the future. So, it is necessary to establish technologies both to use various energy resources and to use them very effictively considering the global environmental protection to maintain a high standard of living

The Institute has conducted various studies on energy technologies such as coal liquefaction and gasification for clean use of coal which has large reserve of resources in the world. liquefaction of Natural Gas and biomass resources and upgrading of heavy oil. Also, new technologies for low emission combustion of coal and thermal energy savings are being developed. For utilization of geothermal energy, effective extraction of geothermal energy and drilling technology for geothermal wells are inder study at present.



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The institute has conducted various studies on energy technologies such as coal indusfaction and gasification for clean use of coal which has large reserve of resources in the world. Industaction of Natural Gas and biomass resources and upgrading of heavy oil. Also, new technologies for low emission combustion of coal and thermal energy savings are being developed. For utilization of geothermal energy, effective extraction of geothermal energy and drilling technology for geothermal wells are under study at present.



循環試験中の楽温岩体実験場(山形県附折温泉) Dverlooking the Hillon, Hot Dry Rock Test Site in Yamagata 石炭を積化し、原植あるいは石油製品の代替として使用可 載なクリーンな想料油を製造するプロセスとして、セレジャ イン計画では褐炭液化技術と豊貴炭液化技術についての技術 開発が進められています。そのなかには解決すべき、多くの 要果技術があります。

当所では主として基礎的な部門を担当し、置貴族から褐炭 までの各種石族について、石炭の物性から始まり、酸化反応、 粗酸化油の水準化設置、設置油の安定性や内燃機関燃料とし ての評価及び酸化副生成物の有効利用等の市広い研究を相互 に連携を保ちつつ総合的に進めています。これらの結果から、 石族の酸化反応条件と酸化生成物の量と置との配体の一般則 を見いだすとともに、生成物の改置処理によって、用途に応 じた性状の製品を製造するための基礎資料を得ています。

The Sunshine Project has developed Brown Coal Liquetaction and Bituminous Coal Liquetaction

The method, which has been carried out at this institute, is one of the coal liquefaction processes conducted by solvent extraction and is essentially a two stage process. The purpose of this research is to provide fundamental data for the selection of optimum reaction conditions for the liquefaction of various coals and the upgrading of z_{i}^{-1} liquid. Product Unitation is also being investigated.

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石皮液化装置 Coal Liguefaction Test Plant

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石炭から液体燃料への変換

石炭の液化反応

石族の離化反応ではまず、石炭を粉砕し、帯剤や触媒と攪 台してペーストとしたのち、高温、高圧水業下で反応させる、 いわゆる一次酸化反応で酸状生成物を出来るだけ高収率で場 ることが必要です。生成物の収率や物性は石炭の種類と反応 条件によって大きく左右されますが、当所では各々の石炭に 適合した常剤や反応条件の濃定をオートクレープあるいは小 型連続装置を用いた実験によって行い、更に酸化酸渣につい ても適切な利用方法を検討しています。

一次務化反応の条件は次の様な範囲です。

反応温度	380~480°C
反応压力	100~300 kg / cg
宿剤/石炭(重量比)	2~5
勉媒	教系

Liquefaction Reaction of Coal

In coal liquetaction, coal is first ground and mixed with a solvent and catalyst to make a feed paste. It is important to obtain c high yield of liquid product in the first stage liquetaction at high temperature under high hydrogen pressure. The yields and properties of the products greatly depend on the reaction conditions and coal ranks and thus selection of optimum conditions and suitable solvents for coal have been studied, at our institute, using autoclave and continuous bench scale plants. The utilization of liquetaction residues is also under investigation.

液化油の改置と評価

一次戦化から得られる粗積化抽は、窒素、酸素等のヘテロ 化合物を多量に含むと同時に、空気や光によって容易に着色 します。二次水素化処理は、粗積化油をクリーンかつ良質な 燃料にするための有効な方法です。

当所では、親化袖の水業化処理方法、最適反応条件を検討 するとともに、灯油、軽拙等の製品評価、貯蔵安定性、着色 防止技術、石油と混合使用する場合の問題点等について検討 しています。

Upgrading and Evaluation of Coai Liquid

Raw coal liquid contains lots of heterolatoms (such as nitrogen), and is readily colored by air and light.

To produce a good quality coal liquid, operating conditions for hydrotreating, evaluation of product oils, storage stability, color stability and various other problems have been studied.







石皮液化反応 Liquefaction Reaction of Coal

液化油からの有効成分の分離

相種化物の中には、化学製品の原料として利用できる化合 物が多種含まれています。これらを分離し、有効に利用する ことは、酸化油を安価に製造するためにも重要なことです。 当所では NEDO とも協力してその分離法について研究を進 めています。

Separation of Valuable Components from Coal Liquid

Coal liquid includes a lot of valuable components, which are useful for chemical feedstocks. It is very important to separate them coal liqued for reducing the price of the production of fuefrom coal.

We are studying their separation methods in cooperation with the New Energy Development Organization (NEDO).



圧力晶析法による石炭液化油中の有効成分の分離 High Pressure Crystalization of Valuable Components



石族を CO, H,, CH,等にガス化する技術は、100年以上も 利用されてきました。今日でも、クリーンなエネルギーへの 転換技術として、サンシャイン計画において営民会同による 開発が進みられています。このプロジェクトでは、石炭を1600°C 以上の発気により O, でガス化する技術を目標としています。

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○、職業でガエ化するため、(1)どんな機類の石炭でも使える、 (2) R₁, CO 以外のガスが生成しにくい。(3)小さな炉で大量の 石炭を処理できる。(4)石炭中の灰分はガラス状の固体になる ので、目然環境に与える影響を最少にできる、などの特徴を 持ったプロセスです。

当期では、効率よく右旗をガス化するための条件を詳しく 調べたり、効率よく生成ガスから硫酸化合物を除去する方法 を整礎的に解明しながら、このプロジェクトを支援していま す。この中で、右提の持っているエネルギーをガスのエネル ギーに載も効率よく転換するためのガス化条件を、理論と実 膝の両面から明かにして来ました。また、これまで運転され たガス化プラントの成績を世界中から集め、これを総合的に 解析するためのデータベースも作成しています。脱硫技術と してはガス化と同時に硫黄化合物を輸出する炉内脱硫技術や、 ガス化炉から出たガスを冷さずに脱硫する高温乾式脱硫剤の 開発を進めています。

Although the gasification of coal has been carried out for more than a century ago, the development of an advanced gasification process in which the problems of efficiency, reliability and environment have been solved, is still being anticipated. The R & D activities associated with coal gasification are being studied by government and private enterprises in cooperation with each other as a part of the Sunshine Project.

At this institute, a fundamental study for coal gasification at a tigh temperature has been carried out. One of the objects of this investigation was the accumulation of basic data to improve the gasification efficiency of coal. Fundamental studies ars in progress on gasification phenomena and the influence of molten slag on gasification reaction above the melting temperature of the ash using a bench scale moving gasifier, plas tometer, and other apparatus.





バイオマスから液体燃料の製造

バイオマスは非化石系エネルギー資源の中で、再生産可能 (リニューアブル) かつ唯一の炭素系の資源です。本研究の 目的は、木質系バイオマスである木材や樹皮、あるいは都市 廃棄物の下水汚況、アルゴール製造工程から排出される有機 汚況から液体燃料を製造することにあります。

このために、これらの原料を高温、高圧下で処理すること により最化反応を進行させます。現在、 5水汚泥の油化実験 に関してはペンチスケールの装置により、連続運転を行って います。

Production of Liqued Fuels from Biomass

Biomass resources are unique because they are renewable and carbonaceous among non-fossil energy sources. The purpose of this study is to produce liquid fuels resembling heavy fuel bil by thermochemical conversion.

In this process, woody material is mixed with an aqueous solution of alkali salt and treated at high temperature and under high pressure to achieve liquetaction. At the present time is learch is being done for sewage sludge liquetaction using a pilot plant.



隆化反応の原料 (木材チップ) Feedstocks for Liquefaction (Wood Chips)

この酸化方法は、(1)水素や一酸化炭素などの還元性ガスが不 要である。(2)反応が木相で進行するため。原料の脱木。乾燥 工程が不要である。(3)木材などのバイオマスは重油に比べて。 硫黄,塩素、重金属などの環境汚染物質が少ないためにクリー シな燃料が得られるなどの利点があります。一連の実験結果 によれば、適切に反応条件を選択することにより、7,000~8,000 kcal/kgの発熱量を有する液体燃料を約50%の収率で得ること ができます。

This process has the following advantages: (1) No hydrogen or carbon monoxide is needed; (2) No drying or dewatering of starting material is needed since liquetaction proceeds in an aqueous phase; and (3) Clean fuel is obtained since woody material hardly contains any pollutants such as sulfur, chlorine and heavy metals. According to a series of experiments, liqued



バイオマス液化装置 Biomass Liquefaction Test Plant

我が国では、未利用の木質系バイオマスが年間で約3,000万 mP程度が廃棄処分されています。また、下水処理場からは年 間5,000万mPもの下水汚死が排出され、同様に各製造工場から も膨大な量の有機性汚死が不可避的に排出され、これらの処 理、処分が大きな社会問題になりつつあります。これらは高 含水状態で排出されることが多く、水の存在下で反応を進行 させる本方法に適した原料であるといえます。当所で開発し た酸化法は、コストをかけて処理、処分しなければならない 廃棄物から逆にエネルギーを取り出すことができ、その成果 が大いに期待されています。

In Japan, roughly 30 million cubic meters of waste wood and 50 million cubic meters of sewage sludge are being generat ed anually. However, the difficulty of finding the available space for disposition and other environmental reasons are making the matter of waste disposal a serious social problem.

Since this type of liquefaction is carried out in the presence of water, it can be applied to many kinds of organic material containing water. For example, sewage sludge, pulping sludge, and peat are desirable potential candidates.





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我が国は原油のほとんどすべてを海外に依存していますが、 近年、輸入原油の質は次第に悪くなり、原油中に含まれる軽 質能(ガソリン、灯油、軽加など)の割合が少なくなる傾向 にあります。一方、これら中間三品の需要増、C 重油の需要減 にみられるように、国内石油製品の需要動向は軽質化の傾向 にあります。このギャップを埋めるために重質油から軽質油 を製造する技術を早急に開発する必要があります。

本研究で開発を進めている高分散触媒は、重要相とよく接 触するので少量でも重要相を効率よく分解して軽要相を製造 することができます。また、重要用中に含まれるパナジウム、 ニッケル、アスファルテン等の触媒被毒物要に対しても強い 抵抗力を持つなど、既存の触媒にない長所を有しています。

Aimost all crude oils consumed in Japan, are imported from foreign countries. Recently, the yield of light fractions in crude oils imported is going down. On the other hand, the demand for light oils (such as kerosene and diesel fuei) is increasing

To compensate for the lack of light oils, it is necessary to produce light oils from heavy oils

A homogeneous catalyst is useful for the hydrocracking of heavy oils which contain a great deal of vanadium, nickel and asphaltenes. Homogeneous catalysts have very high efficiency for contact between a catalyst and heavy oil molecules. Even a small amount of a homogeneous catalyst is capable of reducing asphaltenes in heavy oils. 天然ガスは、石油に比べて比較的広く駅存していますが、 取扱いが不便な気体燃料であるため、これまで世界の生産量 の20%が現地で焼却されています。我が国では、~160°Cに冷 却きれた酸化天然ガス(LNG)として商外から輸入され、大力 発電や都市ガスに利用されています。天然ガスを化学反応に よって報体燃料に変換する技術は、常温での輸送を可能にし、 自動車用・民生用液体燃料や化学工業用原料にまで利活用の 幅を拡大でき、さらに硫酸分や窒素分のほとんどを含まれな い良質燃料を供給できると期待され、我が国のエネルギー酸 の多様化に資献することができます。

Supportion -

OL Natural Gas

本研究では、天然ガスを供化水素や燃料アルコールに化学 変換するプロセスの開発を目的として、転編、反応形式及び 生成液体燃料評価を中心にした研究を行っています。

Although natural gas reserves in the world are next to those of oil, so far about 20% of the world production has been burned off at production sites because of difficulty in storing or handing gaseous fuels. Liguefied Natural Gas (LNG), cooled at 160°C, is imported to Japan for limited uses such as electric power fuel and town gas.

The chemical conversion of natural gas to liquid fuels is desirable. (1) to expand its utilization to automotive fuels, herting fuels, and chemical feedstocks. (3) to easily transport the fuels at ambient temperature, and (3) to supply clean fuels containing no sulfur or nitrogen compounds. Aiming at chemical conversion of natural gas into hydrocarbons and alcohols, research has been carried out on (1) catalysts for the conversion, (2) reaction processes, and (3) evaluation of product fuels.



生成油と触媒 Product Oil and Homogeneous Catalysts









石炭の高効率低公害燃焼

埋蔵量の多い石炭の利用を拡大することは、エネルギー政 量上不可欠のことですが、固体で灰分多く、また燃焼すると NO₄、SO₄を多く摂出する等石炭には多くの防懸があります。 この対策としてガス化・液化1 ンの方法ですが直接燃焼の 改善し有力な手段です。

循環流動層燃焼は最も新しい燃焼法の一つで、合催石炭を はじめ多様な固体燃料の低 NO.,低 SO.での燃焼が可能です。 現在、ペンチスケールのプラントで、燃焼通程の基礎研究を 行っています。

●高級式燃焼法は石炭を高温で燃焼し、炉内で溶融した灰を達心力の作用でガスから分離する燃焼方法で、テストプラントによる研究が進行中です。

従来の敷粉炭燃焼についても、炭種による燃焼性、NO、生 成特性の違いを調べています。このように総合的に石炭の燃 焼法の研究を進めています。

High Efficiency Low Emission Coal Combustion

Coal utilization must be expanded for stable primary energy supply. However coal has many problems, such as handling and pollutant emissions. Gasification and liquefaction are the countermeasures for them, and direct combustion is more efficient in energy utilization. Circulating fluidized bed combustion is one of the newest combustion methods. It can burn many kinds of coal and various soled fluels with flow pollutant emissions. Basic research on combustion procedures is carried on by using bench scale equipment. High temperature slagging combustion method is also under development by using a testing equipment. Conventional pulverlized coal combustion method is being studied to know combustibility and NOx emission characteristics of various coals.



衛價流動層燃燒試験装置 Circulating Fluicized Bed Compussion Equipment



憲温澄式燃焼試験装置 High Temperature Slagging Combustion Equipment

高効率な燃焼制御技術

- 産業用エメルギーの大部分は燃料の燃焼によって産み出き れていますが、地球規模環境問題に対処するためには高効率 な燃焼を達成し省エネルギー化を図ることが重要な課題です。 そのため、CARSなどレーザー応用計測、衝撃投管、レーザー ドップラー流速計などにより複雑な燃焼過程を解明し、様々 な運転条件のもとで燃焼状態を最適に制調する技術を開発し ています。

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Advanced Combustion Technology

In this research program, basic research on reaction kinetics and flame structure is conducted to develop optimum combustion control techniques for stabilizing gobal climate.

CARS (Coherent Anti Stokes Raman Spectroscopy) method facilities, laser pyrolysis laser fluorescence equipment, and shock tubes are used for reaction kinetics research in high temperature. Furthermore, the laser doppler anemometer is used for flame structure analysis.

This basic research is also being conducted to develop combustion simulation modeling.



レーザー加熱/レーザー誘起蛍光実験装置 Laser Pyrolysis / Laser Fluorescence Equipment



高温ヒートバイブ

本研究は高値ヒートバイブ(蒸気温度範囲350~1200°C) に おける最大熱輸送能力等と材料適合性を調べることを目的と して行っています。作動流体には、4種類の溶融金属(木銀、 カリウム、ナトリウム、リチウム)を、容器材料としてステ ンドス、インコネル等を用いています。下の写真は蒸気温度 800°Cで作動中のチトリウムヒートバイブ縦縮肥を示してい ます。

High Temperature Heat Pipes

This study consists of analyzing the maximum heat transport capability etc. for the high temperature hept pipes which is available for vapor temperature range from 350 to 1200°C and testing the container material compatibility. In the heat pipes, mercury, potassium, sodium, and lithium are used as working flued, and stainless steel tubing, inconel steel tubing, etc. are used as the container materials. The following photograph shows the condenser section part of the sodium heat pipe at the vapor temperature of 800°C.



ナトリウムビートパイプ Socium Heat Pipe

電気較正式熱量計

現代産業において、特に熱を加熱・動力用として利用する 分野においては、エネルギー夢となる物質を燃焼させたとき にどれだけの熱が発生するかが重要な尺度となります。そこ で、その尺度の基準となる標準物質の燃焼熱量を電気的なエ ネルギーを用いて精度良く値付ける方法を研究しています。

Electrically Calibrated Calorimeter

From the industrial point of view, in the field where the heat is utilized for heating or producing power in particular, how much heat the material for an energy source generates in burning is an important measure. So the method of determining the quantity of combustion heat of some standard samples presisely with electrical energy is under investigation.







髙温岩体の開発・利用技術

天然の熱水系を有していない裏撮の岩盤を裏握岩体といいます。裏温岩体から熱エネルギーを抽出する方法として2本の抗共を地下採筋の裏混岩体中に掘削し、これらの抗共間をフラクチャ(人工貯留層)で運転した後、1本の抗共へ冷たい水を注入して人工貯留層で暖め、もう1本の抗共から熱水あるいは蒸気として回収する方法が考えられています。

高温岩体を効率的に開発するには、地下の標節かつ高温の 岩盤に十分な熱交換面積を有する人工貯留層をつくらなけれ ばなりません。一般に、氷圧破砕法によって人工的にき数を 造成し、これを人工貯留層として利用しますが、き裂の規模 や方向は、岩盤に加わっ地圧、既存のき数等の天然条件と坑 井に注入する液体の性質、流量、圧力などの作業条件に依存 します。そのため、室内において、き数を含む岩石ブロック に地圧に相当する封圧を加え、水圧破砕によってき数を発生・ 成長させて、き数の発生条件や成長方向を明らかにしていま す。このときの破壊音(AE)を計測し、AEの発生値所と流 体の通路との関係も検討しています。

さらに、山形県射折の高温岩体実験場で得られた定方位コ ア(方位が明らかなコア)を用いて、天然き裂の方向や頻度、 岩盤の物性を調べるとともに、DSCA(差歪曲瞭解析)法 等により地圧を求めています。また、人工貯留層から抽出で きるエネルボー量を予想するため、人工貯留層の規模を知る 必要があります。そのため、現場においてヨウ化カリウムな どを用いたトレーサー試験を実施するとともに、コーピュー タシミュレーションによる貯留層評価を行っています。



水圧破砕実験装置と花崗岩中に発生した水圧破砕によるき殺(左上) Apparatus for Hydraulic Fracturing Experiment and Hydraulically Initiated Fractures in Granite Rock

Development and Utilization of Hot Dry Rock

The heat extraction concept from hot dry rock geothermainsystem requires drilling two wells into hot crystalline rock, connecting them at a depth through large hydraulic fractures, and then circulating pressurized water through this closed connected system to recover heat from the rock. The volume and orientation of hydraulic fractures depend on both natural condition such as earth stress and natural joints in rocks and experimental condition such as flow rate and water pressure. The water pressure required to initiate fracture and the orientation of fracture are investigated using rock blocks containing an existing joint in a laboratory. During these tests, acoustic emission (AE) is measured to map the fractures.

At Hijibri hot dry rock test site, we are investigating the fracture orientation and physical properties of rock using oriented cores. The earth stress is estimated by differential strain curve analysis (DSCA) method. Computer simulation and tracer teses are being conducted to evaluate a hot dry rock reservoir.



地圧測定のための DSCA 試験装置と試験試料 Apparatus for DSCA Stress Determination Method and a Specimen



憲温岩体貯留層評価のシミュレーション結果 A Result of Simulation for HDR Reservoir Evaluation





地熱エネルギーを利用するには地中に存在する熱水や蒸気 を地上に導くために、数千 m の坑井を掘削しなければなりま ん。地熱井銀剤は基本的には石油井銀剤技術が利用されてい ますが、石油井に比較し、岩盤が高温であると同時に硬質で あるため、これに適するように掘削方式や掘削工具(ビット) を改良する必要があります。このため地熱井の掘削乾率向上 を目的として、掘削方式の検討、ビットの性能評価及びビッ トの研究開発を行っています。

写真は搬削方式やビットの性能を基礎的に検討するための 掘削実験装置と実験用のビット類です。開発した新型ビット の方先には人造の多結晶型種ダイヤモンド (Polycrystalline Diamond Compact) を用いています。

Drilling Technology for Geothermal Wells

In order to drill geothermal wells economically and efficiently, improvements in the drilling techniques have been conducted. Aerated mud drilling methods have been introdu- ed to improve the drilling efficiency of rock.

New bits with polycrystalline diamond compact cutters are also being developed for the drilling of hot and hard formations. The photos below are of a drilling test facility and drill bits for the tests.

マグマ近傍からの熱抽出

低生産性の地熱貯留層やマグマ近後の超高温地層、きらに はマグマそのものを開発できれば、我が国の地熱資源をきら に活用することができます。高温地層中に捆削された机井を 同軸熱交換方式は、上記の地熱資源の開発に最も適した切 法と考えられています。これまでの検討により、新熱性能が 優れた内管を用いることにより熱交換器の性能が大幅に向上 することが明らかになりました。そこで、高性能の創熱新新 内管の開発を行っています。また、この方式の熱出力には地 層の熱伝導率や地層中の対抗が大きく影響します。このため、 知石の高温、含水状態における熱伝導率の側定や対抗が熱出 力に及ぼす影響の検討を行っています。

Downhole Coaxial Heat Exchanger System

The geothermal resources such as hot wet rocks which contains insuficient amounts of steam or hot water for power generation, very high temperature formations adjacent to magma bodies and magma itself are considered difficult to develop by ordinary heat extraction techniques. In order to realize the development of those geothermal resources, research on the new heat extraction system, the downhole coaxial heat exchanger system, has been carried out



經濟演算發展 Drilling Test Facility





短行了中国人民中留户建立







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数が国の環境問題は昭和40年代前半には探到な状況を呈し ていました。こうした状況に対処するため当所においては工 葉技術院における公害的止研究のセンターとして積力的に研 究を進めてきました。その結果。我が国は今日では公害対策 先進国とまで言われるようになりました。しかし、公害問題 の危機的状況を脱したとはいえ、問題系内痛や描における富 栄養化現象にともなう木質汚傷や自動車排出ガス等による大 気汚染はまだ未解決の問題として残されています。さらに最 近では CO,等による気候温暖化、フロンガスによるオブン層 破壊など地球規模環境問題が重要となっています。これらの 問題に対処するため当所においては発生源対策、計測技術、 汚染メカニズム、環境アセスメント等の各分野において研究 を進めています。

Air and water pollution problems were serious in the late 1960's in Japan. The National Research Institute for Pollution and Resources, operating under the Agency of Industrial Science and Technology, has conducted research and development on industrial pollution control technology.

As a result air and water conditions in Japan have been greatly improved over the last 10 years. However, there are still many pollution problems to be solved, and recently global environmental problems such as climate changes duo to green house gases such as CO₂, ozone holes caused by chlorofluorocarbons have become world problems. To solve these problems, the institute is carrying out research on the measurement and control of pollutants, their behavior and environmental assessment technology.



贈贈保全

她球環境問題

Global Scale Environment Problems

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她球規模環境予測技術

最近、環境が地球的規模で問題にされています。例えば地球温暖化であり、オゾン層の破壊、散性肉、砂模化、熱帯林の減少、海洋汚染などです。従来の大気汚発等の環境問題は発生原用辺の現象でしたが、産業活動の進展、新規化学物質の登場、過去の排出の累積により影響範囲が地球的スケールにまで拡大したと云えます。これらに共通しているのは原因と結果の間に時間的にも空間的にも距離があることです。

特に地球温暖化は人類共通の問題として解決がせまられて います。二酸化炭素は太陽光線には透明ですが、地表面から の赤外線放射を吸収する温室効果を持っています。太気中の 二酸化炭素の増大は、太陽エネルギーの収支に影響し、対流 幅での気温の上昇を、ひいては気候の変化をもたらします。 このような地球規模での環境問題を科学的に解決するため

このような地球現後での環境の場合が必要です。地 には、まず対象現象の正確な計構技術の確立が必要です。地 球的規模での環境のモニタリングには人工衛星や航空機を利 用した御定報と範囲方法の開発が進められています。

この大気中の二酸化炭素濃度の増大は人間活動による化石 燃料や木質系燃料の大量使用に起因しています。大気中の二 酸化炭素濃度の増加量は、化石松料の使用量から推定した二 酸化炭素の排出量の約半分で、残りがいずこへが吸収されて います。その機構は末解決で今後の研究課題です。解明への 手掛かりとして、二酸化炭素の地球的規模での収支・循環に 関する研究を行っています。これには、発生額での処理、燃 焼方法の改善や省エネルギー技術による燃料使用量の減少、 二酸化炭素の低排出燃料や新エネルギーの開発と転換等の発 生寂対策の有効性の評価も含めて考慮する必要があります。 温室効果を持つ気体は二酸化炭素のほかに、フロン、メタン、 亜酸化窒素等があり、これらの寄与し無視できません。フロ ン類は二酸化炭素と異なる赤外線領域に吸収帯があるため、 フロン濃度の増大は温室効果に影響が大きくなります。

これらを含めて全体的に考える必要があり、発生状況や大 気中における挙動を研究しています。

Studies of Global Environmental Pollution

The most update research subjects are global scale environmental problems. Those are climate changes, the depletion of stratospheric ozone layer, acid rain, deforestation, desertification and ocean pollution. The warming up of the surface temperature is an urgent and commonplace problem for humans. The probable cause is the increase of CO, gas in the atmosphere. This is assumed to be due to human activities in their consumption of coal, petroleum and biomass. CFCs are responsible for the ozone layer depletion and the greenhouse effect.

To investigate the global environmental subject this group is developing simulation models for materials and energy distribution on a global scale. The basic concepts are circulation (advection and diffusion), chemical reaction and balance of materials and energy including the different spheres. Monitoring systems using satellite and airplane are developing further the groundlevel measurements.



日本から放出した粒子の拡散 Particles Released from Japan





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環境汚染のリモートセンシング

人工業業や航空機などを利用したりキャトセンシングによ り、環境汚染を広域に、しかも同時に把握することができま 1.

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現在ではこの手法によって梅表面の水晶分布やクロロフィ ルの分布がある程度観測できるようになっています。しかし ながら、リモートセンシングを簡単現象の解明に用いるには モンサーと梅田の間の大気の条件、商表面の液や生物活動が データに大きく影響を及ぼします。そのため、ランドサット (TM-5) で得られるようなデータと簡単現象を対比させ、 統計的かつ画像解析的に現象を解明しています。また、大気 中の現象としては、森林火災の煙や、火山の噴煙などの拡配 状態を人工衛星データから解析することができます。今後は、 リモートセンシング手法を御祥と大気の両方に適用し、地球 殺権の環境汚染の潮淀法として研究していきます。

Remote Sensing Techniques for Marine Pollution Analysis

Remote sensing data like multi-band spectra of radiation are quite useful for environmental assessment of marine and atmospheric environmental studies.

Measurements of seawater temperature and chlorophyll in coastal area have been already developed. For utilization of remote sensing to marine environmental problems, we have to study the relationship between physical factors and biological activities in the sea through statistical and image analyzing techniques.

It is also quite effective for atmospheric environmental problems, like diffusion of forest fire smoke, volcanic eruption, etc. We will continue the study of remote sensing data analysis for further utilization of it to environmental studies.

二酸化炭業温度・フラックスの 野外立体観測による拳動の解明

糖化炊業、フロンなどは地表面から放出される赤外線を 吸収して対流圏大気を加熱する作用,すなわち處室効果を持っ ています。二酸化炭素、フロンなどの現状の増大傾向が続く と、今後50年間に1~2℃の気機上昇があると予想されてい ます。しかし、二酸化炭素濃度の将来堆移については、産業 革命前の濃度の二倍になる時期を2,040年代とする手種から。 将来にわたって二倍に遭しないとの予想まであり、未確定で す。このような将来農産予選が確定しない原因として二酸化 炭素の循環過程が解明されていないこと、将来のエネルギー 使用見込みに幅かめることなどがあげられます。

本研究では、二酸化炭素の大気、海洋、植物圏各貯蔵庫間 の豆換過程を飛行機、地上野外観測により解明します。観測 は棘崖、発生療状況の異なる北海道から沖縄に至る陶上、針 業樹林、亜熱帯樹林などで四季にわたり行い、二酸化炭素菌 環過程の季節・鱗度変化について検討していきます。

Study of CO, Behavior in the Environment by the Field Observations

Trace gases such as CO, and CFC (Chlorofluorocarbons) absorb the infrared radiation from the earth surface and have possibility of changing the climate.

If the tendency of OO, and OFC concentration continues to increase, the air temperature at the surface will be increase by 1.2°C in the following 50 years

The assessment of the CO, concentration in future is still not clear due to the uncertain behavior of CO, in the environment and the uncertainty of the fossil-fuel comsumpution in IUTURA.

Field observations using airplanes and towers are being carried out over land and sea inside and around Japan. The behaviors of CO, and CFC under the various surface conditions and latitudes are being investigated as a result of these observations











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汚染機構解明

人間の活動に伴い二酸化硫黄や窒素酸化物のような無機が ス、ガソリン等の炭化水素や種々の有機化学物質、金属化合 物やススを含む粒子状物質など多様な汚染物質が様々な発生 費から環境中へ放出されています。これらは大気、水及び土 環に分配され、光、水、微生物等の作用をうけてその形を変 え、人体、動植物さらに自然環境、気候などに大きな影響を 与えることが懸念されています。

Mechanisms of Environmental Pollution

Various kinds of primary pollutants, emitted into the environment from many sources, find their way into the atmosphere, waters and soil and are transformed decomposed by complex chemical reactions. The products and primary pollutants both have serious adverse effects in the environment.

酸性雨生成機構解明及び監視技術

酸性雨の原因は二酸化硫量と窒素酸化物が硫酸と硝酸にな るためと考えられています。この大気中での酸化過程には気 相反応、需需雨磨などの水漏中反応あるいは固体粒子状物質 上での反応があり、酸素、オブン、過酸化水素など様々な物 質が関与します。酸性雨が何故、どのように起こるかを明ら かにするために、これら反応の様子や反応に保わる種々の物 質を分析、概定する方法、装置開発の研究をしています。

Mechanisms of Acid Precipitation and Development of Monitoring Technology

Acidification of environment is considered to arise from sulfuric and nitric acids formed from SO, and NOx in the atmosphere. Many kinds of oxidants (oxygen, ozone, hydrogen peroxide) and chemical species (ammonia, metal ions, aldehydes) are concerned with the atmospheric oxidation reactions taking place in the gas phase, in the liquid phase (clouds, fogs and rain) and in the solid phase (particles). In order to know why and how the acidification of environment occurs, the rate and mechanisms of atmospheric chemistry and development of the instruments and methods measuring the various kinds of chemical species are being investigated.

化学物質の対流圏での変換・分解

フロンなど安定な化学物質は対流圏では分解されにくく、 温室効果をもつとともに、成層圏では強い姿外線で光分解し て成層圏オジン層の破壊に関与すると考えられています。対 流圏に存在する種々の固体粒子状物質(エアロジル、土壌など)はフロン、フロン代替物質等を変換、分解する新たな化 学反応の場となる可能性があり、反応速度や機構を明らかに する研究をしています。

Transformation and or Decomposition of Chemicals in the Troposphere

Stable chemicals such as freens and their alternatives which cannot be decomposed by tropospheric gas phase photochemical reactions are considered to have a reaction to the greenhouse effect and the destruction of the ozone layer. They may possibly be decomposed by the photocatalytic action of the solids in the troposphere. The rate and mechanisms of this transformation and decomposition are upnet to action the transformation and decomposition are upnet.



酸性雨の被害 Forest Damage Due to Apid Precipitation



環境中へ放出される汚染物質(SO₂, NOx, 粉じん, 化学物質 その他)とそれらによりひきおこされる様々な環境汚染問題 Environmental Pollution Problems Resulting From Various Kinds of Primary Pollutants (SO₂, NOx, Particles, Chemicals and Others) and Secondary Dries, Formed by Complex Chemi cal Restricts



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有機ハロゲンの大気放出抑制技術

フロンガス、トリクロロユチレン等の揮発性有機ハロデン 化合物は、オゾン層破壊、発ガン性などが懸念され、広域的、 地球規模の大気汚染問題を引き起こしています。このため、 有機ハロゲン化合物を大気へ放出させないための防止技術の 開発が望まれています。

発生類において気体状有機へロゲンを多孔性固体に吸着し て回収する技術は、その再使用が可能となるため有望な防止 技術と考えられています。本研究では、新規吸着剤の開発、 吸着有機、ロゲンの高効率な回収方法、並びに回者を組合せ た新設着システムの開発を研究しています。

また、上記の吸着法では抑制できない低濃度な廃ガスや廃 撃物中の有機ハロゲレについては、新たな技術を開発する必 要があります。このため、低濃度、難分解性等多くの問題点 を克服すべく、各種高エネルギー並びに触媒による分解法を 研究しています。本研究では分解法の確立とともに、分解し カハロゲンの固定化、並びに微量初生物の分析こも注意を払っ ています。

Emission Control of Halocarbons

Chlorofluorucarbons and trichloroethylene have brought about serious problems of air pollution. Effective technologies for emission control of the halocarbons are greatly needed

Recycling of the halocarbons by adsorption is suivable for the emission from many industries. Research efforts are being made on novel absorption materials, efficient desorption from adsorbed phase, and optimal systems for the adsorption desorption cycle.

Moreover, decomposition of the halocarbons in dilute exhaust gases and wastes is also developed by means of high level energies and catalysis. In addition to the decomposition methods, emphasis is placed on fixation of decomposed halogen and analysis of trace by products.



有機ハロゲン系産業廃棄物の焼助処理

有機ハロゲン化合物は、部品洗浄用や密剤として多量に僅 用されていますが、これらは自然界では分解されにくく、放 出されると環境に悪影響を与えるものが多いため。使用後に 奥酸物を適切に処理する必要があります。焼却処理を行なう 場合は、ダイオキシンのようにさらに有害な物質ができない ように注意する必要があります。

これらの廃棄物を安全に規却するために、処理条件が悪い とどのような化合物が生成する可能性があるか、また、有害 物質を生成させないためには、どのような条件で燃焼させる べきかについて、実験排気の処理装置を完備し、安全性の配 薄された施設で研究を行っています。

Treatment of Industrial Types of Waste Containing Halogenated Organic Compounds

Recently, the amount of hazardous waste containing halogenated organic compounds is increasing. These type of waste should be decomposed properly to avoid environmental pollution. Incineration is the most feasible treatment for their disposal. However, it is important to clarify the conditions of safe treatment because there is the possibility of forming more toxic compounds during the incineration of some halogenated compounds. In this investigation, the possible products and mechanisms of the thermal reactions of halogenated compounds are going to be studied to predict the optimum conditions for their combustion. Bench scale combustion enserts will also be performed to investigate the practical aspects.



有機塩素化合物の熱分解反応例

Thermal Decomposition Behavior of Chlorinated Organic Compounds



産業立地環境予測

大気汚染予測

大気中に産遊している粉じんは、一次粒子と呼ばれている 剤のから粒子として大気中に放出されるものの他に、大気中 で気体から粒子に変化した二次粒子と呼ばれるものがありま す。これらの粒子の拡散や、二次粒子がどのように生成され、 構成して行くかを調べるために、用行機やヘリコプターにパー ティクルカウンター、NO。計などを搭載し粒子状物質の濃度、 粒径分布、NO。濃度を構定し、同時に風速、罹窒、気磁など 気象の立体分布を確定します。また、地上付近でも数十回の広 い範囲の中に覆定点を配置し、濃度や気象の時間的な変化な どを調べます。これらの離定結果を用いて粒子状物質の拡散 シミュレーションモデルを開発しています。

また、先端技術関連の産業から排出される環境市築物質の 拡動予測手法についても研究していますが、工場が内陸の盆 地や谷間など複雑な地形の場所に立地している場合、気気が 複雑なためにSF,などの無害なガスをトレーサーとした拡散実 触や、地形模型を用いた風刷実験などを行っています。

Simulation Model of Diffusion for the Particulates

Particulates flowing through the air are composed of primary particles released from sources as particle phase and secondary particles changed from gas phase to particle. Diffusion of these particlates including the formation and removal process is investigated by particle counter. NOx analyzer and other equipment on an airplane or helicopter, with the measurement of wind, temperature and humidity. Near the ground surface, beyond several tens kilometers in horizontal, the concentrations of contaminants and weather are observed. The data are utilized to develop simulation model of diffusion for the particulates.

Diffusion experiments with SF, as a tracer and wind tunnel experiments over complex terrain are being made to clarify the effects of configuration on the diffusion, since some factories concerning advanced technological industry are located within very complex terrains.

山を越す気流についての風潮実験

大気拡散場構造

※築物質の移機と拡散は主に地上から2000m 程度までの次 気境界層中で行なわれます。そのため閲聴風や都市のヒート アイランドなども汚染物質の拡散に大きく影響します。この ような大気拡散場の構造を調べるために、飛行機や低層ゾン デなどを利用して乱気や血度分布などを広範囲で側定してい ます。また、後間形成される接地逆転層中で汚染濃度が極め て高くなることがあるため、接地逆転層の構造についても小 塔を利用した乱流圏定などにより荒れの場について解明を進 めています。これらの結果を用いて、計算モデルによりコン ビュータで大気の流れを再現する試みを行っています。

Mechanisms of Atmospheric Diffusion Process

Pollutants released near the ground surface are mainly advected and diffused in an atmospheric boundary layer lower than 2000m or so. Under this circumstance, sea/land breeze circulation and heat island near urban areas affect the diffusion of the pollutants. Measurements of turbulence and temperature with an airplane and low altitude sonde are carried out to investigate the mechanisms of airflow in the boundary layer. Concentrations of pollutants often become very high at night in the inversion layer near the ground surface. Turbulence in the inversion layer is also measured by equipment on short masts. Using these results, computer simulation of airflow can be carried out.




Environmental Protect

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海洋污染予测技術

大都市近くの内穫地帯の米質汚染は工業地帯からの初水、 都市頂水、近郊の農地から利用を経由して際に流れ込む農業・ 畜産の排水等の複合した形態で起こっています。これらの約 岸地帯は廃止の交通の重要な場所として、産業活動の拡大に 伴う埋め立ての場所として、また一方では市民の釣り、ボー ドセイリングなどのレジャーなど多方面に利用されています。 近年、ウォーターフロント計画の進行にともない水辺の重要 さが叫ばれています。また、新しい埋め立てや工場の立地に 伴って生じる地形の変化や初水の質・量の変化は水質や御路 の汚染に大きく影響を及ぼします。

これらの影響を予想するために数値シュミレーションを用 いて水質や読賀の将来予測を行い、増め立て地形や工場の立 地の適正化を認ります。

Prediction Methods for Marine Environmental Pollution

In general, the water pollution in coastal areas is caused by mixed wastewater from industries, agriculture, stockbreeding and sewerage. We easily recoginize sea phenomena with physical factors such as fidal currents and wind waves, chemical factors such as water quality and salinity, and biological factors such as the abundant growth of phytoplankt, " and marine organisms. Nowadays, we found to realize the problems of water pollution in open ocean caused in oil spill, heavy metals and chemical breahic compounds. These problems are closely related to the economical and industrial activities of mankind, so we must protect the marine ecosystem and more adequately manage the social behavior of activities of humans in future. Our laboratory is investigating predictive methods for pollution effects on natural coastal waters. For instance, an attempt is being made to develop an ecological and hydraulical numerical simulation model by a computer. For this purpose, we are developing instruments for field serveys and studying a methodology for field observation to obtain accurate data from the 56 ă

沿岸環境保全技術

治学の永賀、底賀、生物の理境は彻水の流動、波波、潮位 あるいは流入してくる種々の排水(工場、都市、農業、新船な ど)や同川の水の量→質によって大きく影響を受けます。 沿岸の際域特有の沢山の種類の生物(梅藤、魚介類、鳥類) が生命を習んでいます。これらと季節、年月を問わず人間生 活とパランスよく生活をしなくてはなりません。近年の産業 構造の変化にともなって分解されにくい物質、蓄積性のある 物質(農業,ハイテク産業物質)などの毎への流入が問題となっ ています。

本研究では、梅に生息する生物(ブラシクトン、ベントス、 パクテリア)と木質,底質との関係の解明に重点を置いていま す。これらの徹定結果を利用して梅の環境保全を図らればな りません、そのために生態系のシュミレーションの開発とそ の精度の向上を図っています。

Development of Marine Environmental Management Techniques

Coastal marine environment such as water quality, properties of sediment and life of organisms is affected by physical factors as tidal current, wave motion, meteorological conditions and chemical factors as discharge of waste water. We must develop an observation instrument and systems for maintaining a clean coastal environment to protect many organisms.

We are studying the relationship between environmental phenomena and marine organisms which enfocussed on bacteria, plankton and bentic organism for ecological study in coastal areas. Moreover, we have developed a numerical model for eutrophication and using it to simulate the environmental changes in the coastal waters.





環境汚染と毎洋生態系

1 11 12 1

Environmental Protection

地下水汚染の予測技術

地下水は、費量な水質層として、量的にも質的にも管理及び監視 を行わなければなりません。しかし、最近になって、有害な化学物 質による地下水汚染が広範囲に生じていることが判明しつつありま す。その対策としてはまず、地下水がどのように汚染されていくか の予測の方法を確立する必要があります。

CARLON CROSS.

右上の回は、典型的な地下水汚袋を示した例です。地表面に汚袋 類があるとすると、師商によって地層中(不飽和層)を浸透し地下水 面に達します。その後に、地下水のある層(帯水層あるいは飽和層) を地下水の流れによって移動し広がって行きます。これらのプロセ スを解明することによって、地下水汚袋の予測手法が確立されます。

種々の汚染物質は、地層中で蔵養したり変質したりします。この 研究では、変化する物質としてトリクロロエチレン等を選び、その 移動や広がりの機構解明に努めています。

Prediction of Groundwater Pollution

We must manage and monitor the quality and quantity of ground water as precious water resources. However, in recent years, it has been found that groundwater pollution by many dangerous chemical substances is becoming a wide occurrence. For appropriate treatment, it is necessary to develop a prediction method of groundwater pollution processes.

The illustration above shows a typical pattern of groundwater pollution. Contaminants on the ground surface inlitrate through the soil (unsaturated layer) to the groundwater surface and migrate to the groundwater (aquifer or saturated layer). A prediction method of groundwater pollution is being developed on the basis of the observation of these processes. Various substances decay and change in the soil. This project examines substances decay and change in the soil and examines trichloroethylene et al. as non-conservative tracers and follows those migrations.



|地下水汚染モデル Groundwater Pollution Model



地下水流模擬装置 Groundwater Flow Simulator

計測・分析

環境保全, 当害防止施策を推進するためには大気汚染物質 あるいは木質汚産物質を高精度で、しかも迅速に搬定する計 期技術が重要です。これにより、汚染の原因及び影響が解明 されると同時に、種々の防止対策はもちろん, 法的な対応も 可能となります。発生熱対策、環境監視、予測あるいは計測 の標準化などの多様な目的に対応し得る高度な計測技術の開 発を先導的に行っています。あわせて、今後のエネルボーの 多様化、産業界の動向をも見極めながら、新しい環境汚染物 質を予測し、これらに対処する計測技術についても研究を進 めています。

The development of accurate and rapid analytical methods for air and water pollutants is essential to preserve a clean environment, and make possible the identification of the sources of pollution and estimation of their effects on the environment. Furthermore, the establishment of various pollution control and regulation systems owes much to the development of these measuring methods.

This institute conducts studies to develop new analytical methods useful to control monitor and simulate pollution. In addition, analytical methods for new pollutants which may emerge from change in future energy sources and industrial.

Measurement of Pollutants





発生源ダストの新しい測定法

昭和47年に環境基準が設定されてから、大気中の存着粒子 状物質 (SPM) の基準達成率は、全国的にみて極めて低いレ べルで推移しており、現在も大気汚染行政上の大きな問題と なっています。この対策として、SPM の発生豪寄与東解明の 研究は重要となっています。現在の公的側定法によって得ら れる発生類データを基に展開されるソースモアルでは、ばい 煙が大気中に放出されたのち生成する粒子に関する情報が十 分でなく、SPM の現状を正確に把握することが困難になって います。

本研究は、現状の発生数ダストの測定方法を見直し、排ガ スが大気中に放出されて凝縮あるいは酸化過程をへて、大気 中で二次的に生成するダストの評価をも含め、SPM との相関 を考慮した新しい側定方法の確立を目指しています。

New Measuring Method of Source Dust

Since an environmental quality standard for suspended particulate matter (SPM) was enacted in 1972, the compliance rate with the standard has remained at low level in Japan, and it is a very serious administractive problem to improve the compliance rate. The identification of emission source is important for this purpose, but it is difficult to determine the origin of SPM because the data about particle, which is produced from exhaust gas after it is emitted from stack to the uir, is insufficient in the source model based on current official measuring method.

At our Institute, research is being carried out to develop a new evaluation and measuring method for source dust in consideration of the relation with SPM by improving the current official method. The new method can take the contribution of secondary particle into consideration, which is produced through condensation and oxidation from exhaust gas in the air.



地下水汚染物質の計測

地下水は豊重な水管要として工業用水,生活」 されていますが、最近になって有機塩業系化合物による広範 な汚染が判明しています。地下水は一度汚染されるとその回 復が非常に困難なので未然防止が重要です。したがって、地 下水の水質監視や、事業所、埋め立て処分場における水質管 理を十分に行なう必要があり、このため地下水汚染物質を迅 速に棚定する技術が不可欠です。

本研究では、地下水汚染計像技術の高度化を目的として、 光ファイパーやレーザー等の先端技術を応用した汚染物質の 遠隔計算法や高感度分析法の検討を行っています。すなわち、 有機汚染物質についてはレーザー蛍光法、光ファイパー化学 センサー法、無機汚染物質についてはレーザー励起イオン化 法を適用し、汚染物質の地下への欄出を早期に検出し、また、 地下水質を的確に把握することが可能な計測技術の開発につ いて研究を進めています。

Measurements of Pollutants in Groundwater

Groundwater is regarded as one of the precious water resources. However, in recent years it has been found that groundwater is widely polluted by toxic organo chloro compounds, etc. Since groundwater is difficult to clean if it is once polluted, prevention of the pollution is of extreme importance. Rapid analytical methods for pollutants, therefore, are essential to sufficiently monitor the quality of groundwater. The object of this investigation is to establish rapid and remote measuring methods for pollutants in groundwater. This investigation consists of two parts 1) the development of remote optical fiber fluorimetry for the determination of organic pollutants; and 2) the development of laser enhanced ionization methods for the determination of trace inorganic pollutants.



レーザーを用いる地下水汚染物質の検知システムの構成 Block Diagram of Measuring System for Groundwater Contaminants





環境保全

職場を守るためには、公害はその発生際において防止する ことが大切です。我が国においては大気汚染防止、水質汚傷 防止について厳しい対策がとられてきた結果、公害が減少し、 大気、水ともきれいになった際は明らかです。しかし近年汚 染の発生薬、発生形態には変化がみられ、都市用辺や幹線道 路における交通公害や開沼などの閉鎖水域での水の汚染、新 素材や電子技術関連産業からの新しい化学物質による汚染な ど、複雑、多様化に対するきめこまやかな対策が必要となっ てきました。

当時ではこれらの問題に対応するため、大気汚染の発生養 対策として、燃焼に係わる低公害燃料、低公害燃焼、排気ガ ス浄化、並びに化学物質の分離・分解について、水質汚染防 止としては活性スラッジにより廃水を処理する生物処理法。 分解、分離、吸着などの適用を図る物理化学処理法について 幅広く研究開発を行っています。

In order to protect our environment from pollution, it is important to control pollutant er usion sources. In Japan, we have taken stringent measures against air and water pollution As a result, environmental pollution is decreasing year by year and the air and water are returning to their former clean state. These days, however, emission sources and the character of pollutants have changed in various ways. We need a further detailed investigation on complex pollution systems, such as

流体化燃料用クリーンコールの製造

石炭を利用する際の最大のホックは低黄及び灰分等の環境 汚染を起こしやすい不純物を多量に含有し、また固体である ために取扱いに手間がかかることです。従って、石炭を利用 する前にあるかじめ不純物を除去し、クリーンで利用しやす い石炭・油スラリー (Coal Oil Mixture: COM)、石炭・ホ スラリー (Coal Water Mixture CWM) 等の流体燃料用の 石炭を製造する粉紙・脱灰技術が重要となります。石炭から 灰分・硫黄分等の不純物質を除去するコールクリーニング法 に関しては多くの技術開発が各国で進められています。

当所では、石炭粒子と不純物質粒子の表面性質の差を利用 して両者を分離するカラム浮遊連度法及び裏剤を用いて処理 する化学的タリーニング法の研究を行っています。カラム浮 適ではカラム中の石炭スラリーに微細気泡を導入し、気泡に 付着した石炭粒子(緑水性粒子)をカラム頂部から回収し、 不純物質(観水性粒子)を底部から排出して両者を分離しま す。化学的タリーニング法は不純物を水に可溶な物質に変え て餘去します。

Ultra Clean Coal Processing for Fluidized Fuel (CWM, COM)

Using coal as a substitute for oil poses serious air pollutant problems due to ash and sulfur emitted when coal is burned. Therefore, it is important to develop techniques to remove these substances from coal prior to combustion.

Coal cleaning processes offer economical means for reducing the sufficient and sub-content to environmentally screptsbir

Pollution Control

traffic air pollution around big cities and along main roads, water pollution in semi-closed water areas such as lakes and marshes, and pollution paused by new types of chemical compounds from new material and electronic industries.

To cope with these problems, we are conducting research to develop better techniques that will ensure high quality fuel, low emission combustion systems, exhaust gas purification techniques and separation decomposition of chemical compounds for air pollution reduction, biological treatment systems of wastewater using activated sludge and physical chemical treatment systems applying various decomposition, separation and adsorption techniques for water pollution reduction.





" 明境保重



Environmental Pro

高燃料比炭の燃焼技術

石油危機を変優として、ポイラーや工業炉などにおいては、 石油代替燃料としての石炭の利用拡大が積極的に推進されて います。この石炭利用拡大のほとんどは毎外からの輸入の増 加でまかなわれる予定です。そして、この商外炭の中でも高 燃料比炭が増加する傾向にあります。

Lobe wants

高燃料比炭は燃焼性が悪く、NO、排出濃度が高いという欠 点があります。そこで微粉飲ガス化燃焼や循環決動層燃焼の ように、低公害で高効率な燃焼技術の開発を行っています。

Combustion Control Techniques for Low Volatile Coal

Coal has been utilized positively in commercial steam boilers and furnaces as one of the main alternative energy sources since the 1970 oil crisis. Therefore, it is forecasted that the demand for imported coal will increase in Japan. Especially, the percentage of low volatile coal among the imported coal is on the increase.

Low volatile coal has poor combustibility and high NOx emission level in comparison with high volatile coal. Therefore, combustion techniques for low volatile coal, such as a pulverized coal combustion with # gasification process or a circulating fluidized bed combustion have been studied to control the elission of pollutants and obtain high combustion efficiency



微粉炭燃烧炉 Pulverized Coal Combustion Furnace

NO. 低減化のための触媒燃焼技術

燃焼反応を触媒表面で低温で行わせる触媒燃焼は、火炎燃焼に比べ本質的に低 NO.,低公害の特性を持っており、これを小型燃焼器用に実用化できれば、大都市域での群小発生夢からの NO.排出を大幅に減らすことができます。そこで、燃焼触媒の評価や触媒中での燃焼反応等の基礎研究と触媒燃焼装置の高性能化のためのシステム研究を行っています。

Catalytic Combustion Techniques for the Small Scale Stationary Sources

Since the low combustion temperature on a catalytically active surface, the use of catalytic combustion has shown significant advantages in the control of emissions. The purpose

ディーゼル自動車排出物質対策

現在,環境中の二酸化窒素及び原遊粒子状物質の環境基準 の達成状況には改善の傾向が認められません。そのため、こ れらの主要排出都であるディーゼン自動車に対する排出規制 の強化が望まれています。しかしながら、ディーゼル機関の 燃焼特性として窒素酸化物に対する低減対策は粒子状物質の 排出を増加させる一方,粒子状物質に対する低減対策は窒素 酸化物の排出を増加させるという二律背反の傾向があるため、 新しい観点からのアプローチが必要とされています。

当所では、ディーゼル機関から排出される窒素酸化物の低 減のため、酸化等囲気下で一酸化窒素を低減できる触媒の開 発、粒子状物質低減のためのフィンタートラップ等による単 増低減技術、窒素酸化物と単雄を同時に低減するためのディー ゼル燃料の改良に取り組んでいます。

Development of the Control Techniques for Diesel Exhaust Emissions

There is no indication that compliance rate with environmental quality standard for NO; and suspended particulate matter (SPM) has been improved. It is, therefore, desired to strengthen the emission standard for diesel powered vehicles which are one of the primary emission sources of NO; and SPM. For the contrary tendency between NOx and particulate emission that the control techniques for NOx increases particulate emission and vice versa, it is necessary to develop other approaches for the control techniques rather than engine modifications.

At our institute, the control techniques for diesel exhaust emissions are under study, especially in the survey of the catalyst for NO reduction, development of trap systems for particulate reduction and improvement of diesel fuels for simultaneous reduction of NOx and particulates.



排出物質特性を評価するための試験装置

超臨界流体による排水中の離分解性化学物質の 処理・回収

化学物質の多様化、使用量の増大に伴い、環境中への化学 物質の拡散、蓄積が採刻な問題になっています。本研究は根 水中の微生物により分解しにくい化学物質を、超越界液体を 利用して効率よく処理する技術の開発を目的としています。 臨界点以上の状態にある物質は、わずかな圧変化で化学物質 の溶解度が急変する等、特徴ある性質を持っています。これ を超臨界流体と呼び、流体の持つこのような性質を利用して 根水中の化学物質を分離、回収することが可能です。根水中 の化学物質の濃度が低い場合は、吸着肥等で根水を処理し、 使用ずみの吸着剤を超臨界流体で再生する方法を、根水中の 化学物質の濃度が高い場合には、肌水と超越界流体を向流接 触させて処理する方法を研究しています。

また、これらの方法を実行するときに必要となる、超結界 状態での相平衡データの収集も行っています。

Treatment and Recovery of Biological Refractory Chemicals in Wastewater with Supercritical Fluid

According to the production of various chemical substances, the dispersion and accumulation in environment of these chemicals have become a serious problem. The purpose of this research is to develop supercritical fluid technology for removing chemical pollutants that are not amenable to biological treatment from wastewater.

A subtance that has been brought beyond a critical point, has the great solubility change with relatively small changes in operating conditions. This substance is referred to supercritical fluid (SCF). SCF can be used to separate and recover pollutants from wastewater by two processes. One is SCF regeneration of adsorbents that have become saturated with pollutants, allowing the adsorbents to be recycled. This method is advancageous in the concentration of dilute pollutants. The other is direct counter current contact of SCF with wastewater for recovery of pollutants. This one step process is applicable when the concentration of a pollutant is relatively high. The data on phinse equilibrium for many SCFs is also collected. This data others the key to useful applications of SCF technology.

微生物による化学物質含有廃水処理

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際水による水質汚染防止は、従来、BOD(生物学的酸素素 家童)やCOD(化学的酸素要求量)といった包括的な影像に 薬づいて行われてきました。生物処理を中心とした多くの廃 水処理法が開発され、現在では、これらの基準はほぼ慮足さ れています。しかし、近年、これらの基準で捉えることが困 難な、有害化学物質による環境の汚染が防酸となってきたこ とから、傷々の化学物質についての廃水募集が起められるこ とになりました。本研究では酸生物の化学物質に対する馴化 能力を利用して廃水中の有害化学物質を勉強する方法を開発 することを回信しています。

そのため、有家化学物質に対する衛生物の動化方法、分離 活性の高い教生物の検索、分解の動力学、分解数生物の生現、 処理装置内での分解微生物の制御法界の研究を行っています。

Biological Treatment of Hazardous Chemicals in Wastewater

The regulatory emphasis in the Water Pollution Control Law in Japan shifted from regulation based on conventional pollutants, such as BOD or COD, to regulations based on specific hazardous chemicals and conventional pollutants. Hence, the need for innovative wastewater treatment technology to mansige hazardous chemicals in wastewater has been strengthened in this work, biological treatment process for hazardous chemicals in wastewater has been investigated based on the adapta bility of microbes to the chemicals. Kence, acclimatization of activated sludge to specific chemicals, isolation of microbes degrading the chemicals, kinetics of degradation, physiology of the microbes, and the behaviour of the microbes in activated sludge process, are under study.



超越界站平衡测定装置 Apparent of the Draws For an on the strengthere







世界に産する資源の膨大な量を輸入かつ角優している数が 国にとって、資源・エネルギーの開発、利用及び貯蔵為にと しやう安全確保は重要な課題です。さらに、産業活動を円滑 にし、快速で安全な社会をつくるために強力に保安研究を推 速する必要があります。

当所においてはご果、優発災子や、破壊の発生機構の解明、 それら災害の予知・予測、防災 ステムの()発酵を基盤とし、 創山保安はもとより大陸度地下 1間の安全・防災技術、制関 発破による機築物破壊活所の発度化、放射性廃棄物処分空隔 の安定性評価等の研究課題に取り組んでいます。 As a major importing country 63 natural resilurces extract ed from all over the workd, it is a very important subject to achieve safety during development of resources and energy and also their utilization.

For the purpose of safety achievement in mines, industrial sites, underground openings for moustrial use and others, basic studies on the occurrance mechanism solution of fire, explosion fracture and on the development of foreseeing or estimating technique and prevention techniques of industrial disasters are carrying out in the Institute. The new demolition technique of old facilities by blasting and the technique for the stability estimation of underground openings for disposal of radioactive woste are also the subjects being carried out.



自動水幕による危険性ガスの急速拡散 Rapid Diffusion by Automatic Water Spray



POメンプレン式度化ガスタンク

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產業火災対策

変要、エネルギーを取り扱う各種産保は、常に可想物い工 堅用資材などによる火災の危険性にさらされています。この ような産業火災を防止し、安全な作業環境を維持するため、 可燃物の火災危険性評価、難燃性試験の標準化、燃焼生成が スの副生機機などに図して検討を進めています。

また、大探度地下空間や鉱山などの閉鎖系空間における火 災性状の解明や退避システムの開発についても取り起んでい ます。さらに、実規構の実験施設を使用し、水増工や防火構 道による火災拡大防止技術や各権消火技術に関して研究を進 めています。

Industrial Fire Safety

Industries handling of many kinds of resources and energy always britigs about the risk of fire related to mineral products and industrial materials. Studies to prevent industrial fires and maintain a safe working environment are being conducted Main research subjects are the flammability of combustible materials, the standardization of flame resistance tests and characteristics of combustion products.

Studies on fire spread characteristics and the escade system at an emergency in great depth underground spaces and coal mines are also in progress. Further, full scale experiments on fire spread prevention techniques such as water spray or fireproof method and fire lighting techniques have been carried out by using large-scale tist galieries.

消火技術

地下空間、鉱山及び大規模建築物などの閉鎖系空間では、 火災が発生すると僅や有害ガスが充満し、初期消火を困難な ものにします。また、空間内の熱的フィードパックにより火 勢が激しくなったり、フラッシュオーバー現象によって被害 を大きくするなどの特徴があります。

このような火災を連腸的に制御し、短時間での増火を図る ために、高整張抱体や不活性ガスを火災区域に往入する技術 を開発しています。また、液体窒棄の急速気化法や液体窒素 と泡沫の檀合消火法などの新しい消火技術についても検討を 進めています。

Fire Prevention and Extinction

Direct fire-fighting to confined spaces such as at a great depth in underground spaces, underground mines and largescale facilities must be much more difficult compared with on ground surface because of the concentration of combustion products, the effect of thermal feedback and explosions caused by the flash over phonomena.

In order to suppress fires underground directly and then to establish the extinction at thort notice, researches have been conducting development of extinguishing techniques using high expansion foam and inert gas. Experimental studies are in progress on new extinguishing techniques such as a rapid evaporation system of liquid nitrogen and a contoined system of inert gas generator and foam generator.



地下環境は地上に比べ閉創性の点で将異性があります。そ い閉ざされた環境から、緊急時の最適進避システムを確立す るため、教命器、緊急通信、教急設備に関する研究を行うほ か、災害時の人間の行動様式の解明に取り組んでいます。

石炭鉱山を主な対象とし、事故教命物の性能向上、性能試 験方法及び緊急時の抗内情報の収集,指令伝達手法について、 また、抗内避難所の安全性の向上についての研究を進めてい ます。さらに熱・光・音の地下環象要素の下での人間の行動 を探り、退避・誘導技術への人間工学的な取り組みを行って います。

Emergency Refuge System

Studies to devise an optimum refuge system in an emergency in underground space or mines are being carried out. The main research subjects are improvement and test methods for self rescues, information processing lechniques and suitable wireless instruction methods for underground communication systems and safety enhancement and structure intensification methods for refuge stations underground. Behavior of the people in an emergency situation underground is also studied to develor optimum refuge system.



試験炭鉱での火災実験 Fire Experiment in the 400m Full Scale test Suriery



人工時による教命器の性能向上 Improvement of Emergency Rescuers by an Antina Ling







爆発対策と火薬類の利用転構

Explosion Prevention and Utilization of Explosive Energy

爆発事故につながる物質は、化学工場で使用される木業・ エチレン・酸化石油ガス、家庭で使用される天然ガス・LPガ スなど数多くあります。また、葉・食品など粉体を取り扱う 工場における粉体爆発や、炭鉱における炭壅爆発による事故 等があります。これら事故を減少させるため、爆発現象の機 構解明をはじめ、着火類対策、爆発伝播防止技術などの研究 を行っています。さらに、新火薬類も市場に出回っており、 より一層の安全性対策も望まれてるいると共に、都市発破・ 制御発破等、より高度に火薬類の利用技術に対する研究にも 着手しています。

There are many kinds of materials like hydrogen, ethylene, liquefied petroleum gas used in chemical plants and natural gas, propane used in homes, which give rise to accidents. Also another explosion hazard in process industries which are han dling foods, chemicals, plastics, metals is dust explosions. This type of an explosion is very common in coal mines.

In order to decrease these types of accidents, clarification of the mechanism of such explosions, ignition sources and controlling propagation of explosion is under study.

Recent explosives are fairly well safe, and number of accidents due to explosives is on the decrease. However, new safety inspection technology for new explosives developed in future, new construction blasting in the city and controlled blasting for new frontier of reo space is also being studied.

爆発対策

可燃性液体・ガス・粉体といった物質は多くい支筆活動の 場で扱かわれており、それらの爆発災害の危険性はあらゆる ところに潜んでいます。

この爆発の抑制対策として、爆発現象(圧力,火炎等)の 詳細についての研究,水袋をはじめ爆発抑制剤の効果と設置 方法等についての研究を行ってきています。また、爆発災害 の未然防止の観点から、漏洩ガスの洗動拡散の解析,着火限 界・静電気着火の究明、機器の防爆技術等についても検討を 重ねています。

Explosion Prevention

Concerning inflammable gases or dust explosions, the way to lessen explosions and to prevent explosions are are being researched. For lessening the occurrance of explosions, gas explosion phenomena, such at hlast wave development and thame prot tightion, are under stully. The effect of the explosion restraint materials is also one of the institute's research subject. Main subjects about the explosion prevention are the diffusion methods for released gas, ignitionability of flammable gas, static electricity and techniques for the design of explosion proof equipment.



吹上式村じん爆発実験装置における爆発

制御発破技術

発破は他の掘削技術に比較すると安価で効率もよいと考え られます。特に地下空間撮削では空間周辺の破壊したくない 部分を破壊しない発敏方法の開発が望まれております。そこ で、AVL 法のように従来の火薬をうまく使用し岩盤に適した 制御発破工法を研究しています。都市におけるビル・道路な どの寿命は限界にきており、その解体のための制御発破技術 の研究が要請されています。都市における発破は、効率は良 いが騒音・振動が大きいため我が国ではあまり利用されてい ません。騒音・振動の制御技術を開発し、積極的に都市でも 発破を使えるよう検討を行っています。

Controlled Blasting

Blasting is far superior in that the cost of blasting is lower and more efficient than other forms of excavating tecchnology. New controlled blasting which can control the breakage area is desired for excavating the new geo-space. So, new controlled blasting like the AVL (Alternate Velocity Loading Blasting) method and construction blasting method which can control the vibration and the noise is being studied.



爆業の落穏感度試験

腹囊保安

地下環境制御·防災技術

地下空間の利用は狭い我が国にとって重要な舞動です。地 下には恒温性、断熱性、遮光性、気密性など多くの利点があ ります。鉱山で培われてきた保安技術のポテンシャルを発展 させ、次のような地下空間利用に隧速する研究を推進してい ます。

06/2

a new part of the state of the

地下の岩盤や構造物を電磁波、湿音波を利用して可視化す る研究と地下岩盤内の力学特性と安全性評価の研究は地,空 間の利用過程における空洞維持の技術指針に役立てられます。 材料の特殊条件下での腐費の研究はより信頼性のある機器の 開発に、センサー開発利用と通気の研究は閉鎖空間内の環境 の情俗化及び維持に、災害と人間のかかわりに関する研究は ヒューマンエラーの防止や安全システムの構整にそれぞれ役 立つことを目指して研究しています。

地下構造物安全性評価

現在,地下空間利用の例として石油や液化石油ガスの備書, ワイン:穀物の備書、排水処理施設などがありますが、さら にさまざまな生産活動、娯楽・スポーツ施設あるいは産業廃 棄物処分など、利用の可能性を拡大するには地下空間特容の 問題が発きれています。安全に関するそれら問題の解決を図 るため、現在、電磁波などの波動を利用した岩盤の裏前評価 技術、緩み領域の低減、地下構造物の劣化、異常予知などの 安全性評価の研究が進められています。さらに、地下空洞へ 及ぼす地下水の影響、地下水位の低下による地表広下、地下 水汚染の研究など地下環境保全技術について研究しています。

Safety Assessment for Underground Structures

As examples of the utilization of geo space in the rock, there are the oil storage tanks. LPG storage tanks in the rock already in order widely to use the geo space in the rock, the rock assessment by suing geo tomography, the effect of under ground water on the structure of rock, the relaxation renge around the cavity, the mechanism of the deterioration of under ground structures, prediction of the deterioration or sometheng unusual, refreshment of underground rooms is now under study.

Underground Environment Controle and System Safety

Land should be more effectively utilized in Japan due to lack of it. In underground openings, there are many advantages such as temperature stability, insulation of heat light, air and others. For highly advanced utilization of underground openings several subjects are being carried out which are as follows, the visualization technique of underground rock body by using waves, rock mechanics and the stability assessment technique for the purpose long-term utilization of underground openings, the corrosion properties of materials under a special environment for the purpose of developing reliable instruments, sensor development and ventilation research for better underground environment and the mechanism of human en ar occurrence to establish a underground safety system.

センサーと安全監視システム

産業施設の大型化や高度化にともないその保安監視や安全 維持のためにより高度化されたシステムが求められます。ま た施設の利用目的次第では、温度、程度、風速、各種ガス通 度など環境内の多くの項目について監視を継続する必要があ ります。さらに、監視警報システムは誤動作の無い信頼性の 驚けるものでなければなりません。これらの目的遠成のため に、高精度で長期に安定性のあるセンサーの開発、それらを 組み込んだ監視システムの構築及び運用ソフトウェアの開発 なごを含めた総合的な保安研究を推進しています。

Development of Sensors and a Monitoring System

A more highly advanced safety monitoring system is being requested to allow for enlarged and modernized industrial facilities. Also depending on the purpose of the facility, continuous monitoring may be required in such environmental criterea as temperature, humidity, wind speed and the concentration of gases. Furthermore, the monitoring and alarm system should be reliable from the print of view of error action. To achieve the purpose above, resparsh on the development of long term stable sensors and the system software or monitoring are being promoted.



AEによる異常診断技術

- 炭鉱集中自動監視システムの開発 - Section mont - Charter Branch Montening and B





当センターは、鉱山保安を主とする課題に取り組んでいま す。特別研究については、産業保安部及び九州石炭鉱山試験 センターと共同研究の形で実施しています。この中では、北 街道という乾燥気候に伴う酸電気の課題が取り上げられ、主 として酵電気火花によるガス着火防止対策に取り組んでいま す。

- 1. 抗内用移動機構の開発:切羽のような上,下盤等,二面 拘束平面での自走ロボットの開発で,面の傾斜及び相さを 問わないという利点を持つ移動機構の要素技術の開発
- 2、新型監視センターの開発:超音波利用センターでは、静 電気の発生頻度及び発生箇所推定システムの開発、光ファ イパーセンサーの開発では、光干波型検出機構を用いた本 質安全な各種センサーの開発
- 3、本質安全防爆の高度化:九州センターとタイアップして 容量回路の思覚圧限界値の開発

The Coal Mine Safety Research Center (Hokkaido) has focussed its research efforts on developing safety technology for coal mines. Major research activities are being carried out as a special project in cooperation with the Industrial Safety Dept and the Coal Mine Safety Research Center (Kyushu).

The project involvas a study on the electrostatic charge in the dry weather of Hokkaido to prevent gas explosions caused by electrostatic discharges. Typical research work at the Center include:

- 1 Locomotive Mechanism for Underground Development
- 2 Advanced Monitoring Sensor Development
- 3. Extension of the Intrinsic Safety Concept



干渉型光ファイバーセンサーの基礎実験 Experiment of Optical Fiber Sensor



売動メタンガスにおける静電気火花樹火

A PARTIN AND A PARTICULAR

当センターは、国及び産炭地の強い要請に対応して、大正 4年5月の重方に設立され、石炭鉱山の災害防止技術開発の 研究を推進し、約75年の歴史と多くの研究成果を蓄積して炭 鉱現場の保安向上に寄与してきました。特に国立研究機関で 唯一の試験炭鉱を有し、この抗道内で大型実験装置を使用する 実規模に近い大型研究を行い,今日まで幾多の成果をあげてき ました。近時、九州産炭地においても、自然条件の悪化や機 業条件が厳しくなってきたことから、遠隔通信、情報伝達技 術や緊急退避所の高度化、ガスの流動拡散の把握あるいは坑 内火災潤火技術等、当センターにおける研究成果を拡大発展 させて、炭鉱現場への適用を図るために努力しています。更 に、長い歴史と伝統のもとに豊富な炭鉱保安技術研究の書種 を活用し、中国との国際産業技術研究事業や産業保安技術、 公害防止技術等の分野にも研究開発の推進を行っています。 試験炭鉱の主な大型設備としては、ガス、炭じん爆発や坑内 火災の実験に使用出来る400mと230m 試験坑道や実際の炭鉱 と同様な構造の黒川坑のほか火薬類の試験坑道などがありま す。黒川坑内では火薬類の試験や坑内通信の試験、坑内退避 所の幕度化の試験などが実施されています。

The Coal Mine Safety Presearch Center (Kyushu) was established by the Japanese Government and Civilian Organization in May 1915. For over 70 years, the Center has provided technology and technical services to the coal mining industry for coal mine safety. The experimental coal mine of the Center, which is a unique facility in Japan has been used for full-scale tests and useful results have been obtained.

Gver the past decades, the working environment of coal mines in the Kyushu coal-mining area has been getting worse. The Centers research focuses on developing advanced technology for coal mines, such as underground communication systems, data transfer techniques in mines, monitoring the firedamp movement and fire-extinguishing method for coal mines. The Center also encourages technology transfer to apply these technologies. In the mining industry



日配列前の増支活動 Explosion Flame of Explosive in Motar



爆薬検定坑通 Test Garery of Explosives

国際研究協力

International Cooperation of Science and Technology

科学技術立国の日本において国立の研究所として国際研究 協力の重要性はとみに高まって来ています。先進国をはじめ 発展途上国からも、フロンの対策問題に代表されるような地 球規模環境問題や、資源エネルギー関連研究、保安関連研究 等の協力要預が年々急増しています。

当所ではこれらの要請にこたえるペイ多様な国際研究協力 の充実を図っています。研究交流事業の内容は次の通りです。 1、先進国との研究交流協力

多国間協力,二国間協力,サミット協力,国際特定共同 研究事業など,例えばOECDとの環境問題研究協力,天然 資源の開発利用に関する日米会議、日米環境保護協力協定 にかかる研究協力,国際エネルギー機関における多国間研 究協力などを行っています。

2. 発展途上国との研究協力、技術協力

国際産業技術研究事業(ITIT事業)として、「中国レア メタルの分離精製技術に関する研究」、「熱帯地域における 大気汚染アセスメント手法に関する研究」を行っています。 3、国連、国際協力事業団(JICA)等からの関連技術協力

国際連合からの技術協力要請(特に研修), JICA からの 要請として「産業公審防止コース」、「鉱山保安コース」の 2つの集団研修をはじめ、数多くの傷別研修、専門家派遣 依頼等にこたえています。

4. その他の国際交流

長期、中期の在外研究、海外の大学、研究機関からの招へい、国際研究集会(国際会職)への出席、海外からの見 学者の対応などを行っています。今後もあらゆる科学技術の面で増加する国際化の波に応えて行きます。



ITIT 魏臼式(インド) The Signing Ceremony of ITIT Program with India



HCA T DRIATAR

Science and Technology have greatly contributed to our health and welfare, but they also have brought about some problems of a global scale regarding natural resources, energy and the environment. Future science and technology should not be indifferent to such problems, and their research and development should be carried out so as to harmonize industrial activities with the global environment. International cooperation is indispensable to cope with such problems.

On the other hand, as recent science and technology have become and complicated, it is difficult for a country to make their further progress only by itself. International cooperation is necessary from this point of view.

The International Research Cooperation Office was set up to promote research cooperation with foreign countries through the offer of the latest information about countries overseas and their research activities.

The following programs are in progress at the National Research Institute for Pollution and Resources.

1. Joint research with the organizations of advanced countries.

- Institute for the Transfer of Industrial Technology (ITIT) projects.
- Cooperation with JICA (Japan International Cooperation Agency) in its projects by dispatching specialists and accepting foreign trainees.
- The interchange of research personnel and guidance for foreign visitors.



MCA 集団研様 JICA Group Training Course



亡州有色会業研究院メンバーとの計議





技術情報サービス Technical Information Services

当所では下記の技術情報サービスの他に、科学技術週間、 又は環境週間に研究施設の一般公開を行っています。

The following Technical Information Services are carried out at this institute.

研究講演会 Technical Symposium

当所の研究成果の広報,着及を目的とし,研究分野を絞っ てテーマを設定し,年2回開催しています。

(担当 業務課)

共間研究 Joint Research

新技術開発のための研究は、基礎研究から企業化への一質 性のある均衡のとれた展開が必要です。その一環として、 民間企業等と技術・知識を交換し、研究及び費用を分担し て共同で研究を実施しています。

(担当 業務課)

官民連帯共同研究 Government and Private Joint Research 我が国が今後技術立国として飛躍するためには、先端的・ 基盤的分野における基礎研究を充実させる必要があります。 そのため、産業基盤確立に必要な基盤的研究で、民間企業 において共同研究についてのニーズの高いもののうち、公 共性の高いと認じられる研究チーマを工業技術院で選定、 予算化し、各研究所で実施しています。

(担当 業務課)

受託研究 Sponsored Research

当所の研究成果の普及、実用化の観点から、現行の研究 内容で直接企業に有効な資料を提供し得る場合に、民間企 業からの委託を受けて実施しています。

(担当 業務課)

技術相談 Technical Advice

当所の研究業務に関連のある募項について、電話、面談 あるいは文書による相談に応じています。 (担当 技術相談所)

技術指導 Technology Transfer

当所の研究業務に関連のある分析・概定・試験・製造法 などの各種技術についての実験指導を依頼に応じて行って います。

(担当 技術相談所)

見 学 Laboratory Visit 当所の設備、機器などを見学される場合はあらかじめ間

い合わせの上、文書でお申し込み下さい。 (担当 技術相談所)

検定・その他 Certification and Others

鉱山坑内用品検定規則に基づく鉱山用機械器具、火薬等の検定及び計量法に基づく熱量計の検定を行っています。 そのほか一般から依頼された試験、分析、鑑定等を行っています。これには試験、分析及び設備の使用規則による手 ただい下できたす。ます。



工業所有權 Patent Licensing

当所の研究を通じてなされた発明及び考案に関する権利は 国に帰属します。これらの通常実施権は閏日本産業技術振興 協会を通じて許諾されることになります。

Patents and Utility Models for Inventions on studies at the Institute belong to the Government.

Licenses of these patents are granted through the Japan Industrial Technology Association.





刊行物紹介 Publications

当所の業務内容や,研究成果,調査報告は,下記の当所に よる出版物のほか,一般学術誌にも掲載されています。

公審資源研究所報告(不定期) Report of NRIPR (Irregular) 研究成果のうち、集大成された研究論文について発行し ています。

公審査源研究所需報(季刊)
 Bulletin of NRiPR (Quarterly)
 研究成学型び調査結果の報文,並びに一般学術誌に掲載
 された論文い要旨を収録掲載しています。

公審實源研ニュース(月刊) News of NRIPR (Monthly) 業務の目的、内容、成果等の広報及び関連技術ニュース

等の紹介を行っています。

資源(月刊)(旧「採鉱と保安」を1989年1月に改題) SHIGEN (Monthly)

雪額・エネルギー分野の研究報文、調査結果等を掲載しています。

公審資源研究所年報(年刊) Annual Report of NRIPR (Annual)

組織、予算、研究内容等の単年度における業務報告を掲 載しています。

截山坑内用品検定公報(隔月刊)

White Mary's

Bulletin of Inspection of Underground Articles Used in Mines (Bi-monthly)

鉱山坑内用品の機定結果の報告を掲載しています。

◎当所刊行物の入手等については、業務課広報係までお問い合せ下さい。



最近の科学技術の発展は著しく、情報量も整大で、昭和62 年に世界中で発表された論文数は約80万編にもなり、研究上 必要な論文、資料の効率的な収集・整理は重要な課題です。 情報管理室は、これらの文献・特許を収集・整理し、データ ペース化するとともに、必要に応じデータを提供することを 目的としています。

具体的には、(1)オンライン情報検索の強化、(2)購入・寄贈 図書・雑誌の案内、(3)工業技術院内各研究所図書・資料室蔵 書目録のデータベース化などを進めています。

At Technical Information Office, literature and patents are collected, databased, and provided when requested. The following services are provided.

Reinforcement of on-line literature search
 Listing of new books and journals

3 Databasing of book catalogues in AIST

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本所では石炭・石油関係の古い費重な図響・雑誌を多数保 有し、また産業保安、廃水処理及び大気汚染関係の資料、特 に生化学と大気・海洋の拡散関係の充実を図っています。 赤外額、NMR、X線などのデータカードを所有しています。 九州石炭鉱山技術試験センターでは次気の変遷史関係の費 風な図響のほか、廃水処理、騒音 写劇関係を所有していま す。

北海道石炭鉱山技術試験センターでは炭鉱保安のうちでも ガス関係のものを多く所有しています。

The Library of the Central Laboratory contains about 30, 200 volumes. The main collections are books related to coal, petroleum, industrial safety, wastewater treatment and air pollution.

The Library of Coal Mine Safety Research Center, Kyushu has about 4,000 volumes. The main collections are books related to wastewater treatment, prevention of noise and vibration in addition to those on coal mine history.

The Library of Coal Mine Safety Research Center, Hokkaido has about 3,000 volumes, most related to mine safety and the safe use of gas in particular













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〒305 茨城県つくば市小野川16番の3 Tel(0298)54-3000(代表) FAX(0298)54-3038, 3049 Telex 3652570 AIST J

企画室	(0298)54-3026,3027
技術相談所	(0298)54-3036
国縣研究協力室	(0298) 54-3060
照務課	(0298)54-3022
果钙酶	(0298)54-3024.3025
情報管理室	(0298)54-3163,3050

北海道石炭鉱山技術試験センター

〒003 北海道札幌市白石区平和通3丁目北1番25号 Tel(011)861-2191(代) FAX(011)864-3469

九州石炭鉱山技術試験センター

〒822 福間県直方市頓野1541 Tel(09492)6-5511(代) FAX(09492)6-5518

同確并分室(試験炭鉱)

〒820-05 福陽県嘉穂郡雄井町西郷1142 Tel(0948)62-2057(代) FAX(0948)62-5280 National Research Institute for Pollution and Resources 15.3 Onogawa, Tsukubaishi, Ibaraki, 305 Japan

Research Planning Office Technology Advice Office International Cooperation Office General Affairs Division Research Service Division Technical Information Office

- Coal Mine Safety Research Center, Hokkaido Kita 1.25, Heiwadori 3, Shiroishi, Sapporo, 003 Japan
- Coal Mine Safety Research Center, Kyushu 1541 Tonno, Nogata, Fukuoka, 822 Japan

Experimental Coal Mine, Urul 1142 Saigo, Usul, Kaho, Fukuoka, 820:05 Japan.



TECHNOLOGIES DISCUSSED AT JGC CORPORATION

Radioactive Waste Management Technologies

- 1. Treatment of High Conductivity Liquid Waste
- 2. Treatment of Liquid Waste Containing Ammonia
- 3. Microwave Dryer for Spent Resins
- 4. New Vacuum Conveying System for Radioactive Liquid Waste

- Incinerator Technologies

- 1. SIAS
- 2. Gasification Furnace Incinerator
- 3. Hazardous Waste Incinerator
- 4. Energy- Recovery of Liquid Injection
- 5. Radioisotope Carcass Incinerator
- 6. Radwaste Incinerator
- 7. Medical Waste Incinerator
- 8. High Temperature Waste Incinerator
- 9. CyForMelt
- 10. Induction Heating Melting System
- Tritium Separation/Concentration
- Wet Oxidization
- Reprocessing Facility Real-Time Gas Monitoring System
- Mixed Waste and Reprocessing Management Technologies
 - 1. Advanced Cement Solidification
 - 2. Induction Heating Melting System
 - 3. Incineration of Spent TBP Contaminated U and Pu
 - 4. Liquid Waste Treatment

Hazardous Waste Management Technologies

- 1. Recovery of Solvent from Off-Gas by Activated Carbon Fiber Filter
- 2. In-situ Stabilization of Cd and Pb Contaminated Soil
- 3. Hazardous Waste Incinerator
- 4. PCB and VCM Waste Incinerator
- 5. Energy Recovering Type Liquid Injection Incinerator
- 6. Gasification Furnace Type Incinerator
- 7. Treatment of Heavy Metal and Organisms Contained in Liquid Waste
- 8. Regeneration of Spent Activated Carbon

BIBLIOGRAPHY OF LITERATURE RECEIVED FROM JGC CORPORATION "Advanced Waste Management Technologies", JGC Corporation, 100 pages.



ADVANCED WASTE MANAGEMENT

NOVEMBER, 1990











URANIUM - SELECTIVE CHELATE RESIN

- Reasonable Cost Good Performance
- Six years of Operating Experience at Ningyo Touge Uranium Processing Facility
- Proposed for Use at Weldon Spring (Competing technology not yet demonstrated)
- Proposed for Limited Use at Portsmouth Gaseous Diffusion Plant
- Under Consideration at Private Uranium Contaminated Sites
- Could Substantially Reduce the Cost of Removing Uranium from Waste Liquids

JGC'S ACHIEVEMENTS IN THE NUCLEAR FIELD IN U.S.A.

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ON - SITE STABILIZATION PROCESS (OSSP)

- Utilizes a Promising Stabilization Compound
- Proposed in Response to INEL's PRDA

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- TCLP Test Results Indicate Good Results
- Test on Uranium-Containing Soils Required
- Structure of Stabilization Agent can be varied to improve
 Stabilization Characteristics of Particular Contaminates

JGC ADVANCED CEMENT SOLIDIFICATION

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Waste to be Conventional treated cement method		JGC technology	Remarks	
Incinerator ash	No pretreatment	Pretreated by the Ca(OH) ₂ and NaOH	A retarded cement hydrating reaction problem are solved.	
Spent resin	No pretreatment	Pretreated by the cement and water	Swelling phenomenon of the immersion test are protected.	
Boric acid No pretreatment waste		Hydrate calcium metaborate are generated by pretreatment	Volume reduction and stable products are provided	







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Leach Test Results - Treated by fixing agent and bentonite -

Fixing agent(I) and bentonite			Fixing agent (II) and bentonite				
Method Sam (mg	EP - Toxicity			TCLP			
	Sample (mg/l)	Criteria (mg / l)		Sample (mg/l)	Criteria (mg / l)		
Cd	0.63	1.0	0	0.04	0.066	0	
Cr	< 0.01	5.0	0	0.11	0.084~5.2	0	
Hg	0.20	0.2	0	1)	0.025	1)	
Pb	<0.01	5.0	0	0.33	0.18~0.51	0	

1) Analysis is now in progress

Method EP - Toxicity			TCL2			
Sample (mg/l)	Criteria (mg / l)		Sample (mg / l)	Criteria (mg / l)		
Cd	4.86	1.0	x	0.10	0.066	Х
Cr	0.68	5.0	0	0.11	0.084~5.2	0
Hg	0.20	0.2	0	1)	0.025	
Pb	0.71	5.0	0	0.36	0.18~0.51	0

Leach Test Results - Treated by fixing agent (I) and Cement -

1) Analysis is now in progress



.

JGC INCINERATION TECHNOLOGIES

800 - 900 °C

- SIAS (Formerly Labopherix)
 - Submerged flue Incineration and Stabilization
 - Treats organic and inorganic liquid wastes simultaneously
 - Laboratory use (6 operating units)
- Gasification Furnace Incinerator
 - Pyrolizer
 - Municipul industrial wastes (Used tires, refusefuel pellets)
 - Industrial facility (Many operating units)
- Hazardous Waste Incinerator
 - Rotary kiln type
 - Industrial wastes or sludges
 - Waste management facility (8 operating units)
- Energy Recovering Liquid Injection Incinerator
 - Liquid injection type
 - High concentration COD liquid waste
 - Industrial facility (1 operating unit)

AUTOMATED WASTE CONTAINER MANAGEMENT SYSTEM

- Current system can include up to 11 types of fully automated inspection operations non - destructively
- Complete computerized record of Inspection
- Modularity allows flexible design options
- Compressive strength & voidage inspection units currently on location at customer sites
- TRU radioassay technology available, but not yet incorporated. Subject of JGC proposal in response to INEL's PRDA
- Free liquid detection technology similary not yet incorporated. Also part of response to INEL's PRDA
- Would likely reduce DOE cost & manpower requirements for dispositioning "Grouted" drums at Oak Ridge





JGC INCINERATION TECHNOLOGIES (Con't)

Greater than 1400 °C

- High Temperature Incinerator
 - Thin film melting furnace
 - Combustible & Incombustible Wastes
 - Nuclear Power Facilities (2 operating units)
- CyFurMelt
 - Cyclone furnace sludge incinerator / melter
 - Municipal Sewage Sludge
 - Municipal Sewage Facility (2 operating units)
- Induction Heating Melting System
 - Uses Electric Induction to melt ash
 - Treats (vitrifies) ash
 - Nuclear Facility Use (Under development)

JGC INCINERATION TECHNOLOGIES (CONT.)

- Radioisotope (RI) Carcass Incinerator
 - Cyclone Type
 - Carcasses, Spent Solvents
 - Medical Industry (70 operating units, Most are not for RI Waste)
- Radwaste Incinerator
 - Hearth Furnace, Wanson, SGN Type
 - Combustible Radwaste (α, β, γ contamination)
 - Nuclear Facility (3 operating units)
- Medical Waste Incinerator
 - Hearth Furnace
 - Medical, Infections Waste
 - Hospital (1 operating unit)





OTHER TECHNOLOGIES

- Tritium Separation / Concentration
 - Counter flow catalytic column
 - Laser stripping

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- Wet Oxidation
 - Well Tested
 - May soon be available at SEG's facility in Oak Ridge
 - Treats resiris, filteraids, and chelate agents
 - Low temperature & pressure system
- Reprocessing Facility Real Time Gas Monitoring System
 Will detect ¹⁴C, I, & ³H in stack gases

HIGH TEMPERATURE INCINERATOR

- Efficient
 - High Temperature Combustion Assures Complete Destruction
 - Accepts a Wide Range of Combustible and Incombustible Radioactive Waste
 - Accepts Wastes Containing Up to 20% Water
- Safe
 - Low Flue Gas Content/Simple Filtration System
 - Very Stable Vitrified Product
- Cost Effective
 - Simple Operation
 - Stable, Easily Solidifiable Product
 - Compact Facility
- Experience
 - Demonstration Plant
 - Operating Plant
 - Second Operating Plant now under Construction







PART 1

INTRODUCTION TO JGC COPRORATION





Reaction Mechanism of Laser Tritium Separation







JGC'S BACKGROUND

O Established :

October 25, 1928

O Employees :

2,800 (Including) 18 in the U.S.A 120 in other foreign countries

O Contracts completed :\$ 1, 80 Million (fiscal year 1989)

O Contracts awarded : \$3,800 Million (fiscal year 1989)

Memo ;

JGC



JGC'S ACTIVITIES AND FIELDS (1) Fields of Activity



- Radioactive waste management
- ⁾ Spent fuel reprocessing plant
-) Electric power generation plants
-) Environmental conservation facilities
-) Robotics
-) Petroleum refineries
-) Natural gas processing plants
-) Natural gas liquefaction plants
-) Town gas manufacturing plants
-) Petrochemical plants
-) Gas chemical plants

- O Coal chemical plants Olnorganic chemical plants
- O morganic chemical plar
- OPipelines
- **OAirport** facilities
- OPort and offshore facilities
- **ODesalination** plants
- OMunicipal sewage treatment facilities
- O Synthetic rubber resin fiber plants
- Ofood processing plants
- Opharmaceutical plants
- O Medical facilities
- **OOil terminals**

O Development and application of new technologies

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JGC

JGC'S ACTIVITIES AND FIELDS (2)

Services Offered

- O Feasibility studies
- O Project management
- O Planning
- O Basic design
- O Detailed design
- O Procurement
- O QA/QC
- O Transportation

Memo ;

- O Construction
 - O Test and start up
 - O Training
 - Maintenance and post installation services
 - O Financing arrangements
 - O Process license arrangements
 - O Research and development





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JGC'S ACTIVITIES IN THE NUCLEAR AND ADVANCED TECHNOLOGY FIELDS

O Nuclear Project Division established in 1965

O Project achievements - More than 200 projects completed

- O Japan's first spent nuclear fuel reprocessing plant at Tokai
- O R/W treatment facilities

O R & D achievements - More than 120 individual programs

O Nuclear Research Center opened in Oarai in 1984

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JGC'S LEADING ROLE IN THE WASTE MANAGEMENT AREA



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JGC'S PROJECT MANAGEMENT PHILOSOPHY









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JGC'S ACHIEVEMENTS IN THE NUCLEAR FIELD IN U.S.A.





MAJOR ACHIEVEMENTS IN THE NUCLEAR FIELD (1)

JAPAN'S FIRST SPENT NUCLEAR FUEL REPROCESSING PLANT AT TOKAI

- O Main Reprocessing Plant
- O R / W Treatment Facilities
- O Hot Functional Test Operation of Reprocessing Plant
- O Plutonium Storage Unit
- O High Active Liquid Waste Storage Facility
- O Replacement of Acid Recovery Evaporator
- O New Reprocessing Dissolver
- O Storage Facility for Bituminized Products



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MAJOR ACHIEVEMENTS IN THE NUCLEAR FIELD (2)

INDEPENDENT R / W TREATMENT FACILITIES

- O Fukushima Daiichi No.6 Total R / W Treatment Facilities for Tokyo Electric Power Co. (TEPCO)
- O Tokai No.2 Total R / W Treatment Facilities for the Japan Atomic Power Co. (JAPC)
- O Fukushima Daini No.3 & 4 Independent R / W Treatment Facility for TEPCO
- O Tokai No.2 Expansion of Total R / W Treatment Facility for JAPC
- O Large sized Incombustible Solid Waste Treatment Facility for Government Sector
- O North Anna / Surry New R / W Facilities for Virginia Power and Electric Company (VEPCO)

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JGC MAJOR ACHIEVEMENTS IN THE NUCLEAR FIELD (3)

		COMPLETION	CONSTRUCTION
ACKA	GED SYSTEMS		
(1)	Bitumen solidification	13	
(2)	Plastic solidification	2	1
(3)	Cement solidification	2	
(4)	Advanced cement solidification		2
(5)	Incineration	4	2
(6)	Filtration	16	
(7)	Compaction	7	
(8)	Laundry (dry and wet systems)	11	3
(9)	Wet oxidation of resins	1	
(10)) Robotics	2	1

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JGC'S ROLE IN MAJOR NUCLEAR PROJECTS

		FUKUSHIMA DAINI UNITS 3 & 4 R / W	TOKAI - 2 R / W EXPANSION	TOKAI REPROCESSING PLANT	NORTH ANNA SURRY R/W
1.	BASIC DESIGN	JGC	JGC	SGN	JGC
2.	DETAILED DESIGN	JGC	JGC	JGC/SGN	JGC
3.	BUILDING DESIGN	JGC		JGC	JGC
4.	BUILDING CONSTRUCTION	JGC (SUPERVISION)	JGC (SUPERVISION) JGC	JGC
5.	PROCUREMENT	JGC	JGC	JGC/SGN	JGC
6.	INSTALLATION	JGC	JGC	JGC	JGC
7.	TEST AND STARTUP	JGC	JGC	JGC/SGN	JGC

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PART 2

OVERVIEW OF JGC WASTE MANAGEMENT TECHNOLOGIES



INTRODUCTION

 JGC Can Provide Total Engineering/Construction & Test Operation Services in the Environmental Conservation Field & Various Other Fields Based on its 60 Years of Experience



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JGC

RELEVANT JGC TECHNOLOGIES

- Radioactive Waste Management
- Mixed Wastes & Reprocessing Management
- Hazardous Waste Management
- Pollution Control

List of Radioactive Waste Management Technologies



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List of Mixed Waste & Reprocessing Management Technologies



List of Hazardous Waste Management Technologies



VCM : Vinyl Chloride Monomer

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List of Pollution Control Technologies



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R.W. : Radioactive Waste RP/UR : Reprocessing Uranium Refining M.W. : Mixed Waste H.W. : Hazardous Waste P.C. : Pollution Control

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PART 4

MIXED / HAZARDOUS WASTE MANAGEMENT TECHNOLOGIES





TABLE OF CONTENTS

<u>No.</u>	Title				
1.	Automated Waste Container Management System				
2.	RASEP / Uranium - Selective Resin				
3.	On - Site Stabilization Process (OSSP)				
4.	Advanced Cement Solidification				
5.	Wet Oxidation				
6.	High Temperature Incineration				
7.	Other Technologies				





JGC

JGC's Automated Waste Container Management System (AWCMS) Proposed in response to INEL's PRDA

The JGC Automated Waste Container Management System (AWCMS) builds on JGC's drum inspection system already tested and currently being commercially installed in Japan to inspect, classify and label drums of solidified power reactor waste. The current system conducts a series of automated checks to determine compliance to packaging requirements, including the presence of surface contamination, and then scans the package for gamma radiation to perform a fully automated radioassay of the contents. The system then classifies the waste according to Japanese requirements and applies appropriate color-coded labels. Finally, the system records the results of all the inspections, and a video image of the drum on a micro floppy disk. The research and development work proposed in response to the PRDA at INEL would extend the technology to include checks for internal free liquids and perform a radioassay for alpha-emitting isotopes, as well as to determine which technologies would be economically appropriate to include in a system to process the tens of thousands of DOE waste drums awaiting inspection, classification, and disposition.

The demonstration facility at the JGC Nuclear Research Center in Oarai includes four stations, one each for:

- Recording a video image of all surfaces of the package
- Performing a surface contamination smear test
- Performing a gamma radioassay of the drum
- Classifying and labeling the inspected drum

The entire process is automated and computerized to reduce error, limit operator exposure, and to minimize manpower requirements.

In addition to the systems in the demonstration facility, JGC has developed and implemented additional systems to non-destructively check the compressive strength of the waste form, and to check for voids in the drum. These systems are being incorporated into systems being implemented in Japan, and prototypes demonstrating their operation have been built. These prototypes are currently on location at customer sites to verify their performance on actual waste drums scheduled for future shipping.

The technology to perform non-destructive radioassays of alpha-emitting waste containers is still emerging. While basic research to perform this kind of inspection has been completed, it has not yet been adapted to an automated system, nor has it been used for large inspection campaigns like those required by DOE. Non-destructive/non-invasive methods for detecting free liquids within the drums also have not yet been automated, and some basic research remains to determine which combination of technologies might most reliably detect significant free liquids.

However, the ability to reliably and efficiently inspect the undocumented, often poorly stabilized containers of TRU and suspected TRU in the DOE inventory could provide significant cost savings in the final disposition of these wastes. In addition, successful implementation of the AWCMS would provide a very high level of confidence in the actual condition and classification of these wastes.





PRACTICABLE ASSAY SYSTEM FOR RADIONUCLIDE QUANTIFICATION OF DISPOSAL

T. Yagi, T. Kato, N. Hashimoto, H. Kuribayashi, Y. Moriya JGC Corporation

ABSTRACT

The requirements for land disposal of LLW from nuclear power plants and facilities have necessituted a more complete and accurate analysis of the radionuclide contents of waste packages.

JGC has developed a new direct assay technique based on gamma-ray spectroscopy and total gamma-ray counting, combined with a scaling factor methodology for difficult-to-measure nuclides.

The system consists of an HpGe detector, a plastic scintillator, a microcomputer and a waste package handling system such as a turntable.

The radioactivity concentrations of Co-60 and Cs-137, which are key nuclides for difficult-tomeasure nuclides, are calculated from the activity ratio of Co-60 to Cs-137 measured by using an HpGe detector and total radioactivity is measured by using a plastic scintillator.

The concentrations of difficult-to-measure nuclides in a waste package are calculated by combining the radioactivity concentrations of Co-60 and Cs-137 with the waste package data and scaling factors.

The system is simple and enables a complete analysis of all nuclides specified prior to shipment and disposal of waste packages, and also ensures that not only homogeneous solidified waste but also nonhomogeneous DAW (dry active waste) can be measured within a short time.

INTRODU ... ION

In land disposal of radioactive solid waste generated at nuclear power plants, it is necessary to assay radionucludes contained in waste packages for the safe operation of the disposal site.

Importent nuclides from the standpoint of land disposal are Co-60 and Cs-237.

In addition, difficult-to-measure nuclides such as C-14, Ni-63, Sr-90, etc. arc also listed (see Table 1).

In shipping waste packages from nuclear power plants to the disposal site, confirmation of the contents of these nuclides is also required in compliance with disposal package technical package requirements.

Considering such-background needs, JGC has been developing the most suitable radionuclide assay technique for nuclear power plants since 1983 and has developed a new radionuclide assay system which combines the scaling factor method and the gamma scanning and gross gamma counting method.

This radionuclide assay system is essentially simple in nature. It is capable of counting the radioactivity of not only homogeneous solidified waste but also nonhomogeneous waste within a short time, and due consideration has been given to the suitability of the confirmation techniques applied in waste package shipment. The principle, composition, performance, etc. of the radionuclide assay system based on the gross gamma method are explained below.

SYSTEM CONCEPT

In order to assay the radioactivity content specific to nuclides in shipping waste packages from nuclear power stations, a radionuclide assay system suitable for this purpose must be developed with consideration given to the following matters.

- Capable of assaying a and B nuclides which are important in land disposal.
- Applicable to waste which is varied in type, size, and weight.
- Capable of covering the measurement of extensive radioactivity concentrations (most DAW is distributed in the extremely low level region).
- Simple system and short measuring time.

To solve these problems, JGC has developed a new radionuclide assay system based on the following approaches.

- Evaluation of the correlation between difficult-tomeasure nuclides and key nuclides and data verification.
- Simple nondestructive direct measurement of key nuclides.



Walls Processing, Transportation, Storage and Disposal Technical Programs and Public Education

Y 3 RADIONUCLIDE QUANTIFICATION OF DISPOSAL FACKAGES

TABLE 1

Activity Concentration Limits of LLW Burial in Japan

	(7 Ci/g)
C - 14	1 × 10 °
Co - 60	3 × 10 ²
Ni - 63	3 × 10 1
Sr - 90	2 × 10 °
Cs - 137	3 × 10 1
TRU	3 × 10 ·2
the second se	





Fig. 1. Concept of Assaying Radionuclides in Waste Package.

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As shown by Fig. 1, the former approach has been attained basically by establishing a scenario for nuclide behavior in nuclear reactor systems and determining the correlation between key nuclides and difficult-to-measure nuclides (C-14, Ni-63, Sr-90, TRU nuclides, etc.).

The latter approach is based on the method in which the radioactivity of key nuclides is assayed by selecting Co-60 as a key nuclide for CP nuclides (and C-14, TRU nuclides) and Cs-137 as a key nuclide for FP nuclides from among the nuclides important from the standpoint of land disposal, then measuring the ratio between Co-60 and Cs-137 and the total gamma radiation. (Co-60 is used as an empirical key nuclide for C-14 and Cs-137 for TRU nuclides.)

This system is intended to be used for the nondestructive inspection of waste packages before shipment. Therefore, in settling its target performance, due consideration must be given to the trend of applicable laws and regulations, and the actual conditions of the waste.

Trial calculation results have been obtained which indicate that if the radioactivity concentrations of Co-60 and Cs-127 can be measured to $10^{-4} \mu \text{Ci/g}$ or so, there is no effect on the disposal capacity which is important in safety evaluation, when the trend of applicable laws and regulations up to the present is considered.

From the conditions of setual waste at nuclear power plants, it is found that especially the radioactivity concentration of DAW is distributed in a wide range centering around the low concentration region. Therefore, measures to cover every type of waste package are required and the range of the relative concentration ratio between Cs-137 and Co-60 must be considered.

Considering these requirements, the target performance of our system is as follows.

Measuring range	: 10-5 - 10 ⁻¹ µCi/j
Measuring Cs-137/Co-60 ratio	: 1/100
Measuring time	: 5 minutes

BASIC PRINCIPLES

As shown by Fig. 2, in this nuclide assay system, the concontrations of radionuclides contained in waste packages are estimated by measuring the radioactivities of Cs-137 as a key nuclide for FP nuclides and Co-60 as a key nuclide for CP nuclides and considering scaling factors, half-lives, etc.

The gamma-ray counts from any point inside a drum of waste can be express by the following equation on condition that no consideration is given to the background gammaray.

$$\mathbb{E}_{N_{1}}\left(\xi\right) \star \iota\left(\xi\right) \mathbb{E}_{S^{-}\left(x,y,z,\xi\right)} \quad , \quad \frac{\Omega\left(\left(x,z,\xi\right)\right)}{4} \quad , \quad \mathfrak{sep}\left(-b\left(x,y,z,\xi\right)\right) \quad (1)$$

Where, Ni(E) : Detector count of the radiation from the radiation sources Si

E(E) : Detection officiency for each onergy

Si(x,y,z,E) : Intensity of the radiation source inside the drum

 $\Omega\left[(x,y,z)\right]$: Detector solid angle as seen from the radiation source Si

bi(x,y,z,E) : Attenuation distance from radiation source Si to the detector

Eq. (1) indicates that the gamma-ray counts at the detection point depends on the geometrical efficiency at any point inside the package and the attenuation distance between the radiation source and the detector.

In this system, the geometrical efficiency at any point inside the drum is made constant by measuring the gammaray flux radiated from the whole circumferential surface of the drum and correction for attenuation is made based on the mean density of the waste (weight/volume).

Thus, Eq. (1) is expressed as follows.

$$\overset{n}{\sum} Si(x, y, z, E) = \frac{\overset{n}{\sum} Ni(E)}{e(E) \cdot \frac{\overline{\Omega}i}{A} \cdot exp(\overline{b}i(E))} (2)$$

As the measured value Ni(E) is a function of energy, it is necessary to identify the nuclides to be measured.

Assuming that the nuclides to be measured are Co-60 and Cs-137:

$$\sum_{n=1}^{\infty} Ni ({}^{60} Co) = \frac{\sum_{n=1}^{n} Ni ({}^{60} Co)}{e ({}^{60} Co) \cdot \frac{\overline{O}i}{4 n} \cdot exp ({}^{-} \overline{D}i ({}^{60} Co))}$$
(1)

$$\mathbb{DSi}^{122} \mathbb{C}s = \frac{1}{\varepsilon \left(\frac{122}{\mathsf{C}s}\right) \cdot \frac{\overline{\Omega}i}{4\pi}} \cdot \exp\left(-\overline{\mathsf{D}i}\left(\frac{132}{\mathsf{C}s}\right)\right)$$

As the plastic scintillator is incapable of spectroscopic analysis, it is impossible to distinguish Co-60 and Cs-137 from each other. Therefore, the Cs-137/Co-60 ratio must be measured using another detector. This system uses a Ge semiconductor detector for this purpose.

In this system, in order to make the geometrical efficiency constant, the mean geometrical efficiency is raised



Yagi RADIONUCLIDE QUANTIFICATION OF DISPOSAL PACKAGES



. Fig. 2. Schematic Diagram of Nuclide Assay System

to 0.5 - 0.6 by rotating the drum and disposing the plastic scintillator so as to cover the overall height of the drum.

APPLICATION TO WASTE PACKAGES

As described above, the radiation incident on the scintillator from the sources contained in the waste package is expressed by Eq. (1) and varies in accordance with the location of the sources and the effects of the attenuation depending on the waste type.

In order to ensure accurate measurements, it is therefore necessary to make corrections after grasping the condition of the content of the package or to devise a measurement method which is not easily affected by the condition of the content.

In the case of homogeneous solidified waste, the density of the content and the distribution of radioactivity are uniform. Therefore, once the solidifying agent used and the target nuclides are determined, the valiables used in the equations can be made constant and accurate measurements can be ensured.

In the case of DAW packages, however, the kind of waste, packing condition, contamination pattern, etc. are varied and the measured values are affected by these conditions.

The effects of these conditions are observed as the attenuation of radioactivity in waste packages. For instance, the measured values become large when nuclides exist near the drum surface and become small when they exist in the center of the drum.

Such difference in the measured values becomes larger as the density of the content of the drum becomes larger.

In this system, the amount of radioactivity is counted as the gross gamma radiation dose whose energy region is not specified.

Radiation is attenuated due to the scattering of radiation by the contents of packages, but in this system, surscattered rays are also counted together.

It is known that the degree of scattering tends to increase as the density of the content becomes higher and as the transmission distance in materials becomes longer.

Therefore, even if materials having a large attenuation effect unevenly exist in the drum, the influence of such materials can be averaged by measuring the gross gamma radiation dose from various directions but not from a particular direction.

As described above, this system is aimed at correcting measurements for the effects of the uneven distribution of density and radionuclides in waste packages and also iniproving the measuring time and the detection limits by providing several detectors.

Yagi RADIONUCLIDE QUANTIFICATION OF DISPOSAL PACKAGES

BASIC SYSTEM SPECIFICATIONS

The basic system specifications are shown in Table II and each item is detailed below. A photograph of the prototype assay system is shown in Fig. 3.

1. Waste packages to be measured

The standard system is calibrated to measure the radioactivity of 200-liter drum waste packages. The system is also applicable to waste packages of other drum sizes by being calibrated. Concerning the kinds of waste to be measured, the system is also applicable to homegeneous solidified waste and DAW.

2. Nuclides to be measured

In principle, the nuclides to be measured are Co-60 and Cs-137, which are the key nuclides in the scaling factor method. However, an optional device permits the measurement of the key nuclide concentration even if waste packages specific to sites contain interfering nuclides such as Mn-54. The concentrations of difficult-to-measure nuclides such as Ni-63, Sr-90, etc. are calculated based on the data base, using the scaling factor method.

3. Measurable activity

The system is capable of measuring the radioactivity up to approximately 0.1 μ CV/drum as gamma-ray radioactivity (for Co-60 when the bulk density of waste packages is 0.5).

4. Measuring time

The standard measuring time is 3 minutes but selection can be left to the operator's selection.

Data processing requires about 2 minutes.

5. Detectors

A high-purity Ge semiconductor detector is used to measure the Cs-13?/Co-60 ratio, and a plastic scintillator is used to measure the gross gamma radiation dose.

6. Operation control





Yagi RADIONUCLIDE QUANTIFICATION OF DISPOSAL PACKAGES

The operation of the system is automatically controlled by a computer. All waste package information can be inputted by interfacing with the computer using the keyboard. (The system can be connected to a superior computer system.) As-measured data, i.e., MCA spectrum data and scaler counts are stored in a floppy disk to permit easy retrieval of past measured data and easy erasure of unnecessary data.

SYSTEM PERFORMANCE

The performance of this system was evaluated stepwise firstly by a pattern test, then a full-scale simulated waste package test, and finally an actual woste package test. Especially in applying this system to DAW packages, such an approach is important because it is impossible to specify the geometrical configuration and the distribution of density and radioactivity in such a package.

An example of the results of each test is described below.

1. Pattern test

In this test, measurements were conducted by preparing a simulated waste package whose inside was divided into several segments so that the density of waste in each segment could be changed, and placing a standard radiation source in the specified position inside the package.

An example of the measurement results is shown in Fig. 4.

2. Full-scale simulated waste package test

In this test, more realistic measurements were conducted, using full-scale simulated waste packages similar to actual ones in the aspects of configuration and material, which were obtained by extending a model package. A single scaled radiation source was used in this test.

3. Actual waste package test

In this test, a reference simulated waste package was used for calibration purposes, and then performance tests were carried out by using actual waste packages (cement packages and DAW packages).

The tests on the compatt packages were conducted to ascortain the calibration constant, and it was

confirmed that errors in the key nuclides measurements were within 5%.

The results of the measurements of the key nuclides were compared with the activity of difficult-to-measure nuclides, which was estimated by the scaling factor method and also with that of the core sample which was analyzed by a manual method.

The ross on DAW were implemented to check for errors due to changes in the filling pattern of radwaste, and it was established that the deviation factor was within $\pm 5\%$ at 1 σ when the bulk density was 0.5 g/cm³.

CONCLUSION

This system is very simple in mechanism and capable of assaying all radionuclides which are important from the standpoint of land disposal. In addition, it is widely applicable to not only homogeneous solidified waste packages but also nonhomogeneous DAW packages. Therefore, this system can be said to be most suited for nondestructive inspection required in shipping waste packages from nuclear power plants to disposal sites. In future studies, more extensive simulation tests and site verification tests will be continued further and the scaling factor estimation program will be improved.

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FULL-SCALE TESTING OF WASTE PACKAGE INSPECTION SYSTEM

T. Yagi, H. Kuribayashi, Y. Moriya, H. Fujisawa, N.Takebayashi JGC Corporation, Japan

ABSTRACT

In land disposal of low-level radioactive waste (LLW) in Japan, it is legally required that the waste packages to be disposed of be inspected for conformance to applicable technical regulations prior to shipment from each existing power station.

JGC has constructed a fully automatic waste package inspection system for the purpose of obtaining the required design data and proving the performance of the system.

This system consists of three inspection units (for visual inspection, surface contamination/dose rate measurement and radioactivity/weight measurement), a labelling unit, a centralized control unit and a drum handling unit.

The outstanding features of : e system are as follows: The equipment and components are modularized and designed to be of the most compact size and the quality control functions are performed by an advanced centralized control system.

As a result of the full-scale testing, it has been confirmed that this system satisfies all the performance requirements for the inspection of disposal packages.

The treatment capacity, operability and data control functions of this system were also checked and confirmed in the full-scale testing of the system.

The results of the performance tests show that JGC can establish the most rational and economical waste package inspection system applicable to any nuclear power station.

INTRODUCTION

At present, approximately 440,000 drums filled with LLW are stores at nuclear power stations in Japan: of them, some drums contain solidified waste having "barrierability" and some contain untreated miscellaneous solid wastes.

In accordance with the Nuclear Installation Regulation Law revised in 1986, and the Waste Burial Regulation and related notifications established in 1988, various items of work concerning the transportation to and the burial disposal of wastes at the Radioactive Waste Storage Center which will be constructed in Rokkasho Village, Aomori Frefecture, are being conducted by national and private organizations for the commencement of the operation of the Center in 1991.

Waste package shipping inspection facilities are being planned at each nuclear power station in Japan for shipping several thousand to dozen thousand waste packages per year.

Waste package shipping inspection facilities are being planned at each nuclear power station in Japan for shipping several thousand to dozen thousand waste packages per year.

This report will discuss the concept of LLW shipping inspection and introduce one example of the structuring of a nondestructive inspection system.

BASIC CONCEPT OF INSPECTION OF WASTE FACKAGES

If the plan for burial disposal is put into practice, the waste stored at each nuclear power station will be transported to the storage center where it will be buried in a concrete pit and controlled therein for several hundred years. (See Fig. 1.)

Thus, the entirely different circumstances of the waste packages, including other barriers such as the concrete pits and earth strata, will enable their long-term safe disposal

Accordingly the shipping inspection, the preparation of base data is required to ensure the control of the safety of packages as minimum units. Therefore, in the case of the inspection of waste packages at the power station, it is important to not only inspect the external appearance of packages but also the quality of their contents and the history after the generation of the waste. Furthermore, it is essential to evaluate and confirm the results accurately and rationally.

It is necessary to ensure the quality of waste packages by obtaining evaluation data for the technical criteria based on a combination of information from (See Table I.):

(1) Waste and process control program

Operation control and quality control data, such as the waste content code, operation records of the waste treatment facilities and the certifications of solidification materials and containers.

(2) Nondestructive inspection

Data obtained by conducting nondestructive inspections of waste packages (surface contamination density, radioactivity concentration, etc.).

(3) Sampling program

Sampling analysis data such as radioactivity concentration data for each type of nuclide and product property evaluation test data (compressive strength, leachability, etc.).

WASTE PACKAGE INSPECTION DEMONSTRATION PLANT

The commercial-scale demonstration plant constructed at JGC's Nuclear Research Center is an example of waste package inspection facilities based on the above basic concept of waste package inspection. (See Fig. 2.)

This demonstration plant is designed to conduct five types of inspection (visual inspection, surface contamination density measurement, dose rate measurement, radioactivity measurement and weight measurement).

It is a fully automatic module system whose inspection equipment, labelling unit, handling unit and centralized control system have been developed and designed to be rational and practical.

The major components of the system are as outlined below.





Yagi WASTE PACKAGE INSPECTION SYSTEM



Fig. 2. Waste Package Inspection Demonstration Plant.

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Yagi WASTE PACKAGE INSPECTION SYSTEM

Visual Inspection Unit

The surfaces of the waste package are all visually checked by using three ITV cameras; one for the top surface, a second for the side surface and a third for the bottom surface.

The image of the waste package is displayed on the monitor screen of the central control unit and recorded on a video floppy disk. (See Fig. 3.)

Surface Contamination/Dose Rate Measurement Unit

The surface contamination density and the surface dose rate of the waste package are automatically measured.

Smear samples (smear pads with which the surfaces of the waste package are wiped) are taken from four locations, the top surface, upper side surface, lower side surface and bottom surface, and each sample is checked to determine the surface contamination density.

At the same time, the surface dose rate is measured by the sensor built into the smear sampling portion.

This unit is designed to carry out all of these operations (sampling, measurement, calculation and smear , ad replacement) automatically.

"adioactivity/Weight Measurement Unit

This unit determines the radioactivity concentration of the waste package nondestructively for each type of the dide contained therein.

The unit consists of a spiral scanning Ge semi-conductor sensor, a plastic scintillator, calculation software and a



Fig. 4. Surface Contamination/Dose Rate Measurement Unit.



Fig. 3. Visual Inspection.



Fig. 5. Radioactivity/Weight Measurement Unit.





TABLE I Qualification of Waste Package for Disposal



Fig. 6. Labelling Unit.

turntable with a load cell. (See Fig. 5.)

The radioactivity concentration is determined for each type of nuclide from the total dose (plastic scintillator), γ scaling factors for difficult-to-measure nuclides and y nuclides, which is available from the data base.

Labelling Unit

This unit labels the waste package with a color band to classify surface dose rates. It also affixes stickers indicating identification numbers for shipping and disposal management after the completion of the abovementioned five types of inspection. (See Fig. 6.)

Color bands (white, orange and red) are automatically selected based on the surface dose rate measurement results and attached to the waste package. The identification numbers are automatically marked on the yellow base labels by heat transfer printing.

This labelling is conducted automatically by recognition of the radiation warning symbol marked on the waste package, using an optical sensor.

Centralized Control Unit

This unit automatically controls each inspection unit and the drum handling unit.

Data on each waste package are received into this system and inspection data from each inspection unit are processed. Daily reports, inspection reports and shipping management data are then prepared.

TEST RESULTS

Performance tests of each inspection unit were conducted and the capacity, workability and measurement accuracy of each unit were evaluated. As a result, it has been ascertained that each unit is capable of achieving the

Yagi WASTE PACKAGE INSPECTION SYSTEM

performance required for the waste package inspection. (See Table II.)

The capacity, operability and data processing functions the entire system were checked by conducting the overall operation of the system.

It has been confirmed that the system can inspect one waste package every 10 minutes. The system can be used at almost all power stations even if the inspection modules are not expanded.

It has also been confirmed that the data processing functions of the system are satisfactory for the confirmation of the data on the waste packages to be disposed of.

FUTURE DEVELOPMENT

Regarding "package damage" and the "compressive strength" of cement solidified waste shown as disposal inspection items in Table I, since there is a possibility that further precise qualification may be required in the future, JGC is proceeding with the development of the following automatic nondestructive inspection units:

Drum Integrity Inspection Unit

This unit is designed to heat the package uniformly by high frequency inductive heating and to check the package for corrosion and damage, sensing the infrared rays from the package by thermography and inspecting by an ultrasonic thickness gauge. (See Fig. 7.)

Compressive Strength Measurement Unit

This unit utilizes the correlation between the compressive strength of the solidified waste and the velocity of the ultrasonic wave propogation through the waste.

The pulsar and receiver probes are attached to the side surface of the waste package and ultrasonic waves are emitted. (See Fig. 8.)

The compressive strength of the waste in the package is determined from the ultrasonic wave propagation velocity.

CONCLUSION

IGC's waste package inspection system is an overall waste package quality control system for the land disposal of LLW.

By using this system, it is possible to provide the data required to ensure the safety of land disposal of waste packages by means of data management from the nuclear plant to the land disposal facility.

From this standpoint, we have shown one concept of waste package inspection and taken up JGC's demonstration plant as an example.



	Madule	System	Required time	Perform	iance.
1	Visual inspection unit	Remote ITV	3 min	3 mm defects detectable	
2.	Surface contamination density /	Automatic smear method /	8 min	Smear efficiency	90%
	Dose rate measurement unit	Automatic whole surface		Measurement limit	1 x 10 4 µCi/cm
		scanning		Measurement range	: 0.1 ~ 2000 mA /
3.	Radioactivity / Weight	Spiral scanning/	5 min	Measurement nuclides	r nuclides
	measurement unit	Load cell		Measurement limit	1 x 10 4 µCi/g
					(Co.60)
				Fluctuation factor	2 20%
4.	Labelling unit	Automatic lettering and	4 min	Labelling accuracy	± 15 mm
		labelling			

TABLE II Results of Performance Test Yagi WASTE PACKAGE INSPECTION SYSTEM



Ultrasonic senser probe

Fig. 7. Principle of Drum In grity Inspection.

In Japan, at present, various committees and working groups are proceeding with the studies and reviews of the inspection of waste packages for land disposal, and, accordingly, opinions are not uniform.

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Fig. 8. Compressive Strength Measurement Unit.

Under these circumstances and in view of future LLW disposal regulatory changes, JGC intends to establish a further improved, rational, overall waste package inspection system.





HORDESTRUCTIVE COMPRESSIVE STRENGTH INSPECTION SYSTEM FOR CEMENT-SOLIDIFIED WASTE FACKAGES

C. Ragiwara, Y. Jtoh, K. Sucai Tokyo Electric Power Company Tokyo, Japan

T. Ujihara. I. Oda. Y. Yoshida Tokyo Electric Power Environmental Engineering Co., Inc. Tokyo, Japan

T. Yagi, M. Takabayashi. M. Rado JGC Gorperation. Tokyo, Japan

INTRODUCTION

in land disposal of low-lavel radioactive waste (LLW) in Japan. It is legally required that weste packages to be disposed of be verified for conformance to applicable technical regulations prior to shipment from each existing power station to the Radioactive Wasts Storage Center. Concerning the ys, properties of cament-solidified waste packsons chn ... requistions require verification that the compra .ngth of the waste packages exceeds 15kg/cm Altho. . conformity to this requirement may be varified indiractly by reviewing such documents as solidification records, the ultrasonic pulse velocity method is considered to be affective when nondestructive inspection is required. Thus, a condestructive compressive stiength inspection system was developed considering the actual conditions of the cement-solidified waste packages

PRINCIPAL DEVELOPHENT ITEMS

In developing this system, the following points were given special consideration.

A. Transmission of ultrasonic waves into the cement-solidified product

In order to measure the pulse velocity in a comment package it is necessary to transmit ultrasonic waves into the solidified product through the stael drum. A slight contraction of the comment may occur during solidification leaving a small gap between the drum wall and the comment-solidified product. To solve this problem, the receiving and transmitting transducers were closely pressed scainst the drum wall and an appropriate ultrasonic wave frequency was selected so that ultrasonic waves propagated in the solidified product.

In securing the transducers on the drugs well, the contact pressure was set in the ringe of 10 \pm 30 kg/cm 2 and the transducer ends were tapered so as to lower the hydraulic pressure. (See Figure 1.)

In addition. In order to select in appropriate transmission frequency, texts were conducted using three transmitting transducers of 24, 54, and 100kHz A frequency of 54 kHz was selected as the transmission frequency.



Fig 1 Suclass wave allenuation mechanism

B. Surface waves and direct waves

As the puise velocity is measured from the outside of the package, two kinds of waves are received by the receiving trantfucer: waves propagating in the solidified product (direct waves) and waves propagating on the aurisce of the steel drum (surface waves).

The transit time of the direct waves is approximately 150 -160 µ sec and that of the surface waves is about 165 - 170 µ sec. Therefore, if the direct waves are slower than the surface waves, the surface waves or the composite waves consisting of the surface and direct waves will be measured at the wave receiving point. For this reason, it is, necessary to reduce the influence of the surface waves. To solve this problem, a close contact mechanise was provided around the transducers as shown in Figure 1 so that the surface waves could be strenusted in the contact sons by bringing the cement-solidified product into close contact with the steel drum wall. The contact pressure was sat in the range of 10 - 30 kg/cm² which was equivalent to the contact pressure of the transducers.

OUTLINE OF INSPECTION SYSTEM

A full-scale inspection system was constructed in consideration of the above. The main components of the system are as follows. (See Figure 2.)

A. Ultrasonic probe unic

This unit measures the velocity of ultrasonic waves propagating inside a censent-solidified product and consists



of transmitting transducers, receiving transducers, a transducer contact device and a surface wave attenuation mechanism.

8. Rydraulic unit

This unit supplies the hydraulic pressure necessary to press the ultrasonic probe unit spainst the drum wall a d detech it therefrom. This unit consists of an oil tank, a drive motor, a solehold walve, and a remote-controlled hydraulic pressure control velve.

C. Lifting and transporting

This unit sets the weste package in position and vertically moves the ultrasonic probe unit. It consists of a transporter, guide rails, and a drive unit.

D. Data processing unit

This unit performs wave pattern recording, pulsa velocity calculation, measurement accuracy judgment, and compressive strangth estimation on the basis of output from the ultrazonic probes.



ig ? Full-scale inspection system

TEST RESULTS

A. Confirmation of the effect of the surface wave attenuation mechanism

Figure 3 shows an example of wave petterns obtained when a surface wave attenuation mechanism was applied and an example of when not applied.

when the mechanism was not used, the transit time measured was approximately 170 µ sec as a results of the influence of the surface waves reaching the receiving transducer in 160 - 170 µ sec. When the mechanism was used, however, the attenuation of the surface waves eliminated their influence and made it possible to measure the transit time of the direct waves (approximately 180 µ sec). Through this test, it was confirmed that the surface wave attenuation mechanism was effective in attenuating the surface waves.





Fig. 3 Effect of surface wave attenuation mechanism

B. Confirmation of the correlation between pulse velocity and compressive strength

Pulse velocity was measured from the outside of the drums of simulated solidified waste packages whose solidification conditions are shown in Table 1, and compressive strangth data were obtained from small test specimen samples from each product of the same lot during drum charging. Figure 4 shows the relationship between the pulse velocity and the compressive strength measured in such a manner.

As a result, it was confirmed that there is a correlation between the pulse velocity and the compressive strength and that this inspection system enabled the pulse velocity in comment-solidified products to be succentically measured in a wide range of compressive strengths.

C. Hot test

Using this inspection system, measurement investigation was carried out on about 1% comment waste packages stored at TZPCO's Fuxushims Dalichi Ruclear Power Station. Of these waste packages, about 60 packages were subjected to core sampling. Destructive inspection tests ware conducted to measure their compressive strength.

Pulse velocity messured from the outside of the pachages and that of core samples

Figure 5 shows a comparison between the pulse velocity measured form the outside of the packages and that of core samples. As seen from this figure, a correlation of approximately 1 to 1 (standard deviet.on 80 m/sec) is obtained in a range of 3.200-3.800 (m/sec).















from this fact. It was confirmed that the pulse velocity in cament vests packages could be measured nondestructively from the outside of the drums using this compressive scrength inspection system.

Correlation between pulse relacity and complexitive attength

Figure & shows the correlation between the pulse velocity measured by this inspection system and the compressive strength obtained from core samples. The following equation was obtained as a regression line of the plats.

do = 204.5 X Vp = 244.4

- de + Compressive scrength (Ag/cm²)
- Vp + Vitrasonic pulse velocity (Km/sec)



Fig 6 Pulse velocity vs Compressive strength

Estimation of comptessive strength

The compressive strengths of 270 sctual waste packages were estimated using the above equation and its frequency distribution was shown in figure 7. The mean compressive strength and standard deviation were 445 kg/cm² and 58 kg/cm², respectively.



CONCLUSION

To establish a bighly reliable nondestructive compressive strength inspection system. this paper reported the design and the fabrication of a full-scale demonstration unit, results of the basic tests and the actual tests, and the practical applicability of this system.

Regulations and standards relating to nondestructive testing such as ASTM, 85, and CIN, show a problem of accuracy of nondestructive inspection techniques as a means of estimating the compressive strength of concrete. Nowaver, as described above, astisfactory accuracy was obtained in the case of cement-solidified waste packages generated at nuclear power plants where very stringent quality control was achieved.

In the future, a more reliable nondestructive compressive strength inspection system will be established through the study of the correlation between the pulse velocity and the compressive strength on the basis of the data obtained from the measurements using actual cement-solidified products.





JGC's Uranium-selective Chelate Resin

To be used in the RASEP processing equipment to remove uranium contamination from liquids

JGC has proposed the use of a proven uranium selective chelate resin for removing uranium contamination from liquid wastes containing other contaminating materials, thereby allowing the wastes to be classified as hazardous rather than mixed wastes. This resin, developed in Japan, is currently being used in the treatment of liquid wastes at the Ningyo Touge uranium processing facility in south central Japan. This successful experience was the basis for its proposed use (in collaboration with Dominion Energy) at Weldon Spring; and it is currently being considered for use at non-DOE uranium contaminated sites. The resin cost is reasonable, and excellent performance has been attained in practice. The resin is highly selective for tranium, and the ion is easily eluted from the resin if required, to obtain a concentrated solution of uranium. The resin has actually been used (for demonstration purposes) to process ordinary seawater to obtain a ten gram sample of yellowcake.

This resin, called UR-3100, has been tested with a solution containing zinc, copper, and iron. It was found that the resin will adsorb virtually no zinc or copper, and only very little iron. Well over 150 column volumes of 71.1 ppm uranium solution could be processed before initial breakthrough of uranium. After initial breakthrough, the effluent uranium concentration increases only very slowly, indicating that significant adsorbtion capability remains for the processing of well over 200 column volumes at this concentration.

It was found that using NailCO₃, the uranium could be eluted almost quantitatively, and that the concentration of zinc, copper, and iron in the eluate were lower than 0.1mg/liter. In other words, the eluate of this resin contained substantially no metal ions other than uranium and sodium.

JGC believes that this resin could substantially reduce the cost of remediation at We'don Spring, and at other sites both within the DOE complex and at privately owned sites. It is our understanding that demonstration tests of the conventional ion exchange system selected at Weldon Spring have not yet begun. However, the process system proposed by JCC through Dominion Energy Inc. may not require further development testing, as it has been in operation for over six years. JGC considered the use of conventional ion-exchange methods and activated alumina treatment process, and concluded that the UR-3100 sesin would provide the most cost effective, reliable treatment to meet the requirements of the RFP. The introduction to that proposal methods appropriateness of the chelate resin technology to this application, has been included in this compendium. We believe that there is a continuing need for this technology at several other DOE locations.



Breakthrough curves of metal ions
1. INTRODUCTION

This Proposal is offered in response to the Request For Proposal No. RFP-3589-VP 140 issued by Contractor, MK-Ferguson Company, Inc. (hereinafter referred to as MK-Ferguson).

Dominion Energy, Inc. (DEI) is a subsidiary of Dominion Resources, Inc., located in Richmond, Virginia. Dominion Resources, Inc. is the holding company for Virginia Pover, the major utility company serving the state of Virginia. It is the intention of DEI to commercialize various technologies and services as a result of its in depth technical and operating experience related to nuclear power plant and fossil power plant vaste creatment. Furthermore, a major strategy of DEI is to enter into partnerships with other companies which possess the technology which compliments the operating experience of DEI.

DEI has selected JGC Corporation (JGC), Tokyo, Japan as a lover-tier Subcontractor for the design of the water treatment facility based on JGC's significant experience in the design of facilities for treating radioactive wastes and hazardous materials.

With respect to the treatment of uranium that currently exists in the Weldon Spring quarry, it is DEI's technical position based on extensive testing and operating experience that conventional water treatment techniques will not, because of known inefficiencies, be successful in removing uranium from the quarry water in accordance with the requirements of the technical specifications of the RFP. Therefore, the use of conventional techniques will be highly inefficient and will not meet the environmental objectives of MK-Ferguson with respect to the discharge of the effluent to an uncontrolled environment. In this proposal, DEI recommends the use of a chelating resin that has been demonstrated to be effective in the removal of uranium impurities from waste water in Japan.

The following specific technical information is provided to support these statements:

1. Use of conventional ion exchange methods -- Based on testing and opprating experience in Japan, the use of ion exchange resin in this spplication will be highly inefficient. The ion exchange resin used in this system should be a non-regeneration system. The reasons are as follows. In the case of a regeneration system of ion exchange resin, the regeneration solution (Na_SO, solution) should be treated by an additional solidification process. For the SO, ion, it is appropriate to add Ca in order to cause CaSO, precipitation. Bowever, in that case, the quantity of Ca(OE), to be added will increase and Na which is not removed by precipitation will accumulate. Therefore, the regeneration interval will be significantly decreased.

For the volume of water and the concentration of uranium contained in the quarry, it is estimated, in the case of a non-regeneration system, that it vill require approximately 20,000 ft3 of ion exchange resin to treat the waste stream. Operating with such a large quantity of resin will be very inefficient and vill result in significant maintenance and operating expenditures in order to operate the treatment system.

The following discussion provides the technical basis for the above statements. In a conventional system, the ion exchange resin will adsorb all ions present in solution. In addition to the uranium contamination, the quarry also contains relatively large concentrations of calcium, manganese and sodium. Since the ion exchange resin will not have a preference for uranium versus other contaminated material, conventional non-regenerative resins will quickly saturate and a very large quantity of resin would be required.

On the other hand, a regenerative ion exchange system will not be feasible since the large quantity of regenerated effluent as well as the resin chemicals will accumulate in the equalization basin. This situation would eriously interfere with the discharge of effluent to an uncontrolled environment.

A chelating resit will selectively remove uranium ions from the vaste stream and therefore require significantly less resin and vill be easily regenerated. DEI estimates that the volume of chelating resin will be 1/240th the volume of the ion exchange resin that would be required. Table 1 demonstrates the efficiency of uranium separation under various conditions.

2. Uranium removal using activated alumina -- DEI predicts that the activated alumina vill not provide the reduction in concentration required by the RFP technical specifications. Based on extensive laboratory testing, it is known that a removal efficiency of approximately ninety percent for uranium using activated alumina adsorption.

Based on an influent concentration of 1750pci/1 stated in table 11300-1 of the specification, it is predicted that an effluent concentration of 30pci/1 would not be achieved.

Furthermore, for the use of a non-regenerative system, the volume of activated alumina that would be required is very large. According to the information from the manufacturer, it is estimated that approximately 6500 cubic feet would be required. On the other hand, in a regenerative system, the frequency of regeneration will be once per eighty hours and a large amount of chemical could be required. Hence, the chelate resin has distinct advantages over activated alumina.

3. Uranium removal using coprecipitation and chelate resin -- Based on JGC's testing and operating experience, DEI/JGC propose to use a system that is based on the use of coprecipitation and chelating resins. The details of this system are described in section 7 of this proposal. In addition to having conducted significant R&D to substantiate the design described in this proposal, JGC has experience in the operation of the Vaste Treatment plant at Ningyo Touge, Japan. This facility which is operated by the Pover Reactor and Nuclear Fuel Development Corporation routinely processes vaste vater contaminated with uranium. The nominal flow rate of this plant is 23 gpm. Figure 1 provides a summary of the demonstrated removal efficiencies for uranium. Please note that although some samples exceed 30pci/1, the average value of the effluent would meet the 30pci/1 limit.

The Ningyo Touge plant was commissioned in January 1984 and the vater treatment plant has operated in a consistently reliable manner over the last five years. The commercial purpose of the plant is to provide water treatment for liquid waste generated from the refining of uranium one into UF. Therefore, the operation of this plant has provided JGC with extensive operating history and operations experience with respect to uranium removal. This type of operating experience does not exist in the U.S. Therefore, JGC has a unique base of experience that has been factored into the design described in this proposal. As a result of the experience gained at this Japanese plant, DEI can state, with a high degree of confidence, that the other competitors who are bidding on this RFP do not have this extensive experience base.

In summary, DEI has developed sufficient test and operating data to demonstrate that the chelating resins and coprecipitation techniques described in this proposal will provide an effective means of removing uranium from the Veldon Spring quarry in accordance with the effluent discharge limits specified in this RFP.

JGC has extensive experience in integrating the various technologies required to treat the Weldon Spring quarry vaste water into an integrated turnkey system that operates in a highly reliable manner.

Additionally, DEI has selected NUS Corporation located in Gaithersburg, Maryland, to be responsible, as a lower-tier Subcontractor, for the detail design work to ensure compliance with the laws, regulations and standards in the U.S.A. and also for the engineering coordination between the U.S. fabricators from whom all of the equipment and materials will be ordered, except the new chelating resin which is only manufactured in Japan.

DEI has selected Corrigan Company as the local construction contractor for site work. Figure 1.1 provides a summary of the work scope of the companies in this team. Figure 1.2 provides a summary of proposed equipment suppliers.



Concerning the remainder of this Proposal, Chapter 2 provides the administrative text of this Proposal in accordance with MK-Ferguson's format, and Chapter 3 provides clarifications and modifications to the commercial conditions proposed in the RFP.

The submittals required in the Technical Instructions to Proposers in the "Request for Proposal" are provided in Chapters 4 though 7.

Chapters 8 through 11 provide the Proposer's technical description of its processes.

Chapter 12 indicates the extent of items which will be imported and which will be limited to one consumable essential for the above-mentioned innovative processing.

Table 1

Adsorption parameters of UR-3100 for Mⁿ⁺ at 25°C

 $Q = KC^{1/n}$ Q = mg M/I-Res. C = mg M/I-Sol.

HCL	1.0	N	0.1	Ν
M ⁿ⁺	К	1/n	К	1/n
U ⁶⁺	2800	0.53	8400	0.39
Fe ³⁺	380	0.60	700	0.59
Cu ²⁺	< 1		220	0.56
V ⁵⁺	< 1		780	0.63
Zn ²⁺	< 1		< 1	
Ni ²⁺	< 1		< 1	
Mn ²⁺	< 1		< 1	

Definition of terms:

M - applicable metal, for example Fe, U, etc.

Q - amount (mg) of adsorbed metal per liter of resin

C - amount of influent metal per liter of solution

K - constant







OPERATION DATE

Figure 1.1 Demonstrated Removal Efficiencies

Contract I



RADIONUCLIDE SEPARATION PROCESS (RASEP)

H. Kuribayashi, Y. Koshiba, K. Suzuki, M. Shibuya JGC Corporation, Japan

ABSTRACT

Liquid radwaste generated from nuclear power plants or other nuclear facilities consists of a small amount of radioactive nuclides and a large amount of non-radioactive matter. By separating radioactive and nonreleased as non-radioactive waste. In addition, by fixing the separated radionuclides according to their nature, it will be possible to effectively and efficiently meet the waste disposal requirements. With this approach, JGC liquid waste and fixed in an inorganic adsorbent. As a result, maximum reduction of waste via a simple and economical method, plus safe discharge of the treated (decontamination) liquid waste to the environment can be achieved.

INTRODUCTION

Liquid radwaste generated at nuclear power plants and other nuclear facilities usually contains insoluble CP nuclides, soluble CP/FP nuclides and nonradioactive substances. The CP and FP are contained in the waste with a considerable amount or nonradioactive substances, which occupy substantial volumes in waste packages produced by conventional immobilization processes.

By separating and fixing only the trace amounts of radioactive nuclides from the large amount of nonradioactive substances present in liquid radwaste, the quantity of final waste packages can be greatly reduced since the resulting decontaminated effluent may be discharged to the environment. By using inorganic material and metal as long term and radioactivity-fixing media, JGC developed a process which solely and selectively separates radioactive nuclides from liquid radwaste. This process fixes separated nuclides in the form of stable packages for safe, long-term storage and disposal.

In cooperation with Tokyo Electric Power Inc., and other five Japanese utility companies, JGC has established this radionuclide separation (RASEP) process which mainly consists of filtration, adsorption and electro-deposition.

RASEP PROCESS

- Filtration: The filtration step separates suspended solids including insoluble CP nuclides, such as Co-60, Mn-54, Fe-59 and Zn-65, from liquid radwaste.
- Adsorption I: The soluble CP nuclides such as of Co-60, Mn-54, and Zn-65 are then selectively adsorbed and removed by chelating resin from the liquid radwaste.
- Adsorption step II: Here, Sr nuclides are adsorbed by chelating resin from the liquid waste in k' ch soluble FP nuclides remain.
- Adsorption step III: Zeolite, an inorganic material with a high adsorption selectivity for Cs nuclides, is next used to separate the Cs nuclides.
- Electro-deposition step: The CP nuclides adsorbed in adsorption step I are now eluted from the chelating resin and fixed on a metal cathode, such as a stainless steel plate, by electro-deposition.
- Auxiliary adsorption step: Sr nuclides concentrated in adsorption step II are eluted from the chelating resin and fixed by synthetic zeolite which has a high selective adsorption performance with regard to Sr.



Fig. 1. Major Steps of RASEP System.

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BASIC PROCESS FLOW

Figure 2 shows the basic process flow. Principal specifications of the filter materials and adsorbents are summerized in Table I.

Moreover, Sr nuclides adsorbed in the chelating resin during adsorption step 111 are eluted by hydrochloric acid for re-adsorption onto inorganic adsorbents which are suitable for disposal.





TABLE 1

Principal Specifications of the Filter Materials and Adsorbents.

Unit Operation	Material
Filtration	Hollow fiber filter
Adsorption I	Chelate resin (Unicellex UR-10)
Adsorption II	ditto
Adsorption III	Natural mordenite
Auxiliary adsorption	Synthetic zeolite (A-7)

Major steps for the treatment of liquid waste and filtration, and three adsorption steps from (1) to (111).

Liquid radwaste is fed to the RASEP process by a pumm under slight pressure. Radioactivity level of the treated liquid radwaste is below the detectable limit of conventional monitoring method. The adsorbed radioactive nuclides in the chelating resin at the adsorption step I are eluted by sulfuric acid then electro-deposited under certain electrochemical conditions and the nuclides are finally fixed on the metal cathode. RADIOACTIVE NUCLIDE SEPARATION PERFORMANCE TEST

- Simulated liquid radwaste solutions: Radwaste solutions simulating the high conductivity liquid waste generated at a RWR power plant were used to test the RASEP process performance.
- Filtration step: A High Efficiency (HE) filter, capable of removing particles larger than 0.04 microns, thoroughly separated suspended solids consisting mainly of Fe, and most Co-60, Mn-54, Zn-65.
- 3. Adsorption step 1: Chelating resin, Unicellex UR-10, was adopted in the test. All soluble CP nuclides present in a simulated solution were removed to the extent that the radioactivity level of the treated solution was below the detectable limits. The bed volume defined as a ratio of volume of treated liquid waste to resin volume, was found to be extremely large. Figures 3 and 4 show adsorption curves and the bed volume of the resin respectively.



Fig. 3. Radionuclide Adsorption Performance of UR-10 Chelating Resin.





- Adsorption step 11: As shown in Fig. 3. Sr nuclides passing through adsorption step 1 were efficiently removed by this additional UR-10 column.
- Adsorption step III: Natural mordenite selected from various inorganic adsorbents was used to selectively remove Cs nuclide from the waste. Radioactivity level of the treated effluent was below the detectable limit.

Adsorption steps II and III are auxiliary steps when soluble FP nuclides coexist in the liquid radwaste.

6. Electrodeposition step: The eluted solution was obtained by treating spent UR-10 by sulfuric acid. Co-60 and Mn-54 as major CP nuclides were tested in the presence of a small amount of Ni ion with the application of DC voltage. The Ni deposition on a metal cathode was observed with simultaneous incorporation of these nuclides.

Figure 5 shows the removal ratio of CP nuclides vs time under Ni deposition conditions. Figure 5 indicates that more than 90% of the CP nuclides was removed and the metal cathode could be repeatedly used.







- Fig. 5. Metallic Ion and Radionuclide Removal Characteristics by Electro-deposition Unit.
 - Auxiliary adsorption step: For the purpose of concentrating Sr nuclide from original liquid waste, chelating resin is superior to zeolite because of higher bed volume even for high conductivity liquid waste. However, zeolite is a more stable long termymatrix material for fixing Sr nuclides.

VOLUME REDUCTION

High conductivity liquid waste is conventionally concentrated by evaporation, then solidified using cement, asphalt or plastic.

In the RASEP, however, liquid radwaste can be treated to obtain almost non-radioactive liquid effluent which renders the solidification of such liquid unnecessary as a rule. As a result, a very high volume reduction is achieved. Figure 6 compares volume reductions achievable with the RASEP process and with conventional processes at BWR plants.





COST-EFFECTIVENESS OF THE RASEP SYSTEM

The RASEP consumes minimal energy since it can be operated at normal temperatures and pressures. Cost can also be reduced since its simple construction eliminates the need for any special, expensive components or equipment.

Compared with an evaporation concentration system (without a solidification process), the RASEP facility and utility costs are below one-sixth of those of the evaporation system, as shown in Fig. 7. Since a costly solidification unit is usually installed for conventional evaporation systems, the RASEP process can be expected to be far more economically advantageous than conventional processes.



Fig. 7. Cost-Benefit Comparison between the RASEP and Evaporation Systems.

ACKNOWLEDGEMENTS

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JGC's Wet Oxidation Process

1. General

The Wet Oxidation Process is applicable for volume reduction of organic radioactive solid and liquid waste. It is especially effective for the treatment of liquid waste containing organic chelating agents which may be generated from the decontamination of nuclear facilities. Through an oxidation reaction with hydrogen peroxide, chelate compounds are decomposed into CO_2 and H_2O . BWR cellulosic filter aids can also be treated using this process.

JGC's Wet Oxidation Process consists of the following:

- (1) Waste feed
- (2) Chemical feed
- (3) Reaction and distillate condensation
- (4) Distillate neutralization

The auxilliary equipment for the Wet Oxidation system includes an Ultraviolet oxidation system to complete the decomposition of chelate agents, and a solidification system to solidify the sludge produced by the system.

2. Process Conditions

2.1 Plant Capacity

The design basis of the existing Wet Oxidation Process (at JGC's Oarai Research and Development Center) is as follows:

)	NTA concentration	3 %
)	EDA concentration	0.7 %
)	Feed rate of liquid waste:	
	Initial charge	1.4 m ³
	Continuous feed (8hrs.) Total	0.7 m ³ /hr
	1.4 m ³ + (0.7m ³ /hr)(8hrs.) (7 m ³ /batch)(2 batches/day	= 7 m ³ /batch) = 14 m ³ /day = 3,700 gals/day

2.2 Operation Schedule

The duration of one batch operation for the treatment of liquid waste is 12 hours. The operation schedule is as follows:

(1)	Initial charge of liquid waste	45mins.
(2)	Adjustment of pH in the reactor	45 mins
(3)	Feed of catalyst into the reactor	15 mins.
(4)	Heating up the reactor	(begin with (2)
(5)	Continuous reaction in the reactor	8 hrs.
(6)	Residual liquid reaction and concentration	1 hr.
(7)	Neutralization of reactor bottom sludge	45 mins
(8)	Discharge of reactor bottom sludge	30 mins.





2.3 Chemicals

- A 50% H2O2 solution is used as an oxidizing agent during the reaction. (1)
- A 98% H2SO4 solution is used to maintain an acidic pH-value. (2)
- (3) A mixture of 10% FeSO4 solution and 10% CuSO4 solution is used as a catalyst.
- (4) A silicone-emulsion type anti-foaming agent is used under the additive condition of 1,000 - 5,000ppm aqueous solution.
- 25% NaOH solution is used to neutralized reactor bottom sludge after the residual (5)liquid reaction and concentration.
- 3. Description of Each Process Unit
- 3.1 Waste Feed

Decontamination liquid waste is directly charge into the reactor from the receiving tank by a pump.

3.2 Chemical Feed

The chemical feed unit consists of the following storage tanks and pumps:

- One storage tank for the 50% H2O2 solution and one pump. ö
- One storage tank for the 98% H2SO4 solution and one pump. Ō
- One storage tank for the 10% FeSO4 and 10% CuSO4 solutions and one pump. 0
- One silicone-emulsion type anti-foaming agent tank and one pump. 0 0
 - One storage tank for the 25% NaOH solution and two pumps.

3.3 Reactor

The reactor unit consists of one reactor, one demister and one condenser.

(1) Initial charge

The effective reactor volume is 111.4m³. After the initial charge of liquid waste into the reactor from the receiving tank, the pH-value shall be adjusted to between 3 and 4 by adding sulphuric acid.

A 10% FeSO4 and 10% CuSO4 mixture solution is fed to the reactor, where the concentrations of FeSO4 and CuSO4 are 0.01mol/liter.

(2) Heating up the reactor

A steam heating coil and an agitator are installed inside the reactor. Chemically adjusted liquid waste is steam heated up to 95%.

(3) Continuous reaction in the reactor

The oxidation reaction proceeds under an evaporative condition by the continuous feeding of liquid waste and 50% H2O2 oxidizing agent solution accompanied by steam heating.

The silicone-emulsion type anti-foaming agent is constantly added.

Vapor and produced gas, mainly consisting of CO2, are evacuated into the condenser through the demister. The vapor is condensed and introduced into the distillate tank.





(4) Residual liquid reaction and concentration

After the continuous waste feed is stopped, heating is required in order for the residual liquid reaction to proceed with the supply of H_2SO_4 and H_2O_2 .

(5) Neutralization of reactor bottom shudge

The residual liquid in the reactor consists mainly of SO₄-2 and suspended solids. Before discharge, a 25% NaOH neutralizing solution is added.

(6) Discharge of reactor bottom sludge

The reactor residue liquids and solids are discharged and transferred to the sludge dewatering unitt.

3.4 Distillate neutralization

The distillate neutralization unit consists of one tank and one agitator. Before discharge of the distillate, the pH value is monitored by the pH sensor and the distillate is neutralized by adding 25% NaOH solution.

- 4. Control System
- 4.1 Process Parameters

During operation of the Wet Oxidation Process, the following parameters shall be controlled in order to ensure the correct reaction conditions:

- (1) Initial Conditions CuSO₄ (catalyst) : FeSO₄ (catalyst) :
- (2) Liquid Waste :
- (3) Chemicals 50% H₂O₂ : Anti-foaming agent : 98% H₂SO₄ :
- (4) Reaction Conditions Operating temperature : Operating pressure : Operating pH range :

0.01 mol/liter of reaction liquid 0.01 mol/liter of reaction liquid

Constant feed

Constant feed Constant feed Constant feed

Boiling point (approximately 100°C) Atmospheric 3-4

4.2 Process Control

The Wet Oxidation Process is controlled remotely and automatically from the control panel. The main indicators and recorders are mounted on the control panel to provide information on the operation status. During the reaction, liquid waste and chemicals are fed at a constant flow rate. Heated steam is supplied at a controlled flow rate based on the evaporation flow rate and the reactor temperature.







(1) Feed Rate of the Liquid Waste

The liquid waste feed rate to the reactor must be kept constant in order to maintain the vaporizing capacity. The flow rate is controlled by presetting the pump discharge rate.

(2) Feed Rate of Chemicals

The 50% H_2O_2 oxidizing solution and silicone-emulsion type anti-foaming agent are fed at a preset flow rate to maintain a normal reaction.

The concentrations of $CuSO_4$ and $FeSO_4$ catalysts in the reactor are 0.01 mol/liter of liquid waste in order to maintain a normal reaction. Prior to the start of the reaction, the liquid level in the reactor is checked and an estimate made of the volume of catalyst required. The flow rate of the pump is then set.

A 98% sulphuric acid solution is intermittently fed at a fixed flow rate so as to keep the pH value in the reactor between 3 and 4.

(3) Liquid Level and Temperature

In order to prevent excessive concentration of liquid, overflow and foaming in the reactor, the level of liquid is maintained at a constant level.

Additionally, the temperature in the reactor must be kept at boiling point to maintain good reaction conditions. In order to maintain the reaction temperature, heating steam consumption is controlled. If the liquid level falls below the "Low Level" set point, subcooled distillate reflux the from condenser is added until the liquid level regains the "Normal Level".

(4) Interiocks and Alarms

For the maintenance of safe operating conditions, interlocks and alarms are installed in the Wet Oxidation Process.

- o Liquid level in the reactor
- When the liquid level in the reactor rises to the "High" set point, feed of liquid waste and chemicals is stopped a: omatically and an alarm sounds.
- o Foaming level in the reactor
- When the foaming level in the reactor rises to the set point "High", the feed of liquid waste and chemicals are stopped automatically and an alarm sounds. • Pressure in the reactor
- When the pressure difference between the inside and the outside of the reactor rises to the set point "High", an alarm sounds.
- o pH value in the reactor
 - When the pH value in the reactor decreases below 2, an alarm sounds.
- (5) CO₂ Concentration Sampling System

The reaction operation is stopped by the time controller. When the decomposition is almost completed, the CO_2 concentration in the off-gas decreases rapidly. Therefore, CO_2 sampling is required.

(6) Radiation Monitoring System

To discharge the distillate, a radiation monitoring system is required.



METHOD OF TREATING RADIGACTIVE WASTE WATER OCMTAINING EDTA

Yasuhiro Moriya, Norimitu Kurumada, Fukuzo Todo and Hiroshi Kuribayashi

JGC Corporation 2205, Naritacho Oharaimachi, Higashiibaraki-gun Ibaraki Pref. 311-13 Japan (0292)66-3311

I. INTRODUCTION

In the decontamination of radioactive waste from nuclear installtions, there is a considerable discharge of radioactive waste water containing decontaminating agents. Decontaminating agents often contain disodium ethylenediamine tetraacetic acid (EDTA), formic acid, citric acid and other organic acids. The radioactive waste water is concentrated by evaporation to reduce its volume, and the resulting residue is solidified by the use of a solidifier such as cement. Nowever, when EDTA and other organic acids are present in the residue, the properties of the solidified product are unfavorably affected, particularly the mechanical strength and the leachability of nuclides thereof. Therefore, it is essential to emove EDIA and other organic acids from the radioactive waste water prior to the evaporation-concentration processing. Here we report a method, the wet oxidation method, of decomposing EDTA and other organic acids present in the radioactive waste water.

II. PRINCIPLE

Wet exidation is a method of exidizing organic compounds by the use of hydrogen peroxide as an exidizing agent in the presence of catalyst under atmospheric pressure. This method has been developed to decompose solid organic compounds such as ion-exchange resins or filter sludges suspended in aqueous solutions¹. Here we apply this method to decompose water-soluble organic acids present

in radioactive decontaminating agents".

III. REACTION CONDITIONS

The decomposition reaction depends on the concentration of organic compounds, the reaction temperature, the concentration of catalyst, the pH of the reaction mixture and the supply rate of hydrogen peroxide.

 Types and Concentrations of Organic Compounds 1-15% EDTA, which is a decompositionresistant material, was mainly tested, and 1% formic acid, 1% citric acid and 1% oxalic acid were also tested as reference materials.

B. Reaction Temperature

The reaction temperature was changed in

the range of 80 to 100° C to study the effect of temperature on the decomposition of organic acids.

C. Types and Concentrations of Catalysts

Fe and Cu are effective catalysts for the present decomposition reaction of organic acids. We added 0.01 to 0.02 mol/liter of Fe, Cu or a mixture of the two as the catalyst.

D. pilof the Reaction Mixture

The catalyst becomes inactive when the pill of the reaction mixture is alkaline. By adding various amounts of sulfuric acid and changing the pill of the reaction mixture, we investigated the effect of the pill of the reaction mixture on the rate of decomposition of organic acids.

E. Supply Rate of Hydrogen Peroxide

When the supply rate of 35wt% hydrogen peroxide is too high, the oxidizing efficiency of the hydrogen peroxide falls. On the otherhand, when the supply rate of hydrogen peroxide is too low, the rate of decomposition of organic acids becomes impracticably slow. We studied the effect of the hydrogen peroxide supply rate on the rate of decomposition of organic acids where the longest reaction time was set at 5 hours.

IV. PROCESS DESCRIPTION

A schematic flow sheet, of the wet oxidation process is shown in Fig. 1. The operation is performed as follows.

A. Initial Feeding

The radioactive waste water is supplied from the storage tank to the reactor. FeSO₄ and/or CuSO₄ is added to the reactor as a catalyst. Sulfuric acid is added to the







Fig. 1 Schematic flow sheet

reactor, if necessary, to make the pilof the reaction mixture acidic.

B. Heating

The reaction mixture is heated to 80 to

 100° C by the use of an electric heater or a steam-coil heater.

C. Reaction

After heating the reaction mixture, the oxidation reaction proceeds by continuously supplying 35% $\rm H_2O_2$. When the reaction mixture

is boiled, the vapour is condensed by a condenser. The condensate is transferred to the distillate treatment unit, and the produced gas to the off-gas unit.

D. Neutralization

When the reaction mixture is acidic at the end of reaction, 25% NaCH is added to neutralize the reaction mixture.

E. Evaporation Concentration

After neutralizing the reaction mixture, it is concentrated by evaporation to make the concentration of Na_2SO_4 25%. The concentrate

is drained to the concentrate receiving tank and then transferred to the solidification system.

V. RESULTS

Tests were performed by the use of beakerscale test system with a glass-made reactor having an effective volume of 1 liter, and also by the use of a pilot-plant test system with a titanium-made reactor having an effective volume of 150 liters. Firstly using the beaker-scale test system, optimal reaction conditions for decomposing EDTA and other organic acids were determined to study the decomposability of various organic acids. Then using the pilot-plant test system, the decomposition of EDTA as studied, as a typical organic acid. Table 1 Eaction conditions

	1	1	1	4		
Despusie mutarial apresentration (west	1.1.18	3	3	3	,	
Reaction Lamparations (⁸ 0)	1.00	80-100	(20	(00	1.045	80-100
Deluivet concentration(epi/)	Pa 0.51 De 0.01	Pa 0.01 Cu 0.01	Pa 0.01-0.07 Cu 0.91-0.07	2.67	78 0 01 Qu 0.01	78 8.51 Cu 0.01
Assettion time (hr)	5	1	5	5	4	5
neture of %3804 addeel (9)	10-50	15	15	6-15	15	15
Hy0; lead resa (g/hr)	30 - 279	70-90	76 - 90	76	30-140	30-40

A. Beaker-Scale Test

1. Decomposition of EDTA. Using a beaker-scale test system, we studied the effect of the EDTA concentration, the reaction temperature, the type and concentration of the catalyst, the pH of the reaction mixture and the supply rate of hydrogen peroxide on the decomposition degree of EDTA.

a. Effect of the EDTA concetration. One liter of test waste water, containing 1, 3 and 15 with of EDTA, was treated under the reaction conditions shown in column 1 of Table 1. The relations between the decomposition degree of EL:A and the consumption of 35% ${\rm H_2O_2}$

were investigated. Here the decomposition degree is defined as:

Decomposition degree(%) = [(initial TOC-TOC after reaction)/initial TOC)*100

where TOC represents the total amount of organic carbon. At 90% decomposition of EDTA, the amount of H_2O_2 consumed was about 4.6g per

ig of EDTA decomposed, which did not depend on the concentration of EDTA. The decomposition of EDTA reached 99% at the completion of reaction when EDTA concentration was 1% and 3%, but it leveled off at 94% when the EDTA concentration was 15%. The latter results is due to the raising of the pH of the reaction mixture in the later phase of the reaction, as will be described later.

b. Effect of the reaction temperature. One liter of 3% EDTA aqueous solution was treated under the reaction conditions shown in column 2 of Table 1. At 90% decomposition of EDTA, the consumption of H_2O_2 per 19 of EDTA decomposed was 4.69 at

 100° C, and 6.7g at 80° C. `t lower reaction temperatures, the consumptive f H_2O_2 increased substantially, i.e. the oxidizing efficiency of H_2O_2 is low. Therefore, it is recommended to treat EDTA at as high a temperature as possible.





c. Effect of the catalyst concentration. One liter of 3% EDTA aqueous solution was treated under the reaction conditions shown in column 3 of Table 1. The decomposition of EDTA was compared in the presence of FeSO₄, CuSO₄ or a mixture of the

two as a catalyst. We found that the mixture of $FeSO_4$ and $CuSO_4$, each concentrantion being 0.01 mol/liter, is the most effective catalyst.

d. Effect of the pH of reaction mixture. One liter of 3% EDTA aqueous solution was treated under the reaction conditions shown in column 4 of Table 1. When H_2SO_4 was not added, the decomposition degree of EDTA was 75% at best. However, it reached 99% when H_2SO_4 was added. In the decomposition of EDTA sodium ions are dissociated, which increases the pH of the reaction mixture if H_2SO_4 is not added. As the catalyst precipitates and becomes inactive in the range of alkaline pH, the decomposition degree of EDTA levels off unless H_2SO_4 is added. Therefore, it is recommended to keep the pH value of the reaction mixture acidic by adding H_2SO_4 .

e. Supply rate of H_2O_2 . One liter of 3% EDTA aqueous solution was treated under the reaction conditions shown in column 5 of Table 1. In the range of 0.90 to 4.8g of H_2O_2 per 1g of EDTA per hour for the supply rate of 35% H_2O_2 , the consumption of H_2O_2 did not depend on its rate of supply. Therefore, one can select any H_2O_2 supply rate mentioned above as long as the reaction rate is within a practicable range.

2. Other organic acids. Other than EDTA, we selected 1% formic acid, 1% citric acid, and 1% oxalic acid and treated them under the reaction conditions shown in column 6 of Table 1. The decomposition degree of all these organic acids reached 97 to 100% within one hour. The consumption of 35% H_2O_2 was less than 2.5 to 3.6g per 1g of organic acid decomposed. In the case of volatile formic acid, the decomposition reaction was performed

at 90°C. The decomposition degree of formic

acid was the same at 90° C and at 100° C. Therefore, it is recommended to treat volatile organic acids at lower temperatures in order to reduce loss by evaporation.

B. Pilot-Plant Test

A volume of 150 liters of 3% EDTA solution was treated by a pilot-plant test system based on the reaction conditions indicated in Fig. 2. The relations between the decomposition

degree of EDTA and the consumption of $\rm H_2O_2$ were studied and the results are shown in Fig. 2. At 90% decomposition of EDTA, the consumption of H_2O_2 was 4.3kg per 1kg of EDTA decomposed,





which was comparable to consumption of 4.6kg of H_2O_2 per 1kg of EDTA decomposed observed in the case of the beaker-scale test. Therefore, we consider that EDTA is similarly decomposed in the pilot-plant system as in the beaker-scale system.

VI. CONCLUSION

Almost perfect decomposition of the organic acids is achieved by the wet oxidation method when the reaction temperature increases up to the boiling point. In order to avoid decay of the catalyst, a small amount of sulfuric acid must be added to the reaction mixture before starting the reaction so as to maintain the pH value of the reaction mixture within the allowable low range. More than 90% decomposition of EDTA was achieved when the reaction was performed at the boiling point and at a low pH value. A quantity of 5 to 10kg of 35% H_2O_2 was consumed per 1kg of EDTA



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IMPROVEMENT OF THE HIGH TEMPERATURE SLAGGING INCINERATION SYSTEM

H. Kuribayxshi, N. Kurumade, S. Shibata and K. Kural UGC Corporation Yokohama, Japan

ABSTRACT

The objective of the paper is to describe the High Temperature Slagging Incinerator (HTSI) application study to reduce the volume of low level dry active weste (DAW) generated at nuclear power stations in Japan. The system, originally developed by SCK/CEN in Belgium, is being practically applied for the treatment of radioactive waste in Europe. JGC has been roducting the R&D work with a view to improving overall performance of the system and ennancing its safety and reliability. The performance tests of a plot plant with a design capacity of 100 kg/nr revealed a number of positive attributes. It also ensures stable products which can be readily solidified when necessary for disposal.

INTRODUCTION

The High Temperature Slagging Incineration (KTS1) system characteristically volume-reduces combustible and non-combustible dry active waste (DAW) at the same time. The system was developed at the Belgian Nuclear Research Center (SCK/CEN) where the system shown in Fig. 1 is presently operating. To inter use HTSI as a main subsystem into the DAW treatment ...tem, UGC has advanced improvement of the SCK/CEN system and related technology development through cooperation with Belgonucleaire, with the following goal:

- Improvement of the pretreatment subsystem to enlarge composition range of waste to be treated, and enhance system component durability.
- 2. Scaling up of the incinerator.
- Improvement of the off-gas cleaning subsystem to raise the DF value, relieve the corrosive environment and raise heat recovery.
- Development of a solidification process which ensures easy solidification of incineration residue.



Fig.1. HTS1 system at SCK/CEN

In order to achieve these goals, JGC constructed a 100 kg/hr pilot plant at the JGC Nuclear Research Center in Dharai in July 1985 by making principal improvements of the pilot plant in Belgium. Operation records after those improvement are described below.

BACKDROUND OF IMPROVEMENTS

The following DAW is generated at nuclear power stations mainly through periodical inspection and facility improvement:

- Hard materials such as concrete, thermal insulation material, wood blocks, etc.
- Soft materials such as waste cloth, plastic sheets, etc.
- Comparatively easily shreddable small metals such as wire, thin iron plates, small diameter pipes, etc.
- Unshreddable large metals such as large diameter shick pipes, valves, etc.

To ensure stable incinerator operation, DAW should be shredded into 5 cm or smaller size pieces, while the waste listed in Item 4 above should be excluded for treatment by the HTSI system. The DAW (hard and soft) listed in Items 1 to 3 differ in properties from each other; when they are treated by a shear shredder only, the abrasion of shredder edges lowers the throughput of the pretreatment subsystem.





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In order to solve this problem, the pretreatment subsystem was modified as shown in Fig. 3, so that the subsystem could perform its anticipated function; shredding waste into small pieces for prolonged period of time.

The modified pretreatment subsystem characteristically treats hard and soft materials by a crusher and shredder, respectively.

After waste sorting, acceptable hard and soft DAW is directly fed to the crusher and shredder, respectively, through Line 2. When perfect waste sorting is not expected, mixed DAW is fed to a shorting unit through Line 1. Hard DAW is then crushed into small pieces by the crusher and separated from the mixed DAW by a trommel screen before remaining soft DAW is fed to the shredder. As a result, abrasion of the shredder edges is avoided and durablity of the shredder is improved. In order to allow the HISI system to function satisfactorily, hard DAW must be completely crushed into small pieces by the crusher. Thus, a rotary hammer type machine was selected for

In the SCK/CEN off-gas cleaning subsystem, off-gas is released after being cooled by a water injection cooler (WIC) and then treated by a bag and HEPA filter. The following were considered in modifying the off-gas cleaning subsystem:

- Replacement of the bag filter with another high efficiency filter system aiming * a prolonged life of HEPA filter elements.
- Elimination the WIC since some measures must be taken to prevent dew condensation due to high water content when the WIC is used.
- High temperature off-gas treatment to prevent condensation of HCl and SO3".

HALO DIAN COMPOSED

Soli DAN

LINE 1

Fig. 4 shows a modified off-gss cleaning subsystem in which these improvements were incorporated:



Fig. 3 Modified pretreatment subsystem.









The HEPA filter load is reduced by adopting a high-efficiency ceramic filter, and the entire system can be operated at high temperature by adopting a high temperature HEPA filter. Water content in the off-gas is reduced by adopting a heat exchanger and fuel saving is ensured by using heated air for combustion air.

PILOT PLANT DESCRIPTION

The main process flow diagram of the HTSI is shown in Fig. 5. The simulated non-radioactive waste consists mainly of wood, paper cloth, sheet, rubber, small metals, concrete and thermal insulation material. The waste is crushed or shredded, then fed to a mixing bin. The mixed waste is then fed to the incinerator at a normal feeding role of 100 kg/hr, with intermittent operation of a conveyer at 20 minutes per hour. The incinerator residue, falling from FLK in the form of droplets into a water bath, is cooled in water. dried by heated air, then packed into 200 liter drums.

The incinerator off-gas is cooled to about 500°C in a shell-and-tube countercurrent air heat exchanger and further cooled to 300°C by an air diluter, then filtered by the ceramic filter. To prevent the off-gas from leaking, the entire facility is kept below atmospheric pressure by an off-gas blower.

Pratreatment unit

The crusher is a heavy duty industrial machine equipped with a vertical rotating shaft with hammers capable of crushing hard materials into 1 to 3 cm pieces, having a crushing rate of 2 ton/nr for average composition waste.

The shredder consists of a single low-speed rotating shaft equipped with disk-shaped blades, stationary edges fixed on the outer wall, and a 5 cm perforated screen below them. Its shredding rate is S ton/hr.



Fig. 5. Flow scheme of the HTSI Pilot Plant

Cff-cas cleaning system

To avoid corrosion problems, the heat exchanger tube sneet and tube inlet portions are coated by ceramics. To confirm filtering performance and duratility, two types of ceramic filters are equipped just for performance evaluation purposes. The ceramic filters are classed on-line by blowing compressed pulse jet air on each element.

OPERATING EXPERIENCE

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Cumulative operating time has reached approximately 2,000 hrs, and about 40 start-up and shut down procedure have been experienced. Soundness of all refractory has been verified, though small cracking caused by thermal cycling has been observed. In the highest temperature range, the refractory inside the main combustion chamber has shown no evidence of significant surface attack by adhered molten slag.

In the performance test, non-combustibles such as iron, concrete, thermal insulation material, HEPA filter elements, etc., and combustibles such as paper, wood, plastic sheets. On exchange resin, etc., were treated in the pilot plant. Although waste composition effected incinerating and shredding performances and volume reduction, the residual carbon was kept at approximately zero. Continuous and constant composition feeding ensured very small fluctuation of the residue properties as shown in Fig. 6.



Fig. 6. Incinerator residue properties in a campaign.

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	Waste Compo	\$(1)D+++#1#1	Operation	Treeted	Capacity
RUG ND	Unburnable	Burnable	(n / Run)	(Ion/IRun)	Ckg Zh 3
ei~ 0.8	5.0	5.0	3.5	2.8	
H=0 0 - 1 0	30	7 0	60 90	5 ~ 8	90 ~ 100
si≈1 2 × 1 8	4.0	6.0	60 - 100	6 ~ 9	90 - 10
si = 1 8 - 2 0	6.0	4.0	80 ~ 50	8 ~ 8	(00
R-21-32	7.0	3.0	08 ~ 03	1 ~ 3	120
H = 3 3 = 3 8	6.0	2.0	30 - 50	3 ~ 5	120
H-33-39	7 0	3.0	30 ~ 65	5 ~ 8	120
H-40-41	60	40	5.0	5	110

Table 1. Typical operating results



The HTS1 system achieves high volume reduction since combustibles are almost completely burned, and the residue hulk density is high. DF value of each component was then measured, using $Co(MO_3)_2$ and Cs_2SO_4 as chemical tracers; results are shown in Table 1. Of measurement indicated a DF of over 10⁴ for the cershic filter, and above 10⁵ for the entire system including the incluerator.

In commercial plants, additional HEPA filters may be installed to the system for safety purposes so that radioactivity released to the environment can be reduced to near zero.

Run No.	(H = 3	0)	(8+22,	23,24)
Element	Ċo	Cs	Co	C s
Incinerator	167	۱.5	229	2.8
Ceramic filter	\$1.74×10 ⁴	1.1×10 ⁵	\$1.74×10 ⁴	1.1×10 ⁵
Total	\$2.9x10 ⁶	1.7×10 ⁵	\$4.0x10 ⁶	3.1x10 ⁵

* Calculated from the "stection limit.

Table 2. DF value of each component

Duration	Leaching rate*
(day)	(g/cm ² day)
3	1.6 × 10+7
3	8.4 × 10 ⁺⁸
3	1.5 x 10*6
• 20	8.4 x 10*9
2.0	** 8.9 x 10"8
	3 3 20 20

**Calculated from the detection limit Non-combustibles /combustibles * 70/3D (wt/wt) * Measured by the MCC-55 method

lab's 3. Leaching rate of the residue

INCINERATION RESIDUE SOLIDIFICATION

In the main combustion chamber (FLK), the residue becomes molten slag at a high temperature of 1,500%C. For easy handling, the molten slag is granulated by quenching in water. The average granule size was approximately 3 mm, apparent specific gravity approximmately 3, and leaching rate based on the Soxhlet test below 10⁻⁶ g/cm² day as shown in Table 2. Although the residue can be easily solidified with conventional cement by out-drum mixing, this method generates secondary waste liquid from washing the mixor. The in-drum post packing method was found to be suitable for eliminating secondary waste. In this method, the residue is poured into a 200 liter drum, half filled with cement grout, which sits on the vibrating table.

CONCLUSION

The incinerator was scaled up actually attaining the design treatment capacity, and the HTS1 system having wide range of throughput can ben designed.





Outline of Laser Tritium Separation Process



Reaction Mechanism of Laser Tritium Separation



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- After treatment, soils exhibit very low leach rates for me, ury and heavy metals, as well as for other contaminates.
- Uranium and TRU contamination can be effectively statilized.
- Excessive volume increases are eliminated.

In addition, the system can easily be adapted to a mobile system, providing economical treatment *it* diverse locations without excessive set-up costs.

Successful Implementation of the concept will:

- Allow more effective stabilization and leach resistance for heavy metals.
- Provide the first effective method of on-site stabilization of low levels of mercury contamination.
- Reduce the cost of remediating sites contaminated with mercury.
- Allow the stabilization of soils contaminated with a combination of hazardous metals, uranium, and TRU.
- Allow efficient remediation of smaller contaminated sites.
- Reduce volume of low concentration TRU required to be sent to WIPP
- Provide a method for stabilization of soils containing heavy metals and TRU fixed to chelate agents such as EDTA and DTPA.

JGC is optimistic that this process could substantially improve the effectiveness and cost of remediation of these DOE sites. In addition, it is likely that many EPA-designated Superfund sites could benefit from the successful implementation of the concept.





Method	EP	- Toxicity			TCLP	
	Sample (mg/l)	Criteria (mg / l)		Sample (mg/l)	Criteria (mg / l)	
Cd	4.86	1.0	x	0.10	0.066	х
Cr	0.68	5.0	O	0.11	0.084~5.2	0
Hg	0.20	0.2	0	1)	0.025	-
Pb	9.71	5.0	0	0.36	0.18~0.51	0

Leach Test Results - Treated by fixing agent (I) and Cement -

1) Analysis is now in progress











Leach Test Results - Treated by fixing agent and bentonite -

	Fixing agen	t(I) and ben	tonite	Fixing age	nt(II) and ben	tonite	
Method	Aethod EP - Toxicity				TCLP		
	Sample (mg/l)	Criteria (mg / l)		Sample (mg/l)	Criteria (mg / l)		
Cd	0.63	1.0	0	0.04	0.066	0	
Cr	<0.01	5.0	0	0.11	0.084~5.2	0	
Hg	0.20	0.2	0	1)	0.025	_1)	
Pb	< 0.01	5.0	0	0.33	0.18~0.51	0	

1) Analysis is now in progress

Comparison between the Conventional Cement System and JGC's AC - Process

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Waste to be treated	Conventional cement method	JGC technology	Remarks
Incinerator ash	No pretreatment	Pretreated by the Ca(OH) ₂ and NaOH	A retarded cement hydrating reaction problem are solved.
Spent resin	No pretreatment	Pretreated by the cement and water	Swelling phenomenon of the immersion test are protected.
Boric acid waste	No pretreatment	Hydrate calcium metaborate are generated by pretreatment	Volume reduction and stable products are provided





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Table 1 Typical Product Properties

		Regulatory	Position (NRC)			JGC's Data		
No.	Test Item	Standard	Estimation value	Boric auid waste	Spent resins	Laboratory drain	Laundry drain	Incinerator ash
1	Free liquid	ANS 55.1	No more than 0,5 vol%	0	0	0	0	0
2	Compressive strength	ASTM C 39	More than 60 Psi	3,150	3,070	8,380	7,250	3,600
3	Gamma irradiation	ASTM C 39	More than 60 Psi	2,800	2,550	6,760	7,190	3,400
4	Leach test	ANS 16.1	More than Co	1 12.1	13.0	14.6	11.1	14.1
			(90 dyas) Cs	8.4	7.9	8.5	7.9	8.2
5	Immersion	ASTM C 39	More than 60 Psi	3,090	2,650	8,620	7,790	3,700
6	Thermal degradation	ASTM B 553	More than 60 Psi	2,810	1,970	8,670	4,830	3,100
7	Homo - geneity	ASTM C 39	More than 60 Psi	2,950	2,140	6,510	4,390	3,900
8	Bio -	ASTM G 21	More than	3,740	2,060	7,140	5,970	3,700
	degradation	ASTM G 22	60 Psi	3,510	2,940	6,370	8,280	4,600

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ADVANCED CEMENT SOLIDIFICATION PROCESS

Teruo Iji, Hideo Kodama Kyushu Electric Power Co., Inc. 1-82, Watanabe-Dori, 2-Chome, Chuo-ku, Fukuoka, B10 Japan.

René Pierlas, Claude Jaouen Société Générale pour les Techniques Nouvelles (SGN) 78184, Saint Quentin Yvelines Cedex, France.

> Hiroshi Kuribayashi, Norimitsu Kurumada JGC CORPORATION 2205, Naritocho, Oharai-machi, Higashi-Ibaraki-gun, Ibaraki Pref., 311-13 Japan.

ABSTRACT

The Advanced Cement Solidification Process, which features improved volume reductivity and properties of solidified wastes, has been developed to establish a better radioactive waste management system. Cement has been widely used as an inorganic solidification agent for the treatment of radioactive wastes generated at nuclear facilities. With current technology, borate waste solutions generated at PWR plant are neutralized with caustic soda and solidified directly with cement. This causes an increase in the volume of waste. Moreover, since borates retard the hydration of cement, the properties of solidified waste are not always such that meet the final disposal requirements. In order to eliminate these defects and to improve conventional cement solidified ment of borate. As a result of numerous experiments, a new process has been developed, by which solidified products with high volume reductivity and excellent physical properties can be produced. On the other hand, research and development work on a pilot plant have been carried out in co-operation with Kyushu Electric Power Co., Inc., including small scale hot tests of actual wastes at its Genkai Nuclear Power Station. A commercial-scale cold pilot plant was constructed and several proving tests have been conducted successfully.

INTRODUCTION

An increase in the amount of radioactive wastes generated at nuclear power plants along with an increase in their construction and operation, has created significant problems of radioactive waste management. Under such circumstances, the development of practical treatment processes for such wastes is an urgent task facing the world's nuclear power industry.

In order to store and transport medium and lowlevel radioactive wastes generated at nucle r power plants easily and economically, it is essential not only to reduce their volume to the maximum extent but also to transform them into a solidified product so that it is suitable for final disposal and will maintain its chemical and physical soundness over a very long period of time.

For this purpose, the Advanced Cement Solidification Process with high volume reducibility has been developed. Cement has been most commonly used as an inorpanic solidification agent for radioactive wastes at nuclear facilities, since it has many advantages as follows:

- It is standard, low cost embedding material.

- It is compatible with wet waste.

 Its hardened product is stable, and has high density and high mechanical resistance.

However, it has been generally observed that the volume of solidified products produced by conventional

cement solidification processes becomes larger than the original volume of radioactive wastes. Noreover, the high quality solidified products are not always produced in cases where borate wastes generated at PWR plant are treated by such processes, because borates present in the wastes retard the hydration of cement and impede the progression of the hardening of cement paste.

In order to eliminate these defects of most of the conventional cement solidification processes, mainly related to borate wastes, SGN had been developing a new volume reduction and cement solidification system, in which a borate waste solution is pretreated and then overconcentrated. Independently of SGN, SGC was conducting basic research on the protreatment of borates and on the hydration of cement.

In 1982, SGN and JGC agreed to begin co-operative work on the research and development of a volumereducing cement solidification process, based on their past experience. After a series of basic experiments, particularly on the pretreatment of borates. SGN and JGC developed the Advanced Cement Solidification Process, by which the volume of wastes can be highly reduced and solidified products with good physical properties can be produced.

Small scale hot tests of actual wastes based on this process, with the co-operation of Kyushu Electric Power Co. Inc., were carried out at its Genkai Huclear Power Station. In this hot test, the results of basic research were reconfirmed and the practicability of the process was proved. Based on preliminary performaance test data on selected equipment, commercial-scale



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pilot plants were designed and constructed in both JGC and SGN laboratories, the former spontured by Kyushu Electric Power Co., Inc., to carry put various demonstration tests on the operability, durability and optimal operating conditions of all equipment and systems. These tests have almost been completed, with the result showing that the process can be put into commercial service.

An outline of this process mainly relating to treatment of borate waste, is described in this paper.

PROBLEMS WITH MOST OF THE CONVENTIONAL CEMENT SOLIDIFICATION PROCESSES

In most of the conventional cement solidification processes, the concentrated borate waste solutions generated at PWR plants are directly solidified with coment. This causes two main problems. These are:

- Borates ions present in the waste solution retard the hydration reaction of cement. This makes it difficult to obtain sufficiently hardened products with good physical properties.
- The volume of hardened products is larger than the original volume of wastes.

The first problem may be solved by the following mechanical or chemical means:

Mechanical means when cement particles come in contact with a solution containing soluble borates, calcium borstes is formed on their surfaces. It remarkably retards the hydration reaction of cement clinker mineral, such as calcium stilicate, and also consequently setting time of cement. In order to make the hydration proceed, the calcium borate must be removed from the surfaces of the cement particles. This can be unnived, for

Chemical means
 Borates present in the waste solution can be
 transformed into almost nardly soluble borates.
 This eliminates the formation of calcium bo rates on the surfaces which retard the hydra tion and setting of cement.

In order to solve the second problem whereby the volume of solidified products must be further reduced, some means that would enable further concentrating of such solutions had to be developed.

example, by using a high-shear minute.

SGN and JGC decided to solve these two problems by chemical methods and carried but extensive basic research on various pretreatment methods for borate waste solutions. SGN and JGC finally succeeded in overcoming the problems by developing an effective pretreatment method, by which insoluble borates are formed from such solutions.

BASIC RESEARCH

Basic Research on Pretreatment

An objective of the pretreatment is to transform soluble borates present in the waste solution into hardly soluble borates so that their retarding action on the hydration of cement can be eliminated.

Some alkali earth metal salts of bonic acid such as calcium borate are known to be very insoluble in water. For examples, calcium borates exist in nature as born containing minerals. It is anticipated to be stable in the hardens: cement matrix which also consists of calcium compounds. Therefore, in the basic research, SGN and JGC's efforts were directed toward finding methods by which insoluble calcium borates could be formed when a calcium compound is added to the borate waste solution.

In cases where the pH of concentrated borates waste solutions was on the acidic side, the rate at which calcium borates were formed was extremely slow, whatever calcium compounds were added.

When an alkaline calcium compound was added to an acidic waste solution, the reaction rate was extremely slow, so it seemed that no reaction was taking place. The alkaline calcium compound is solid. When its particles came in contact with a boric acid solution, a thin film of insoluble calcium borates was formad on their surfaces, and it prevented the particles from being dissolved in the solution and stopped their reaction.

In order to further advance the reaction, two methods were considered.

- To enlarge the lattice structure of the thin film of the insoluble salt so as to allow calclum ions to easily pass through the film.
- To promote the dissolution of the calcium compound under specific conditions.

It is well known that in cases where cement is mixed directly with a borate solution, the mixture does not harden neither develop strength even after curing for one month, but the addition of sodium hydroxide to the solution to transform boric acid into sodium tetraborate or sodium metaborate causes the mixture develop strength. This fact was effective in solving the problem.

Sodium hydroxide was added to a borate solution, and the solution was then mixed with a calcium compound. The mixture was continuously stirred. As the reaction advanced, a thixotropic paste was produced. The higher the concentration of boric acid, the more viscous it became. And it sometimes stiffened and remained in a nearly pasty condition only when the stirring was continued. When the stirring was stopped, it became a gel and was difficult to handle. By this pretreatment method, it was possible to obtain low soluble calcium borates but it was difficult to concentrate the solution.

In order to improve reaction conditions in the pretreatment step, such as reaction temperature, stirring conditions (stirrer type, blade shape, revolution speed), dosage of cremicals, method of dosage, etc., further detailed studies were conducted.

It was found that pretreatment under adequate stirring conditions prevented the formation of the pasty substance and produced precipitates of insoluble calciumborate. The precipitates are crystals having good sedimentation tendency and can exist stably in a cement matrix.

Basic Research on Mixing Ratio

The solid/water/cement mixing ratio to prepare a cement paste is determined according to the procedures shown in Fig. 1. The cement paste must have a sufficient consistency at which it can be poured into a container. The consistency or workability can be measured as flow values defined in JIS R 5201 or ASTM C124-71.







Fig. 1. Procedure of Design of Mixing Ratio.

Experiments showed that the consistency or workability of cement paste increased proportionally as the volume fraction of liquid in a mixture increased. The volume of liquid in a mixture can be determined if the target flow value is set.

Meanwhile, the strength of solidified products depends upon cement/water ratio in a mixture. The cement /water ratio can be determined if the target strength of solidified product is set. Since liquid volume determines the quantity of water, the mixing ratio can be determined if the consistency of a mixture and the strength of solidified products are set.

Experiments or "onsistency and strength showed List a mixing ratio o: 50/30/20 (solid/cement/water) by weight was adequate in cases where the targe: flow value was set at 200 mm or larger, while the target one month compressive strength of solidified product was set at 20 MPa or larger.

HOT TEST

A serie, of small-scale hot tests of this process were carried out at the Genkai Nuclear Power Station of Kyushu Electric Power Co., Inc. The experimental results are summarized below.

- Calcium borate precipitates were successfully obtained in a pretreatment step.
- Pretreated waste solutions were concentrated. The concentrates were solidified with cement. Sufficiently hardened products were produced.
- In cases where the concentration of boric acid in the waste solutions was 12 wt 1, the volume of solidified product became approximately 1/4 of that of their original waste solution.
- Solidified products showed good water resistance.

The not tests confirmed the results of the basic research and proved that the process is effective in solidifying actual borate waste solutions.

BASIC PROCESS FLOW

The basic flow of the Advanced Cement Solidification Process is shown in Fig. 2. This process consists of the following three main steps.

Fretreatment Step

The pretreatment step, the key to the successful operation of the process, is indispensable for ensuring



Fig. 2. Basic Flow Diagram of Advanced Cement Solidification Process.

the hardening of cement to produce sufficiently solidified products, with good physical properties in a reproducible way.

in the pretreatment step, calcium borate is formed and precipitated. The precipitates are stable crystals and have good sedimentation properties.

Since dissolved borate, deterimental to the hydration of cement, can be converted into insoluble calcium borate precipitates, it has become possible to obtain sufficiently hardened products with good physical properties, which are suitable for waste storage, transportation and final disposal.

Concentration Step (Volume Reduction Step)

In this step, a borate waste solution is concentrated to such an extent that only a minimum quantity of water necessary for mixing with cement is left. This operation has become feasible by removing dissolved borate in the form of inscluble calcium borate showing good sedimentation.

From the viewpoint of reducing the volume of a borate waste solution, the most common method that enables the greatest volume reduction is to evaporate such solution to complete dryness. In the process, the borate waste solution is not evaporated to dryness but is concentrated to an optimum volume, because cement must be mixed with water so as to harden it.

The concentration step avoids technically difficult dry powder handling operations.

Mixing Step

An out-drum mixing method is employed in the mixing step. In order to produce solidified products having high strengt: and highly reduced volume, it is important to prepare a cement paste of good consistency with nimum quantity of water. It is also important to case the packing efficiency of such paste in a consiner. For these reasons, it is preferable to adopt the out-drum method, which can provide high mixing and filling.

Different kinds of cement can be used depending on requirements of the final product qualities. The results presented hereinafter are based on the ordinary portland cement (corresponding to ASTM Type I).

Advantages

The advantages of the Advanced Cement Solidifica. tion Process are:

- Based on 12 wt % of boric acid, the volume of solidified products produced by the process is 1/7 = 1/8 of that produced by conventional cement solidification processes.
- Solidifies products having good physical properties are produced.







- The process is safe. The process does not need any flammable material. Moverover it is a wet process, air-borne contamination du to radioactive nuclides resulting from a dry , "ocess can be avoided.
- All materials to be used in the process are readily available and inexpensive.

PILOT PLANT TEST FOR COMMERCIALIZATION

Examples of Typical System

As shown in Fig. 2, the process basically consists wee steps: pretreatment of borate waste solution. of three steps: concentration of slurry, and cement mixing. With regard to the concentration methods and operating modes, several systems are feasible. Examples of a typical system are shown in Fig. 3.

System-1 is a continuous process and consists of three steps: pretreatment, overconcentration and mix-ing. The slurry, obtained in the pretreatment step, is evaporated and concentrated in the step of overconcentration. The concentrated slurry is cooled and mixed with cement. This system is of adva tage in cases where there are frequent discharges of borate waste Since the slurry is concentrated by evapora. solution. tion, the concentrating vessel is equipped with a specially designed scraping mechanism so as to keep the heat transfer surface exposed at all times to the The mixer is also equipped with a similar slurry. SGN has designed and built a pilot plant mechanism. based on this system and has been conducting experiamnts.

System-2 is a batch wise process and consists of four steps. The concentration step in the basic process, which is shown in Fig. 2, is divided into two steps: solid-liquid separation and concentration of separated liquid. This system is advantageous in cases where discharges of borate waste solutions are less Calcium borate precipitates obtained in the frequent. pretreatment step have good sodimentation properties and easy to separate. This separated solid portion is and easy to separate. sent to the mixer and the separated supernatant is concentrated by evaporation. Since borate precipitates can be almost removed in the separation step, a conventional type concentrator can be used.

With the co-operation of Kyushu Electric Power Co., Inc., JGC has built a commercial-scale pilot plant based on this system and has been carrying out various tests to put this System-2 process into commercial SETVICE.

System-3 is also a batch wise process, using pretreatment, concentration and cement mixing-depending on the basic data. These three steps can be operated in one or two stages. The system is based on the concentration under reduced pressure condition of the slurry, during and/or after the pretreatment step. concentrated slurry is then mixed together with the The concreted product is poured into a drum in binder. This system has been developed with a the last step. view of reducing the number of equipment and making the entire system design compact. SGN intends to design a compact mobile unit based on this system, the corresponding tests are going on.

Pilot Plant Based on System-2

The effectiveness of system-2 process has been proven in pilot plant, which is outlined below.

Pretreatment:

A reaction vessel, equipped with a specially designed stirrer, was used. The stirrer served to effectively mix the borate waste solution with an added calcium compound. Crystalline precipitates of insoluble calcium borate were formed.

Solid-Liquid Separation:

A conventional type separator was used to separate the slurry into the supernatant and solid portions. The separated solids were sent to the mixer. The separated supernatant was first stored in a tank and then fed to an evaporator.

Concentration:

A small conventional type evaporator was used to concentrate the supernatant. Concentrated liquid was sent to the mixer and used as mixing water. Therefore, the supernatant was concentrated to the minimum



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possible volume by which the separated solids could be mixed with cement.

Mixing:

The pilot mixer used has blades which rotate at high speeds. Weighed quantities of the separated solids and the further concentrated liquid were premixed. The pre-mixture was well mixed with a weighed quantity of cement for a few minutes so as to obtain a workable cement paste. The paste was poured into a container. After curing the mixture at room temperature, a sufficiently and uniformly hardened product was obtained. After the mixing operation was completed, the mixer was easily washed.

A series of pilot plant tests on the performance, operation and durability of the equipment used in each step proved that the System-2 process successfully set all pre-set targets. Also, a large volume of engineering data including the operability and optimal operating conditions of the whole system was obtained. All experimental results showed that it would be feasible to put this process into commercial service. A conceptual design of an actual plant was carried out, using the experimental results. It was known from the conceptual design that a sufficiently economical and compact system could be constructed.

PHYSICAL PROPERTIES OF SOLIDIFIED PRODUCT

Several typical physical properties of solidified product obtained by this system-2 process are shown below.

Strength Development and Specific Gravity

The strength development of a solidified product obtained is plotted in Fig. 4. Aft. It was cured for about one month, it showed a strength of more than 20 NPa. Even one year later, measurements showed that its strength was steadily increasing.

An electron microscopic observation of a crosssection of the solidified product proved that the calcium borate formed in the pretreatment step was also stable in cement matrix for a long period of time.

The specific gravity of the solidified product is about 1.8.

Nater Resistance

The solidifed product does not lose its strength, as shown in Fig. 5, even if it is impursed in water for a long period of time. Its volumetrir change was equal to or less than 1% in one year, with result that there was no appreciable chamge in its shape. It has good water resistance.

Leachability

The leaching rate of two typical nuclides, Co⁴⁴ and Cs^{13*}, measured by using a specimen, 45 mm in diameter and 44 mm in height, was as follows.

 $Co^{4.0}$: 10^{-4} cm³/cm², day or less

Cs^{11*}: 10^{+*} cm¹/cm², day or less

The leaching rate of Cs13* can be further improved, if necessary, by adding zeolite or using blended cement.

Uniformity

A 200-liter (S5-gallon) solidified product was









produced. Samples were taken from the core to measure the specific gravity and strength. The measurements showed that the specific gravi. and strength of the solidified product were uniform throughout product.

VOLUME REDUCIBILITY

The volume reducibility obtained by the Advanced Cement Solidification Process is shown in Fig. 6. When treating 1 m² of concentrated waste solution containing 12% boric acid, conventional cement solidification processes produce about 2 m² of solidified product, with the result that the original volume is increased. On the other hand, this process produces 0.25 m² of solidified product, resulting in the volume being 1/4 of the original volume. When compared with most of the conventional process, the volume of the solidified product obtained by this process is about 1/8. Thus, this process attain an extremely large volume reducibility.

Accordingly, this process is very effective inmany respects, i.e., reductions in storage space, transportation costs, etc., when treating radioactive waste containing borates.

CONCLUSION

A method by which radioactive wastes containing bonate generated at PWR power plants can be chemically treated in such a manner that insoluble borates can be formed has been established, and based on the method, the Advanced Cement Solidification Process has been developed.







Fig. 6. Comparison of Volume Reduction.

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- The volume of solidified products produced by the process is about 1/8 of that obtained by conventional cament solidification processes.
- The process produces solidified products having good physical properties. Such products are suitable for storage, transportation / d disposal

- Cement, used as a solidifying agent. is inexpensive.
- Radioactive wastes are processed in liquid or slurry form, so no airborne radioactive powders are produced. Thus, the process can be safely operated.
- The system is highly reliable and consists of practical equipment. Proving tests were successfully carried out on a commercial scale pilot plant.
- The pretreatment step of the process was proved to be practicable by numerous pilot plant tests and also by hot tests.





ADVANCED CEMENT-SOLIDIFICATION PROCESS FOR SPENT ION-EXCIIANCE RESINS.

K. Sauda, F. Todo, T. Nakashima, T. Kagawa, H. Kuribayashi JGC Corporation

Yokohama, Japan 232

AUSTRACT

JGC has developed an advanced cement-solidification process (AC-Process) for the treatment and stabilization of radioactive spent ion-exchange resins generated at nuclear power plants. The AC-Process can produce excellent products in comparison with other existing cement solidification processes. In addition, this process requires a lower operating cost than that of the HIC system.

In general, cement-solidification products derived from spent ion-exchange resins tend to swell in water. Such exclining is caused by the expansion of resins in water due to the absorption of water and by the sdsorption of soluble contents in the cement matrix. In order to solve this problem, JGC has developed a new predesatment technique for obtaining coment-solidified products which will meet the requirements for final disposal. Extensive tests were conducted to determine pretreatment conditions. The properties of obtained products were evaluated to verify that they met the requirements for final disposal. The results of the tests and evaluation are reported below.

INTRODUCTION

JGC has carried out research and development work on coment solidification technologies for radioactive wastes for a long time. Such wastes as PWR evaporator concentrates, incineration ashes, etc., were difficult to cement-solidify using existing technologies. To stabilize such wastes, JGC has already established a new technology for obtaining highly volume-reduced solidified products excellent in water resistivity and other properties by using a JGC-developed unique pretreatment method (USP 4800042).

Also for ion-exchange resins, JGC has succeeded in solving problems associated with existing coment-solidification processes by pretreating ion-exchange resins before ixing them with coment.

The advanced cement-solidification technology for spent ion-exchange resins, developed by JGC, is described below with respect to the pretreatment method, process control program, Technical Position tests for cementitious waste forms.

PRETREATMENT OF SPENT ION-EXCHANGE RESINS

A problem associated with existing well-known coment-solidification processes for spent ion-exchange resins is that the matrix of the coment-solidified products obtained is disrupted when immersed in water for a long time. Coment-solidified products derived from spent bead resins particularly exhibit such a phenomenon. For this reason, the pretreatment of spent ion-exchange resins has been actively discussed at the Workshop on Coment Stabilization of LLRW held by the U.S. Nuclear Regulatory Commission (1).

This report describes the concept of the spent ion-exchangeresin pretreatment method developed by JGC in comparison with other pretreatment methods. The mechanism of the matrix disruption of coment-solidified spent ion-exchange resin products in water has not yet been clarified. However, such a disruption is said to be caused by the adsorption of soluble coment components by resins during the curing and by the swelling and contraction of resins due to the reactions between coment-water and resin-water.

Various methods have been proposed to prevent such a disruption of the matrix and they can be classified roughly as follows:

- Adsorption of such ions as Na, Ca, etc., by cation exchange resins (2).
- 2) Improvement of central, binders (3).
- Coating of resins with polyester or similar materials (4).
- 4) Pretreatment to raise the water content (1).

Concerning method 1, the following problem exists: When spent ion-exchange resins are pretreated by NaOH, the adsorbed Na ions are replaced after a long period of time by Ca ions contained in cement. As a result, the matrix of the cement-solidified product is disrupted. The pretreatment of spent ion-exchange resins by lime (CaO) cannot prevent an increase in the anion exchange resin volume, though it is very effective in preventing an increase in the cation exchange resin volume.

According to our test results, it is revealed that when a coment-solidified product derived only from anion exchange resins is subjected to a water immersion test, the "sume of the product gradually increases and finally disruption of the matrix and cracking occurs. Figure 1 shows the immersion test results.

On the other hand, the following problems are associated with methods 2 and 3: As methods for improving cement binderc, examples exist of using slag cement, alumina cement, polymer gypsum cement, or sulfur cement. However, these methods do not differ from the use of

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Sauda CEMENT SOLIDIFICATION PROCESS



Fig. 1. Swelling of Solidified Spent Resin Products During Immersion Tests.

ordinary Portland cement. In method 3, spention-exchange resins are coated with an organic high-viscosity liquid such as polyester which requires a special chemical or heating for hardening to occur. This complicates the solidification process. Therefore, this method is undesirable although is effective.

Alternatively, as listed as method 4, spent ion-exchange resins can be coment-solidified under water-containing conditions. However, this method is not very effective because spent resins become dry during the curing.

As described above, the spent ion-exchange tesin pretreatment methods reported up to the present time have both advantages and disadvantages. The development of a new coment solidification te cology has therefore been desired. Considering such circumstances, JGC conceived the following pretreatment.

After the addition of a certain amount of coment and water, dewatered spent ion-exchange resins are agitated at a high speed in a highly alkaline coment slurry. Coment hydration of the spent resins after mixing with coment can thereby be prevented.

On the basis of this concept, JGC studied the spent ion-excluance resin protreatment conditions, especially the amount of cement to be added, pretreatment time, and agitator revolutions. As a result of the study, it was revealed that satisfactory pretreatment effects could be achieved by high-speed agitation at 350 rpm using a high shearing mixer, coment addition of 20 with on a dry resin basis, agitation for shown in Fig. 1, application of the coment pretreatment could also prevent matrix disruption of coment-solidified anion reain products.

TECHNICAL POSITION TESTS ON SOLIDIFIED PRODUCTS

Technical Position tests were conducted to verify that cement-solidified products obtained by applying this new pretreatment method met the technical requirements of 10 CFR 61, Technical Position (1983). The tests were comprised of a compressive strength test, radiation stability test, biodegradation test, leachability test, immersion test, thermal cycling test, free liquid test, and a full-scale test.

Preparation of test products

Non-radioactive spent ion-exchange resins were used to prepare solidified products for laboratory and fullscale tests. For leachability tests, the radionuclides, Co and Cs, were added to resins. For field tests, spent ion-exchange resins actually generated at a PWR plant in Japan were used.

Spent bead and powdered resins were both used as ion-exchange resins and the standard mixing ratio of cation to anion exchange resins was 1:1.

Test procedure

Compressive strength tests were conducted in accordance with ASTM C 39. Test specimens were cured in water-saturated air for 30, 60, and 90 days.

For radiation stability tests, specimens were irradiated with gamma rays of 10⁵ rads at the Japan Atomic Energy Research Institute. Biodegradation tests were conducted in accordance with ASTM G21 and ASTM G22.

Leachability tests and immersion tests were conducted __coordance with ANSI 16.1 Test specimens cured in water-saturated air for 30 days were used for all the tests.

Thermal degradation tests were conducted in accordance with ASTM B553 and free liquid tests were conducted using full-size (55-gal drum) solidified products prepared by a demonstration pilot plant (Fig. 2) at JGC Oarai Research and Development Center.

Full-scale tests were conducted using specimens taken from a 55-gallon drum size solidified product prepared by the pilot plant, using a core boring machine.

Physical properties of products

The physical properties of cement-solidified produets obtained are shown in Table I. All the data satisfied the criteria required by the Technical Position on Waste Form. From the test data, it is concluded that the cement-solidified



Sauda CEMENT SOLIDIFICATION PROCESS

products obtained by this process much the requirements for Class B and Class C waste forms.

System Description

JGC's advanced cement-solidification process, for the effective pretreatment of spent ion-exchange resins, is shown in Fig. 3. Spent ion-exchange resins are received in a spent resin receiving tank, then sent to a dehydrator or centrifugal separator for dehydration purposes. The supernatant is returned to a spent resin storage tank and the dewatered resins are received in a mixer, where the resins are mixed with the specified amount of cement and water. After the specified amount of cement is added, the pretreated resins are sufficiently mixed, then filled into drums.

Control parameters

The following parameters are controlled to obtain satisfactory cement-solidified products. The amount of cement to be added for pretreatment is more than 20 wt% on a dry resin basis. Resins are mixed for more than 10 minutes at an agitation speed of 350 rpm and cured for more than 3 hours. In addition, when the pretreated resin is mixed in the mixer, the weights are controlled so as to enable the follow-







Fig. 4. Relation Between Mixing Ratios (R/W/C) and Bounding Limits.

ing mixing ratio:

Resin/Water/Cement = 18/36/46

Resin : Dry resin (lb)

Water: Water contained in resin + Free water (lb)

Cement: Portland cement (lb)

When the amount of water increases beyond the No. 1 line in Fig. 4, bleeding occurs, and when the amount of water decreases below the No. 2 line, the dischargeability from the mixer is lowered. An increase in the amount of resin lowers the compressive strength of centent-solidified products and causes disruption of the matrix in immersion tests. The No. 3 line indicates the condition under which cement-solidified products cured in water-saturated air for 30 days show a compressive strength of 700 psi. When the amount of resin is increased above the No. 4 line, cracks develop on the surface of cement-solidified products in 90-day immersion tests.

ECONOMICAL EVALUATION

This advanced cement solidification process eliminates the need for using expensive HIC liners because spont ion-exchange resins are stabilized by cement. A large reduc

tion of direct expenses such as container cost, disposal cost, etc., can therefore be achieved.









CONCLUSION

In the advanced cement solidification process developed by JGC, solidified products obtained are excellent in physical properties because spent ion-exchange resins are sufficiently pretreated before being cement-solid fied. Therefore, the products can satisfactorily meet the waste form criteria required by 10 CFR 61 and Technical Position. In addition, direct expenses can be 'argely reduced because there the use of expensive HIC liners is not required.

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INCINERATION CONFERENCE 1990

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Technology of Stabilization for Incinerator Ash Wastes

K. Yokoyama, K. Suzuki, F. Todo and Y. Moriya JGC CORPORATION, (Tokyo, Japan)

ABSTRACT

This report concerns the stabilization of incinerator ashes; advanced cement solidification and high frequency induction melting.

(1) AC Solidification

Most important points in this process are pretreatment of ashes by $Ca(OH)_2$ addition and use of high shearing mixer to give efficient homogeneity. The products contain 35wt% ashes, and have high compressive strength (200kg/cm²) and low leachability.

(2) HFIM

Ashes are melted at 1,200 - 1,600°C by the induced heating using a 25KHz high frequency, in a melter made of carbon material. The vitrified products have good chemical-physical properties. Furthermore, the process is capable of treating concrete, heat-insulator materials and other noncombustible DAW.

INTRODUCTION

The stabilization of incinerator ash wastes is classified into two main processes ; one is embedding in a binder matrix(1)(2), and the other is melting into hard blocks(3). Cement-, bitumen, plastic-solidification belong to the former process. These are easily operated by simple equipment, though both volume and weight of solidified products are increased. The latter is the melting treatment by method such as Joule-heat, high frequency induction, microwave and so sy. These processes have higher volume reduction but involve considerably more simplicated systems.

The application of these schniques to actual wastes should be reasonably selected with consideration given to several parameters ; waste-volume, -weight, -form, contamination levels of radicactive or/and hazardous wastes, required capacity and foreign materials in ashes.

In this paper, advanced cement (AC) solidification and high frequency induction melling (HFIM), developed by JGC, for radioactive ash wastes are reported.

ADVANCED CEMENT SOLIDIFICATION

I. Concept of AC solidification

AC solidification process for incinerator ashes consisted of two main procedures : pretreatment and homogeneously mixing by high shearing mixer. Figure 1 shows the basic flow diagram. Pretreatment is required to effectively prevent a cement-hydration reaction due to Zn. PD metal compounds. Also, this can avoid the deterioration of product-block strength due to H₂ gas generation reacting with A1 metal. The above mentioped metal compounds are generally contained in incinerator ashes. The process proceeds with the direct addition of Ca(OH)₂ and NaOH into ash-wastes is the presence of appropriate amount of water.

II. Properties of cement product

To obtain the fairly homogeneous mixture of cement the pretreated ash waste, a high shearing mixer, as shown in Figure 2, has been used at the mixing condition of 35wth ash-waste content and weight ratio of water to cemental:1. The pretreatment requires about one day.

The physical-chemical properties of products are summarized in Table 1. The density and compressive strength are range from 1.7 - 1.9 and 200 -400 kg/cm2, respectively. The incinerator ashes are effectively embedded in cement matrix by AC process.

Leaching index(LI) values, defined by ANS16.1 are obtained for radioactive Co and Cs, based on the leaching tests results over 90day. These LI values satisfy the requirement that LI>6.

- III. Features of the process system
 - The pretreatment of additive Ca(OH)2 to incinerator ashes is sufficiently effective to obtain stabilized cement products.
 - This process system has very simple equipment and a easy to operate. although the weight and volume of wastes finally increase.
 - The homogeneous mixture of cement and ash waste (35wt% content) is given by high shearing mixer.

HIGH FREQUENCY INDUCTION MELTING

1. Concept of HFIM

Figure 3 illustrates main system-components in the RFIM test pilot.

The noncombustible wastes, which are shredded or crushed, if necessary, are fed to the melting pot made of carbon material. The outside of this pot is surrounded with high frequency induction coils. The wastes are directly melted in the pot without any additives, and blown down into the capister to storage vitrified products. The off-gas is cleaned through the combination system of cyclone, elecrostatic precipitator and HEPA filter. The main wastes to be treated are the following: The main wastes to be treated are the following:

- Incinerator ash
- Heat-insulation material (containing asbestos)
- HEPA filter material
- Concrete debris

The melting temperature of these waste are ranges from 1200 to 1600°C, so that the wastes can be melted by HFIM method in either respective or mixed forms.

- II Feature of pilot plant

The mair features of the pilot plant to be used in experiments are shown below;

capacity	1	15 kg/h (based on incinerator ash)
melting pot	1	5 liter
electric power	1	4.2 KWH
cooling water	1	3m ³ /h
off-gas	1	12 - 13 Nm ³
canister	1	50 liter



Figure 4 to Figure 6 correspond to the photographs of induction heating melter main part, canister/holder, and off-gas treatment, respectively. The melted wastos are blown from the bottom of the carbon material pot, by controlling the carbon rod stopper setting in the pot. A sample of vitrified product is shown in Figure 7.

III. Characteristics of the process

(1) Melting pot material

In this experiment, tests were conducted on the applicability of carbon and silicon-ceramic materials to a melting pot in the HFIM process. As a results, the ceramic had severe damage due to the complicate reaction with a little amount of iron metal in melted wastes. On the other hand, although carbon material has easily reacted with oxygen and decreased the weight in high temperatures, the long operation time, 60 - 100 hours could be done under the nitrogen atmosphere. Then, carbon material was selected. The effective procedure will be expected by complete decxygen-pretreatment of wastes and shielding the inside of whole processes by nitrogen.

(2) Off-gas treatment

In order to confirm the performance of off-gas treatment system, the wastes doped radioactive materials, 60_{CO} , 134_{CS} were malted in the melter. Table 2 shows results of DF in the off-gas system. More than 10^4 is obtained as the total DF values.

(3) Physical-chemical properties of vitrified products

Table 3 indicates the volume reduction factors of noncombustible wastes treated by the pilot plant. High Volume reduction was achieved. Table 4 summarized the density, compressive scrength etc., of the vitrified products. The leachability of Fe and Na tested by the MCC-55 method (100°C, '/days) ranges from 10⁻³ to 10⁻⁶ g/cm² day, being almost same as that of general vitrified glass melted products.

IV. Features of the process system

- Various popcombustible wastes can be treated simultaneously

- Final products are vitrified form with excellent physical-chemical properties.
- High volume reduction can be obtained, for *xample, 5 9 for inciderator ashes.
- The melter is heated from outside usin; high frequency induction coils.
- The inventory volume of melted wastes in melter is small, thus
- allowing the maintenance of the melter to be readily carried out.

CONCLUSION

As a result of the experiments, it was confirmed that both processes of AC solidification and HFIM can be applied to treating incinerator ash wastes. The AC solidification process is suitable for treating large amounts of ash waste which is less contaminated by radioactive and/or hazardous materials. The HFIM process can be applied to treating small amounts of ash waste which is highly contaminated, in order to obtain stable vitrified products.

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Table 1 Characteristic of Ce	ment Products for Ash Wastes
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Waste	Content	Density	Compressive Strength	Leaching Index value
Incinerator	35 wt%	1.8	420 kg/cm2	60C0 14
Filtered ash	35 wt%	1.6	620 kg/cm ²	134Cs 8

Compressive Strength : at the curing time = 30 days Leaching Index : the required LI value > 6, at 90 days, in ANS 16.1





Figure 6 Off-gas Treatment System

Figure 7 Vitrified Product Block



Table 2 DF in Off-gas Treatment

Nuclide	Cyclone	Electric Pricipitator
6000	8 × 102	10
134Cs	2 × 102	6 × 102

Table 3 Volume Reductivity of Wastes

	Waste	VR Factor
	Ash	5~9
	Heat-insulator	9~53
	HEPA - filter	13 ~ 54
*	Concrete debris	1 ~ 3

Table 4 Characteristic of Vitrified Wastes

Waste	Density	Compressive Strength	WeightLoss	Leachin Fe	g Ratio* Na
Ash	2.7	>4000 kg/cm2	0.02 ~ 2wt%	10-4-10-6	10.2-10.5
Ash/Heat-insulator (50/50)	2.6	> 2000	0.2 - 2wt%	10-4~10-5	10-3-10-4
Ash/HEPA-Filter (50/50)	2.5	> 104	0.2 - 2wt%	10.3-10.6	10.3-10.4

^vg/cm²· day MCC-55 (100·C, 7 days)





PART 5

DESIGN AND CONSTRUCTION OF VIRGINIA POWER LOW LEVEL RADWASTE FACILITIES PROJECT



PROJECT OVERALL SCHEDULE



Memo ;

-



OUTLINE OF NRF BUILDING





The second



NRF MAJOR GOALS AND OBJECTIVES

- 1. On-site storage requirements to be one (1) year.
- Radioactive effluent discharges not to exceed
 0.1 Ci/year/site, and chemical discharges less than
 50% of NPDES limits.
- Waste shipped off site not to exceed 16,000 CF / year / site.
- 4. No inadvertent radioactive gaseous release.









PROJECT FEATURES - SCOPE OF WORK





PROJECT CONFIGURATION - SUBCONTRACTING



PLASTIC MODEL





3-D CAD MODEL



PROJECT ORGANIZATION AND OFFICE LOCATIONS









OWNER INVOLVEMENT

E	 PERIODIC MEETING DOCUMENT REVIEW MODEL CHECK
Ρ	VENDOR SELECTIONSHOP INSPECTION
с	 FIELD INSPECTION AUDIT, MONITORING PERFORMANCE TEST TRAINING

QUALITY ASSURANCE





SURRY CONSTRUCTION SITE





TECHNOLOGIES DISCUSSED AT PNC, TOKAI WORKS

- Vitrification of HLLW
- Nuclear Fuel Cycle and Reprocessing
 - Receiving and Storage of Spent Fuel
 - Mechanical Treatment
 - Dissolution, Clarification and Adjustment
 - Separation
 - U-Purification, Concentration, and Denigration
 - Pu-Purification, and Concentration
 - Analysis and Operational Testing Laboratory
 - Trea ment of Waste
- Microwave Melter
- ESR Furnace
- Assessment of Long Term Durability of Engineered Barrier Materials

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P. Marias





Reprocessing Plant in Japan

Since Japan has asside energy resources and depends for most of as energy on other countries, it's a very important problem how to secure energy resources. Our deare is to aubstitute nuclear energy for oil energy. To adhere this, an independent nuclear fuel cycle must be established so they the valuable fuel con be used effectively. The reprocessing plant for open tuel is the most important component of the multian tuel cycle since the nuclear fuel can not be used without it.

The reprocessing plant built by PNC at Tokan Mora in the fast step in the establishment of an independent mulear fast cycle.

Eperocenning Plants in the World

-	FRC	ł	india	Fame	U.K.	0.5.A	Nation
Tester	Karlanube	Mod	Transbey Tempone	Marcoule La Hagor	OrmCoray Wunducate	idahe Haniord Savaniah River West Valley Barnwell	Plant
D Yeldaw	351/y	1004/r for EWR research reactor fael, awaring permission to restant	30r/y vor netword U, military purpose 0.5s/day for EWR H9KR fuet	900-1 "X00k/y mulatasy purpose 8600/y. For natural wramum fuel 460/ly, for LMMR fuel	flox remeanch reactor fixed 3. SORity: Fox tractural unanhum fuest 400/r/y, fox 1.WB fixed, created operation	for restarch reactor had several tona/day, military purpose 300t/y, for LWR fuel; crawed operation 1.500t/y, for LWR, ebandoned	Capability and Status





Nuclear Fuel Cycle and Reprocessing

Nuclear Fuel Cycle



Development of the nuclear fuel cycle and PNC

PNC is developing most parts of the nuclear fuel cycle, from prospecting for uranium to reprocessing of spent fuel, as well as development of new reactors such as FBR and ATR. Reprocessing plant construction began in June, 1971, and after finishing blank test, chemical test, and hot test, operations began in January, 1981.

This plant can reprocess 0.7 ton-uranium of fuel per day and has a key role to promote the future reprocessing industry in Japan.



Establishment of the nuclear fuel cycle

In the operation of a reactor, uranium-235 content decreases and fission product (FP) content increases as the fuel burns. At the same time nonfissionable uranium-238 is converted to plutonium by neutron absorption. After about three years of fuel irradiation in the reactor the fissionable uranium-235 content is reduced and the FP content increased so that the reactor can not maintain criticality, and it is necessary to refuel the reactor.

A one million kWe LWR plant consumes about 30 tons of fuel per year on the average. Spent fuel contains about 1% uranium-235 (new fuel contains about 3%), about 95% uranium-238 (new fuel, 97%), about 1% plutonium, and $2 \sim 3\%$ FP. The reprocessing plant function is to recover the plutonium produced and the uranium remaining from the spent fuel and to separate and treat the FP as waste. Therefore, the nuclear fuel is not disposed of after only one use but is used many times as nuclear fuel because of the reprocessing.

The recycle of nuclear fuel in this way is called the nuclear fuel cycle and the reprocessing plant is the key to the nuclear fuel cycle.



ie of main plant

About 0.7 Metric ton of uranium per day it capacity nt fuel to be processed Enriched uranium fuel Cladding material Zin aloy or scuicles stael About 28/000 MM/D/t Average) About 35 MW/t Mvereje) Burn-up Specific power 180 days (Minimum) Cooling time Enrichment 4% (Aaximum) xe of process Purex process with chop and leach duct Uranium trioxide (UO3) Plutonium nitrate (Pu(NO3)4)

- Truck airlock
- Cask decontamination room 2
- 3 Fuel unloading pool
- 4 Fuel Storage pool
- 5 Mechanical treatment cell
- 67 Dissolver loading cell
- Feed adjustment cell
- 8 Separation cells
- 9 U-purification cell
- 10 Pu-purification cell
- 11 Utility room 12 Control room



3



mocessing process

maiving and Storage of Spent Fuel

e cask containing spent fuel is receiv-I the truck airlock. Then the cask is ed into the cask decontamination n, the water in the cask is exchanged. sc on. Then, the cask is moved into unloading pool and the spent fuel oved. The spent fuel assemblies are into fuel baskets, sent to a storage , and cooled there.

is cooled storage purpow is to await decay of relatively short-lifetime FP ents in the spent fuel assemblies. h decreases the activity and decay

After the fuel is cooled over a in period of time, the assemblies are to an intermediate pool, discharged the fuel basket one at a time, and to a mechanical treatment cell.

e storage pool of this plant can about 100 tons of spent fuel.



Storage pool





Reprocessing process Marchanical Treatment

The spent fuel assemblies sent to the mechanical treatment cell from the storage pool have the end parts of the fuel assemblies removed, and then the remainder is chopped into pieces about 5cm (3 inches) long. These fuel pieces are sent to a dissolver in the dissolver cell. Solid wastes, such as the end parts of the fuel assemblies which have been off first, and the hulls (fuel element clad) which don't dissolve in the dissolver, are put into a waste container and stored in a high active solid waste storage facility.

The mechanical treatment cell is surrounded by a concrete wall about 150cm thick. Viewing windows and manipulators are set up for remote operation of the chopping work. A decontamination cell and airlock cell are also set up for and maintenance of the machinery.







Chopping machine



Hulls

processing process

ssolution, Clarification and Adjustment

ent fuel elements chopped by the hanical process are transferred to the sliver in the dissolver cell. Of the ded fuel pieces, only oxide fuel is blived in nitric acid and the cladding e of stainless steel or zircaloy reis undissolved.

the dissolution, oxygen is introduced te dissolver, which oxidizes the nitrooxides, which originates in the dision.

er dissolution, dissolved solution is iferred to the buffer vessel, diluted adjusted with nitric acid. Undissolved rials such as solid particles are read by a pulse filter and the acidity is solution is adjusted in the adjust-

vessel of the feed adjustment cell. I this vessel, after accounting for the inium and uranium, this solution is ferred to the 1st extraction bank in 1st so iration cycle. Undissolved s'' are rinsed, removed with a pered basket, put into a waste container sent to the high active solid waste ge facility. Off gas originating in the lution is sent to the off gas treatment 'ss.











Reprocessing process



Dissolved solution contains U. Pu and FP. The purpose of the separation process is to separate U. Pu and FP from each other in the solution. This process can be roughly classified as the 1st separation cycle (co-decontamination), and the 2nd separation cycle (partition). The 1st separation cycle is the process to separate fission products from dissolved solution and the 2nd cycle is to separate Pu from U.

1st separation cycle (co-decontamination)

This process has the function to extract U, Pu in the solution into solvent and to separate them from FP. U and Pu extracted into solvent at mixer-settler extraction 1, are turned again into the squeous phase by stripping the solution at the settler extraction 2 and they are the insferred to the 2nd cycle. On the other, side, FP remaining in the aqueous phase at mixer-settler extraction 1 are vashed at the diluent washer and transierred to the high active waste evaporator via the buffer vessel.

() 2nd separation cycle (partition)

At the extraction 2 of the 1st separation cycle, U and Pu are stripped into the aqueous phase and this aqueous phase is introduced to the extraction 3. And there, U and Pu are extracted is the organic phase to separate them from FP



Extraction maintenance area

remaining in the aqueous phase by the same process as the extraction 1. Of U, Pu in the organic phase, Pu is reduced and stripped by the mixture of uranous nitrate and hydrazine at the extraction 4, and resulting in separation of uranium in the organic phase from Pu in the aqueous phase. After this, the aqueous phase containing Pu is transferred to the Pu purification process. The organic phase containing U is transferred to the corraction 5 and there U is stripted in 6 the aqueous phase and transferred to the U purification process.



processing process

purification, -concentration and Denitration

he solution containing U that is strippinto the aqueous phase at the extrac-1.5 of the 2nd separation cycle, is isferred to the extraction 6 of the U dication cycle. There, U is separated in the trace of FP by the solvent as in extraction 1 and they are purified. parated U is transferred to the extrac-1.7, stripped again into the aqueous se by dilute acid and then transferred the U concentration-denitration pro-

containing solution (uranyl nitrate) isferred to the U concentration-deation is concentrated in the 1st evapor. This concentrate, keeping a cont concentration, is removed from the porator and after cooling it is transed to the dilute vessel via the contrate receiver. At this vessel, inspecis done and if within specification it ansferred to the feed vessel.

e concentrate transferred to this feed el is re-concentrated at the 2nd evapor, transferred to the denitrator via buffer vessel and there it is decomid into UO_3 powder.

D₃ powder is continuously removed a the bottom of the denitrator, packnto the container as final product and to UO₃ storage.



U-purification cell



Uranium nitrate and uraium trioxide

inciple of Solvent Extraction Method d Mixer-settler

the reprocessing plant, the main technique covering U and Po from spent fuel is the ni extraction method.

a method has the function of extracting al elements from the nitric acid solution ting from dissolving spent fuels, conversion organic solvent or stripping elements from irganic phase into the aqueous phase and ation of U and Pu from FP.

example, at the 1st separation cycle that he function of separating U, Pu from FP. d Pu is extracted into the organic phase see the higher acidity makes it them easy to we in the organic phase. Under this cont, FP remain in the aqueous phase because solubility.

i FP are separated from U and Pu. And U, the organic phase are able to be stripped into the aqueous phase because the lower acidity makes it easier to dissolve in the aqueous phase than in the organic phase. Furthermore, tetravalent Pu in the organic

Furthermore, tetravalent Pu in the organic phase is reduced to trivalent Pu by a reduction reagent. Trivalent Pu is so insoluble in the organic phase and is stripped into the aqueous phase. So Pu is separated from U remaining in the organic phase. Thus, repeating the extraction, stripping and Pu stripping make it possible to separate U, Pu and FP from each other.

• The mixer-settler consists of the mixing part and the settling part.

mixing part : contacting the organic solvent with acid solut.

settling part : dividing the aqueous phase For example, at the extraction 1, the extracion operation proceeds as in the figure

tion operation proceeds as in the figure. Real mixer-settlers differ in size from each other, but a model is shown in the photo. Solvent Weir P Mixing Settler section FP + S

Reprocessing process

Pupurification and -concentration

Pu solution that is stripped at the extraction 4 of the 2nd separation cycle is transferred to the extraction 8 of the Pu purification cycle and there Pu is separated from a small amount of FP by solvent extraction and purified.

The aqueous solution containing a small amount of FP is transferred to the medium level liquid waste treatment process as waste. The organic phase containing Pu that is separated at the extraction 8, is stripped to the aqueous phase by the mixture of uranous nitrate and hydrazine at the extraction 9 and they are transferred to the evaporator in the Pu concentration cell.

The concentrate, plutonium nitrate, is transferred to the storage tank as a final product.



Plutonium nitrate



Pu-purification cell





Model of mixer settler
analytical laboratory and OTL are central building which is the largest ig next to the main plant.

role of the analytical laboratory is ovide the necessary information the process and the material balance ning uranium and plutonium. Samken by means of sampling vehicles ansferred through the pneumatic is system inside the plant. These is are at different activity levels, so are analysis cells for high and m activity samples and glove boxes v activity samples, all with modern nents.

is a miniature of the main plant, on O scale, of the chemical treatment, ing the fission product concentrais a tium denitration, plutonium itration and acid recovery.

role of the OTL is to reproduce the operation, review the problems and ositive proof of operation before ng the process.

ples of irradiated fuel are transfrom the plant to the laboratory in ad cask "Cendrillon".





Operatinal area of OTL

Atomic absorption spectrometer



Sectace ionization mass spectrometer



Transfer of spant fuel to OTL





Tuestment of Waste



In the plant, nuclear material such as uranium and plutonium is recovered, but also, radioactive waste must be treated safely. Airborne radioactivity from mechanical treatment and dissolution processes are washed and filtered several times. Radioactive liquid wastes are reduced in volume by evaporation and stor ed in a stainless steel vessel. Condensate removed from the evaporator is closely monitored for radioactivity and the concentration of radioa, tivity is confirmed to be within acceptable limits before release to the sea.

Gaseous waste

st of the radioactive gaseous waste from the dissolution process and nechanical treatment process, with a amount from various cell and l ventilation systems. These gaseous is are assayed and filtered several according to their activity level and released to the atmosphere after

ul activity monitoring.

Liquid waste

th active level liquid waste (HALW) is ly the aqueous raffinate from the 1st ration cycle, which contains nearly the nonvolatile fission products. W is reduced in volume by evaporaand stored in underground tanks.

dium active level liquid waste LW) is the raffinate from the 2nd ration cycle, uranium purification e and plutonium purification cycle the nitric acid solution recovered absorber of the uranium denitration ess and HALW concentration process. W is treated by the acid recovery ess. The distillate is reused in the ess as recovered nitric acid and the entrate in the evaporator bottom is sferred to the HALW evaporator. active level liquid waste (LALW)

n other processes and facilities is sferred to the auxiliary active facility F) and stored temporarily according ictivity level. The comparative high ve liquid waste such as the raffinate n the solvent regeneration cycle is iced in volume by evaporation and its densate is transferred to the buffer is and the other concentrate is stored essels.

te other liquid waste such as laundry te is treated by flocculation, and the other water is transferred to buffer ks and the sludge is stored in vessels.

reduce the sea-discharge activities, itionally two evaporator stations for LW (E and Z facility) were constructed is start of hot operation. For the rained TBP, the de-oiling station (C lity) was installed before sea-discharge nitoring vessels.

he water is carefully monitored and ifined, and the concentration of radioivity is determined to be within accepte limits before release to the sea.

Solid waste

ligh active level solid waste (HASW) hsists of the end plugs of spent fuel emblies sheared in the mechanical atment process and insoluble hulls in the dissolving process. HASW is ntained in stainless steel drums and ired in HASW storage.

ow active level solid waste is sorted as mbustible waste or non-compressible ste, which is either incinerated, comcted or placed in drums.

These treated wastes are stored in the NSW storage area.













Treatement of LASW



bety Desigh and Safety Administration

The reprocessing plant must be designed and administered considering first the mfety of the facility, process design and influence on the circumstances.

(1) Confinement

The main parts of plant have triple confinement barriers where the primary confinement barrier is the building, the secondary is the interior wall and the third is the reaction vessel. If leakage of a vessel containing radioactive material occurs, the plant is designed to restrict radioactive release to the environment.

(2) Aseismic design

Aseismic design is based on the basic conditions that the buildings, structures. equipment and piping have sufficient strength to assure safety of the employees and public of the nearby area during and after earthquakes.

(3) Criticality control

The following basic control parameters asidered.

uipment geometry limitation resionable material mass limitation · concentration limitation

The nuclear safety of the system is maintained by exercising control over one or more of the above parameters and using neutron poison supplementally.

(4) Maintenance

The mechanical cell equipments treating the high active material are maintained remotely and are able to be remotely decontaminated prior to repai. The equipment in the chemical process is all to be repaired by direct maintenance after decontamination. The key cells and equipment such as HALW concentration process are installed in parallel.

In addition to the above safety design. the health physics of employees is controlled by a PNC system. PNC establishes the monitoring station, ground monitoring posts, sea and coast monitoring, monitoring cars and one monitoring boat.



Environmental Monitoring System at Tokai





Monitoring post





Present Status of R & D Activities on HLLW and TRU Waste Conditioning in Tokai Works

November 1990

POWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION(PNC)



Major RD and D Activities on HLLW and TRU Waste Conditioning in Tokai Works.

R & D phase

- HLLW : Vitrification by LFCM Process
- TRU Wastes: Nuclide Separation from Low Level Liquid Waste

Decomposition and Nuclide Separation from Spent Solvent

Demonstration phase

- PWTF : Pu Contaminated Waste Treatment Facility
- AspF : Bituminization Demonstration Facility
- STF : Solvent Waste Treatment Facility







HLLW Conditioning Technology

C

Vitrification Technology Development

Process Development





Cold Test (ETF, MTF, EDF-III) Hot Labo. Test (CPF)









- Two - armed manipulator · Rack system Ventilation System, ISI, Instrumentations











TRU Waste Conditioning Technology



Low Level Liquid Waste Treatment Process









Present Status of Waste Treatment Facilities in Tokai Works

November 1990

POWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION(PNC)











Major TRU Waste Treatment Facility in Tokai Works

Demonstration phase

- PWTF: Pu Contaminated Waste Treatment Facility
- AspF : Bituminization Demonstration Facility
- STF : Solvent Waste Treatment Facility

Design phase

LWTF : Low Level Radioactive Waste Treatment Facility HWTF : Hull Waste Treatment Facility

Pu-Contaminated Waste Treatment Facility (PWTF)

Objectives

- 1. Demonstration of the volume reduction and conditioning processes developed by PNC
- 2. Characterization of the conditioned waste











Sep. 1990



× simulated waste

w Level Radioactive Waste Treatment Facility (LWTF)

Objective

- (1) Volume reduction and conditioning of low level solid wastes from Tokai Reprocessing Plant
- (2) Demonstration of the nuclide separation processes
- (3) Characterization of the conditioned wastes









Reduction Ratio of Waste (design)

	Combustible Incineration	CI Contained Materials	Nuclide
Waste [kg / year]	30,600	19,800	912,000
Product \sh, sludge) [kg/yeer]	2,780	2,480	45,600
Weight	. 11	8	20 •

	High Pressured Compaction	Metal Melting
Vyaste [m³/year]	. 342	: 116
Product (m ³ /year)	69	17
Volume reduction	5	7

0



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Present Status of Other Activities on Nuclear Fuel Cycle in Tokai Works

November 1990

POWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION(PNC)







Fig. Sampling Points for Surface Water and Sediment

Table-1. Analysis and Measurement Methods and the Detection Limits of Radionuclides in the Terrestrial Environmental Monitoring Program at PNC Tokai Works.

Sample	Nuclide	Analysis and Measurement Method	Detection Limit	
Airborne Gross Alpha		Direct alpha counting of filter paper	0.02	mBq/m3
Particulates	Gross Beta	Direct beta counting of filter paper	0.7	mBq/m3
, untreduced	Sr-90	Radiochemical analysis and beta counting	0.004	mBq/m3
	Cs-137	Gamma spectrometry	0.007	mBq/m3
	Pu-239,240	Radiochemical analysis and alpha spectrometry	0.0001	mBq/m3
Airborne Iodine	I-131	Gamma spectrometry	0.2	mBq/m3
Airborne Rare Gas	Kr-85	Continuous measurement of air	7	kBq/m3
Rain Water	H-3	Liquid scintillation spectrometry	2	Bq/l
Fallout	Gross Beta	Beta counting	4	Bq/m2
Drinking	Gross Beta	Evaporation and beta counting	0.04	Bq/l
Water	H-3	Liquid scintillation spectrometry	2	Bq/I
Leafy	I-131	Gamma spectrometry of chopped samples	0.2	Bq/kg fr sh
Vegetables	Sr-90	Radiochemical analysis and beta counting	0.04	Bq/kg fresh
	Cs-137	Gamma spectrometry of chopped samples	0.08	Bq/kg fresh
	Pu-239,240	Radiochemical analysis and alpha spectrometry	0.00008	Bq/kg fresh
Polished Rice	Sr-90	Radiochemical analysis and beta counting	0.04	Bq/kg fresh
Surface	Sr-90	Radiochemical analysis and beta counting	0.08	Bq/kg dry
Soil	Cs-137	Gamma spectrometry of dried samples	0.8	Bq/kg dry
	Pu-239,240	Radiochemical analysis and alpha spectrometry	0.04	Bq/kg dry
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Cample	Nuclide	Analysis and Measurement Method	Detecti	on Limit
Airborne	Gross Alpha	Direct alpha counting of filter paper	0.02	mBq/m3
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Airborne	1-131	Gamma spectrometry	0.2	mBq/m3
Airborne Rare Gas	Kr-85	Continuous measurement of air	7	kBq/m3
Rain Water	H-3	Liquid scintillation spectrometry	2	Bq/I
Failout	Gross Beta	Beta counting	4	Bq/m2
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	Cs-137	Gamma spectrometry of dried samples	0.8	Bq/kg dry
	Pu-239,240	Radiochemical analysis and alpha spectrometry	0.04	Bq/kg dry



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Present Status of Other Activities on Nuclear Fuel Cycle in Tokai Works

November 1990

POWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION(PNC)





Reduction Ratio of Waste (design)

	Combustible Incineration	CI Contained Materials	Nuclide Separation
Waste st	30,600	19,800	912,000
Product sh, sludge)	2,780	2,480	45,600
	. 11	8	20 •

	Pressured Compaction	Metal Melting
Waste [m³/year]	* . 342	116
Product (m ³ /year)	69	17
Volume reduction	5 States 19	7







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w Level Radioactive Waste Treatment Facility (LWTF)

Objective

- (1) Volume reduction and conditioning of low level solid wastes from Tokai Reprocessing Plant
- (2) Demonstration of the nuclide separation processes
 - (3) Characterization of the conditioned wastes
Weight Reduction Ratio of the Wastes (PWTF)

Sep. 1990



% simulated waste













²u-Contaminated Waste Treatment Facility (PWTF)

Objectives

- 1. Demonstration of the volume reduction and conditioning processes developed by PNC
 - 2. Characterization of the conditioned waste

Major TRU Waste Treatment Facility in Tokai Works

Demonstration phase

- PWTF: Pu-Contaminated Waste Treatment Facility
- AspF : Bituminization Demonstration Facility
- STF : Solvent Waste Treatment Facility

Design phase

LWTF : Low Level Radioactive Waste Treatment Facility

HWTF: Hull Waste Treatment Facility











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Present Status of Waste Treatment Facilities in Tokai Works

November 1990

POWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION(PNC)





Spent Solvent Oxidative Decomposition Process







Low Level Liquid Waste Treatment Process



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9.



Vitrification Technology Development

Process Development





HLLW Conditioning Technology





R & D phase

- HLLW : Vitrification by LFCM Process
- TRU Wastes: Nuclide Separation from Low Level Liquid Waste

Decomposition and Nuclide Separation from Spent Solvent

Demonstration phase

- PWTF : Pu Contaminated Waste Treatment Facility
- AspF : Bituminization Demonstration Facility
- STF : Solvent Waste Treatment Facility

Present Status of R & D Activities on HLLW and TRU Waste Conditioning in Tokai Works

November 1990

POWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION(PNC)





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CEC 4th Natural Analogue working Group Meeting

Pitlochry, Scotland, 17-22 June 1990

(CEC EUR REPORT)

SOME ASPECTS OF NATURAL ANALOGUE STUDIES FOR ASSESSMENT

OF LONG-TERM DURABILITY OF ENGINEERED BARRIER MATERIALS

- RECENT ACTIVITIES AT PNC, JAPAN -

(YUSA, Y., KAMEI, G. and ARAI, T.)

5. Framework of our analogue studies: Our natural analogue programme has three components: 1) investigation of alteration phenomena, 2) examination of environmental conditions, 3) support experiments. The support experiments are an essential part of our study in order to enhance the wider applicability of the natural analogue.

1. INTRODUCTION

1.1 COMPONENTS OF ENGINEERED BARRIERS

The Components, candidate materials, and functions of various types of engineered barriers are as follows:

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Components	Candidate materials	Function expected
· Vitrified waste	· Borosilicate glass	· Restricts release*
• Overpack	• Carbon steel or Cast iron	 Retards water penetration * Provides favourable chemistry *
• Buffer materials	• Bentonite	 Restricts water penetration * Delays commencement of release* Restricts radionuclides release *
• Backfill materials	• Concrete (Cement)	 Minimizes water access to package f Alters groundwater chemistry f Retards solute transport f
		* :NAGRA [1985], † :Chapman et al [1987]

As a part of the study on engineered barrier materials and systems for geological disposal of radioactive waste in Japan, analogue studies for the assessment of long-term durability of engineered barrier materials are conducted at PKC Tokai Works. This paper describes the state-of-the-art on the studies, specifying their main purposes and framework, and demonstrating our emphasis on natural materials. The results obtained to date will be summarized. Some parts of studies on natural glass and bentonite were already presented at the MRS Symposium (Arai et al., 1988; and Kamei et al., 1989), although revised and expanded data are shown here.

1.2 DEFINITION OF "NATURAL ANALOGUE"

One of the most critical aspects in the evaluation of the durability of candidate materials for engineered barriers is the extrapolation of the results of short-term experiments over a long time scale. Natural analogues currently provide the only means by which such extrapolated long-term behaviour can be confirmed.

SOME ASPECTS OF NATURAL ANALOGUE STUDIES FOR ASSESSMENT OF LONG-TERM DURABILITY OF ENGINEERED BARRIER MATERIALS -- RECENT ACTIVITIES AT PNC TCKAI, JAPAN ---

YUSA,Y., KAMEI,G. and ARAI,T. Geological Isolation Technology Section, Tokai Works Power Reactor and Nuclear Fuel Development Corporation 319-11 Yokai Ibaraki JAPAN

SUMMARY

This paper contains an overview of analogue studies for the assessment of longterm durability of engineered barrier materials at PNC Tokai.

Materials of young age and with simple history are the most suitable for study as: 1) properties of the materials tend to deteriorate over longer historical time intervals; and 2) detailed quantitative data on time intervals and environmental conditions are more likely to be available. The following materials and their alteration phenomena were selected: 1) weathering alteration of basaltic glass (as vitrified waste form), 2) corrosion of iron in soil (as overpack), 3) illitization of smectite associated with contact metamorphism (as buffer material), 4) alteration of cement (as buffer or backfill material).

1. Weathering alteration of basaltic glass: Basaltic glasses, from the Fuji and the Izu-Ohshima pyroclastic fall deposits were studied. The observations were made: 1) Climatological conditions have not varied significantly during the last three thousand years. Therefore, values for temperature, amount, and chemistry of ground water are quantified. 2) The cases studied could be regarded as leaching experiments in groundwater, using mass balances in waterglass interaction. 3) Although the groundwater is of Ca(Mg)-HCO₃ type in the Fuji area and of Na-Cl type in the Izu-Ohshima, similar alteration ratios (2 \sim 3μ m/1000yr) were obtained.

2. <u>Corrosion of iron in soil</u>: Industrial materials, such as gas/water service pipes of carbon steel or cast iron embedded in soil for 20 \sim 110 years, were selected for an analogue study of corrosion of iron in bentonite. The maximum corrosion rates obtained so far fall in the range of 0.04 \sim 0.09 mm/yr.

3. Illitization of smectite associated with contact metamorphism: In the Murakami bentonite deposit in central Japan, lateral variation of smectite to smectite/illite mixed-layer minerals are found in the aureole of the rhyolite intrusion body. Conversion of smectite to the mixed-layer mineral composed of 40% illite was found to have occurred in a period of 2.4 Ma over a temperature range of above 240 (\pm 50) °C to 105 °C.

4. <u>Alteration of cement</u>: Concrete components of fabrications, such as estuary walls, with a known age were studied. Chemical alteration of the cement were detected to a depth of few centimeters by EPMA, SEM, TEM and XRD.





1.4 SELECTION OF SUBJECTS FOR THE STUDIES

Cases of younger age and simple process, therefore, are regarded as more suitable subjects for the studies, as quantitative data on time and environmental conditions are probably available. Many previous analogue studies consisted of descriptions of the results of natural experiments without incorporating data on well-defined environmental conditions.

We selected subjects for the analogue studies according to the following criteria: 1) analogy of materials with candidates, 2) analogy of environmental conditions with simulated repository conditions, 3) simplicity and availability on environmental conditions, and 4) availability of chronological data. Table 2 shows the subjects of our analogue studies on engineered barriered materials.

Engineered	Candidate	Assummed phenomena in	Analogue Phenomena in
Barriers	Material	Repository Conditions	Analogous Conditions
1. Waste Form	Boro- silicate Glass	Leaching of Waste Borosilicate Glass with Groundwater	Weathering Alteration of Basaltic Glass with Goundwater
2. Overpack	Carbon	Corrosion of carbon	Corrosion of Iron
	Steel	steel in Bentonite	in Soil
3. Buffer	Compacted	Illitization of	Illitization with
Materials	Bentonite	Smectite in Bentonite	Contact Metamorphism
4. Backfill	Concrete	Alteration of cement	Alteration of Cement
Materials	(Cement)	with Groundwater	with Groundwater

Table 2 The subjects of our analogue studies on engineered barrier materials

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The term "natural analogue" can be defined as "natural phenomena which resemble those assumed in geological disposal scenarios". The selection of an appropriate natural analogue is the key issue which will determine whether the natural analogue study will be successful.

1.3 PROPERTIES OF NATURAL ANALOGUE

First, consideration is given to the properties of the natural analogues. In order to extrapolate the results of short-term experiments to the long-term, it is desirable that the natural phenomena can be individually and quantitatively described in terms of three constituents: 1) starting materials, 2) environmental conditions (including time scale), and 3) results. These are the three normal constituents of all "experiments".

However, there are some intrinsic difficulties in regarding such phenomena as experiments. Most naturally occurring materials, from which an relevant analogue must be selected, have complicated histories resulting the overprint of different processes, as shown in Table 1.

	Laboratory Experiments	Natural Phenomena
(1) Materials	Candidates	Analogue
() Number	Many	Solitary, few
(2) Environmental (Experimental) Condition	Simple, Uniform Constant, Controlled Common Small Scale	Complicated Variable Individual Large Scale
(3) Period	Short-term	Long-term
(4) Results	Independent variables Discrimination among conditions is possible	Overprinting of actors Restoration is difficult

Table 1 Comparison between laboratory experiments and natural phenomena

Secondly, geological and historical records are often incomplete, and errors in the determination of time scale and environmental conditions are not small. Although such disadvantages differ case to case and sample to sample, as materials age, their histories generally become more complicated; the factors with which alteration phenomena were related become overlapped, and quality and quantity of available data decreases. Thus, estimation from present observations becomes virtually impossible with very old samples (Figure 1).



Sample				Oxi	de (I	rt%)			
o unip i o	S102	TiO2	A1202	Fe:03*	MgO	CaO	Na ₂ O	K20	Total
Fuji HS ZS	52.9 50.7	2.3 1.6	12.1 15.0	16.5 13.8	3.9 4.9	8.5 8.8	1.8 2.9	1.2	99.2 98.6
Izu-Ohs N1 N4	hima 53.7 53.0	1.4	13.0 13.1	15.6 15.3	3.4	8.8 8.4	2.3	0.5	96.9 98.6

Table 3 Chemical compositions of glasses.

*: Total Fe as Fe203

Alteration layer

The alteration layer is optically isotropic and X-ray amorphous. The surface form of the alteration layer of the HS is grainy and that of ZS, N1, and N4 is flaky.

Despite the difference in morphology, the chemical composition of the alteration layer of the Houei Scoria is similar to that of the Zunazawa Scoria. The morphology of the alteration layers of both scoria is strikingly similar to that observed on the surface of experimentally altered borosilicate glasses (Hirose, unpublished data). Alteration layer thicknesses are summarized in Table 5. The elemental concentrations in the alteration layer are characterized by greater depletion of Mg, Ca, Na, and K, as compared to Si, Al, Fe, and Ti.

Table 4. Chemistry of pore water, spring water and rainwater.

		Chemical composition: (mg/l)				- H	Eb.					
Samp	e	Na	ĸ	Ca	Mg	Fe	HCO3	SO.	Cl	SiO,	pn	(mV)
Fuji												
HS F	.W.	4.4	1.7	5.3	1.3	87100A	24	6.5	4.1	34	-	-
ZS F	.W.	8.4	3.3	4.6	1.3	5.0	35	6.4	4.3	218	-	-
5	5.W.	5.0	1.4	8.9	6.9	2.9	67	4.4	2.6	41	7.0	178
Izu-(hshima											
N1 F	.W.	78	2.9	27	12	3.2	7.6	27	176	52	(6.0)	-
NA F	.W.	86	3.7	40	15	4.0	5.1	26	222	48	(6.0)	- 1
Rain	water*	1.1	0.3	0.4	1.0	0.2	. inclusion	1.5	1.1	0.8	-	-

Water chemistry

2. Weathering alteration of basaltic glass

2.1 SCOPE OF STUDIES

Many analogue studies of the alteration of natural glasses indicate that the alteration rates at low temperatures of natural glasses vary from $0.001 \,\mu$ m/1000 yr to 30 $\,\mu$ m/1000 yr (Hekinian et al. 1975; Bryan et al. 1977; Allen 1982; Lutze et al. 1985 & 1987; Grambow et al. 1986; Ewing et al. 1987; Jercinovic et al. 1988). This variation is interpreted as the result of variations in environmental conditions. However, few detailed studies on environmental conditions have been reported.

Described below are the effects of alteration by weathering of basaltic glasses with well established environmental conditions and ages. The alteration is a long-term leach test carried out by nature with rainwater as the leachant and groundwater as the leachate. The young-aged (280 \sim 2800 years) samples were selected to investigate environmental conditions during alteration based on present meteorological data.

Samples

Volcanic glasses constituting scoria of pyroclastic fall deposits were studied. Scoria samples were collected at the foot of the Fuji and Izu-Ohshima volcanoes, on both of which the stratigraphy and chronology of pyroclastic fall deposits have been studied in detail.

The samples collected were Housi Scoria (HS, 280 years ago) and Zunazawa Scoria (ZS, 2800 years) from the Fuji, and N1 (880 years) and N4 Scoria (1240 years) from the Izu-Ohshima volcano.

All of the scoria samples contained pore water, and spring water was found about 2.5 m below the Zunazawa Scoria bed.

Methods

Glass compositions were determined by Electron Probe Microanalyser (EPMA). Alteration layers were studied by optical microscope, EPMA and Scanning Electron Microprobe (SEM). The thickness of alteration layers was measured from SEM photos of the sections oriented nearly perpendicular to the layers.

In the field, the pH and Eh of the spring water were measured by portable meters. The spring water was filtrated through a $0.45\,\mu$ m filter and the filtrate was analyzed by absorption spectrophotometry, flame spectrometry and atomic absorption spectrometry.

2.2 RESULTS

The chemical compositions of the glasses are shown in Table 3. These are within the range of basalts.

The relation between alteration layer thickness and age is shown in Figure 2. The two kinds of alteration rates, the forward rate of alteration (3 \sim 20 μ m/1000 yr, under silica-unsaturated conditions) and the final rate of alteration (0.1 μ m/1000yr, under silica-saturated conditions) by Grambow et al.(1985), are also shown in Figure 2. The alteration rates estimated in this study are near or below the forward rate of alteration.

Mass Balance Between Alteration Layer and Spring Water

Spring water can be regarded as the leachate. In order to discuss the leaching behaviour of glass, it is necessary to clarify the relation between the elemental concentration in the leachate and the elemental loss from the alteration layer. Elemental concentrations in groundwater have previously examined (Arai et al., 1989) and the results indicate that the calculated composition of groundwater is in fair agreement with the composition of spring water (Figure 3). The discrepancies in the concentrations of Fe and SiO₂ can be explained by the precipitation of iron hydroxides and silica gel respectively among scoria grains.

	F	UJI	IZU - OHSHIMA		
(1) MATERIALS STUDIED	HP	HS	ZS	N1	N4
(2) GLASS COMPOSITION [Si02wt96]	64	53	51	54	53
(3) ENVIRONMENTAL CONDITIONS ① TEMPERATURE (°C) ② WATER CHEMISTRY ③ WATER SUPPLY RATE (2/cm²/yr)	Ca(Mg)	14 - HCO, 0.20	type	15 Na - (0	5 C1 type .21
(4) PERIOD (yr)	280	280	2800	880	1240
 (5) RESULTS ① ALTERATION RATE (μm/1000yr) [Alteration Layer Thickness:μm] 	<0.2 <0.05	1.6 0.44	3.1 8.8	1.7 1.5	1.8 2.2
 ALTERATION PRODUCTS Amorphous Materials Goethite Smectite 	N.D. N.D. N.D.	00×	000	000	000

Table 5 Summary on alteration behaviour of volcanic glasses and their environmental conditions

N.D. : Not Determined, O : Present , X : Absent

2.4 CONCLUSION

The chemistry of the pore and spring water is listed in Table 4, together with the average of rainwater in Japan (Sugawara, 1968). The elemental concentrations in ZS pore water are higher than those in HS pore water. This implies that elemental concentration in groundwater increases with depth.

2.3 DISCUSSION

Environmental Conditions

Analyses of paleo-sea level variations (Sugimura, 1977) and paleo-climatological data (Yamamoto, 1980; and Maejima, 1984), indicate that the climatic conditions in Japan have not varied significantly for the last 2800 years. Therefore, the temperature and the water supply rate are estimated from meteorological data such as mean annual temperature, annual rainfall, and evapotranspiration. The samples were situated in the unsaturated zone; accordingly, percolating meteoric water is the only source of pore water. The pore water flows downward in the deposits and dissolves the components of scoria. This natural phenomenon can be regarded as a leach test being constantly renewed fresh rainwater.

Alteration Rate

In natural alteration systems, it is generally difficult to know the exposure age of a sample, that is the time that the glass has actually been in contact with water (Jercinovic et al., 1988). The exposure ages of the samples in this study are equivalent to the samples ages as their surfaces were always in wet conditions and were always in contact with renewed pore water.





Fig. 2 The relation between age of samples and thickness of alteration layer

Fig. 3 The comparison between elemental concentrations of spring water and calculated groundwater









Fig. 5 Maximum corrosion depth as a function of time intervals

In conclusion, the maximum corrosion rates of cast iron and carbon steel embedded in soils were estimated in the range of $0.04 \sim 0.09$ mm/yr. Corrosion of industrials materials in soil is a useful analogue and further studies are planned.

 MATERIALS STUDIED Site Sample Material 	Yokohama	Nagasaki	Tokyo	Tokyo
	Gas S.P.	Water S.P.	Water S.P.	Watar S.P.
	Cast Iron	Carbon Steel	Cast Iron	Cast Iron
(2) ENVIRONMENTAL	Sandy Clay	Gravel with	Cohesive	Cohesive
CONDITIONS		Org. Comp.	Soil	Soil
(3) PERIOD (yr)	110	50	56	20
(4) RESULTS ① CORROSION RATE (Uniform Corrosion Pitting Corrosion	пип/ут) 0.03 0.08	0.01 0.09	N.D. 0.04	N.D. 0.06
© CORROSION	FeCO3	Not	FeCO3,	FeCO₃,
PRODUCTS		identified	a-FeO(OH)	α-FeO(OH)

Table 6	Corrosion	behaviour of	firon	in soil
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S.P. = Service Pipe; Org. Comp. = Organic Compounds

1) It was possible to determine the alteration behaviour of volcanic glasses from three experimental constituents: 1) starting materials, 2) environmental conditions and time scale, and 3) results (Table 5).

2) Calculation of the mass balance between the elements depleted from the glasses and the chemical composition of groundwater permitted us to regard the cases studied as experiments in the leaching of glasses by groundwater.

3) The natural alteration products of the volcanic glasses were very similar to those of laboratory experiments with simulated waste glasses.

4) Although the ground water is Ca(Mg)-HCO₃ type in the Fuji area, and of Na-Cl type in the Izu-Ohshima area, similar alteration rates ($2 \sim 3 \mu m/1000 \text{ yr}$) were obtained.

3. CORROSION OF IRON IN SOIL

Industrial materials such as water service pipes, were studied for the following reasons: 1) iron or steel is one of the candidate materials for waste package, 2) soil environment is probably similar to the bentonite fill environment, 3) samplavailability, and 4) chronology and environmental data are fairly assessable in ccomparison to those of archeological artifacts. One of the purposes of this analogue study is to validate whether the results of corrosion rates and models derived from the results of laboratory experiments can be extrapolated to a few tens of years (Figure 4).





The samples studied were gas or water service pipes, composed of cast iron or carbon steel. The soil or clay adjacent to the pipe was examined in order to avoid the influence of the macro-cell effect. The corrosivities of the soil environment at each site were estimated as not being very severe from the viewpoint of both electrochemical and chemical characteristics of the soil. Corrosion rates were derived from the measurements of the thickness of the pipe, and chemical composition of the material were determined. The corrosion products





4. ILLITIZATION WITH CONTACT METAMORPHISM

4.1 SCOPE OF STUDIES

The research on illitization of smectite in the natural environment affords indispensable information on the long-term durability of bentonite.

Geological processes associated with smectite-illite conversion can be classified as follows:

Diagenesis, 2) Regional metamorphism, 3) Contact(or thermal) metamorphism,
 Hydrothermal alteration

Among theses, contact metamorphism has been selected as being a suitable analogue because of the prevailing temperature and the water/rock ratio. Furthermore, a study of contact metamorphism has potentiality to give clear-cut data on the reaction term and the thermal conditions of illitization of smectite, provided that: 1) the bentonite bed is distributed, and 2) simple history and simple geology can be recognized.

One such case of contact metamorphism is the Murakar - entonite deposit in central Japan, where a homogeneous bentonite bed and rhyolitic intrusive rock are present. Geological, petrological and geochronological studies have already been presented at the MRS symposium in Boston, 1989 (Kamei et al. 1990), so only a brief description of this deposit is given below:





Tuffaceous sedimentary rocks

+ Granite (basement)

- Fault

Fig. 6 Geological map of the Murakami deposit



Fig. 7 An idealized geological section of the Murakami deposit

Geology

A geological map and an idealized section of the Murakami deposit area are shown in Figure 6 and 7, respectively. Rhyolitic lava and tuff are distributed in a graven with a width of approximately one kilometer. The reported age of deposition of this unit ranges from 18 to 14 Ma (Muramatsu, 1988). The tuff is regarded as being deposited in a marine environment, and was converted into bentonite probably due to diagenetic reaction. Subsequent intrusions of biotite rhyolite are found in the bentonite bed. The contact between the intrusive rock and the tuff dips about 30° near the surface of the ground, and the inirusive rock body is assumed to form a funnel with a diameter of less than 200 meters (Figure 7).

Samples

Sample A was collected from a point 30 meters distant from the contact between the intrusive body and the bentonite bed. X-ray diffraction showed that sample A contained illite-smectite mixed layers with an illite ratio of about 40 %.

4.2 RESULTS

Thermal History

The cooling rate of the intrusive rock was determined from combining radiometric mineral ages and each closure temperature. The cooling rate of sample A was estimated by the "TRUMP" thermal analysis code. The results are shown in Figure 8. The cooling rates of the intrusive rock and of sample A were 70 °C/Ma, and 60 °C

/Ma, respectively. (Figure 8 and the values for cooling rates are newly revised, therefore those reported in the MRS Proceedings (Kamei et al. 1990), should be ignored.)

The Illitization Period

In the Murakami deposit area, a minimum temperature of illitization is regarded as

(1) Material	Smectite in marine sediment
 (2) Environment ① Water chemistry ② Temperature 	Modified seawater > 240 ℃~ 105 ℃
(3) Period	2.4 Ma
(4) Result	<pre>I/S mixed layers mineral (Illite;approximate 40%)</pre>
(5) Activation energy	Approximate 27 kcal/mol

Table 7 Summary of a study on illitization of smectite associated contact metamorphism — A case study at the Murakami deposit

4.3 CONCLUSION

Once again returning to the three-part concept of starting materials, environmental and chronological conditions, and results, it was possible to describe the illitization of smectite associated with contact metamorphism in terms of 1) material studied, 2) environment, 3) period, and 4) results (Table 7).

A more precise estimation of activation energy is possible through an estimation of the overall thermal history during contact metamorphism, using a thermal analysis code. This work is in progress.

5. ALTERATION OF CEMENT

Concrete components such as tunnels or estuary walls with known ages were studied. Environmental conditions such as temperature, surrounding materials, water content, and water chemistry ware either measured or estimated. The alteration of cement materials has been analyzed by EPMA, SEM, TEM(Transmission Electron Microscopy), and XRD. Results obtained to date are shown in Table 8.

The following alteration features of cement materials was able to be traced:

- (1) Decrease in pH of pore water,
- (2) Decrease of CaO/SiO2 ratio of C-S-H gel,
- (3) Partial dissolution of C-S-H gel,
- (4) Formation of CaCO3,
- (5) Permeation of Cl, resulting in formation of Friedel'Salt.
- (6) D lution of Calcium hydroxide,
- (7) Diss ution of Calcium which cause dissolution of CaCO3,

Such alteration phenomena were detected within a range of a few centimeters. Further studies are necessary to permit any definite conclusions. 105°C, because this was the temperature estimate made by Oda et al. (1985) for the appearance of illite-smectite mixed layers in Japanese oil fields, of which the Murakami deposit form a part. In the vicinity where sample A was collected, the temperature at 6.4 Ma was presumed to be 240 \pm 50°C. Therefore, a period of about 2.4 Ma was required to cool these rocks from 240 to 105°C.

In short, smectite was converted into illite-smectite mixed layers, in which the illite ratio is approximately 40 %, in the period of more than 2.4 Ma.





Water Chemistry

The chemical composition of the rocks distributed in the Murakami area is mostly rhyolitic, and the tuff, now converted into bentonite bed, is of marine origin. The geological evidence leads to the idea that the chemistry of the water related to illitization was very similar to that of seawater after it was modified by interaction with rocks of rhyolitic composition.

The hydrogen isotopic composition (D/H) of water, in the form of hydroxyl groups, in the illite and the smectite-illite mixed layers, were measured, and from this an assessment of the water involved in illitization was made. The results supported the idea noted above.

Provisional Calculation of Activation Energy

The activation energy for illitization at the Murakami deposit was provisionally calculated on the basis of the estimated thermal history. The calculation procedure was already described in Kaunei et al. (1990). Using the revised cooling rate of 60 °C/Ma and a period of 2.4 Ma. the activation energy is approximately 27 kcal/mol. This value is close to that obtained by Roberson & Lahann(1981) of approximately 30 kcal/mol. The water used for their experiments contained 400 ppm K * and 9400 ppm Na * , the chemistry of seawater. A similar water chemistry can be inferred at Murakami.

6. SUMMARY AND FUTURE PROSPECTS

Table 9 summarizes the present state of PNC analogue studies on engineered barrier materials, and Figure 9 shows the framework of our analogue studies.



Natural Analogue Studies (Long-term)

Fig. 9 Framework of analogue studies on engineered barrier materials

Our natural analogue studies have three components:

- 1) investigation of alteration phenomena of analogue materials,
- 2) examination of environmental conditions (time intervals, water chemistry etc),
- 3) support experiments

The validity of the analogue study is determined by the selection of alteration phenomena of analogue materials out of various natural phenomena acting on historical materials from the point of view of best analogical fit.

An examination of environmental conditions occupies an inevitable part of the study. Time intervals, prevailing temperature, and water chemistry related to the alteration are key items.



(1) MATERIALS STUDIED ① SITE ② SAMPLE MATERIALS	Kanagawa-Manazuru Concrete of Railway Tunnel Wall	Yokohama-katabira Concrete of Estuary Wall
 (2) ENVIRONMENT Temperature Surrounding Materials Water Content Water Chemistry 	1 3 °C Lapilli tuff 4 0 % Ca - ND ₃ (HCO ₃)	1 5 °C Soil 3 3 96 Na - Cl
(3) PERIOD (yr)	6 7	6 1
(4) RESULTS OF ALTERATION	Ca depletion <few mm<="" td=""><td>Cl permeation >10 cm CaCO, formation >8 cm CaCO3dissolution >5 cm</td></few>	Cl permeation >10 cm CaCO, formation >8 cm CaCO3dissolution >5 cm

Table 8. Alteration behaviour of cement fabrics

Table 9 The present state of PNC analogue studies on engineered barrier materials

Barrier Components (Candidate Materials)	Materials and Mode of Occurrence	Period (yr)	Estimation of Period	Estimation of Environmental Conditions
Waste Form (Borosilicate glass)	Scoria (Pyroclastic Fall Deposit)	10 ² ~10 ⁴	Tephro- chronology	From Recent Cliumatological Conditions
Buffer (Compacted Bentonite)	Contact Metamorphosed (Natural) Bentonite	106~107	Radiometric Age Determination	Closure Temperature of Radiometric Ages Geological and Geochemical Data
Overpack (Carbon Steel)	Industrial Materials (Pipe)	101~102	Documents	From Present Embedded Conditions
Backfill (Concrete)	Industrial Materials (Components of Fabrics)	10 ¹ ~10 ²	Documents	From Present Embedded Conditions

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Support experiments are indispensable to the study in order to enhance the wider applicability of the natural analogue. Comparison of differences in composition or condition is the key issue for laboratory support experiments. Such experiments for the comparison of compositional differences between basaltic glasses and candidate waste glass have already started, and the results to date indicate that there is no significant recognizable difference in the leaching rates.

From a combination of the natural analogue studies outlined above and laboratory experiments on the candidate materials, an integrated evaluation of the long-tem durability of candidate engineered barrier materials can be conducted.

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5) Laboratory of Cement Mineralogy, Nagoya Institute of Technology, especially Professor K. Suzuki as technical advisor concerning hydration of cement materials;

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Tokai Vitrification Facility (TVF)



Power Reactor and Nuclear Fuel Development Corporation

Tokai Works



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Pu-contaminated Waste Treatment Facility (PWTF)



Power Reactor and Nuclear Fuel Development Corporation

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Cross-section view of melted ash



Power Reactor and Nuclear Fuel Development Corporation Head Office 1-9-13 Akasaka Minato-ku Tokyo Japan Tel 03-586-3311

Key Technical Issues for Commercialization of FBR Power Plants

Key Technical Issues		Target of Development	
Long-life high performance fuels		Reliable fuel with its burn-up above 200.000MWd/t	
High performance core for large- scale FBRs	1	Optimization of a high performance core for 1.500MWe plant	
Plant service at high temperature	k	Structural material resisting higher temperature (over 550°C)	
Optimization of heat transport and fuel handling systems		Optimization of systems' layout to reduce the size of reactor building Development of compact and reliable components	
Optimization of reactor contain- ment design	1	Realistic estimation of a rise in pressure at severe accident	
Seismic isolation	•	Licensable evaluation model and standards of seismic isolation design	
Elimination of secondary heat transport system	•	Establishment of safety logic and corresponding protection system Development of reliable double-walled tube type steam generator	
Reliable decay heat removal system	•	Passive decay heat removal system of natural convection	
Autonomous plant operation	•	Fully automatic plant operation using the artificial intelligence (AI) technology	
Optimization of safety logics		Safety design and evaluation with an adequate margin	
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Artist's View of a Commercialized FBR

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		ptilene.
	Reantor vessel	
2	Core	
3	Shield plug	
4	Control rod drive mechanism	
5	Refueling machine	
6	in-cell crane	
7	Spent fuel storage rack	
8	Reactor coplant pump	
9	Reactor coplant pipe	
10	Sleam generator	
11	Air cooler	
12	Dump tank	
1.4	Water steam pipe	PRESE LONG
15	HVAC system	
16	Seal bellows	
17	Advanced shut down mechanism	S ALL
18	Heat exchanger for residual heat removal	
19	Ceramics liner	To a more than the second seco
20	Seismic isolation device	
6.4	A ala lisa	

Developing Plutonium Utilization Technologies is a Key Role of PNC



In order to realize the full potential of nuclear power, it is essential to utilize plutonium and establish the nuclear fuel cycle. To utilize plutonium as an energy resource, Japan is pursuing a strategy of shifting from light water reactors (LWRs) to fast breeder reactors (FBRs). PNC is carrying out extensive R&D programs in various elements of the fuel cycle.



Power Reactor and Nuclear Fuel Development Corporation

Head Office Sankaido Bldg. 1.9-13. Akasaka. Minato-ku. Tokyo. Japan 107 Tel. 03.586-3311 Telecopy: 03.505-5125 (International Division) Tix J.27654 PNRD

OLEMERS OF CH

PNC Paris Office 4-8. Rue Sainte-Anne, 75001 Paris, France Tel: 1-4260-3101 Telecopy, 1-4260-2413 Tix: 42-240750

PNC Washington Office 2600 Virginia Avenue, N.W., #715. Washington, D.C., 20037 U.S.A. Tel: 202338-3770 Telecopy 202-333-1097 Tix, 23-089-2777

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PNC Exploration (Canada) Co., Ltd. 650 West Georgia Street. #2401, Vancouver, B.C., V6B4N8 Canada Tel: 604-681-6151, Telecopy, 604-682-3452, Tix: 21-4507862

PNC Exploration (Australia) Pty. Ltd. 16th Floor. Royal Exchange Building, 56 Pitt Street, Sydney, N.S.W. 2000 Australia. Tel. 2.241-1594. Telecopy. 2-251-1584. Tix. 71-25912

TECHNOLOGIES DISCUSSED AT PNC-OARAI

- Decommissioning Technology for Nuclear Fuel Cycle Facility in WDF
- Separation and Removal
 - Decontamination with Melting
 - Electopolishing I
 - Electropolishing II
 - Ice Blasting Decontamination
 - REDOX
- Radiation Control
- System Engineering
- Waste Treatment
- Remote Handling
- Dismantling- Plasma Arc Cutting
- Monitoring- Radiation Image Display
- R&D of Fast Breeder Reactors (FBR)
- R&D of Advanced Thermal Reactors (ATR)
- R&D on Fuel Recycling
- Development of a Heat resistant and Angle Beam Type Electro-Magnetic Acoustic Transducer
- R&D on Plutonium Fuel
- Low Level TRU Bearing Waste Management Technologies
 - Incineration
 - Acid Digestion
 - U-Contaminated Waste Incineration
 - TRU- Contaminated Waste Incineration
 - Cyclone Incineration
 - Waste Incineration
 - Decomposition
 - Dismantling
 - Plasma Arc Cutting
 - Laser Arc Cutting
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 - Electro-Slag Remelting
 - Hip Solidification
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 - Hypothermal Solidification
 - Cementation
 - Plastic Solidification
 - Bituminization
 - Krypton Immobilization

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DEVELOPMENT OF A HEAT RESISTANT AND ANGLE BEAM TYPE ELECTRO-MAGNETIC ACOUSTIC TRANSDUCER

K. ARA, H. RINDO, K. NAKAMOTO

O.zri Engineering Center.

Power Reactor and Nuclear Fuel Development Corporation

JAPAN

T. DOI

Advanced Technology Research Center.

Mitaubiahi Heavy Industries Ltd.

JAPAN

K. MORIMOTO

/ Takasago Technical Institute,

Mitsubishi Heavy Industries Ltd. /

JAPAN

T. SAKAMOTO

Kobe shipyard & Machinery Works

Mitsubishi Heavy Industries Ltd.

JAPAN



ABSTRACT

The invervice inspection (ISI) system of fast breeder reactor (FBR) is required to be miniaturized and rationalized, because of the severe environmental conditions around the reactor vessel of an FBR during ISI. A new ultrasonic testing methods using an electro-magnetic acoustic transducer (EMAT) has been developed to apply for the ISI of the FBR reactor vessel in a practical manner. The new method can generate directly ultrasonic waves in the materials without the couplant such as oil and water, through the interaction of a magnetic field and eddy currents. The high performance EMAT has been developed. So the inspection system with the EMAT will be able to get compact, because it needs no couplant supplying and collecting equipment.

The developed "EMAT" can be available up to 240 °C and detect 20% slit in depth artificial flaw in austenitic stainless ageel which wall thickness of 50mm under 240°C in laboratory. Transmission cable of forty meters was applied for the EMAT to consider the actual interval between an inspection point and the signal processing equipment.

For presentation in IAEA Specialist's Meeting on "Experience and Further improvement of In-Service Inspection Methods and Programmes of NPPs with Particular Emphasis on On-Line Techniques"

1. INTRODUCTION

The ISI will be done in the narrow space between the reactor vessel and the guard vessel of a FBR under the condition of high temperature and high radiation field as shown in Table 1. The probe is required to be compact for remote inspection and to have the excellent property under high temperature without cooling. If conventional UT probes are used in this case, the ISI squipment might be large because it needs the additional couplant equipment. The inspection system applied DMAT satisfies all of the above requirements. The DMAT is a couplant free sensor and has potential of high temperature use, as described in the next section.

The basic design conditions for the EMAT are shown in Table 2.

Table 1: Environmental Conditions

Temperature (*C)	1	applox.200	
Radiation dose rate (R/hr)	8	max.1000	
Access space (IND)	1	300	
Signal transmission length (m)		max. 40	

Table 2: Basic Design Conditions requirements

Subject :	Heat affected zone of weld (austenitic stainless steel)
Detectability :	50% slit in depth at wall thickness of 50mm (In laboratory, 20% slit)
Operating temperature and operating time	: 200 °C, 100 hr

2. PHINCIPLE AND STRUCTURE OF EMAT

A principle of the EMAT is shown in Figure 1. The EMAT consists of a set of magnets array and coil, and the high frequency current in the coil generate eddy current in the surface layer of the materials. Lorentz force \vec{F} is generated by the interaction between the magnetic field and the eddy current. And Lorentz force \vec{F} is defined by the following equation,

5 . J × B

where \overline{J} is eddy current and \overline{B} is magnetic flux density. The direction of Lorentz force depends on the frequency of eddy current, and vibration of a material results in generation of ultrasonic waves. The thickness (t) of one magnet piece is given by the following equation.

where C is sound velocity f is frequency 6 is refraction angle

The ultrasonic wave generation and detection mechanism of shear horizontal (SH) wave angle beam EMAT are shown in Figure 1. In the opposite way, in detection, ultrasonic waves near the surface of the test sample besides the coil and electronic oscillations are produced. By the interaction between external magnetic field and vibration force, electrons flow in the direction of the force generated and produced alternative curve. In the surface of the materials. The current induces a voltage in the detection coil on the test piece.



Figure 1: -Schematic diagram of ultrasonic generation by SH wave angle beam EMAT

3. DEVELOPMENT OF EMAT AND EVALUATION

(1)INVESTIGATION OF SPECIFICATIONS OF EMAT AND SELECTION OF OPTIMUM CONDITION

The specifications of EMAT were investigated under the actual environment conditions and basic design requirements for the EMAT are shown in Table 3.

able 3: Specification of EMA	T
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Ultrasonic wave mode	1	SH wave
Refraction angle (deg)	1	60
Frequency (kHz)	1	700 (at 240 ℃)
Operating Temperature ($ hinspace ($	4	200 (max. 240)
Transmission cable length (m)	1	40

ULTRASONIC WAVE MODE: In the case of the inspection for the heat affected zone (HAZ) of welded part in austenitic stainless steel, ultrasonic wave mode is one of the most important factor on designing of the probe, because that acoustical anisotropic materials exist in HAZ. SH wave was selected by the results of wave mode characteristics test. SH wave keeps its sound velocity in HAZ. And the loss of wave mode transformation is lower than that of shear vertical (SV) wave, because SH wave has no transformation at reflection on the boundary of a flaw.

REFRACTION ANGLE: The EMAT is required to detect primarily a defect in the inner surface of the stainless wall. Because of the wide heat affected zone of welded part, the angle beam function is necessary for a probe, especially applied for heavy wall thickness such as reactor vessel. The refraction angle of 60 degree was selected for the EMAT by experiment.

FREQUENCY: Generally, the signal level obtained from an EMAT is very low. Especially, a flaw detection ability of an EMAT in applying for inspection of nonmagnetic materials is inferior to that in applying for magnetic materials. The reactor vessel of FBR 'MONJU' was applied for non-magnetic stainless steel. Therefore, the optimum frequency was investigated. As a result, it was confirmed that the EMAT had the highest signal level at the frequency of 700kHz.

HIGH TEMPERATURE APPLICATION: The EMAT for high temperature use of 200 °C has been developed. High performance heat resistant parts and components were developed shown in Table 4. The sheet type coil is coated poly-imide film. The magnet assembly is an array of thin Sm-Co magnet pieces. In this case, the thickness of a zagnet (t) is 2.4mm.









Figure 2 : - Prototype EMAT





Table4: Materials of Elements of EMAT

ELEMENTS	MATERIALS	
Coll	Poly-imide sheet c	011
Magnet	: Sm-Co(Curie point;82	0*0)
Cable	: Teflon insulato	r



ANGLE INCIDENT OF SH-WAVE EMAT: The sound field of the EMAT was measured and confirmed its angle incident function. Figure 4 shows the sound field at the distance of 100mm from the center of the EMAT. The points marked "*" show the measured sound intensity of transmitter and "o" show that of receiver. And the line shows its simulation result. Close agreement between measured and simulation value was obtained, and the center angle of the sound field was 60 degree.



Figure 4: Sound fields of EMAT

(2) DETECTION OF FLAW

The flaw detection ability of the developed EMAT was confirmed by experiment. As shown in Figure 5, the EMAT was placed on the outer surface of the test specimen, which has the artificial slit type flaw at the inner surface. The test specimen is made of the austenitic stainless steel as same as the reactor vessel of FBR.

One of the typical results of flaw indication pattern is shown in Figure 6. A slit type flaw, 35mm in length and 20% of wall thickness in depth, in the heat affected zone could be clearly detected with the signal to noise ratio (S/N) of over 2.0, under the both conditions of room and high(240 °C) temperature. Sensitivity obtained was wept continuously for 100 hours at the temperature of 240°C.

(3) SIGNAL TRANSMISSION PERFORMANCE

The EMAT applying for FBR reactor vessel is required to have excellent property of the signal transmission from itself to signal processing equipment at intervals of ROM. The output signal strength from the EMAT is very low, furthermore it



Figure 5: Test configuration for detecting the flaw



Figure 6: CRT traces for detecting the slit type flaw in laboratory



Figure 7: . Test configuration for signal transmission

attenuate through the long transmission cable. Therefore, signal impedance matching between the DMAT and the r gnal processing equipment is needed. Test configuration for signal transmission is shown in Figure 1. The attenuation ratio in transmission line under high temperature of 240 °C was higher than that in r om temperature as shown in Figure 8.

(4)EVALUATION

HIGH TEMPERATURE OPERATION: Minimum Requirement of operating temperature is 200°C for 100 hours. It was confirmed by the experiment that the EMAT could be operated up to 240 °C for 100 hours over. The element with the low heat resistance are coil and cable, especially electrical insulator. The maximum operating temperature of the EMAT depends on that of the electrical insulation materials. It is necessary to improve these materials for realizing higher temperature use.

DESIGN PARAMETERS: The specifications of the EMAT for FBR reactor vessel were decided through some experiments. It was confirmed that design value agreed with measured value well. Design parameters depend on the conditions of an inspected material. For example, in the case of applying for the thin wall inspection such as piping, refraction angle and frequency of ultrasound should be selected optimum value.

DETECTABLEITY: The requirements of flaw detection ability is 20 % slit in depth in austenitic stainless steel under high temperature of 240°C. The results of the research satisfied with the target. We could get the result that the flaw signal level in high temperature of 240°C was about 3dB lower than that in room temperature.

APPLICATION: In remote operation, the flaw signal level is attenuated through the long transmission cable. But impedance matching is effective, it could transmit the flaw signal without a amplifier.







(Time 20µs/div, Amp, 50mV/div)



Figure 8: CRT traces for transmitting signal at intervals of 40m in labe atory

4. CONCLUSION

SH wave, angle beam EMAT with high temperature use was developed which is applicable to the reactor vessel inspection of FBR. It was confirmed in laboratory that the EMAT could be operated up to 240°C for over 100 hours, the flaw detection ability was 20% slit in depth artificial flaw in austenitic stainless steel wall thickness of 50mm, and forty meters of signal transmission cable could be applied.

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MASAO SHIOTSUKI. SATOSHI IKECA. and HIDEHIKO MIYAO Power Reactor and Nuclear Fuel Development Corporation (PNC)

1. INTRODUCTION

Nuclear facilities such as power reactors, reprocessing plants and fuel fabrication plants are generally said to have a limited life of up to 30 or 40 years. When they are superannuated, these facilities have to be dismantled and removed safely, and wastes from such dismantling must be treated under appropriate control. These operations are comprehensively termed as "decommissioning".

Power Reactor and Nuclear Fuel Development Corporation (PNC) has so far dedicated itself to the technical development of fast breeder reactors, reprocessing techniques and MOX fabrication techniques. Programs are based on national policy of plutonium fuel recycle. PNC is now developing fuel cycle facility decommissioning techniques.

At the Waste Dismantling Facility (WDF) located in O-arai Engineering Center (OEC). PNC is eager to validate its technical development efforts aimed at the treatment of surface-contaminated large size wastes from post irradiated FBR fuel and material examination (PIE) facilities.



Photo.1 View of Waste Dismantling Facility (WDF)

2. OUTLINE OF MDF

The plan and the waste stream of WDF are shown in Fig.1 and 2. The WDF is a ferromonomete building with three stories and one basement. The building area is 1,700 s² and the total floor area is about 5,400 s².

The wastes are classified into three categories such as high radiation level α wastes (surface radiation level: $\geq 50 \text{ mrem}/h$), low radiation level α wastes (surface radiation level: <50 mrem/h), and β , γ wastes. The high radiation level α wastes are received through the overhead hatch of the α waste loading cell.

The wastes are then sent to the decontamination cell via an air lock chamber and unpacked by means of master slave manipulators. After measurement of the dose rate and surface contamination, the wastes are subjected to surface decontamination by an ice blasting process. Thereafter, they are transferred to the dismantling cell and are cut into pieces with a plasma cutter and a hacksaw (Photo.2 and 3). Compressible wastes are further subjected to a compressing process (Photo.4) and then are packed in metallic containers.

The low radiation level α wastes are brought into the acceptance hall and then are transferred by a cart to the decontamination hall, where the wastes are unpacked and dose rate and surface contamination are measured directly by the workers wearing airline suits. Then the wastes are decontaminated by an electropolishing process. Thereafter, the wastes are dismantled with the plasma outter into small pieces and packed into containers. The hall is constantly monitored from the 2nd floor control room during these operations.

The β , γ wastes are introduced directly into the β , γ dismantling cell from the overhead hatch of the cell and out into small pieces by remote outting techniques. The high radiation level wastes are packed into metallic containers and stored into casks for transportation. The low radiation level wastes are sent to the β , γ loading cell, and after sorting and classifying, they are packed into drums. The process flow sheet is shown in Fig.3.



Photo. 2 Plasma cutting

Photo. 3 Hacksaw cutting









Photo. 4 Compressing Machine

Photo. 5 Operator wearing Frog-man suit



Fig.1 #DF Plan

		e Dissantling Cell	# Decontagi- betion Gell	a Loading Cell	A r Dismantling Cell	F + Loading Cell
Dimension (m) L × W × H		13.4 × 4.0 × 6.1	3.5 × 4.0 × 6.1	6.8 × 4.0 × 6.1	4.8× 4.2 × 6.6	4.6× 4.6 × 6.6
Shimiding	Front	1000 :	1000 : Heavy Concrete		750 : Concrete	550 : Concrete
(8.8)	Roof	1050 : Conprete		500 : Concrete		
	Floor	1150 1	Concrete	A second s	550 : Conorel	Le .
Lining	Wall/ Roof	EIE 3/A		Epoky Resin	Ероку Певіп	Taxan Basis
	Floor			SUS 304 SUS 304		They wante
Air tightness		< 0,1 WD	18 /Hr	Contraction of the Address of the Instance	Negative Pressure	

Table, 1	Specifi	1780	no	6f	Celi	\$
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Table. 2 Specification of equipment

EQUIPMENT	SPECIFICATION
(PROCESS) 1) Pleame Cutting	· Cas · Current : Max. 250 A · Current : SUS 70
2) Hacksaw Cutting	· Capacity : SUS 200 cm
3) Compression	· Type : Uniaxial Press · Object : 259 # × 320 mm · Capacity : 70 Ton
4) Press Cutter	· Object : 200× 1000× 8 mm · Capacity : 400 Ton
(HANDLING) 1) Roller Conveyor	Type : Motor Drive with Chain Capacity : Max. 2.0 Ton
2) Transfer Car	 Type : Self-Drive with Motor Self-Drive with Linear Motor Cepacity : Max. 2.0 Ton
(AIR LINE SUITS)	<pre>* Type : Fixed , Free * Number : 2(Decon.),3(Dismantl.) ?(Beta-Campa) * Temp : 122 ~ 35 %</pre>
(REMOTE HANDLING) 1) Master-Slave Manipulator	 Type Handling Max. 23 Kg Load Number 5 Pairs 3 Pairs
2) Power Manipulator	 Handling : 67.5 Kg (All Position) Load Shoulder : 450 kg Hoek Load Number : 3
) In-Cell Crane	· Capacity : Max. 2.0 Ton



alle.

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Fig. 2 Waste Stream in WDF

- Contraction

-

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3. R & D STRATEGY OF D/D TECHNOLOGIES

Technology needed for areas on nuclear fuel cycle decommissioning operations have been identified and prioritized using the results of past power reactor decommissioning studies for each major decommissioning activity. (Fig.4)

In comparison with reactors, the decommissioning of nuclear fuel cycle facilities has distinctive features that objects to be removed are contaminated with TRU nuclides and their contamination conditions, structure, configuration and materials vary.

These factors have been considered in developing decommissioning techniques for PNC's nuclear fuel cycle facilities. Safe and effective decommissioning of nuclear fuel cycle facilities with minimum generation of secondary wastes and cost would be achievable with an integration of the techniques discussed bellow.



Fig. 4 Development of Technologies on D/D of Nuclear Fuel Cycle Escilities

4. DEVELOPMENT OF DECONTAMINATION TECHNIQUES

Decontamination techniques are classified into two groups, namely, "Primary decontamination techniques" and "Complete decontamination techniques". (See Fig. 5.)

The former envisages the removal of loose contaminants to reduce the exposure dose rate involved in handling nuclides and to prevent spread of contamination. The latter aims at the absolute reduction of radioactivity down to background level.

The WDF is now developing an "Ice blast decontamination process" as a means of primary decontamination as well as an "Electropolishing process" and a "REDOX process" as a means of complete decontamination.

CATELLEY	MELTAWICEM	REMEWAL STZE		TECHNIQLE
Primery -	D Extaliation of Contaminants	Particle	+	Spray Ice-Blasting
	(2) Solution of Munlides	1 Molecule	******	Cheminal Decon
	3 Surface Removal	-		Electro-Polishing
Camplete		Atom		REDOK
Decon.	3 same	-		Electro-Slag -Remelting

Fig. 5 Decontamination Techniques and Mechanism

4-1 Ice Blast Decontamination

Ice blast decontamination is a surface decontamination process using ice and dry ice mixed particles made by pelletizer, which are blasted onto an object to be decontaminated using a carrier medium such as compressed air.

This process features the utilization of blast impact energy and low temperature to remove nuclides, coatings and oils for improved decontamination efficiency in comparison with that of a spray process with far less secondary waste generation.

The impact energy producted by a blaster in the WDF (with compressed air rated at 6kg /cm²) come up to hundreds of kg /cm².

Fig.6 is a conceptual illustration of ice blast system. The system consists of a pelletizer and a blaster. Only a maintenance-free flexible pressure hose and a blasting mozzle are installed in the cell.





The pelletizer is capable of producing blast particles at a rate of about 200kg /h.(Photo.6) The particles are made by converting liquefied carbon dioxide into a fine powder through adiabatic expansion and then compacting these into column shaped particles

of 4mm diameter and 5mm length. The pelletizer is provided with a water supply system, which can mix up to approximately 20% water (ice) into blast particles. The objective of mixing water into blast particles is to increase the hardness of blast particles and to facilitate transfer of nuclides to the liquid waste stream during decontamination.

The evaluation of test results obtained so far has verified the validity of this decontamination process. In comparison with pressurized water processes, this method holds down the secondary waste generation to the order of one tenth and achieves higher decontamination levels.



Photo, 6 The Pelletizer





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Hence, this process will be given increased ispetus for development as an effective primary decontamination process that promises wider applicability and further improved decontamination efficiency.

4-2 Electropolishing Decontamination

This is the application of electropolishing, a common industrial technique for surface processing of setsis, to nuclear decontamination.

In this process, as the surface of contaminated metal dissolves, nuclides will move into the electrolyte. Theoretically, a decontamination efficiency as high as BC level can be expected.

The WDF started developing this process in 1982 and selected a 5% sulfuric acid solution as the electrolyte in consideration of its electrolytic properties such as polishing efficiency and uniform dissolubility as well as after treatment of spent electrolyte. WDF sets up a demonstration decontamination system (Photo.7) in a Decontamination Hall.

The basic concept of electroplishing system is illustrated in Fig.7.





Fig. 7 Schematic Diagram of Electropolishing

Photo.7 Electropolishing Decon. system

Decontamination will take place with the application of positive charge to metallic wastes in a conductive electolyte because the charge will displace the metal surface into the electrolyte as cation. Also, successive electropolishing operations will lead to an increased metal ion concentration in the electrolyte, decreasing the polishing efficiency down to 1/3 to 1/4 of the initial value when the concentration rises to 30 g metal ion/liter or more. Finally the spent electrolyte itself becomes waste. In order to recover such a spent electrolyte for reuse, an electrodeposition technique which is reverse to electrolysis is used to recover metal ions from the spent electrolyte. A critical element of this technique is pH control, which can be achieved by providing an electrolytic diaphrage between the regenerative cathode and anode which selectively allows the permeation of the fulfuric acid ion (SO. ?-).

From the decontamination of wastes derived from irradiated FBR fuels. WDF has

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confirmed the high efficiency of this decontamination process (Fig.8). The WDF's oceparative evaluation of this process versus ultrasonic cleaning has verified excellent efficiency of this process (Fig. 9) as demonstrated by the distribution of contamination and the SEM observation of polished metal surface. The sulfuric-acid electrolytic decontagination which can remove effectively the contagination embedded in the grain boundaries of metals, is particulary effective.



Fig. 8 Relationship A.R.T. and DF

4-3 REDOX Decontamination

This is an electro-chemical decontamination process, where the dissolution of metals and contaminated wastes is accelerated by the addition of quadrivalent cerium ion (Ce(IV))to nitric acid to form a strong exidizing agent. Reduced cerium ion (Ce (11)) is exidized into quadrivalent cerius ion by electrolysis to regenerate the oxidant. Its principle is shown in Fig. 10.



Unlike electropolishing decontamination, this process has its electrolytic and decontamination steps clearly separated. Since its decontamination is based on the electrochemical reaction between solution and metallic surface, it resulted in a high DF and unifors dissolubility.

Rinsing and Electropolishing

The WDF has been operated as a cold decontamination experiment system (with a 100liter decontamination bath) mince 1985 and has verified the validity of its highly unifors __ssolubility in polishing tests on plates and valves with different surface roughnesses :

This process, however, is not free from problems due to the strong acidity of quadrivalent cerius ion. Such problems include the selection and evaluation of the equipment component materials and the treatment of spent decontamination solutions. These will be solved through subsequent studies plus hot tests to be conducted on actual wastes.

5. DEVELOPMENT OF DISMANTLING TECHNIQUES

Facilities, machines and equipment used in the nuclear fuel cycle are diverse in construction, configuration and material. Thus, it is essential to evaluate the applicability, safety and efficiency of dismantling techniques under development.

In its efforts to develop dispantling operations, the WDF places greater exphasis on plasma outting technique which has wide applicability to many components. In the WDF, large-size wastes generated in OEC have been out by a plasma are or a hacksaw, and operation of the plassa torch with a master slave manipulator and preliminary plasma outting robot have been demonstrated. In the same way, various methods, peripheral techniques and remote control techniques for dismantling are being developed.

5-1 Plasma Cutting Robot

In order to dismantle large-size equipment and machines of complex configuration installed in high-radiation and high-contamination areas, it is essential to use remote control techniques for automatic, efficient and safe dismantling and removal operations.

In 1984 a plasma cutting robot (Photo.8) was installed in WDF, as a modified version of industrial "Nots. as a link in the development " resole control technology to verify its usefulness in the dismantling of wastes from operating plants. This robot was based on a teaching playback method, in which a outting path on an object is preliminarily taught to the robot and outting is made to the given outting path. If the object to be cut has a complex configuration, its teaching procedure takes such time. To solve this problem, some notable improvements have been made.i.e. the addition of a voltage are sensor which will feedback voltage fluctuations to the robot during outting for automatic operation. In addition a non-contact type laser distance sensor, a joy stick and a master arm (Photo.9 and 10) have been added.



Photo. 8 Plasma Cutting Robot





Photo. 9 Laser Distance Sensor

Photo. 10 Controller

Based on the dismantling method, algorism and many other design factors obtained from the WDF robot, a small-size portable robot for decommissioning use is now being developed. Its design concept is shown in Fig.11.



Fig. 11 Concept of D/D Plasma Robot

6. DEVELOPMENT OF MONITORING TECHNIQUES

Efficient and safe decommissioning operation addresses importance to monitoring techniques by which to determine the quantity and distribution pattern of contaminated nuclides.

PNC is developing a radiation image display (RID) which could be replaceable the conventional spear method and direct survey method and provide a reliable and quick means of evaluating the distribution of nuclide contamination by remote control.

6-1 Radiacion Image Display (RID)

To improve measurement and evaluation efficiency for decommissioning, to decrease radiation exposure during the work and also to improve the reliability of measurement data. PNC has been developing a radiation image display capable of remote and automatic measurement and image display of the distribution of radioactive substances.

Its operating principle is to run a colimated γ ray detector and distance meter to scan across a contaminated object to be measured and obtain radiation information and distance information, from which a computer will create a picture of the distribution of radioactive substances (evaluation picture) composed of 1,500 to 9,000 plots divided into ten color levels and then will display the picture on a TV monitor as a synthetic image.

The measurement principle is illustrated in Fig. 12. Photo.11 shows the prototype equipment No. 1 manufactured in 1986.



Table 3 Specification of RID No. 1

ITEM	RID (1)				
	CSI(T ()+PD				
γ Detector	¢ 28 × 50(mm)				
Shielding Material	Tungsten				
Shielding Thickness	50mm				
Shielding Power (7 Energy:1MeV)	17100				
Detector Weight	~50kg				
Detector Dimension	300(W) × 350(D)				
Cable Numbers	5				
Measurement Time	10,15,30,60 min				
Calculation Time	3min				

Fig 12 Composition of Radiation Image Display



Data processing section

Detecting section

Collimator. (2) TV camera. (3) Upper box .
 (4) Personal computer. (5) Image display equipment



Photo.12 shows the results of measurements on a liquid waste tank. The TV monitor shows the entire measurement range. The evaluation picture gives a counting value at each plot in ten color levels (red-yellow-green-blue-non color) with the largest counting value of the plots in the picture placed as the upper limit. The synthetic image of both the screen and the picture provides information about radioactive substances deposited on the bottom of liquid waste tank.



Photo. 12 Example of Liquid Waste Tank Measurement

The applicability of this equipment was evaluated in a test made on wastes and a liquid waste tank. The results obtained from the test are shown below:

(1) Measurement of wastes

The measurement results of a 200-liter drum containing processed wastes are shown in Photo.13.

The evaluation of the result clearly indicates the location of radioactive substances existing in spots.



① Sealed position of radioactive substances Photo.13 Result of 200-liter Drum Measurement

(2) Measurement of a liquid waste tank before and after decontamination

A tank containing liquid vaste from the cleaning of FBR fuel assemblies was decontaminated with high-pressure water. This decontamination process was evaluated by this equipment. Images obtained before and after decontamination are shown respectively in Photo.14. The measurement was done over 30 minutes and from a distance of 3.5 meters. It is noted that the high-pressure water decontamination carried highly contaminated

It is noted that the high pressure sector tank before the decontamination, to the bottom deposits, which is observed at the top of tank before the decontamination, to the bottom of tank.



Before decontamination

After decontamination

The display of high-level contamination ① shifted to the display of low-level contamination. ② and ③.

Photo. 14 Evaluation of Liquid Waste Tank before and after Decontamination

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7. CONCLUSION

We will continue the development of decontamination and dismantling technologies undertaken by WDF to establish techniques that can validate the safety and economy of various nuclear fuel cycle facilities.

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FBR Development In PNC for Commerciality.



Construction of "Monju", A Prototype Fast Breeder Reactor

The construction of Monju is now 85% completed at the end of June 1990 All the components is scheduled to be installed by April 1991 The target date for initial criticality is October 1992







searanten completes in may race

Reactor containment vessel completed in April 1987

CONSTRUCTION SCHEDULE OF MONJU

FISCAL YEAR	184	185	165	187	80	180	190	- 191	- 19.2	. 93
KEY	EXCAVATION START OCT 85		BASEMAT CONCRETE START FEB'86		REACTOR VESSEL			INITIAL CRITICALITY OCT 192		



Research and Development on Plutonium Fuel

PNC's Tokai Works is mainly developing and manufacturing plutomum test, developing reprocessing technology, treatment and dispsed of high lavel to an active waste and uranium enrichment.



Arrial view of Tokar Works

An essential element of fuel for new reactors such as the ATR and FBR is plutonium - a byproduct of nuclear reactor operation that can be recovered from spent fuel. The plutonium is combined with uranium to form mixed or ide (MOX) fuel.

Since MOX fuel consists of both uranium and pli tonium, an effective non-proliferation measure is to mix the two and convert them to oxide immediately after extracting the plutonium.

PNC developed its own process, called the "microwaveheating direct denitration process," to carry out this coconversion. It has been used since 1979 at the Tokai Plutonium Fuel Fabrication Facility (PFFF). The Tokai Plutonium Conversion Development Facility, with a daily co-conversion capacity of 10kg MOX, went into operation in 1983, using plutonium nitrate and uranium nitrate from the reprocessing plant to produce MOX powder for ATR Fugen, FBR Joyo, and FBR Monju.

O The Tokai Roprocessing Plant Plutorium Fuel Facilities at Toka



Uranium-plutonium mixed oxide (MOX) powder
"Monju" is a 280 MWe Reactor Cooled by Liquid Sodium in Three Loops





Reactor vessel installed in October 1988

MONJU Plant Parameters

Reactor type

Thermal power Electrical power Fuel material Discharged fuel average burnup rate Breeding tatio Number of loops

Mixed oxide fuel, sodium cooled. fast neutron breeder reactor, loop type 714MW 280MW PUD, UD,

80000MWd/1 1.2

Secondary sodium temperature (IHX outlet/inlet) Type of steam generator Steam temperature (Tu bine inlet) 483°C Steam pressure (Turbine inlet)

Refueling system Refueling interval 505/325°C

Helical coil, once through unit type

127 kg/cm2g Single rotating plug with fixedarm fuel handling machine 6 months



R&D for Advanced Power Reactors



Although conventional light water reactors will continue to provide the majority of Japan's nuclear-generated electricity into the next century, the government has designated the development of fast breeder reactors as a top priority. PNC is working toward this goal with the development of the Joyo and Monju FBRs.

Because they utilize uranium resources so efficiently. FBRs represent the ideal nuclear power source for Japan's future. The FBR, which can be fueled by plutonium recovered from spent fuel, is often called the "ultimate reactor." It not only generates electricity, but actually creates more fuel than it consumes by converting U-238 in the fuel to Pu-239. After the year 2030, the FBR is expected to take over as Japan's chief source of nuclear energy.

PNC's role is to carry out R&D to develop commercialscale FBRs that are competitive with LWRs in terms of safety and economy.

This work involves many new technologies, such as the use of uranium-plutonium mixed oxide (MOX) fuels and the use of efficient yet chemically active liquid sodium coolant.

Airial view of O-arai Engineering Cente



Development of computer analysis code for major structures as a reactor vessel and related components





Our work in this area began with basic research on plutonium handling. MOX fuel properties, and fuel fabrication at the Tokai Plutonium Fuel Development Facility (PFDF). Technology developed here was confirmed on a larger scale at PFFF, which began producing fuel in 1972 and has supplied fuel for experimental FBR Joyo and prototype ATR Fugen. Initial core fuel assemblies were completed for Joyo in 1975 and for Fugen in 1978.

The next step is the demonstration of mass production technology in the Tokai Plutonium Fuel Production Facility (PFPF), which will supply large quantities of fuel to FBRs Monju and Joyo and to the demonstration ATR. The FBR fuel production line at PFPF started operation in 1988 with a capacity of 5 tons of MOX fuel per year. Another line now under construction will supply 40 tons of MOX fuel per year for the demonstration ATR and other plants. It is scheduled to start operation in 1993.















Comparison of Dose Equivalents from Natural Radionuclides and Radionuclides Discharged from Tokai Reprocessing Plant

> The height of poles means the amount of radiation exposure to an individual in a year.

The radiation exposure from radionuclides discharged from the reprocessing plant is less than one-hundredth of the exposure from natural radionuclides including radon and thoron and these daughters.





Radionuclides Discharged from Reprocessing Plant



Inhalation of Radon

Natural Radionuclides except for Radon and

RESEARCH AND DEVELOPMENT IN OARAI ENGINEERING CENTER

November 7

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OARAI ENGINEERING CENTER, FOWER REACTOR AND NUCLEAR FUEL DEVELOPMENT CORPORATION

RESEARCH AND DEVELOPMENT IN OARAI ENGINEERING CENTER

1. Mission of Oarai Engineering Center ··· P. 1
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1 Mission of Oarai Engineering Center

- © Research and Development of Key Technologies Associated with FBR & ATR Power Plants (Plant Systems, Fuels and Core, Sodium Technologies, Components, Safety, etc.).
 - (1) Design Studies for Framing Plant Systems with Safety and Economic Competitiveness
 - (2) Research and Development on Base Technologies and Innovative Technologies
 - (3) Research and Development Using Experiences Obtained through Construction and Operation of Joyo a: Monju, and Fugen.
 - Research and Development on Fuel Recycling Research and Development on Decommis- sioning, Nuclear Criticality, etc. of Fuel Re-cycling Facilities



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(2) Organization of Oarai Engineering Center



(3) Chronology of Oarai Engineering Center

- Oct.1967 Foundation of Power Reactor and Nuclear Fuel Development Corporation
- Dec.1969 Initial Nuclear Criticality in the Deutorium Criticality Assembly (DCA)
- Mar.1970 Establishment of Oarai Engineering Center
- May.1970 Start of the Fast Experimental Reactor "Joyo" Construction
- July 1974 Start of 50MW Steam Generator Test Facility Operation
- Apr.1977 Initial Nuclear Criticality of "Joyo"
- Nov.1982 First Nuclear Criticality of "Joyo" Irradiation Core(MK-II Core)
- Sep.1984 Completion of the Fuel Cycle
 'Loop' Using "Joyo"



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- Feb.1987 Agreement between "Joyo" and "Phenix" on Exchange Irradiation of Fuel Assembly
- Mar.1989 Contract of Technical Cooperation Agreement between PNC and JAPC for Research and Development of the Demonstration Fast Breeder Reactor.
- DEC.1989 Achievement of 40,000 accumulating hours' Operation for "Joyo"

- 3. Research and Development of FBRs
 - Extensive Preparations for the Protoype FBR "Monju" Operation on Items As Follows
 - > Planning of Start-up Test Program
 - Research and Development for Operation, Maintenance and Repairing
 - Training for "Monju" Operators
 - (2) Cooperation for the Demonstration FBR

Cooperation Based on Technical Agreement between PNC and JAPC

- Application of Operation Experience on "Joyo"
- ② Application of R&D Results Obtained in Oarai Engineering Center
- ③ Conduct of R&D Associated with the Demonstration FBR by Utilizing Technical Capabilities of Oarai Engineering Center





- (3) Research and Development for Commercial Type FBRs
 - Framing of Concepts for Commercial Type FBR Plants
 - Conceptual Design Studies for These FBR Plants
 - ② Research and Development of Key Technologies
 - i) Development of High Performance Fuel
 - ii) Development of Large Reactor Core with High Performance
 - iii) Development for Higher Temperature Service of Plant System
 - iv) Rationalization of Component, Piping and These Layout
 - v) Development of Rationalized Reactor Containment System
 - vi) Development of Seismic Isolation Structures

- vii) Development of Plant System without Intermediate Heat Transport System
- viii) Development of Decay Heat Removal System with High Reliability
 - ix) Reduction of Radiation Exposure during Maintenance
 - x) Establishment of Rational Safety Logic
- ③ Utilization of "Joyo"

i) Modification to a highly Efficient Irradiation Facility

ii) Development of Advanced Technology and Concepts





- 4 Research and Development of Advanced Thermal Reactors(ATRs)
 - Cooperation to the Demonstration ATR (Oma Nuclear Power Station in Aomori Pref.)
 - Conduct of Tests to Ensure Designing of the Demonstration ATR under Contract with Electric Power Development Corporation
 - (2) Research and Development of Base Technologies
 - Conduct of High Performance Fuel Development, and of Safety Research Associated with Severe Accidents



5 Research and Development on Fuel Recycling

- (1) Research and Development on Fuel Recycling
 - Conduct of Development concerning Decommissioning Technologies Common to Fuel Recycling Facilities
 - ② Conduct of Research Concerning TRU Transmutation Treatment Technologies by Utilizing FBR
 - Planning of Transmutation Demonstration Tests in the Fast Experimental Reactor "Joyo"
 - ③ Conduct of Research for Waste Disposal Technology
- (2) Research on Nuclear Criticality Safety
 - Conduct of Research Concerning Nuclear Criticality Safety Necessary for FBR Recycling Facilities by Using DCA

6. 'Frontier' Research

Conduct of the Following 'Frontier' Research Subjects for Innovation and Improvement of Advanced Power Reactors

- (1) Research for Nuclear Reactor Materials
 - New Materials Resistant to High Temperature
 - New Materials Resistant to Neutron Irradiation
- (2) Research on Artificial Intelligence Technologies for Nuclear Reactor Plants
 - Conduct of Research and Development for Nuclear Reactor Plants with Arti-
 - ficial Intelligence Systems
 - Demonstration Tests of Artificial Intelligence Systems in Joyo.



- (3) Diversification of Plutonium Utilization
 - Conduct of Research on FBRs Using Advanced Fuels, Small- and Medium-Sized FBRs, Inherent Safe FBRs, etc.







7. Facilities for Research and Development in Oarai Engineering Center

Present Facilities under Operation ;

- (1) Reactor Facilities (Joyo and DCA)
- (2) Test Facilities for Post-Irradiation Examination
- (3) Test Facilities for Developments of Components and Structures
- (4) Test Facilities for Safety Research
- (5) Test Facilities for Fuel Recycling
- (6) Administration Facilities for Health and Safety

◎ Total No. of Facilities : 34

New and Modified Facilities under Planning ;

- Expansion of the Fuel Monitoring Facility for MONJU and Other Large LMFBR Fuel Assemblies
- (2) Modification of Joyo for Improvement of Irradiation Capabilities and Demonstration of Innovative Technologies (Joyo MK-III Program).
- (3) Modification of DCA for Research of Nuclear Criticality Safety
- (4) Construction of a Information Center for Advanced Utilization of Computers
- (5) A Safety Test Reactor "SERAPH"^{**} for FBR (Under Investigation)

* Safety Engineering Reactor for Accident Phenomelogy



Technical Draft for Comments R D & D Program

on

Low-Level T R U Bearing Waste Management Technologies

March 1, 1990

Power Reactor and Nuclear Fuel Development Corporation

(PNC)

Translated into English and Prepared by Radioactive Waste Management Project, PNC in February 1990 for the U.S. Department of Energy



Technical Draft for Comments R D & D Program

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Low-Level T R U Bearing Waste Management Technologies



Radioactive Waste Management Project Power Reactor and Nuclear Fuel Development Corporation

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(4) Entrusted Research of

1. Incineration

Incineration technology is to oxidize combutibles such as paper, clothe, etc. and Clcontained wastes such as PVC by heating or other means, in order to reduce the waste volume and to transform into inorganic materials.

(1) Acid Digestion (Tokai)

Cold engineering test has been conducted at Pu-contaminated Waste Treatment Facility (PWTF) for volume reduction of Cl-contained wastes generated at MOX fabrication facilities etc., and for recovering Pu with acid digestion process characterized by low-temperature treatment. These test results will be reflected to future TRU waste treatment facilities.

(2) U-contaminated Waste Incineration (Waste Plants Operation Division at Tokai)

<u>Vertical fixed-grid fueled incinerator</u> is under hot operation, which object is to reduce the volume of combustible U-contaminated wastes gamerated at U handling facilities. The operational experiences and know-how will be utilized to the design, construction and operation of U-contaminated Waste Treatment Facility (UWTF).

(3) TRU-contaminated Waste Incinerator (Waste Plants Operation Division at Tokai)

<u>Moving-grid incinerator</u> is under demonstration operation at PWTF, which object is to reduce the volume of TRU Wastes generated at TRU handling facilities. The results will be reflected to the design, construction, etc. of the Low-level Waste Treatment Facility (LWTF).

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U-contaminated Wastes Incinerator	Study on Facility Concept	Basic Design	Detailed Design		Construc	tion Cold	Test	Opera	ition	
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TRU Wastes					Demonstr	ation Test				
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LWTF. etc.	Design	5	Design				1	Cold Te	st Opera	tion
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(4) Cyclone Incinerator (Tokai)

<u>Self-burning cyclone incineration</u> technology which is characterized by using goodcorrosion-resistant electro-melted Alumina for refractories, and non-moving parts, is under demonstration test at PWTF, in order to establish the technology of the volume reduction with incineration for Cl-contained wastes generated from MOX fuel fabrication facilities, etc.

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(5) B y Waste Incinerator (Tokai)

Moving-grid incinerator is under construction to reduce the volume of B 7 wastes from Tokai Reprocessing Plant.





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2. Decomposition

Decomposiontion is to transform organic materials into stable inorganic form except oxide by heating, etc.

(1) Immobolization of Spent Solvent into Inorganic Stable Form (Tokai)

RI tracer tests of the advanced technology characterized by incorporation of synthetic mica or hydrated lime and pyrolysis are being carried out to inorganize spent solvent.





Fy 1997 1998 1990 1991 1992 1993 1994 1995 1996 1989 lten RI Test Evaluation 1111 Immobilization of Spent Solvent into Inoganic Stable Form] -] -... 1 -]

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1. 18

37 Separation and Removal

Redioactive nuclides in liquid waste shall be separated and removed by chemical methods such as filtration, precipitation, etc., in order to change the liquid waste into nonradioactive or lower-level waste.

(1) Pu Recovery from Incinerated Ash (Tokai)

In order to reduce the volume of Cl-contained waste from MOX fuel fabrication facilities, etc. and to recover Pu, etc., adhered on the waste, tests of acid digestion characterized by low temperature treatment are currently being carried out at PWTF. The test results will be reflected to future TRU waste treatment facility.

(2) CP Separation/Removal Technology (O-arai)

Radioactive corrosion product(CP) separation/removal process with hollow film and reverse osmosis is under design at "Joyo" Waste Treatment Facility. The objective of the process is removal of CP contained in liquid waste from fuel washing. The results will be reflected to commercial FBR project.

(3) Separation of Long-lived Nuclides from Low-level Liquid Concentrate (Tokai)

In order to dispose of bituminized wastes from Tokai Reprocessing Plant in shallow land burial, long-lived nuclides contained in waste concentrate are planned to be separated and removed as pretreatment of Bituminization by ferrite treatment, precipitation, ion-exchange, etc. The results of cold tests obtained are currently evaluated, and the achievements will be fed to Bituminization Facility (AspF) for practical use.



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(4) Nuclide Separation from Spent Solvent (Tokai)

This technology is to reduce the radioactive concentration in the liquid waste below the regulatory level of sea discharge, by separation and removing the radionuclides in phosphoric-acid liquid waste with coprecipitation-ultrafiltration, etc., after inorganization of spent solvent into phosphoric-acid waste using hydro peroxid. The results will be reflected to the design of LWTF.

(5) Nuclide Separation from Low-level Liquid Waste (Tokai)

The objective of the technology is the volume reduction of low-level concentrated liquid waste, etc., generated in Tokai Reprocessing Plant. The technology will enable to reduce the concentration of the radioactive nuclides in liquid waste below the regulatory level of sea discharge, by adsorption, coprecipitation-ultrafiltration and ion-exchange. Cold basic test using simulated waste is currently implemented, and the results will be reflected to the design of LWTF.

(6) Iodine Removal from Sperit Solvent (Tokai)

In order to reduce iodine concentration discharged to the environment, hot test of removal(adsorption, etc.) of iodine contained in spent solvent, TBP and dodecame is currently being conducted. The results will be reflected to Solvent Waste Treatment Demonstaration Facility (STF) for practical use.

Fy 1990 1991 1992 1993 1994 1995 1996 1997 1998 1989 lien C/R Basic Nuclide Separa-tion of Spent Solvent ∇ Engineering Test Test EDLINTE Cold Hot Engineering C/R Basic Basic Test ∇ Nuclide Separa-Test Test tion of Lowlevel Liquid Waste =>LITF Development Test lodine Removal from Spent Solvent Design/Reconstruction of STF Evaluation Test

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4. Decontamination

Decontamination technology enables to remove radioactive nuclides adhered on solid wastes by chemical or mechanical methods. The objective is to change the radioactive solid wastes into non-radioactive or lower-levc: wastes.

(1) Melt Decontamination (Tokai)

Melt decontamination can achieve the volume reduction and Gocontamination of metal wastes, simultaneously. The electro-slag melting is currently under cold and hot testing at PWTF for volume reduction and stabilization of TRU-contained metal wastes. The test results will be reflected to the future TRU waste treatment facility projects such as LWTF, Hull Waste Treatment Facility (HWTF), etc.

(2) Electropolishing 1 (Tokai)

Electropolishing will be util'zed to decontaminate the surface of metal wastes for volume reduction of TRU contaminatec metal wastes generated at reprocessing plants and MOX fuel fabrication facilities. Hot test is currently being carried out using active metal wastes, and the results will be reflected to the practical use for TRU waste volume reduction hereafter

Fr 1989 1990 1991 1993 1994 1995 1996 1997 1998 1992 lien Melt Decontamination Practical Test PWTF (Hol) Evaluation Test Cold Test C/R (Nuclide Removal method Off-gas Bahavior) Cold Conceptual Operation Detailed Design Construction Test Design LUTT Conceptual Detailed Design Construction Basic Design Design HITF Hot Test of Electrodeposition Pu Recovery Test Regeneration Electropolishing Decontamination morespective waste treatment

(3) Electropolishing I (O-arai)

Electropolishing is removing the surface layer of metal wastes electrochemically as the wastes for anion in electrolyte. Cold plant-scale test and hot basic test of electropolishing is currently being implemented so as to develop the reuse technology of depleted electrolyte and the electrodeposition technology separating eluted metal ion from acid, as increasing the concentration of eluted metal ion in electrolyte. The aim of the tests is to make life of electrolyte longer.

(4) Redox (O-arai)

Redox decontamination is expected to be advantageous technology for miscellaneous waste decontamination, because of its applicability for complex shaped metal wastes and high decontamination effect. Utilizing Ce(IV) in nitric acid as strong oxidizing agent, it is currently under development through cold engineering tests, etc.



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(5) Ice Blasting (O-arai)

Ice Blasting has been developed as the decontamination technology in which water and dry ice are sprayed on to waste surface using air as the transfer media. It is regarded as good primary decontamination technology because of generating small quantity of secondary waste, and hot test has been conducted using actual waste, at the Waste Dismantling Facility (WDF) since Fy1984. Ice blasting will be applied more extensively for decontamination use of in-situ equipment surface and concrete, etc. aiming establishment of decommissioning purpose, hereafter.







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Expansion of application			o'					
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5. Dismantling

Dismantling is the cutting or sectioning technology for large solid wastes by the use of thermal and or mechanical techniques, in order to facilitate the handling of solid wastes.

(1) Plasma-arc Outting (O-arai)

Plasma-are cutting method, and related recovering methods of aerosol and dross, etc. are under development as remotely dismantling technology for large TRU-contaminated solid waste to obtain good efficiency of work, execellent dismantling capability and extended applicability.

In particular, automatic arc-cutting robot has been designed and fabricated for dismantling in-cell equipments. The achievements will be reflected to R&Ds of TRU waste treatment and decommissioning technologies.

(2) Laser Outting (O-arai)

Laser cutting technology, which is suitable for remote operation, because of generating less amount of aerosol or dross and being applicable to broad materials for cutting, is under development as the advanced, promising dismantling technology for large equipments, etc.

Especially, the basic research has been implemented on the cutting performance of COlaser, power transmission characteristics, etc. and application for the decommissioning of fuel cycle facilities is planned, increasing laser power step-by-step, hereafter.

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6. Immobilization (Stabilization)

Immobilization is waste stabilization technology by which solid waste including incineration ash and cut scrap, liquid waste and gaseous waste are solidified by wastes mixing with matrix, melting and so on.

(1) Micro-wave Melting (Tokai)

Micro-wave melting technology is characterized by homogenized heating(melting). Cold and hot demonstration tests of the technology have been currently implemented in order to stabilize and to reduce the volume of incineration ash or residue arisen from incineration and/or acid degestion. The test results will be reflected to the future TRU waste treatment facilities such as LWTF for practical use.

(2) Electro-slag Remelting (Tokai)

Electro-slagging technology is characterized by achieving volume reduction and decontamination simultaneously. Hot demonstration tests are now implemented at PWTF for volume reduction and stabilization of TRU-contaminated metals. The results will be utilized for the future TRU waste treatment facilities such as LWTF and HWTF for practical use.



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(3) HIP Solidification (O-arai)

Cold test of solidification by Hot Isostatic Press (HIP) method is under progress. The HIP technology is planned to be applied for volume reduction and stabilization of hulls and other metal wastes arising at Tokai Reprocessing Plant, and the achievements obtained from the development activities will be utilized for the facility design of HWTF.

(4) Dehydrating and Solidification with Micro-wave (Stabilization) (O-arai)

Dehydrating and solidification technology which converts the fuel washing liquid waste of "Joyo" (high-alkali property including corresion products), into glass-like block safely with simple facility, is under development. The results will be utilized to reconstruct the existent waste treatment facility.

(5) Solidification of Separated Nuclide Residue (Tokai)

In order to stabilize the separated nuclide residue removed from low-level liqued waste and spent solvent, an advanced solidification process is to be developed. In the present, solidification technologies is under investigation, and the results will be reflected to the design of LWTF.



(6) Hydrothermal Solidification (Tokai)

In order to solidify uncombustible residues, including incineration ash, spent silicagel, waste sand and spent iodine filter elements arising from Tokai Reprocessing Plant, solidification process applied hydrothermal reaction(Temp:100°C, Press:300 kg/cm³) is under cold basic testing. The results will be reflected to the design of LWTF.

(7) Cementation (Tokai)

In order to solidify uncombustible residues, including incineration ash, spent silicagel, waste sand and spent iodine filter elements arising from Tokai Reprocessing Plant, Cementation using cement-glass, low-hydrated cement, silica cement and portland cement is under basic testing. The results will be reflected to the design of LWTF

(8) Krypton Immobilization (Tokai)

In order to immobilize gaseous Krypton recovered from off-gas flow of Tokai Peprocessing Plant, Krypton is planned to be ionized and immobilized in metal. Cold basic tests are currently being implemented and the verification of process principle has been completed. The results will be utilized in Krypton Recovery Development Facility (KRF) for practical use.

(9) Plastic Solidification of Spent Solvent (Tokai)

In Solvent Waste Treatment Development Facility (STF), separation of TBP from spent solvent from Tokai Reprocessing Plant and its plastic solidification is under development operation to demonstrate the stable operation of STF.

Fy 1990 1391 1992 1998 1993 1994 1995 1996 1997 1989 lten Cold Basic Test Hydrothermal est WTF Solidification 1.04 Cold Basic Test Cementation SUNTE Basic Design | Licensing Fabrication Test Krypton =+KRF lmmobilization Development Operation Plastic Solidification of Spent Solvent

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00 Flame Retardation of Bitumen (Tokai)

The technology of flame retardation of bitumen by adding reagent before bituminization. is being tested (cold. laboratory-scale). The results will be utilized for the operation improvement of Bituminization Facility (AspF).

ov Bituminization (Tokai)

In order to demonstrate the bituminization process for liquid waste arising from Tokai Reprocessing Plant and the storage of bituminized wastes, developmental operation has been continued at AspF.



7. Solidified Waste Characterization

In order to establish the database for quality assurance of conditioning process and disposal, machanical and chemical characteristics of solidified wastes are planned to be evaluated.

(1) Evaluation of Sythetic Minerals Produced at PWTF (Tokai)

Solidified waste characterization such as leachability, etc., is conducted for performance assessment at PWTF using actual wastes to establish the safe and rational storoge and disposal of molten waste of incineration ash, etc. In addition, nuclide leach mechanism will be evaluated by observation and measurement of nuclide stability, taking account of disposal conditions. The results will be used to determine the conditions of solidification process, and to establish waste characterization database.

(2) Evaluation of Metal Ingots produced at PWTF (Tokai)

In order to establish the proper management of metal ingots produced from electro-slag remelting at PWTF, meterial properties including ingot compositions and distribution of nuclide concentration are being measured in cold and hot experiments. The obtained data will be used to determine the melting conditions of metal wastes and to develop the standards for reutilization of metal wastes.

(3) Compressive Strength Measuring Test of Plastic-Solidified Waste (Tokai)

Non-destructive compressive strength measuring tests are planned to control the spent solvent treatment process at STF, and the obtained data will be reflected to the operation of STF.

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Compressive Strength Measuring Test of Plastic-Solidified Waste		Prototyp) Test	cal ==STF													

(4) Evaluation of Bitumen (Tokai)

Mot evaluation test of bitiminized waste and plastic-solidified waste is being implemented to verify the soundness during the long-term storage at 2nd Bituminized Waste Storage and disposal. The results will be reflected to the plans of storage facilities and repositories

(5) Surface Contamination Measurement Test (Tokai)

Surface contamination measurement tests except smear method are planned to be implemented at 2nd Bituminized Waste Storage for contamination control of bituminized waste. The results will be utilized for practical use in AspF, and Shipping Facility, if necessary.

(6) Verification Test of Solidified Waste Homogeneity (Tokai)

Verification test of solidified waste homogeneity by non-destructive measurement is planned for quality control of bitumen waste at 2nd Bituminized Waste Storage. The results will be utilized for practical use in AspF, and Shipping Facility, if necessary.





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8. Source Term

Source term has to be established for obtaining the basic data for performance assessment of waste disposal.

(1) NDA (Tokai)

Non-destructive assay method incorporating τ activity measurement and neutron measurement is under development at PWTF in order to determine proper classification standard for TRU waste management. The obtained results will be utilized for evaluation τ^{*} nuclide inventory and for establishing classification standard value in TRU waste treatment processes.

(2) Scanner for Solidified Waste (O-arai)

Solidified waste scanner which enables automatic measuring evaluation is currently being investigated at Waste Management Section at O-arai. The objective of the study is quantitative and qualitative evaluation of total radioactivity and nuclides of glass-like solidified waste which incorporates radioactive corrosion products (CP). The results will be reflected to modification of "Joyo" Waste Treatment Facility.



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9. Facility Management

R&Ds for appropriate operation of waste handling facilities.are summarized.

(1) PWTF Operation Supporting System (Tokai)

Operating supporting system on the basis of knowledge database is currently being developed at PWTF. The objectives of the system are the improvement of the safety and reliability, and are the establishment of long-term stable operation of Pu-contaminated waste treatment process by integrating data related to design, operation and maintenance. It is expected that the facilities can be operated and maintained without skilled engineers by the support system. The results will be applied to the future waste treatment facilities such as LWTF and HWTF.

(2) Automation and Remotization of a Hall (O-arai)

By automation and remotization of the facilities in α -Decontamination Hall and α -Dismantling Hall of Waste Dismantling Facility(WDF), flogman work, man-hour and waste treatment cost have to be reduced as well as improving safety.

In FY1988, the sysytem optimization was studied. In FY1989, the basic design has been carried out to take shape of the whole system. The obtained results from this will be reflected to the future decommissioning projects of nuclear facilities.

(3) Hull Retrieval Technology (O-arai)

In order to retrieve the Hull cans piled in water pool of High Active Solid Waste Storage (HASWS) located in Tokai, the remote system comprised of underwater robot and large manipulators is currently under conceptual design. The results will be utilized for Hull treatment in the future Hull Waste Treatment Facility (HWTF).

FY 1995 1989 1990 1991 1992 1993 1994 1990 1997 1998 lien Demonstration Operation Actual Operation PACTE System Ĩ 14 Phase 1 Ш Development of Design Operation Supporting System LWTF. etc. Design Basic Study Desigt. - C/K Detailed Automation and Remotization of Design Lincensing a Hall Operation Decontamination/ Installation =>Decompissioning Design/Fabrication Mock-up Test Technology Development 1 C/R 1 C/R (Reference) Detailed Design/ Basic Safety Review Construction Design MASN'S Modification

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(4) Radiation Image Display (RID) (O-arai)

Radiation image Display (RID) which visualizes the distribution, quantity etc. of radioactive substances is currently being developed, as a part of development of radiation measurement technologies which aim efficient and economical operation of decontamination and dismantling (D&D), measuring contaminants and dose prior to determining D&D method.



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10. Miscellaneous

(1) Treatment of Alcoholate Liquid Waste (O-arai)

Advanced treatment technologies such as the catalytic oxidation method or the membrane treatment method is under investigation in order to treat the alcoholate liquid waste generated at "Joyo" Waste Treatment Facility during washing of fuel gripper. The information obtained will be reflected for practical use at "Joyo" Waste Treatment Facility.

(2) Purification of Recoverd Xenon (Tokal)

Xenon contained in off-gas of Tokai Reprocessing Plant is planned to be separated and purified by adsorption for reutilization. The basic test is currently being carried out and the results will be reflected to KRF for practical use.

(3) De minimis Level Measurement Technique (Tokai)

Since the dominimis level is required to be decided to establish disposal system of TRU waste which is arisen from MOK fabrication facilities, reprocessing plants and decommissioning, related measurement techniques have to be developed. In the present, the measurement techniques are under investigation evaluation. The results will be reflected to planning of the future waste management system.

(4) Entrusted Research of "Hot Treatment Test of Concentrated Liquid Waste" (Tokai)

Hot test of dry pulverization of concentrated liquid waste from reprocessing plant is under authorization procedure at AspF.

The test results will be used at Low-level Concentrated Waste Treatment Facilicy of Shimokita Reprocessing Plant.









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TECHNOLOGIES DISCUSSED AT JAERI- Takasaki Radiation Chemistry Research Establishment

- Amidoxime Group Containing Adsorbents for Metal Ions Synthesized by Radiation-Induced Grafting
- New Type of Amidoxime Group Containing Adsorbent for the Recovery of Uranium from Seawater

BIBLIOGRAPHY OF LITERATURE RECEIVED FROM TAKASAKI RADIATION CHEMISTRY RESEARCH ESTABLISHMENT

- "Amidoxime-Group Containing Adsorbents for Metal Ions Synthesized by Radiation-Induced Grafting", Written by J. Okamoto, T. Sugo, A. Katakai and H. Omichi. JAERI, Takasaki Radiation Chemistry Research Establishment, 11 pages.
- "A New Type of Amidoxime-Group-Containing Adsorbent for the Recovery of Uranium from Seawater", Written by H. Omichi, A. Katakai, T. Sugo and J. Okamoto. JAERI, Takasaki Radiation Chemistry Research Establishment. 16 pages.

Amidoxime-Group-Containg Adsorbents for Metal Ions Synthesized by Radiation-Induced Grafting

J. OKAMOTO, T. SUGO, A. KATAKAI, and H. OMICHI, Japan Atomic Energy Research Institute, Takasaki Radiation Chemistry Research Establishment, Takasaki, Gunma 370-12, Japan

Synopsis

Amidoxime-group-containing fibrous adsorbents for metal locs were synthesized by radiation-induced grafting of acrylonitrile followed by amidoximation of cyano groups with hydroxylamine. The degree of amidoximation and the distribution of amidoxime groups in the fiber were followed by means of electron probe X-ray microanalysis. The efficiency of adsorbing metal ions was increased by alkaline treatment of the adsorbent at high temperature for a short period before use. The order of adsorption for various bivalent metal ions was Hg > Cu> Ni > Co > Cd. From the distribution pattern of metal ions in the fibrous adsorbent, the adsorption was found to be controlled by the diffusion of the solution containing metal ions inside the adsorbent. It was found that confining amidoxime groups superficially and making short chain length of grafts were effective to obtain a high degree of adsorption.

INTRODUCTION

A variety of adsorbents for recovering metal ions dissolved in water or seawater has been reported.¹⁻⁴ Especially, adsorbents containing amidoxime groups which make chelate complexes with uranyl ions are noted for the recovery of uranium from seawater.⁶⁻¹²

These amidoxime-group-containing adsorbents are synthesized through the reaction of acrylic resins and hydroxylamine. The amidoximation, however, often causes a dimensional instability of the resin when used in aqueous solution because of large swelling in water.¹⁹ When the acrylic resin is made from a copolymer of acrylonitrile and a crosslinking monomer such as divinylbenzene to reduce the swelling, on the other hand, the ability to adsorb metal ions decreases tremendously.¹⁴ One of the causes is the decrease in the free movement of amidoxime groups due to the crosslinking among polymer chains. Therefore, both the stability to swelling and the free movement of the functional groups are essential to the adsorbant which is used in water.

The radiation-induced grafting is known as a method for introducing functional groups in a variety of polymers and inorganic substances.¹⁶ As polymer chains containing functional groups are chemically bonded with trunk polymers only at their chain end, a free movement of the polymer chains is maintained by this method. When a hydrophobic polymer is used as a trunk polymer, the part swollen in water can be restricted only to graft chains. Therefore, the two essential conditions mentioned above are satisfied by this synthesizing method. In addition, the distribution of the introduced functional groups is easily controlled by selecting reaction

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conditions such as irradiation dose, dose rate, temperature, and reaction time. However, the studies of synthesizing adsorbents for metal ions by radiation-induced grafting have been little reported.

In the present study, fibrous adsorbents containing amidoxime groups were synthesized by the radiation-induced grafting of acrylonitrile onto fibers and adsorption of heavy metal ions such as copper ions were attempted.

EXPERIMENTAL

Fibers used for synthesizing adsorbents are listed in Table 1. The fiber (ca. 40 μ m in diameter, and 15mm long) rinsed with acetone and dried in a vacuum oven for 16 h was packed in a polyethylene bag under nitrogen atmosphere and was irradiated with an electron accelerator (Dynamitron, Model IEA 3000-25-2, Radiation Dynamics) operating at beam energy of 1.5 MeV and a current of 1 mA at room temperature. In order to reduce the heat accumulation on the sample along with the irradiation, the polyet^{3,1}ylene bag was conveyed back and forth under the window of the accelerator at a rate 2.3 m/min. The dose rate was 1 Mrad/pass.

The irradiated fiber was installed in a glass ampoule and was evacuated for 5 min followed by the introduction of purified acrylonitrile under a nitrogen atmosphere. The graft polymerization was carried out at 25°C. When acrylonitrile was grafted in vapor phase, the fiber was separated from liquid acrylonitrile with a perforated plate. After the grafting, homopolymer and unreacted monomer were extracted with N,N-dimethyl formamide. The grafting yield was obtained from the weight increase based on the initial weight.

The amidoxime group-containing fiber (AO fiber) was obtained by heating grafted fiber with 3% hydroxylamine solution (methanol/water = 1/1) at pH 7 then rinsed with methanol and dried at 40°C under a reduced pressure. The amount of amidoxime groups was measured by elemental analysis.

The distribution of amidoxime groups combined with metals in the cross section of AO fiber was measured by means of a JEOL electron probe Xray microanalyzer (EPMA), Model JXA 733.

About 0.5 g of AO fiber was immersed in 1 L of metal ion (0.1-0.2 mM)containing Clark-Lubs buffer solution in the pH 2-7 region at 30°C for the prescribed periods. Then the fiber was rinsed with water and was dried in a vacuum oven. A concentrated sulfuric acid was added to a p¹2tinum

No.	Materials	Producers
1	Tetrafluoroethylene-ethylene copolymer [poly(TFE-E)]	Arabi Glass
2	Polypropylene (PP)	Ube Industria
3	Polyamide	Teijin
4	Polyethylene (PE)	Mitaui Petrochem
5	Polyester	Teifin
6	Carbon fiber	Toray

			TABLE I	
Fibers	Used	for	Synthesizing	Adapthenta

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crucible containing the fiber and was evaporated to dryness. This procedurs was repeated three times. Sodium carbonate was added to the ash of fiber, which was calcined at 550°C and melted. Then 1N hydrochloric acid was added to dissolve the melt. The amount of metal in the hydrochloric acid solution was measured by means of a Jarrell-Ash atomic absorption spectrophotometer, Model AA-845.

RESULTS AND DISCUSSION

Grafting

Grafting of acrylonitrile was attempted onto the preirradiated fibrous raw materials as shown in Table I. Figure 1 shows the grafting yield at different reaction time. A relatively high grafting yield was obtained with poly(TFE-E), PP, and polyamide. Especially, the grafting yield with poly(TFF E) reached ca. 60% in 24 h. The grafting yield with PE, polyester, and carbon fiber, on the other hand, was less than 5%, which is probably due to the low radical concentration in these irradiated polymers. From the results in Fig. 1, the following study was carried out mainly with poly(TFE-E). Moreover, it has a high heat stability and a sufficient resistance to chemical reagents such as base and acid solution due to C-F bonds in polymer structure.

Figure 2(a) shows the effect of diameter of poly(TFE-E) fiber on the grafting yield. The smaller the diameter, the higher the grafting yield. The rate of grafting obtained from Figure 2(a) was found to have a simple relationship with specific surface area of the fiber which was calculated from the fiber diameter as shown in Figure 2(b). These results are due to the fact that grafting is controlled by the diffusion of monomer into the fiber.¹⁶

As shown in Figure 3, both the grafting rate R_{p} and the final grafting yield G_{f} increase with the increase in the irradiation dose D. If the polymer radical is produced in proportion to the nth power of D, that is, $[\mathbb{R}_{p}]$



Fig. 1. Grafting yield of acrylonitrile onto preirradiated fibrous materials. The numbers are the same as shown in Table I.

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Fig. 2. Effect of diameter of poly(TFE-E) fiber on (a) grafting yield and (b) rate of grafting.
(a): (○) 40 µm; (Φ) 100 µm; (Φ) 150 µm; (Φ) 200 µm; (Φ) 250 µm.

= $k_i D^*$, where [R ·] is the radical concentration and k_i is the rate constant for initiation, respectively, R_i is expressed as

$$R_p = k_p[\mathbf{R} \cdot] \quad [\mathbf{M}] = k_i k_p D^*[\mathbf{M}] \tag{1}$$

where k_p is the rate constant for the propagation and [M] is the monomer concentration in the reaction site, respectively. The final grafting yield G_r at $t = t_f$ is obtained by integrating eq. (1) from t = 0 to $t = t_f$:

$$G_{f} = \int_{0}^{t_{f}} R_{p} dt = k_{i}k_{p} D^{n}[M]t_{f}$$
(2)

Equations (1) and (2) show that both R_p and G_f are proportional to D^* . From the results in Figure 3, n is estimated as ca. 0.5. The result that R_p is proportional to D^* agrees with the previous work in which styrene was grafted onto the preirradiated TFE Teflon.¹⁷

From the dependency of grafting rate on the reaction temperature the apparent activation energy was estimated as ca. 13 kcal/mol, which is



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comparable to the previously reported values for a variety of radiationinduced grafting.¹⁶

Amidoximation

As . own in the experimental section, AO fiber was obtained by converting cyano groups of graft chains to amidoxime groups. Figure 4 shows the effect of temperature on the amidoximation of the grafted fiber with average degree of grafting: 54-58%. At 40°C the conversion after 10 h was still less than 5%. The temperature was raised until it reached the boiling point of hydroxylamine solution (ca. 80°C). The conversion reached more than 60% after 6 h. The activation energy for amidoximation was estimated as 12 kcal/mol from the Arrhenius plots of the conversion as shown in Figure 4.

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The distribution of the amidoxime groups in the cross section of AO fiber was measured by means of EPMA. Figure 5 shows the distribution of amidoxime groups when the period of amidoximation was changed from 1 to 6 h at 80°C. It is clear that the amidoximation gradually proceeds from the surface to the center of the fiber and that it takes more than 6 h to obtain a homogeneous distribution.



Reaction Time (hr)



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Figure 6 shows the temperature effect on the width of distribution of amidoxime groups in AO fiber obtained from EPMA measurement. When the temperature is high and the reaction time of amidoximation is long, the distribution becomes widespread. The almost linear relationship of the width with reaction time indicates that amidoximation in the center of the fiber proceeds at a similar rate as on the surface region. The activation energy for spreading amidoxime groups in the fiber was estimated as ca. 10 | cal/mol. This value is in fair agreement with the above-mentioned activation energy for converting cyano groups to amidoxime groups.

Such thermodynamic similarity between the conversion of amidoximation and the spread of distribution of amidoxime groups inside the fiber is more clearly indicated when these values are plotted against each other as shown in Figure 7. Clearly, the conversion increases in proportion to the increase in the width under all the present experimental conditions. Therefore, it is reasonable to say that the amidoximation occurs homogeneously at least within 20 μ m from the surface of the fiber. Probably, cyano groups of graft chains exist homogeneously in the fiber and as soon as the hydroxylamine solution reaches the cyano groups the amidoximation occurs. In other words, the reaction is controlled by the diffusion of hydroxylamine solution in the fiber.

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width of Amidozime Layer from Surface(um)



Adsorption of Metal lons

The dried AO fiber did not adsorb metal ions at room temperature. One of the causes is the insufficient swelling of the fiber in metal ion-containing solution. The hydrophilicity of poly(TFE-E) fiber itself is virtually negligible and amidoxime groups are not so hydrophilic.¹⁰

Heating of AO fiber in alkaline solutin was found to increase the swelling of AO fiber in aqueous solution. For example, the contact of AO fiber with 0.5N potassium hydroxide solution at 20°C for 8 h elicited 30% augmentation in the water uptake.

The increase in hydrophilicity of the fiber is expected to increase the adsorption of metal ions. Figure 8 shows the effect of temperature of alkaline treatment on adsorption efficiency expressed as the percentage of amidoxime groups used for complexation with copper ions. It is clear that raising the temperature increases the efficiency. Although the efficiency abruptly




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increased when the fiber was treated with alkaline solution for a short period, the elongation of treatment period was not so effective to increase the efficiency. As shown in Figure 8, the efficiency levels off when the period is more than 8 h. In addition, the long contact of AO fiber with alkaline solution at high temperature should be avoided to reduce an undesirable decomposition of amidoxime groups.¹⁰ It was found that the treatment at 80°C for 10 min was the optimum condition to keep a high adsorption capacity with less decomposition of amidoxime groups introduced into poly(TFE-E).

When a sufficient period of adsorption was maintained, the adsorption of copper ion increased in proportion to the amount of amidoxime groups in the alkaline treated AO fiber as shown in Figure 9. From the slope, the number of amidoxime groups necessary for fixing one copper ion by complexation was estimated as, on the average, 3.3. As amidoximes are supposed to be bidentate, two amidoximes may be used for chelate formation with a copper ion which makes a square planar chelate. Therefore, the number 3.3 indicates that more than one extra amidoxime which does not directly participate in the chelate formation exists. Probably, the graft chains containing amidoximes have a loop structure, which makes it impossible that some of the amidoximes which exist in the middle of the loop take part in the chelate formation.

Figure 10 shows the effect of acidity of metal ion- containing solution on the adsorption of Hg²⁺, Cu²⁺, and Cd³⁺. About 0.1 mmol of Hg²⁺ was adsorbed per 1 gAO fiber at pH 2, while no adsorption of Cu³⁺ and Cd²⁺ was observed at this acidity. Cd²⁺ was not adsorbed until pH was beyond 4.

The amount of adsorbed metal ions increased with the increase in pH value. A steep increase was observed in the case of Hg²⁺ and Cu²⁺ when pH was more than 5. However, when pH was beyond 6 a precipitation of Cu was observed. The order of adsorbing various bivalent metal ions was

Hg > Cu > Ni > Co > Cd

which agrees with Irving-Williams order.18

Figure 11 shows the distribution of copper ions complexed with amidoxime groups at various contact periods. It is apparent that the migration of



Concent of Amidoxime Group(meg/g)

Fig. 9. Relationship between the amount of amidozime groups in AO fiber and the amount of adsorbed copper ions.

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copper ion through the amidoxime group layer is slow. It took about 1 day to obtain an almost homogeneous distribution of copper ions in AO fiber. The rapid increase in the population of copper ions on the surface region, on the other hand, suggests that confining amidoxime groups superficially on the adsorbent fiber is effective to obtain a high adsorption ability. This assumption was confirmed by the result shown in Figure 12 where the distribution of amidoxime groups in AO fiber and the adsorption amount







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Fig 12. Distribution of amidozime groups and adsorption amount of copper ions with the adsorbents made by (O) surface grafting and by (@) homogeneous grafting.

of copper ions were compared between two types of adsorbents-one made by surface grafting with acrylonitrile vapor and the other by homogeneous grafting using liquid acrylonitrile. It is clear that amidoxime groups in the surface layer (within 10 am from the surface) provide about twice as much adsorption of copper ions as the homogeneously distributed amidoximes do. Another possibility of obtaining high adsorption ability is shown in Figure

13, where AO fibers with similar grafting yield and therefore an almost constant amidoxime amount obtained at different irradiation doses were



Fig. 13. Effect of irradiation dose for grafting acrylonitrile on adsorption of copper ions.

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used. By increasing the irradiation dose, the number of trunk polymer radicals which can initiate graft polymerization increases. As the grafting yield is the product of the number of initiation radicals and the length of the graft chains, the result that these AO fibers have similar grafting yield indicates that the graft chain lengths are different according to the differthe in the number of initiation radicals. It is clear that the AO fiber of shorter chain length (or irradiated with higher dose) provides a higher adsorption capacity for copper ions. Probably, the shorter chains have more Dexibility of the movement of amidoxime groups for the complex formation

AO fiber has high stability to various kinds of treatment. For example, with copper ions. when dry AO fiber was treated with 2.5% potassium hydroxide solution at 80°C for 10 min, the diameter increased only 4%, which proves that AO fiber sweils little in alkaline solution. No further increase in diameter was observed when the alkaline treated AO fiber was contacted with water at 30°C for 24 h, while a considerable amount of water uptake (ca. 30%) was observed as mentioned above.

Thus, this new adsorbent produced by radiation-induced grafting proves very effective for the use in water because of high adsorption ability for

heavy metal ions and sufficient stability under various conditions.

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A New Type of Amidoxime-Group-Containing Adsorbent for the Recovery of Uranium from Seawater

H. OMICHI, A. KATAKAI, T. SUGO, and J. OKAMOTO

TAKASANI RADIATION CHEMISTRY RESEARCH ESTABLISHMENT JAPAN ATOMIC ENERGY RESEARCH INSTITUTE TAKASANI, OUNMA 370-12, JAPAN

Abstract

We new type of adsorbent containing amidoxime groups for the recovery of uranium from seawater was synthesized by the radiation-induced graft polymerization of acrylonitrile onto polymeric fiber followed by amidoximation with hydroxylamine. When amidoxime groups were introduced superficially on the fiber, the amount of uranium adsorbed by the amidoxime groups was higher than that with the amidoxime groups introduced homogeneously in the fiber. The introduction of the poly(acrylic acid) chain and the increase in temperature and flow rate in the adsorption process were effective in increasing the amount of adsorbed uranium. Although alkali metals and alkaline earth metals were found in the adsorbent, the concentration factors for these metals were less than $1/10^3$ of that for uranium. The present adsorbent had a high stability to various treatments such as contact with alkali and seawater.

INTRODUCTION

Separation of uranium from seawater has been studied with various kinds of inorganic (1-5) and organic adsorbents (6-15). Among them, amidoxime-group-containing polymeric adsorbents are noted because of the high loading of uranium and the rapid adsorption, rate. Recently, a fibrous adsorbent containing amidoxime groups synthesized through the amidoximation of a commercially available acrylic synthetic fiber with hydroxylamine has been used. It has been reported that the fibrous adsorbent has much higher adsorption ability for uranium when compared with a corresponding bead-type adsorbent (16).

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The fibrous adsorbent, however, has poor mechanical stability when it is contacted with alkali. This is supposed to be due to the hydrolysis of residual cyano groups in the polymer chain, which brings about the swelling of the whole fiber and a decrease in mechanical strength. Therefore, in order to synthesize a more stable fibrous adsorbent, a controlled amount of amidoxime groups should be introduced in the prescribed part of the fibrous material without changing the original mechanical strength. Radiation-induced grafting is a convenient method for such a purpose.

When a polymeric substrate is irradiated with 60 Co γ -rays or electron beam, about 10¹⁹ radicals are produced in 1 gram of polymer. The graft polymerization of a monomer is initiated by using these radicals when the monomer is introduced to the irradiated substrate. The number and the length of graft chains are easily controlled by irradiation and polymerization conditions. For example, the number of graft chains has roughly a linear relationship with the irradiation dose. The chain length is affected by the reaction time, the temperature, the presence of chain transfer agents in a monomer solution, etc.

When acrylonitrile is grafted onto fiber followed by amidoximation, a fibrous adsorbent is obtained. This adsorbent has the rescribed number and the length of amidoxime-group-containing graft chains which are connected with the trunk polymer fiber only at their chain ends. It is said that about one graft chain is connected with one trunk polymer on the average (17). In other words, the chemical modification of a trunk polymer by grafting is restricted to a very tiny region of the trunk polymer. Therefore, the chemical structures which were previously possessed by the trunk polymer are well maintained even after grafting.

It is another advantage of the radiation-induced grafting method that the graft chains can be introduced to any part of the substrate—for example, only in the surface region, the inner part, or in the entire substrate. According to the distribution of graft chains containing functional groups, the ability to adsorb metals is expected to differ.

In the present paper, several types of fibrous adsorbents containing amidoxime groups were synthesized by changing the grafting condition of acrylonitrile onto tetrafluoroethylene-ethylene copolymer fiber and were applied to the recovery of uranium from seawater.

EYTRIMENTAL

Fibrous adsorbents containing amidoxime groups (AOF) were prepared by the routes shown in Scheme 1.

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SCHEME 1. Synthesis of AOF by radiation-induced grafting.

Tetrafluoroethylene-ethylene copolymer fiber (40 μ m ϕ , 15 mm length) was irradiated with a Radiation Dynamics electron accelerator, Dynamitron, model IEA 3000-25-2, under nitrogen atmosphere at room temperature. The irradiation dose was estimated as 10 Mrd (100 kGy). The irradiated fiber was introduced in a glass ampule containing purified monomer under nitrogen atmosphere. The graft polymerization was carried

out at 25°C. After grafting, the homopolymer as well as the unreacted monomer were extracted with solvents: N_*N -dimethylformamide for acrylonitrile homopolymers and water for acrylic acid homopolymers. When both acrylic acid and acrylonitrile were introduced to the fiber, acrylic acid was grafted as the first step, then the grafted fiber was irradiated again followed by the introduction of acrylonitrile as the second step of grafting (18). The cyano groups in the grafted fiber were converted to amidoxime groups through the reaction with hydroxylamine (3% methanol/water 1:1 solution) at pH = 7. The conversion was determined by elemental analysis. The details were reported in a previous paper (19).

The distribution of amidoxime groups in the cross section of the amidoximated fiber was measured by means of a JEOL electron probe x-ray microanalyzer, model JXA 733.

The adsorption of uranium from seawater with AOF was carried out by the following four processes:

- Batch process: AOF was mixed with seawater in a vessel under vigorous agitation at 25 ± 1°C for the prescribed period.
- 2. Semibatch process I: The seawater in the batch vessel was intermittently exchanged for fresh seawater.
- Semibatch process II: Seawater was supplied continuously to the stirred vessel.
- Fixed-bed process: AOF was packed in a column (10 mm φ, 10 cm length) and seawater was continuously supplied.

The desorption of uranium adsorbed in 0.1 g of AOF was accomplished by contacting the fiber with 25 mL of sulfuric acid for 1 h at room temperature. The amount of uranium complexed with arsenazo III was measured optically (20) at 665 nm by means of a Shimadzu spectrophotometer, model UV-100-02. Metals other than uranium were determined by means of a Jarrell-Ash atomic absorption and flame emission spectrophotometer, model AA-8200.

RESULTS AND DISCUSSION

Preparation of Adsorbents

As mentioned above, one of the advantages of the grafting method for synthesizing adsorbents is that any amount of functional groups can be introduced to the trunk polymer fiber by selecting proper irradiation and

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grafting conditions. Several kinds of fibrous adsorbents containing amidoxime groups up to 6.5 meq/g were synthesized by changing the reaction time. Figure 1 shows the results when these AOF were used for uranium adsorption from seawater in semibatch process 1. As 2 L of seawater was exchanged with fresh seawater for 5 d, the total amount of supplied seawater was 10 L. The amount of adsorbed uranium increased in proportion to the increase in the concentration of amidoxime groups in AOF. Such relationships have been observed in various amidoxime-groupcontaining bead-type adsorbents (9).

Although the results in Fig. 1 show that the adsorption of uranium is facilitated by increasing the amount of amidoxime groups in the fiber, it should be pointed out that the amidoxime groups used for adsorption of uranium from seawater is only a small portion of the total amidoxime groups introduced in AOF. The molar ratio of the adsorbed uranium to amidoxime groups is of the order of 10^{-4} . Even if four amidoxime groups are necessary to make a chelate complex with one uranyl ion (21), the number of the amidoxime groups used is estimated $25 \sim 1/2500$ of the total amidoxime groups. In other words, the present adsorbents shown in Fig. 1 can, in theory, adsorb 0.5-1.5 mmol uranium per 1 g adsorbent. This value is reasonable when compared with the previous work by Schwochau et al. (21).

Two types of AOF, AOF-L and AOF-V, each containing ~5 meq of amidoxime groups per 1 g of fiber, were synthesized by liquid-phase grafting and vapor-phase grafting of AN, respectively. As shown in Fig 2, the distribution of amidoxime groups in the cross section of AOF-L

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observed by means of EPMA is homogeneous. On the other hand, in AOF-V the distribution is heterogeneous-mostly restricted to the layer within 10 µm from the surface. Because uranium in seawater is adsorbed in the thin surface layer of the adsorbent (21), only the amidoxime groups existing in the surface layer are supposed to be effective for adsorbing uranium.

The concentration of amidoxime groups in the thin surface layer of AOF-L was compared with that of AOF-V. The ratio between the two amounts is given as

$$F_1 \frac{S_0 - S_1}{S_0} / F_2 = (F_1 / F_2) \left(\frac{D_0^2 - D_1^2}{D_0^2} \right)$$
(1)

where F_1 and F_2 are the concentrations of amidoxime groups for AOF-L (5.1 meq/g) and AOF-V (5.0 meq/g), respectively, and Do and D, are the outer and inner diameters of cross section, respectively, as shown in Fig. 2. S_o and S_i are the corresponding areas. By introducing $D_o = 80 \ \mu m$, $D_1 = 60 \ \mu m$, $F_1 = 5.1 \ meq/g$, and $F_2 = 5.0 \ meq/g$, the ratio was obtained as 0.45. This result indicates that the concentration of amidoxime groups in a unit surface layer of AOF-V is twice as large as that of AOF-L. According

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FIG. 3. Adsorption of uranium from seawater with (O) AOF-V and (A) AOF-L in the semibatch process I.

to the results in Fig. 1, therefore, the adsorption of uranium with AOF-V is expected to be twice as much as that with AOF-L. However, Fig. 3 shows that the adsorption of uranium with AOF-V is slightly higher than that with AOF-L. This is a rather unexpected result because we observed that the adsorption of copper ions from a buffer solution at pH = 6 with AOF.V was about twice as much as that with AOF-L (22). In the case of adsorbing uranium from seawater, the functional groups existing on the surface layer are occupied by various kinds of metal ions other than uranium and are quickly covered with organic substances such as seaweeds. Therefore, there are fewer residual functional groups which work for adsorbing uranium compared with those functional groups existing inside an adsorbent like AOF-L. In the latter case, the surface layer may work as a filter for these obstacle substances. The result in Fig. 3 indicates that an adsorbent which is effective in a pure solution is not always effective in seawater. Therefore, the following experiments were carried out with the homogeneous adsorbent AOF-L.

Figure 4(a) shows the amount of uranium adsorbed from 2 L of seawater with 0.1 of three types of adsorbents, AOF-I, II, and III, which were made by liquid-phase grafting. The compositions of the adsorbents are shown in Table 1. AOF-I, containing only carboxyl groups, did not adsorb uranium at all. Referring to the report that poly(acrylic acid) can make a chelate complex with uranyl ions under acidic condition (23), the present result may be partly due to the insufficient pH value of seawater for chelate formation between carboxyl groups and uranyl ions. At pH = 8,



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TABLE 1 Composition of Adsorbents AOF

	Adsorbent	Concentration of functional groups (mmol/g)		
		Carboxyl	Amidoxime	
	AOF-I	5.2	0	
	AOF-II	0	5.1	
	AOF-III	1.9	3.4	

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poly(acrylic acid) dissociates into carboxylate anion, which may be undesirable for chelate formation.

When AOF-II and III are compared, AOF-III adsorbed more uranium in spite of the fact that the amount of amidoxime groups in AOF-III is less than in AOF-II. Therefore, the higher adsorption with AOF-III is supposed to be due to carboxyl groups. The effect of carboxyl groups on uranium adsorption was more clearly indicated when the molar ratio of adsorbed uranium to amidoxime groups was plotted against the adsorption period as shown in Fig. 4(b). AOF-III adsorbed twice as much uranium as did AOF-11. The effect of carboxyl groups on uranium adsorption with AOF is explained as follows: As already pointed out (14), uranyl tricarbonate, which is the major form of uranyl ion in seawater at pH = 8, is converted to bicarbonate when the pH value is lowered. Therefore, it is probable that poly(acrylic acid) graft chains locally decrease the pH of seawater sorbed in the adsorbent, which promotes the conversion from tricarbonate to bicarbonate. This conversion may be accompanied by a new chelate formation with uranyl bicarbonate and an amidoxime group in the vacant site. In other words, the presence of carboxyl groups is thought to have a synergistic effect on chelate formation between amidoxime groups and uranyl ions.



FIG. 5. Effect of temperature of seawater on adsorption of uranium with AOF in the batch process: (O) 15°C, (O) 25°C, and (O) 35°C.



FIG. 6. Effect of exchange rate of 2 L of seawater on adsorption of uramium with AOF-III in the semibatch process I: (@) every 1 d, (@) 2 d, and (O) 4 d.

Adsorption Method

The adsorption of uranium from seawater with AOF was carried out at 15 to 35° C. Although the equilibrium amount of adsorbed uranium did not change, the initial rate of adsorption was increased by increasing the temperature as shown in Fig. 5. From Arrhenius plots, the activation energy for adsorption was estimated as ~8.8 kcal/mol.

The results in Fig. 5 indicate that a higher temperature is preferable for the recovery of uranium from seawater with AOF. Especially when the adsorption-desorption cycle is carried out at short intervals, such a large initial rate is desirable.

As shown in Fig. 4(a), the amount of adsorbed uranium levels off after 7 day's adsorption in the batch process. About 0.1 g of AOF-III recovered more than 80% of the uranium contained in 2 L of seawater. The szturation of adsorption is, therefore, supposed to be due to the depletion of uranium in seawater contained in the batch vessel. In order to clarify this assumption, seawater was exchanged with fresh seawater every 1, 2, and 4 days in semibatch process I. Figure 6 shows the amount of adsorbed

AMIDOXIME-GROUP-CONTAINING ADSORBENT



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FIG. 7. Effect of flow rate and exchange rate of seawater on adsorption of uranium with AOF III in (O) semibatch process I, (Δ) semibatch process II, and (D) fixed-bed process.

uranium thus obtained. When the results in Fig. 6 are compared with those in Fig. 4(a), it is clear that uranium adsorption is facilitated by exchanging seawater. The adsorbed uranium was at most 50% of the total uranium in seawater, even if the exchange rate was small. This result indicates that the depletion of uranium was fairly avoided by this exchange procedure

Uranium adsorption did not increase in proportion to the exchange rate of seawater. Doubling the exchange rate brought about only a 20 to 30% increase in uranium adsorption. This adsorption behavior was observed when the flow rate was increased in semibatch process II and the fixed-bed process. As shown in Fig. 7, the increase in uranium adsorption levels off when the flow rate of seawater in semibatch process II and the fixed-bed process increases. It is interesting that the amount of adsorbed uranium obtained in semibatch process I is virtually the same as in semibatch process II and the fixed-bed process. This is probably due to the slow feed of seawater. For example, 10 L/d of flow, adopted in the present experiment, corresponds to only 8.8 cm/min of line velocity, which means that at least 2 s is necessary for one drop (\sim 0.2 mL) of seawater to get out of the column. Therefore, it is supposed that the flow rate in the continuous processes is slow enough to assure a sufficient contact of AOF with seawater as observed in the batch process with vigorous agitation.

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(6)

(a)





Characteristics of AOF

It has been reported that alkaline earth metals disturb the recovery of uranium from seawater (24). The smaller uptake of these metals is desirable as adsorbents for uranium. Although alkali metal ions and alkaline earth metal ions were reported to have small affinity for amidoxime groups (10),



1



-

Fig. 9. Distribution of (a) Mg and (b) Ca in the cross section of AOF-111 when contacted with seawater for 25 d.

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Based on Adsorption with AOF		
Metal ion	Abundance in seawater (g. L)	CP
U	3×10^{-6}	1.3 × 10 ⁴
Zn	5×10^{-6}	7.0 × 10 ⁵
Ni	7×10^{-6}	2.0 × 10 ³
Ca	0.41	29
ms	1.3	6.3
Na	10.8	0.34
K	0.39	0.13

TABLE 2 Natural Abundance of Various Metal Ions in Seawater, and Concentration Factors (CF)

a considerable amount of these metals was found in AOF after contact with seawater. Figure 8 shows the amounts of (a) alkali metals and (b) alkaline earth metals in AOF-III at different adsorption periods in the fixed bed process. The amount of such alkali metals as sodium and potassium gradually increases with an increase in the adsorption period (amount of seawater). The amount of alkaline earth metals, on the other hand, levels off quite soon, as shown in Fig. 8(b). From the distribution pattern of Mg and Ca in AOF as shown in Fig. 9, it is clear that these alkaline earths locate only in the surface region of the adsorbent even if AOF are contacted with seawater for a long period.

The equilibrium adsorption of Mg was ~2 mg, which is at least 10 times that of uranium. However, this large amount is due to the large content of Mg in seawater. Table 2 shows the natural abundances of various metal ions in seawater, and the concentration factors (CF) indicated are the ratios of the concentration in the adsorbent to that in seawater. It is clear that CF for uranium and some transition metals is over 104. On the other hand, CF for alkalis and alkaline earths is less than 102. These results indicate that amidoxime groups in AOF have a high selectivity for heavy metal ions.

The adsorbent AOF made by the radiation induced grafting method has some characteristic polymer structures when compared with the corresponding adsorbents made from commercially available acrylic synthetic fiber. First, the graft chains which contain amidexime groups are connected with the trunk polymer fiber only at the chain ends. Therefore, the mechanical properties of the trunk polymer itself is mostly unchanged, even after grafting. Second, swelling occurs only in the amorphous region of the trunk polymer because grafting is restricted to this region (17). This heterogeneous swelling behavior brings about a small dimensional change of the adsorbent when it is contacted with various solutions such as alkali,



ANIDOXIME GROUP CONTAINING ADSORBENT

			TA	BL	E 3				
Change in	Diameter	01	AOF	by	Contact	with	Vanous	Liqi	uid:

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2.74

a and the second second second	Liouid	Diameter (um)		
		and the second s		
	Crisinal AOF	66		
	alkalif	20		
	Seawater	72		
	COMPANY AND ADDRESS OF ADDRESS OF ADDRESS OF	In Property Line Construction and the second s		

*2.5% %.O.H solution, 10 min at 10*C.

*24 h at 30*C

acid, and senwater. As shown in Table 3, the increase in diameter of AOF by contact with these liquids is less than 10% on the whole. This dimensional stability is very important when the adsorbent is used repeatedly through the cycle: pretreatment with alkau-contact with seawater—acid desorption. A preliminary study showed that the decrease in adsorption of uranium and the dimensional change of AOF are both negligible up to cycle five. Further details will be presented in future publications.

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TECHNOLOGIES DISCUSSED AT PNC-CHUBU WORKS

- National Analogue Study on TONO Sandstone- Type Uranium Deposit in Japan

BIBLIOGRAPHY OF LITERATURE RECEIVED FROM PNC-CHUBU WCRKS

- "Field Tour Guide for Tono Mine Gallery (Tsukiyoshi Deposit)," PNC Chubu Works, 28 pages.
- "Natural Analogue Study of Tono Sandstone Type Uranium Deposit in Japan," Written by C. Sato, Y. Ochiai and S. Takeda. Waste Management and Raw Materials Division, PNC-Chubu Works, 11 pages.
- "Natural Analogue Study of Tono Sandstone Type Uranium Deposit in Japan," Written by T. Seo, Y. Coniai, S. Take'a and N. Nakatsuka. PNC-Chubu 6 pages.

Field Tour Guide

for

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Tono Mine Gallery(Tsukiyoshi Deposit)

Chubu Works Power Reactor and Nuclear Fuel Development Corporation(PNC)

JAPAN

Field Tour

Meeting Room :

- Brief Overview of R&D Activities of PNC at Chubu Works
- Introduction of Field Tour

Tono mine gallery : (see next three pages)

	STOP :	Engineered Barrier Materials Field Tests
-	STOP 2.3	Hydrogeochemistry of grounwater
-	STOP 4	Geochemistry of Natural U-Th Series Nuclides
		Uranium Wineralization
-	STOP 5	Mine-by Experiments on Excavation Responses
	A GOT2	Shaft Excavation Effect Project Site







Location Map of Test Area in Tono Mine and Adjacent Area





Location Map of Test Sites and Facilities in Tono(Tsukiyoshi Deposit) Mine.

STOP 1 ; Engineered Barrier Haterials Field Tests

Objectives

- (1) Evaluation of chemical durability of waste glass under field conditions.
- (2) Evaluation corrosion behavior of overpack materials under field conditions.
- (3) Establishment of experimental methods of field tests for engineered barrier materials.

Test items

- (1) Hydrological characterization of field test site:
 - Sampling and analysis of groundwater composition.
 - Periodic monitoring of groundwater:
 - pH.Eh.DD, conductivity and temperature measurement.
 - Groundwater flow measurement.
- (2) Engineered barrier materials corrosion tests under field conditions:
 - Leaching test of simulated waste glass: (Glass=P0798, Temperature=19°C, 90°C)
 - Corrosion tests of candidate overpack materials; (Specimen: mild steel, cast steel, copper. titanium alloy(ASTM G-12Ti), hastelloy C, Temperature:19°C)
 - Evaluation of specimens:

Corrosion rate(weight loss), analysis of alteration layer and corrosion products(SEM, XRD)

Result

- (1) Hydrological charaterization of field test site:
 - Tono groundwater chemistry (See Table 1); Na-HOO₂ Type
 - Hydraulic conductivity at Tono test site; 10^{-*}~10⁻⁵cm/sec
- (2) Corrosion behavior of engineered barrier materials under field condititions:
 - Waste grass (P0798) leach rate as a function of temperature (See Fig.2): The activation energy (65KJ) of waste glass alteration caluculated from field leaching tests was quite similar as that obtained from a soxhlet test in laboratory.
 - Overpack materials weight loss as a function of time (See Fig.2);
 Weight loss of test specimens (mild steel, cast steel and pure copper) in field tests at Tono test site were smaller than those in laboratory tests using Tono groundwater.

Very low corrosion rate at the field tests was obtained for Titanium.

	1 st Test Site	2 nd Test Site
Teep (t)	19	19
pH	8.9 ~ 9.1	8~9
Eh (mV)	242 - 296	160 - 260
Cond. (#\$/cm)	244 ~ 260	200 - 300
DO (ppm)	< 1	< 1
Ka'	46.8(mg/L)	65. 0 (mg/ l)
K*	0.32	3.0
C8°.	2.04	11.0
Mg 2 .	0.04	ND
fe?"	0.024	< 0.3
A13*	1. 2	r 6
C1.	1.0(eg/2)	1.5(mg/f)
50.1.	0.7	1.2
F ·	8.7	5.4
PD.	0.07	ND
HCD .	90	140
CO.***	11	< 3
Si	6.5(mg/2)	6.5(eg/2)
В	0.3	0.3
7-Fe	< 0.12	< 0.3

ND : Not Detected



Figure 1 - Waste gloss (P0798) leach rate as a farction of temperature ...





Figure 2 Overpack materials weight loss as a fanction of time

STOP 2; General Programme and Work Scope on Natural Analogue Studies

Objectives

- Contribution to validation of migration models in natural barrier for long-term safety assessment.
 - -To understand geochemical basis related to migration and fixation of

U-series nuclides for long-term prediction models.

(2) Contribution to the site investigation process.

-To develop the methodology and equipments for the characterization of

suitable geological environment for isolation of radioactive wastes.

(3) Contribution to establishment of public acceptance.

-To support the feasibility of geological isolation of radioactive wastes in Japanese geological environment.

Work Scope

The following studies are in progress to investigate fixation-migration

of U-series nuclides, and its relevant geological and geochemical environment.

(1) Migration and retardation studies uranium of series nuclides.

-Uranium series disequilibria in ore zone and around the Tsukiyosi Fault. -In-situ distribution coefficients -Matrix diffusion into granite boulders in ore zone.

-Transportation to biosphere.

(2) Geochemical study of groundwater.

-Geochemical parameters. -Characterization of natural colloids. -Geochemical modelling of groundwater.

(3) Hydrogeological study at Jono area.

-Hydrogeological parameters. -Modelling of groundwater flow system.

(4) Geological and geochronological study.

-Occurrence of uranium deposits. -Geological history of Tono area. -Mineralogy of host rock.

(5) Migration modelling.

-Data base of above studies.

Remarks

- General and economic geology have been studied i.e. stratigraphy, geological structure, geological history, ore distribution, ore grade ore charcteristic, mineralogy of host rock, etc.
- (2) There is no evidence of migration of uranium series nucldes such as 23*U, 23*U and 230Th for a period of at least 1 million years.
- (3) ²²⁶Ra has been migrated over a distance of several meters for recent thousand years.
- (4) The chemical and isotopic composition is charcterized in correspondence with the stratigraphy.
- (5) The tritium concentration in groundwater show that shallow groundwater is directry recharged by rainfall and discharge very quickly and the deep groundwater is stagnant.









Natural Analogue Program in Tono Uranium Deposits





Fig. Chemisrty of groundwater around Tsukiyosi deposit.



(1) Seto Group, (2) Mizunami Group, (3) Toki Group, (4) basement granite, (5) ores, (6) sampling point of waters.

The number represents the sampling point given in left figure. STOP 4; Geochemistry of Natural U-Th seies Nuclides (Disequilibrium of Uranium series Nuclides)

Objectives

- To estimate the time scale and spacial sacle of migration of U-series nuclides in sedimentary ore zone and fault zone.
- (2) To identify the minerals on which U-series nuclides are fixed.
- (3) To understand the geochemical mechanism related to migration and fixation of U-series nucides for long-term prediction model.

Horks

The measurement of U-series disequilibrium and analysis of associated minerals are in progress concerning the following samples.

- (1) Three dimensional grid samples in fresh ore zone of gallery.
- (2) Two dimensional grid samples in fault zone of gallery.
- (3) Drilled core samples in one body along the direction of groundwater flow.
- (4) Drilled core samples in the vicinity of fault zone.

Results

- Uranium has not been migrated over distances of 1 m for at least recent 1 million years.
- (2) ²²⁶Ra has been leached over distances of 1 m for recent thousands years.
- (3) Radioactive disequilibrium is observed within a few meters along fault zone.
- (4) Uranium is associated with various materials such as zeolite, clay titanium compound and organic carbon.



Sampling site for grid survey in the gallery.



Disequilibrium of 238U-234U-230Th-226Ra. D:25cm x 25cm x 25cm blocks. D:Whole block (lm x lm x lm).
STOP 5 ; Mine-by Experiments on Excavation Responses

Objectives

- (1) Preliminary study on the monitoring system of excavation responses.
- (2) Test and evaluation of the instruments and methods which zre currently available for the measurements of rock mass behavior.
- (3) Aquisition of the geomechanical and hydraulic data on excavation responses for the preparation of the further experiments in the actual deep underground research laboratory.

Horks

- (1) Pilot boring for the initial investigations.
- (2) Laboratory tests on the boring cores.
- (3) Geological mapping.
- (4) Measurements of the pourwater pressures.
- (5) Permeability measurements.
- (C) Measurements of the rock mass displacements.
- (7) Measurements of the axial stress of rockbolts.
- (8) Borehole loading tests.
- (9) Seismic tomography.
- 00) In-situ stress measurements
- QU Groundwater level monitoring.
- 0.2 Comparison of the actual rock mass displacements with the predicted ones by the F.E.M.model.



Results

- Rock mass displacement was almost terminated when the excavation face procceeded about 2 times of the drift diameter from the measuring point.
- (2) In-situ stress was not isotropic.
- (3) Permeability measurement was impossible at the zone of 0.5-1.0m from the drift face after the drift excavation.
- (4) Seismic survey suggested the low velocity zone of 0.8m thickness around the drift.
- (5) F.E.M. simulation result was consistent with the actual measurements assuming the excavation influenced zone of 1.On thickness around the drift.









③ Rock Mass Displacement (Extensometer)

@ Rock Mass Displacement (Convergence)

S Axial Stress of Rockbolt

(D) Borehole loading Test

② Seismic Tomography

(In-situ Stress (Hydro-fracturing method)

(In-situ Stress (Over-coring Method)











Rock Mass Displacement (by Extensioneter)



Objectives

- (1) Evaluate the mechanical and hydrological characteristics of rock mass which is influenced by the shaft excavation.
- (2) Evaluate the change of hydrological condition around the shaft.
- (3) Develop the repository design and the performance accessment of geological isolation for nuclide transport.

Works

 Measurement of the mechanical and hydrological changes of zone influenced by the shaft excavation.

(?) Numerical model development of the groundwater flow around the shaft.

(3) Natural analogue study.

Remarks

- (1) Following items are ... epared as the pre-excavation monitoring :
 - ① Tensiometers and piezometers for the monitoring of the subsurface water flow are installed.
 - ② Boreholes of up to 200m depth are drilled and geophysical loggings. BTV-monitoring and permeability measurement are performed.
 - ③ MP systems are installed and the multiple piezometric pressure measurements are being performed in the boreholes.
- (2) Shaft excavation was started in January 1990 and its present depth is 96m. During this period, some property measurements of disturbed zone has been done in the shaft.





SCHEMATIC FIGURE OF PROPETY MEASUREMENT OF DISTUBED ZONE (1)



4.

SCHEMATIC FIGURE OF PROPETY MEASUREMENT OF DISTUBED ZONE (2)



Hydrogeological Investigation for Regional Groundwater Flow

Objectives

- Development of methodology and equipment for analyzing the groundwater flow relevant to mechanism of radionuclide migration.
 - To develop method of hydrological, hydrogeological and hydrogeochemical investigation.
 - To develop equipments obtaining data relevant to hydrological.
 hydrogeological and hydrogeochemical characters
- (2) Development and validation of groundwater flow models for long-term safety assessment.
 - To understand hydraulic and hydrogeological characters related to the 3D-migration model in the regional area (12km×15km×depth 1.000m) including Tsukiyoshi Uranium deposites.
 - To develop regional three-dimensional hydrological models.
 - To validate reional three-dimensional hydrological models by hydraulic data obtained on hill-slope (surface and subsuface) hydrology and in borehole and drift.
- (3) Contribution to establishment of public acceptance.
 - -- To support the feasibility of hydrogeological isolation of radioactive waste in the Japanese rainy environment.

Works

The following studies are in progress to investigate hill-slope hydrology, hydrogeological characters. And it is also to develop the hydraulic equipment and to establish the model.

- (1) Investigation of hill-slope hydrology.
 - Lineament analysis by LANDSAT.
 - -- Vegetational and morphological analysis by aerial photograph.
 - Geological and topographic interpretation by mapping.
 - Zebra map and drainage interpretation by topographic map.

- --- Investigation of surface hydraulic characterization of evapotranspiration, river flow and precipitation.
- Measurement of specific discaharge and electric conductivity.
- (2) Hydrogeological characters.
 - 2-1 Core logging
 - Investigation of fracture characle isation(fracture pattern,filling materials.RQD etc.)
 - Measurement of physical properties (effective porosity, density and hydraulic conductivity etc.)
 - 2.2 borehole hydraulic investigation
 - Measurement of hydraulic parameters(hydraulic conductivity, pore pressure and groundwater flow velocity by tracer test).
 - Geophysical logs (sonic log, neutron log, BHTV and RADER etc.).
 - Measurement of groundwater physico-chemical parameters(Eh, pH, electric conductivity and groundwater composition etc.) and groundwater sampling.
 - 2-3 Hydraulic investigation in drift
 - ·- Geological analysis of fracture system.
 - Measurement of hydraulic parameters (hydraulic conductivity, pore pressure and groundwater discharge etc.).
 - Evaporation analysis on tunnel wall.
 - Geochemical analysis of groundwater composition and filling materials in fracture.
- (3) Development and validation of 3D regional hydrogeological models. (TAGSAC CORD developed by Dr. WATANABE, SAITAMA UNIVERCITY)
 - Data base of above studies.
 - Validation of regional three-dimensional hydrogeological model from hydraulic data obtaining on hill-slope hydrology, and in borehole and drift.
 - -- PNC Tracer Test System.

- (4) Development of hydraulic equipments.
 - PNC Aquifer Test System.
 - PNC Low Pressure Lugeon Test System.
 - Hydraulic Testing Machine(Laboratory Permeability Test).
 - PNC BAT Groundwater Sampling System.
 - PNC Tracer Test System.
 - PNC Geochemical Logging System.

Results

- Based on core observation, most of the fractures are classified as four types as follows. OPlaner type, @lrregular type, @Curved type, @Stepped type(Fig.1).
- (2) In-Situ hydrulic conductivity in the granite is approximately 10^{-*}~10^{-*} cm/s at fracture-predominants parts. 10^{-*}~10^{-*} cm/s at fracture-predominant parts in case of occurence of filling-minerals in fracture, and 10^{-*}~10^{-*} cm/s at fracture-poor parts measured by PNC Aquifer Test Method (Fig. 2).
- (3) In-situ hydraulic conductivity at fracture-predominant parts in the granite tends to descrease with depth as areas in some other countries (Fig.3).
- (4) Hydrogeological models has been developed for groundwater flow in the sedimentary rock and the granite, considering the recharge of water from the overlying high permeability Seto Group, the geochemical analysis of the surface-water and the groundwater, the geological survey, and the in-situ hydraulic test(Fig.4).

Natural Analogue Study of Tono Sandetone Type Uranium Deposit in Japan

C. Sato, T. Ochiai and S. Tekeda Waste Management and Rav Material Devision Power Reactor and Buclear Fuel Development Corporation (FMC)

SURMARY

Sandstone type uranium deposit, located in Tono area, cential mirt of Japan, has been recognized as a potentially useful analogue of geological isolation of radioactive wastes in Japan. The uranium deposit occurs as the erretiform which is less than 150 m in depth and below to water table. The studies area is not far from inhabitant. Preligioury study has been unde taket on migration of matural uranium series muchides and or hydrogeochemistry in Tsukiyoshi ore body. The radioactive disequilibrium study of drill core shows that the equilibrium has been simpst kept within the ore body. The hydrogeochemical study has revealed that there are three types of groundwater classified in correspondences with the stratigraphy and high-alkaline, fluorineriched groundwater is confined into, paleoweathered basement granite and perseable beds in the ore horizon.

The sim of this study is to reveal the geological, geochemical and hydrogical conditions which is favorable to keep uranium series nuclides undistur, of for a certain period of time, and to prove the feasibility of underground dispos i of radioactive wastes in Japan where peological sovironment is complicated and unstable.

1. IntreJuction

Japanese island arc> 's a part of the circum- cific mobile belt and not considered to have protogics' stability as the les such as Precamblian shield. Bowswar, geological isolatic of high level radioactive waste has been recognized to be basically feasible in Japan by means of a combination of angineered and natural barriers. Occurrence of uranium deposit as natural unalogue is thought to support the varification of geological isolation 870.4M.

Uranium deposits including small bodies in Japan, most of which are eacdstone-type bosted in Tertiery system overlying cretaceous granites. have been reviewed from the point of natural analogue. As a result, Tono deposit was chosen as the most favorable mos for the following reasons, (1) it is the only unwined deposit with minable ore grade, (?) it has a size which is roughly equivalent to that of the repository, (3) it is possible to investigate the influence of fault and groundwater on nuclide migration and (4) many information is slready available (1), (2), (3), (4), (5).

A boundary between oxidized and reduced some which is common in roll



3.2 Kineralogy

The mineralogy of samples in the gallery are shown in Table-I. The samples within ore zone mainly consist of quarts, plagioclass, kaolinite, montmortilonite, calcite and heulandite. Low amount of iron oxides as limo-nite are also present. In a few samples(No.5,6,7), pecondary uranium miner-a's as andersonite and sippeite are observed which may have mineralized due to oxidation after construction of the gallery. Sample No.11 contains a

463

The samplas were collected in the exploration gellery, and their locations are shown in Fig. 3,4. The redistribution of radionuclides after construction of the gallery is assumed for samples (Ro. 1- (0) taken from the sur-

radioactive anomaly accompanied with a fragment of carbonized wood (11) Drill core eamples

Drill core samples were taken from the uranium mineralized zone and

its neighboure at upstream and midstream in channel structures of the ors body. The locations of the drill boles, SM-2 and Al-12, are shown in Fig.3.

face of the gallery. Sample No.11, in the outside of the ore body, is a

(1) Samples in the gallery

The preliminary studies was made for ore samples which were taken from the gallery and drill core within the ors body. 3.1 Sample and Location

3. Distribution of Matural Uranium Muclices

Kisano, Utozaka and Jorinji. Tsukiyoshi ore body is roughly 3400 m by 500 m in size with thickness of 1 to 3 m. There is a reverse fault mamed Tauklyoshi Fault, Z-W, 60-70'S, with a throw of 33 m, which cuts the ore body. Shaft and gallery for exploration have been constructed in the middle part of the ore body. Present natural analogue study is undertaken within Taukiyoshi

The deposit is classified into sandstone type. Most of urgalum is characteristically accompanied with smolite, clay minerals, carbonaceous matter et al (2). Very small amount of uraninite and coffinite are observed. Tono uranium deposit consists of four ore bodies namely Taukiyushi,

Tono area is located in 350 km southwest of Tokyo. In this area, sedimentary rocks of Miccens age overlie uncomformably cretaceous granites (Fig. 1). The remerkable structual feature of uranium occurrence is paleochannel control on the plane of unconformity. The most favorable some in the changel for uranium mineralization is in the lower part of tertiary fluvial sediment of Toki Group which is composed of arkosic sandstone, taffaceous samdetone, carbonacaous mudatons and conglomerate. The depth of uranium occur-

2. Gaolosical Outline

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Preliminary study was cerried out on mainly muclides distribution and hydrogeochemistry in Taukiyoshi ore body of Tono deposit in order to discuso the availability of natural analogue study in this deposit.

front type deposit is not observed in this deposit. A reducing environment may have been dominant during and after the deposition. It is particularly important in metural analogue study to it. stigsts the geological, grochemical and hydrogical conditions which is favorable to keep utant - series ical and hydrogical conditions which is revolable to keep trans ... series nuclides utdisturbed. Product of this study also make a contribution to si-ts characts isation as well as to validation of model for performance asse-



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large quantity of sulfate minerals as jarosite and gypsus in addition to montmorillonite, isolinite and carbonized wood. Autoradiography and radiolunograpy abow uranium does not exist in carbonized wood but in clay minerale in this particular sample.

Except for small amount of pyrite, the mineral composition of the dri-11 core samples is similar to that of the gallery samples No.1-4.

3.3 Natural Dranium Series Disecullibrium

(1) Samples from the gallery Assay of D-238, D-234 and Th-230 were carried out by alpha spectrometry. Re-226 and Pb-210 were nondestructively analysed by high pure germanius detector. In the gamma-ray measurement, semie was kept in scaled container for three weeks and U-238 was calculated from the 63 keV gamma ray of the short lived daughter Th-234.

The results of assay are presented in Table-I together with the uranium contents. The relative activities of U-238, U-234 and Th-230 are plotted in ternary diagram Fig.5 (6). According to their radioactive disequilibrium states, the following geochemical processes which cause redistributions of uranium series nuclides can be infered.

No.1: U-238, U-234 and Th-230 are almost in radioactive equilibrium. This means that the age is older than at lisst a million years. On the other hand redium is elightly leached out by reduced groundwater fairly recently No.2,4,10,11: Dramium, preferentially 0-234, and radium is partially leached out probably by oxidited groundwater. While long 'ived daughter Th-230 which is almost immobile is present in excess state.

No.3,5,6,7,8,9: Accumulation of uranium occurred fairly recently by secondary introw of uranium, in which excess of U-234 was contained by alpha reco-11 process.

(11) Drill core samples

The D-234/U-238 and Th-230/U-234 activity ratios are shown in Fig.6. together with uranium content and geologic colum. There are no significant fractionstions between these nuclides in the two drill cores from upstream and midstream of the channel structure of Taukiyoshi ore body. However, the results of SH-2 cores suggest partial uranium leaching had occurred just above unconformity.

3.4 Selective Phase Separation

Sequential extraction techniques were applied to identify the phases of uranium series nuclides in ore. The following five fractions were separstad as idanticable phases: exchangeable, bond to carbonates, bond to iron oxides, bond to organic matter and resudial (7). Leaching procedures and reagants are summarized in Table-II, and the results obtained on samples from the gallery are shown in Fig. 7. A large portion of uranium is associated with carbonatas and iron oxides, even though total of iron and carbonace is less than 6 percent in weight for all of the samples.

Complamentary measurements as alpha spectrum and I-ray diffraction are not yet performed on the individual leachates and on the residuals following each extraction.

4. Hydrogeocuemietry

Uranium in Yopo deposit could be originated from the basement granites and transported by circulated groundwater through permeable host rock. The uranium deposit is assumed to be enriched by repetitions of leaching- and fixing cycle between bost rock and groundwater (3). The present ore horizon lies below water table and mostly in equiclude.

The general stratigraphy of Tomo area is composed of Toki Group, Misunami C., and Seto C. in ascending order. Groundwater samples were collected

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from each geological group for chemical analysis (Table-III, Fig.3.), There is a significant increase in pH, bicarbonate, sodium and fluorihe in correspondence with the stratigraphical depth. The groundwater in Toki Group where uranium is deposited is classified chemically as Na-BCO3 type. The high level of these ions seems to be caused by long term interaction between water and the minerals in host rock (4). The results of triti-

us analysis suggest that the groundwater is stagnate for more than 50 years. Rowever, the groundwater in Toki Group domen't contain higher level of uranium in contrast to higher bicarbonate content. The uranium concentration

These results suggest that the groundwater in ore borizon is stagnate

Therefore, the future studies should include the following subjects to

(1) Detailed study on nuclides distribution within and around the ore body. (2) Detailed geological and mineralogical study with a emphasis on uranium

(4) Further study on hydrogeology and hydrogeochemistry including develops-

The authors wish to thank Frof. Dr. T. Makanishi of Kanazawa Dniversity and Dr. T. Murano and Dr. M. Tamakawa of Power Reactor and Muclear Fuel

contribute the assessment of radioactive waste repository sites:

(3) Investigation on colloid formation and transport.

(3) Validation of developed model for nuclide migration.

Development Corporation for valuable suggestion.

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for a long time and the hydrogeochemical unvironment is reducing.

The results of preliminary study are summarized as follows: (1) & quite minor disequilibrium of uranium series nuclides is observed in (2) & large portion of uranium exist in the three fractions of carbonate, (3) Three types of groundwater is identified in correspondence with the st-(4) The groundwater of ore some is characterized by high content of bicarbonate, sodium and fluorine ion against the low content of wranium and These results suggest that this natural analogue study is available to reveal the favorable environment for the repository and to demonstrate the feasibility of the peological isolation as well as to understand the sucli-

is simpst as same as that of surface water.

3. Conclusions and Future Studies

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Sample RO	110			"The state	U. O. M	Main Minenels (*)
1	1.01	1.01	0.95	1.09	0.175	Qr. Pl. Loo . Cal
2	0. 84	1. 87	0. 90	1.02	0.100	QI. FI. Leo . Leo
3	1.04	0.94	0. 9.5	0.95	0.031	Qu . Pi . Nont . Cai
4	0.84	1. 94	1.05	0.92	0.121	Qa. Pi. Non I
5	1.01	0. 80	1.07	1.00	0.166	Q. PI . Leo . Leo . Cal
6	1. 19	0.75	1.01	1.11	0.091	Qi . Pi . Zeo . Lao
7	1.11	0.84	1.02	1.05	0.060	Q1 . PL . Teo
	1.00	0. 7 5	1.29	1. 08	0.097	Qu . Pi . Cal
9	1.01	0.72	1. 2 2	1.03	0.060	Qu. Pl. Mont. Kao
10	0. 94	1.12	0.87	1.03	0.067	Q FI . Ront . Kao . Zee . Cal
n	0. 96	4.94	0. 86	1.05	0.200	Host , Eao , Fel , Jar ,Gyp

Table-1. Activity ratios, uranium concentrations and minerals in the samples from the gallery

(*) Q: : Buartz - Pi : Plagloclase . Kao : Kaolisite . Ront: Monteorillosite

Cal : Calcita , Zeo : Zeolita (Nevlandita) , Pal : Fuidapar

Jar : Jarosite : Gyp : Gypsum

Tuble-II. Sequential extraction procedures for eelective phase separation

Praction	Extractant	Condition
].Erchaagaable	14 NaOAc 10ad (79 8.7)	Room Temperature Occasional Agitation 247
2. Boad to Carbosa iso	(H NaOAc(BOAc)Own (799 5.0)	b icaal AgiLation 24s
3. Boand to Fe Oxidens	0.041 HN,041 - BC1 12532HO4c 0 ml	90 T. 60 Occasional Agitation
4. Boad to Drganic Patter	(1) 3038.0, Seel +0.0298.0, 3eel (2) 30338.0, 3eel	(1) 85°C 681 Decessional Agilation (2) 85°C 29 Occasional Agilation
	(3) 3. 2M NIL, OAC + 203, MR . Sant	(3) Room Temperature Decasional AgiLation
5. Bond to Silicates	(1) HF 10.mf (HC10, 3.mf (2) 6/1 HC1	(1) Digestion (2) Disolution

466

Initial Weight of Sample : Is



tarrier acres







Table-III. Bydrogeochemical composition of waters around Tsukiyoshi ore body

and themes	1414	1	100a	U	-			*	3	2	2		*	Sample.
	1	12	(844)	1	((164)	(1444)	(844)	(=44)	(246)	(add)	(944)	(1/17)4)	
wrface	- Se	6.3	10	m	*	0.2	4	**	*		0.3	к 13	R	•
ant a grant	-8-	6.0	20	*	30	0.1	*	**	en		1-0	0.2	8	*
Sever-Jac.	8	7.2	\$	*	10	0.1	38	*	2	0.5	0.1	0.1		n
ut stat	200	8.5	8	*	*	•	33	0.3	*	0.1	0.1	0.1	0	*









Fig. 6. Furistion with depth of activity ratios and uranium concentrations for drill core samples from the upstream and midstream of Teukiyoshi ore body.





(w) 4 10+0











NATURAL ANALOGUE STUDY ON TONO SANDSTONE TYPE URANIUM DEPOSIT IN JAPAN

T. Seo, Y. Ochiai, S. Takeda, and N. Nakatsuka Power Reactor and Nuclear Fuel Development Corporation Toki City, Japan

ABSTRACT

The Tertiary Tono sandstone-type uranium deposit is recognized as a potentially useful analogue for the geological isolation of radioactive wastes in Japan. The study of this natural analogue is being carried out to acquire information about the geochemical processes that relate to the containment and migration of uranium series nuclides over geologic time. There is no significant migration for uranium for the samples investigated as the result of radioactive disequilibrium study. The hydrogeochemical study suggests that the goundwater in the ore horizons is very stagnant and reducing.

INTRODUCTION

Research and development of gelogical disposal for high-level radioactive waste have been conducted in accordance with a program announced by the Atomic Energy Commission (1987) in Japan. To implement gelogical disposal, it is essential to take account of the geological features. Japan is located in the Pacific rim unstable mobile belt, and its hydrogeology is characterized by a large amounts of rainfall. However, geological disposal has been considered feasible in Japan by means of a multibarrier system of engineered and natural barriers. In order to validate predictive mode's for safety assessment of a geological disposal system. the natural analogue study provides invaluable information and understanding for similar processes occurring over geological time and large spatial scales in a natural system.

The main purpose of this study is to clarify the

geological, geochemical and hydrogeological conditions that have been contained uranium successfully over geologic time, and to verify the fundamental process of safe isolation of radioactive wastes in a Japanese geological environment

Since 1986, PNC Chubu works has conducted a natural analogue study on the Tono uranium deposit. The main site of this study is limited within the Tsukiyoshi ore body, which is the largest ore body in the Tono uranium deposit. It is remarkable that the ore body is apparently cut by a fault that is available to investigate and to evaluate the effect of faulting on uranium series nuclides migration.

The main areas of the natural analogue study in the Tono uranium deposit are as follows:

- Investigation on occurrence of uranium series nuclides in a geological environment.
- (2) Investigation on migration of uranium series nuclides along a fault.
- (3) Investigation of hydrogeochemistry and hydrogeology of groundwater.
- (4) Evaluation of the role of colloids in the migration-retardation process.

DISCUSSION Geology

The Tono area, the largest uranium deposit, is located in central Japan, approximately 350 km southwest of Tokyo (Fig.1). Since the deposit was discovered in 1964, more than 560 holes of 50 m grids have been drilled. The 130 m-o. — shaft and galleries for the exploration tests hild already been constructed in the middle of the ore body.

The area is underlain by Tertiary sediments on the







basement rocks, which are composed mostly of granitic rocks ranging in age from the late Cretaceous to the early Paleogene period. Tertiary sediments are composed of the Mizunami Group and the Seto Group. The detailed regional geology is given Itoigawa (1974), and study of the uranium deposits are given in Katayama et al. (1974).

The Mizunami Group sedimentation occurred in a basin, which developed in the first stage of the Miocene transgression. The sequence is formed of lacustrine sediment which is overlain by marine sediment and divided into three formations. The lower unit has been correlated with the Toki lignite-bearing Formation, which is unconformably overlain by the Akeyo Formation and the Oidawara Formation. The general dipping of formation within the basin is from 0° to 10°.

The Toki lignite-bearing Formation consists of arkosic sandstone and mudstone with tuffaceous materials, interbedded with granule to pebble conglomerate. The sediments often contain carbonaceous and coal materials. The basal part is conglomeratic and consists of angular to subangular granite and guartz porphyry gravel. The basal conglomerate and several distinct parts of the upper beds are the principal uranium-bearing units.

The Akeyo Formation consists of fine to medium grained sandstone and mudstone containing many marine fossils. The Oidawara Formation, composed of siltstone and mudstone, is deposited in the deeper marine than that of the Akeyo Formation. The Mizunami Group is unconformably overlain by clay, sand and gravel beds of the Seto Group in Plice, he age (Fig 2).



 Fig 2 Schematic cross-section of the Tsukiyoshi ore body and sampling points of rock and water.
 (1) Seto Group. (2) Akeyo & Oidawara Formation.
 (3) Toki Formation. (4) basement granite.
 (5) ore body. (6) sampling site for grid survey.
 (7) sampling points of water.

Uranium Mineralization

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The Tono uranium deposit consists of several cre bodies, the largest of which is the Tsukiyoshi ore body as shown in Figure 1. The Tsukiyoshi ore body is the Tono uranium deposit in a practical sense because other ore body occur as satellites.

The Tono uranium deposit contains 4000 tons of contained U_3O_8 at an average grade of 0.06% U_3O_8 it is about 3400 meters in length and 500 meters in average width, with a thickness of 1 to 3 meters. The ore body is 100 meters to 150 meters in deep.

The uranium deposit is located in the basal part of the Miocene sediment and its distribution is typically controlled by paleochannel structures of the plane of unconformity on the basement. The uranium mineralization occurs as layers and is highly concentrated in the middle part.





types by the difference of host rocks: the conglomerate type and the lignite-bearing tuffaceous sandstone type. In the case of the former, uranium is adsorbed either in clay minerals such as montmorillonite or in calcite within the matrix of fine to coarse sandstone and siltstone. In high-grade ores, the uranium is characteristically associated with zeolite, which belongs to the heulandite-clinoptilolite group in the case of the latter, uranium is adsorbed in carbonaceous material which is rich in coal materials A very small amount of primary uranium minerals such as uraninite and coffinite are associated with pyrite and carbonaceous materials. According to fission track data, this sequence indicates an approximate age of 5-20 million years for the sedimentation. The age of mineralization is estimated to be about 10 million years. It is inferred that the Tono deposit is the egigenetic deposit.

In the Tono area, a reverse fault called the Tsukiyoshi Fault, runs along the central part of the channel and cuts the ore body. The fault strikes E-W and dips 65° to 70°S, with a replacement of 35 m. The Tsukiyoshi fault occurred before the deposition of the Seto Group.

Migration of Uranium Series Nuclides

Sato et (1987) discussed that there is no significant fractionation among 238U, 234U and 230Th for the two drill core samples in upstream and midstream of the channel structure of the Tono uranium deposit.

As shown in Fig.2 and Fig.3, a grid survey was carried out. The large block specimen of 1m³ was sampled by dividing it into blocks 25 cm x 25 cm x 25 cm in the fresh ore zone. The total of 64 samples were analysed individually to investigate the migration of uranium series nuclides



Fig. 3 Sampling site of ore for grid survey in the gallery

The uranium contents of these samples ranges from 0.01 % to 0.049 % U_3O_8 in every block. The thorium contents are less than 10 ppm. The data of 238U/234U vs. 230Th/234U activity ratio are plotted in Fig.4. and the data of activity ratio of the whole-rock (1m³) is also plotted for comparison. Although a slight deposition and leaching of uranium have occurred in some samples, the total activity of these nuclides is completely in equilibrium in the whole rock. It appears



Fig. 4 Disequiligrium diagram of 2380- 2340- 230Th for the sampes in the grid survey.

that these uranium series nuclides have not migrated more than 1m in the last one million years

In the same way, as shown in the disequilibrium diagram of 234U-230Th-226Ra (Fig.5), the depletion of 226Ra is observed in the most of samples and there is no enrichment of 226Ra. The total activity ratio of 226Ra/230Th in the whole block is 0.86. From this result, it might be inferred that the radium has been leached recently by reducing groundwater over a distance of several meters.

The mineralogical and chemical composition for these 64 blocks were also investigated to reveal the mineralogical and the geochemical condition where uranium is preserved for long time. The mineral assemblages of samples have large amounts of quartz, k-feldspar, plagioclase and small amounts of heulandite, kaoline minerals, smectite and others. Uranium minerals are not observed in any separated fractions. Although there is no evidence of correation between uranium and these minerals, chemical analysis show that uranium is correlated to iron, titanium and cation exchangeable capacity. Therefore, it is suggested that uranium is concentrated in clay, iron and/or titanium minerals.





Fig. 5 Disequilibrium diagram of 234U. 230Th. 226Ra for the samples in the grid survey.

This sampling method seems to be effective for investigating three-dimensional radionuclide migration.

Hydrogeochemistry and Groundwater System

The isotopic and chemical characteristics of groundwater and surface water in the area of the Tono uranium deposit were investigated. The results are summarized in Table1. In correspondence with the stratigraphy, there are significant differences in the hydrogeochemical compositions.

Groundwater in the aquifers of the Toki lignitebearing Formation, where uranium is deposited, were sampled from boreholes in the gallery. Some of the boreholes are intersected across the Tsukiyoshi Fault. These groundwater samples have similar characteristics, that is, lower values in 3H, 5D

Table 1 Chemical and isotopic composition of waters around the Tsukiyoshi ore body

бамрія На	-		N a 1 prest			U (MAA)	#203		1	E h (**)	JN	10	1180
Tr.		5	K1	ø	3	0.1	10	0	2		35	-16	
1	8	1	2	¢	1	9.1	20	0	1		35	• • •	12.5
3	,	2	ы	<0	x,	9.1	60	4	i		•	- 5 2	-1-1
•	•	i	3.5	10	1	Q.1~	90	3		- 300	a	- 5.5	-4.5
		4	34	<0	x	0.1-1	60				63	-55	4.4

and \$180 and higher values in pH, bicarbonate, sodium, and fluoride.

As shown in Fig 5, OD and O160 values of groundwater and surface water samples, which are important parameters of a groundwater recharging systems, fall on the line of OD = 8O160 + 13. This relation is in the range of the meteoric origin in the southern part of Chubu district of Japan. The OD and O160 values of the groundwaters in the Toki lignite-bearing Formation are lower than those for surface water and groundwater in the Seto Group over scale of 10 and 1 per mil respectively. Consequently, although the origin of these natural waters is rainfall, the source of groundwater in the Toki lignite-bearing Formation is clearly different from the present rainfall, which is represented by surface waters and groundwaters in the Seto Group.



Fig. 6. 6 D vs. 6¹⁸O of surface and ground voters around the Tsukiyoshi ore bore body. The number represents the sampling point given in Fig. 2.

The groundwater in the Seto Group is directly recharged by rainfall and discharged within a very short residence time through the loose sand gravel beds. The groundwater in the aquifer of the Mizunami Group, except for the Toki lighte-bearing Formation, which is overlain by the Seto Group, is more stagnant and has characteristics intermediate between those of the Seto Group and the Toki lighte-bearing Formation.

On the other hand, the groundwater in the Toki lignite-bearing Formation is stagnant for a long time in the deepest aquifer, and very little recharge of vertical flow occurs along the Tsukiyoshi Fault. The groundwater is reducing due to the presence of



Table 2. Hydrological parameters mesured in a	a chil ho	le.
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	Pore Pr		Perme al	ality	Welosity and	Direction
Depth Rock Type	JTT (m)	Rugeon (#)	JTT (CR/145)	Rugeon (ch/seb)	Velosity (cm/sec)	Direction
6 2-8 5 siltatone (OF)	-71		>10-7	•	>10-6	N D.
9.0-14.0 siltstone (O.F.)	11.50	-7.4		5 x 10-7		fick i g
33 5-38 5 conglomerat	e -31.1	-31.3	1.3 × 10-5	3 x 10-4	2.5 x 10-5	$S \longrightarrow N$
38 5-39 5 sandstone (A.F.)	-28.7		>10-7			
40.5-45.5 conglomera (A.F.)	te -30.9	-28.0	>10-1	>10-7	>10-6	N.D.
48.5-53.0 sandstone (A.F.)	-28.5	-28.4	9 x 10-6	5 × 10-7	>10-6	N.D.
57.0-62.0 sandatone (A.F.)	-27.3	-27.2	>10-7	4 x 10-4	>10-6	N.D.
69.6-74.6 sandstone (T.F.)	-26.9	-27.3	.2 x 10-5	7.1 × 10-4	>10-6	N.D.
SO 0-85 0 sandstone	1.*	-29 9	>70-1	5 x 10-6	>10-6	N . D .

O.F. Oldawara Formation A.F.: Akeyo Formation T.F. Toki Formation, N.D. Not Detectable

organic materials in the Toki lignite-bearing Formation.



A hydrogeological investigation on the regional groundwater movement is also in progress Understanding the dynamics of groundwater is important in evaluating the effects of Japan's heavy rainfall on the migration of the uranium series. It is emphasized that investigation of hill-slope hydrology is necessary as well as the hydraulic characteristics of deep groundwater to establish a regional hydrogeological model of this area; core logging, geophysical logging and hydraulic testing have been under way.

The result of a hydrogeological also indicates that the shallow groundwater in the Seto Group is directly recharged by rainfall and that the groundwater in the Mizunami Group is stagnant for a long time in the deepest aquifer.

The hydrological parameters such as pore pressure, flow direction and velocity measured in the borehole are in Table 2. There are various values of permeability which depend on rock type. Almost all groundwater velocity values are lower than 10⁻⁶ cm/sec.

The Investigation of colloids and other components such as microorganisms in groundwater is important to evaluate the effect on radionuclide migration. The physical and geochemical composition and characteristics of natural colloids are also investigated

- CONCLUSION
 - General geology and geology have been studied i.e. statigraphy, geological structure, general geological history, ore distribution.

ore grade, ore characteristic, mineralogy of host rock, etc.

- (2) There is no evidence of migration of uranium series nuclides among 238U, 234U, and 230Th. On the other hand, 226Ra is inferred to have been leached recentry by reducing groundwater more than several meters.
- (3) The chemical and isotopic data of groundwater show that the shallow groundwater in Seto Group is directly recharged by rainfall and discharge very quickly and the deep groundwater in the Mizunami Group is stagnant.

FUTURE WORK

- Investigation on migration of uranium series nuclides nuclides in order to understand the geochemical migration processes.
- (2) Investigation on distribution of uranium series nuclides along the fault to evaluate effects of faulting on uranium series nuclides migration.
- (3) Investigation of hydrogeology and hydrogeochemistry to understand the threedimensional water transport pathways and how they affect nuclide migration.
- (4) Interpretation of geological events by age determination in order to estimate time scale and paleo-environment.
- (5) Investigation of the role of colloids and microorganisms in nuclide migration. These studies are under way through cooperation with national research institutes, universities and private consultants.

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A SELECTIVE CO SENSOR USING TI-DOPED a-Fe2O3 WITH COPRECIPITATED ULTRAFINE PARTICLES OF GOLD*

TETSUHIKO KOBAYASHI, MASATAKE HARUTA, HIROSHI SANO and MASANORI NAKANE

Government Industrial Research Institute of Osaka, Midorigaoka 1, Ikeda 563 (Japun) (Received January 15, 1987; in revised form June 30, 1987; accepted October 15, 1987)

Abstract

Titanium-doped α -Fe₂O₃ with ultrafine deposits of Au was prepared by the coprecipitation method. A thick-film sensor fabricated from the above semiconductor had high sensitivity and excellent selectivity to CO against ethanol and H₂ in the temperature range 30 °C to 100 °C. This unique sensing property was found to originate from a high catalytic activity of the material, which could catalyse CO oxidation even at a temperature as low as -70 °C. High-resolution electron microscopic observation has revealed that gold ultrafine particles with a mean diameter of 36 Å are homogeneously dispersed and that they are not merely supported on but strongly held by the host oxide α -Fe₂O₃-Ti⁴⁺. The sensing mechanism and the origin of the high selectivity to CO are discussed.

1. Introduction

There are growing needs for selective detection of CO in both domestic and industrial sectors of society, because it is toxic even at a concentration as low as 100 ppm. Selective CO gas sensors are indispensable for the detection of incomplete combustion, the prevention of poisoning in mines, and the detection of early stages of a fire.

Gas sensors based on a conductivity change of n-type semiconductive oxides, such as ZnO [1] and SnO₂ [2], have been widely used in gas-leak alarms. Several oxidation catalysts have been used as sensitizing additives to the above semiconductive oxides in order to control the sensitivity and the selectivity to reducing gases [3, 4]. In the case of CO detection, the addition of Pt [5], Pd [ε -8], CuO [7, 9], or Pd, MgO and ThO₂ [10] can increase the sensitivity. However, since these sensitizers also tend to increase the sensitivity to ethanol and/or H₂ [11 - 13], it has been difficult to obtain sufficiently high selectivity for CO.

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We have recently reported that a novel catalyst composed of a-Fe₂O₃ and ultrafine particles of gold (UFP-Au) with an optimum atomic ratio of Au:Fe = 5:95 can catalyse the oxidation of CO even at a temperature as low as -70 °C [14]. This catalyst exhibits remarkably enhanced activity in comparison with those of Au powder and α -Fe₂O₃; when these single compounds were used, the CO oxidation needed much higher temperatures, 200 °C for α -Fe₂O₃ and 300 °C for Au powder. Furthermore, the UFP-Au/or-Fe2O3 catalyst is so stable that it does not lose its high catalytic activity during continuous oxidation of CO for one week in a moistened atmosphere at room temperature.

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It is reasonable to expect that the high catalytic activity of the UFP-Au/a-Fe2O3 catalyst may lead to a high sensitivity to and high selectivity of CO at low temperatures, if suitable modification of the catalyst can be made. The present study has been undertaken in an attempt to make n-type semiconductors from UFP-Au/ α -Fe₂O₃ and to utilize them as a CO gas sensor.

2. Experimental

2.1. Preparation of semiconductive materials

Haematites $(\alpha \cdot Fe_2O_3)$ doped with tetravalent metal ions were prepared by coprecipitation from a mixed aqueous solution of $Fe(NO_3)_3$ and each of SiCl₄, Ti(OH)₃(OOH), SnCl₄ and ZrOCl₂ with an aqueous solution of Na₂CO₃. The precipitates were washed with distilled water then vacuum dried and calcined in air at 400 °C for 3 h. Titanium-doped α-Fe₂O₃ with ultrafine gold particles (UFP-Au/α-Fe₂O₃-Ti⁴⁺) was prepared by a similar coprecipitation method using a mixed solution of HAuCl₄, Fe(NO₃)₃, and Ti(OH)₃-(OOH). For comparison, the following four kinds of materials were also prepared; (1) Photodepo-Au/ α -Fe₂O₃-Ti⁴⁺ was obtained by the photodeposition method [15] from a mixed aqueous solution of HAuCl4, ethanol and a dispersion of α -Fe₂O₃-Ti⁴⁺ powder under the illumination by a mercury arc lamp (100 W) for 10 h. (2) Au-sol/ α -Fe₂O₃-Ti⁴⁺ was prepared by coprecipitation from a mixed solution of $Fe(NO_3)_3$ and $Ti(OH)_3(OOH)$ with a Na_2CO_3 solution containing Au-sol. The gold sol was previously prepared by the reduction of HAuCl₄ with formaldehyde [16]. (3) Impreg-Au/a-Fe₂O₃-Ti⁴⁺ was obtained by impregnation of a HAuCl₄ solution into α-Fe₂O₃-Ti⁴⁺ particles. (4) Au/SnO2 was prepared by coprecipitation from a mixed solution of HAuCl4 and SnCl4. These materials were calcined at 400 °C for 3 h and were further calcined at 600 °C for 1 h.

2.2. Characterization

X-ray diffraction (XRD) analysis was made by using the Rad-B system (Rigaku Denki Co. Ltd.). The mean particle diameter of α -Fe₂O₃ and Au was estimated from half-widths of the (104) peak of α -Fe₂O₃ at $2\theta = 33.15^{\circ}$ and the (111) peak of Au at $2\theta = 38.18^{\circ}$ using Scherrer's equation [17].

To observe the fine structure of the material, a high resolution transmission electron microscope (TEM) (H-800, Hitachi Co. Ltd.) operated at 200 kV was used.

2.3. Fabrication of thick-film sensors and gas sensitivity measurements

An aqueous paste of the gas-sensing material calcined at 400 °C was deposited on an alumina plate on which a pair of comb-like Au electrodes was previously printed. Both the width of the Au electrodes and that of the gap between the electrodes are 0.2 mm, and the total length of the electrodes is 77 mm. The amount of gas-sensing material used for one sensor device was about 10 mg per cm² of the alumina plate. The thick film was finally heat treated at 600 °C for 1 h in air before measurements. The sensor device thus obtained was set on a temperature-controlled hot plate in a test chamber (7000 cm³) that was filled with clean air humidified to 65% r.h. The change in resistance of the device was measured when a reducing gas, such as CO, ethanol or H₂ was introduced by a gas syringe into the test chamber. The sensitivity is expressed by the ratio of the resistance measured in air alone, *R*-air, to that measured in the presence of reducing gases, *R*-gas.

2.4. Catalytic activity measurements

Catalytic activities of the materials for CO oxidation and H₂ oxidation were measured by using a small fixed-bed reactor with 0.2 g of the materials, which had passed through 70 and 120 mesh sieves. A standard gas consisting of 1.0 vol.% CO or H₂ balanced with dry air to 1.0 atm was passed through the catalyst bed at a flow rate of 66 ml min⁻¹. Conversion efficiencies from CO to CO₂ or from H₂ to H₂O were determined by the gas chromatographic analysis (G-1880, Yanagimoto Co.) of the effluent from the reactor. The catalytic activities were expressed by the specific temperatures, denoted as $T_{1/2}$, at which 50% conversion efficiency was attained, and therefore lower $T_{1/2}$ implies higher catalytic activity for the oxidation.

3. Results

3.1. Doping of metal ions into a-Fe₂O₃

The resistance of a thick-film device fabricated from UFP-Au/ α -Fe₂O₃ catalyst (Au:Fe = 5:95 in atomic ratio) was over 10 M Ω at 250 °C in air. This resistance value is so high that a precise measurement of the resistance change caused by reducing gases is difficult without sufficient amplification. Therefore, several tetravalent metal cations like Si⁴⁺, Ti⁴⁺, Sn⁴⁺ and Zr⁴⁺ were used to dope α -Fe₂O₃ by a coprecipitation method in order to increase its n-type semiconductivity. The resistances of the thick-film devices prepared from the doped α -Fe₂O₃ were measured at 250 °C in air and the results are shown in Fig. 1 as a function of the radius of the dopant ions. Titanium ion, Ti⁴⁺, which has a nearly identical ionic radius to that of Fe³⁺, was found to decrease the resistance of α -Fe₂O₃ most effectively. Figure 2 shows that



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Fig. 1. The conductivit, shange of α -Fe₂O₃ on doping with tetravalent metal cations. The resistance was measured at 250 °C in air in the thick-film device. (a) Si⁴⁺; (b) Ti⁴⁺; (c) Sn⁴⁺; (d) Zr⁴⁺; (e) dopant free (Fe³⁺). M⁴⁺: Fe³⁺ = 1:99 in atomic ratio.

Fig. 2. Dependence of the conductivity of a Fe₂O₃ at 250 °C on Ti⁴⁺ concentration.

the resistance of α -Fe₂O₃ could be minimized at a composition of Ti:Fe = 3:97 in atomic ratio. Based on the above results, Au/ α -Fe₂O₃-Ti⁴⁺ (Au:Fe = 5:95, Ti:Fe = 3:97 in atomic ratio) has been chosen as the most promising gas-sensing semiconductor to be examined.

3.2. Sensing performance of Au/Ti-doped a-Fe2O3

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Figure 3 shows the sensitivities to CO, EtOH and H_2 of the thick-film sensors fabricated from Au-loaded and -unloaded α -Fe₂O₃-Ti⁴⁺ as a function of sensor operating temperature. Gold-loaded α -Fe₂O₃-Ti⁴⁺ exhibits a significantly high sensitivity to CO at temperatures below 100 °C, while having maximum sensitivities to EtOH and H_2 at around 150 °C and 250 °C, respectively. From the comparison with the results obtained for Au-un'oaded α -Fe₂O₃-Ti⁴⁺, it is clear that the presence of Au leads to a shift of the temperatures for the maximum sensitivities towards lower temperatures, and that this shift and the enhancement of the sensitivities are most remarkable for CO detection.







Fig. 4. The sensitivities of UFP-Au/ α -Fe₂O₃-Ti^{4*} sensor operated at 40 °C as a function of gas concentration. ----, CO₁ ----, EtOH₁ ------, H₂.

Fig. 5. The response time of UFP-Au/ α -Fe₂O₃-Ti⁴⁺ sensor to 100 ppm CO. - - -, 50 °C: ----, 40 °C; -----, 30 °C.

At temperatures below 50 °C, an excellent high selectivity to CO against EtOH and H₂ was obtained. Figure 4 shows the dependence of the sensitivities on the concentrations of these gases at 40 °C. The sensitivity obtained for 20 ppm of CO corresponds to those for 1000 ppm of EtOH and for a much higher concentration of H₂. Figure 5 shows that the sensitivity reached 90% of the final steady-state signal after 40 s for 100 ppm of CO at 40 °C. This response at such a low temperature as 40 °C can be considered to be sufficiently rapid.

3.3. Effect of the particle size of gold and of the kind of host metal oxides on sensing properties.

Figure 6 shows the sensitivities to CO and F_2 of α -Fe₂O₃-Ti⁴⁺ with Au deposits of different particle size. The sensit⁴ day as well as the selectivity to CO is markedly enhanced by a decrease in the Au particle size. In addition, CO sensing properties are dependent on the kind of host metal oxide on which the Au particles are supported. The sensitivity of Au/SnO₂ to CO at 50 °C, also shown in Fig. 6, is lower than that of photodepo-Au/ α -Fe₂O₃-Ti⁴⁺, although the particle size of Au on SnO₂ is smaller than that of Au photo-deposited on α -Fe₂O₃-Ti⁴⁺.

Figure 7 shows the catalytic activity of the semiconductors for CO oxidation and for H_2 oxidation. The same tendency as observed in the gas sensitivities can also be seen in the catalytic properties of the materials for the oxidation of CO and H_2 .

3.4. Characterization of UFP-Au/a-Fe₂O₃-Ti⁴⁺

Figure 8 shows an X-ray diffraction pattern of UFP-Au/ α -Fe₂O₃-Ti⁴⁺. Several sharp peaks due to α -Fe₃O₃ and appreciably broadened peaks due to metallic Au can be seen. From these peaks, the mean particle diameters of α -Fe₂O₃ and Au were estimated to be 170 Å and 30 Å, respectively. As TiO₂ could not be detected by X-ray diffraction, Ti⁴⁺ seems to be dissolved in the lattice of α -Fe₂O₃ to form a solid solution [18]. Since the particle size of


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Fig. 6. Variation of the gas sensitivity at 50 °C with the particle size of Au. (a) UFP-Au/ α -Fe₂O₃-Ti⁴⁺; (b) Photodepo-Au/ α -Fe₂O₃-Ti⁴⁺; (c) Au·sol/ α -Fe₂O₃-Ti⁴⁺; (d) Impreg-Au/ α -Fe₂O₃-Ti⁴⁺; (e) Au/SnO₂. Su content in atomic ratio. (a) · (c), Au:Fe = 5:95; (d), Au: Fe = 1:99; (F), Au:Sn = 5:95; (c); 300 ppm CO, (c); 300 ppm H₂.

Fig. 7. Variation of the catalytic activity for the oxidation of CO (CD) and H₂ (WW) with the particle size of Au. (a) UFP-Au/ α ·Fe₂O₃-Ti⁴⁺; (b) Photodeno-Au/ α ·Fe₂O₃-Ti⁴⁺; (c) Au·sol/ α ·Fe₂O₃-Ti⁴⁺; (d) Impreg-Au/ α ·Fe₂O₃-Ti⁴⁺; (e) Au/SnO₂. Au content in atomic ratio: (a) · (c), Au :Fe = 5.95; (d), Au :Fe = 1.99; (e), Au :Sn = 5.95.

 α -Fe₂O₃, *i.e.*, 230 Å, in Au/ α -Fe₂O₃ was larger than that in Au/ α -Fe₂O₃-Ti⁴⁺, the doping of Ti⁴⁺ might lead to the formation of smaller particles in the coprecipitates.

Figure 9 shows the fine structure observed by TEM of Au/α -Fe₂O₃-Ti⁴⁺ calcined at 400 °C. Since both the XRD pattern and the catalytic activity of the material calcined at 600 °C were nearly identical with those for the material calcined at 600 °C, the fine structure of the material calcined at 600 °C might be almost the same. Gold ultrafine particles are homogeneously dis-





persed on the surface of $\alpha \cdot Fe_2O_3$ particles of about 200 Å in diameter. The diameter of 2663 particles of Au in the enlarged TEM photographs was measured by using a slide caliper and their distribution is plotted in Fig. 10. The mean particle diameter of Au and the standard deviation were calculated to be 36 Å and 13 Å, respectively. The perticle diameters of $\alpha \cdot Fe_2O_3$ and of Au determined from TEM coincided well with those estimated by XRD and indicate the validity of the XRD method.

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Figure 11 shows that, typically, at the junction between Au particles and α -Fe₂O₃, the basal planes of hemispherical Au particles are in contact with the surface of α -Fe₂O₃ particles. From the interplanar spacing of the crystallites, it can be seen that the (111) plane of a Au crystallite is grown on a α -Fe₂O₃ (110) plane. Such a crystal orientation is often but not always observed at Au/ α -Fe₂O₃ junctions. Furthermore, ultrafine Au particles are stable on α -Fe₂O₃-Ti^{a+} and did not coagulate under the electron beam radiation in a TEM, while ultrafine Au particles vacuum deposited on SiO₂ change their crystal structure in less than 0.1 s and occasionally move on the SiO₂ surface to coagulate with each other [19].

Neither the strongly bound interface nor the crystal orientation could be observed by TEM for impreg Au/ α -Fe₂O₃-Ti⁴⁺, where spherical particles of Au were grown to a size larger than 200 Å in diameter and were present as a simple mixture with α -Fe₂O₃-Ti⁴⁺ particles.







Fig. 11. The fine structure of the interface be ween Au and α -Fe₂O₃. Note that the interplaner spacings of Au(111) and α -Fe₂O₃ (110) are 2.355 Å and 2.519 Å, respectively.

4. Discussion

4.1. Sensing mechanism

It has been shown that the gas-sensing semiconductor, UFP-Au/ α -Fe₂O₃-Ti⁴⁺, can selectively detect CO with a high sensitivity at temperatures

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below 100 °C (Figs. 3 and 4). Some Pd-added SnO₂ sensors [6, 8] have also been reported to detect CO at temperatures below 100 °C. Gold-loaded α -Fe₂O₃-Ti⁴⁺ has a sensitivity and selectivity comparable to those of the Pd/SnO₂ sensor of best performance reported so far [20]. However, there is a marked difference in the response time between the two materials. The response time with UFP-Au/ α -Fe₂O₃-Ti⁴⁺ is less than 1 min (Fig. 5), while it is over 10 min with Pd/SnO₂ [6].

Another big difference is the catalytic activity for CO oxidation; i.e., over UFP-Au/a-Fe₂O₃-Ti⁴⁺, CO oxidation takes place even at -70 °C (Fig. 7), while it commences only at tround 100 °C on Pd/SnO₂ catalysts [21]. Hence, at the temperatures for which both of the materials have high CO sensitivity, CO oxidation takes place vigorously on UFP-Au/a-Fe₂O₃-Ti⁴⁺ but not on Pd/SnO₂. This difference strongly sup last that there should be a substantial difference in the CO successing mechanism between these two materials.

The activation mechanisms of the gas-sensing semiconductors sensitized by Pd, Pt and Ag have been classified into two types [22]. In the first mechanism, the reducing gases are adsorbed and activated on the sensitizers, and then the activated gases react with the oxygen adsorbed anionically on the semiconductor surfaces. The resultant removal of the surface oxygen causes an increase in the conductivity of the n-type semiconductors. In this case, a high sensitivity is expected to appear as a reflection of a high catalytic activity for the oxidation of reducing gases at the same temperature range.

The second mechanism is an electronic one in which the sensitizers create such surface states as to directly exchange electrons with the semiconductors. The reducing gases are adsorbed on the sensitizers and then donate electrons to the semiconductors via the surface states originated from the sensitizers, which also results in an increase in the conductivity of the n-type semiconductors. In this case, catalytic activity for the oxidation is not necessarily needed for sensing reducing gases.

From the relation between the sensitivity and the catalytic activity at tend, evaluates below 100 °C, the first mechanism would be valid for UFP-Au/- α -Fe₂O₃-Ti⁴⁺, and the second for Pd/S₃O₂. In the latter case, it is likely that CO molecules will be adsorbed too strongly on the Pd surfaces [23] to react with the surface oxygen on SnO₂ at such low temperatures. This may be responsible for the very slow response time of Pd/SnO₂.

4.2. Origin of high CO sensitivity

Carbon monoxide oxidation takes place at temperatures above 200 °C on α -Fe₂O₃ [14], where the rate-determining step is the adsorption of CO onto the α -Fe₂O₃ surface [24]. Such a catalytic nature of α -Fe₂O₃ is reflected in the CO-sensing property of α -Fe₂O₃-Ti⁴⁺ without UFP-Au (Fig. 3). If UFP-Au provides sites for the activation of CO and then facilitates its transfer to the α -Fe₂O₃ surface, the fast CO oxidation at low temperatures and hence the high sensitivity to CO of UFP-Au/ α -Fe₂O₃-Ti⁴⁺ will be explainable.

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Since the surface of pure gold is known to be essentially inactive to most chemicals, including CO [25], it can be suggested that the surface property of Au is significantly changed in UFP-Au held on the α -Fe₂O₃ surface.

The first experimental evidence is the size effect of Au particles shown in Figs. 6 and 7. The UFP-Au particles on α -Fe₂O₃-Ti⁴⁺ are so small that a large part of the constituent Au atoms is exposed to the surface. The number of constituent Au atoms in an Au sphere with a diameter of 36 Å is calculated to be about 10³, and about 40% of the Au atoms exist on the outermost surface of the particle. The catalytic properties of such ultrafine metal particles are expected to differ significantly from those of bulk metals [26], due to an increase in the surface dangling bonds [27] and/or a change in the electronic structure due to quantum size effects [28] with a decrease in the particle size. In fact, it has been reported that atomic Au can react with both CO and O₂ even at 10 K and that the resultant Au(CO)₂O₂ complex decomposes to produce CO₂ at 40 K [29].

Secondly, it can be pointed out that 'metal-support interaction' [30] may govern the chemical properties of UFP-Au. Electron microscopic observation showed that UFP-Au particles were not spherical but hemispherical and were strongly held on the host oxide of α -Fe₂O₃-Ti⁴⁺. The activity to CO at low temperatures was appreciably dependent on the kinds of host metal oxides; on SnO₂, instead of α -Fe₂O₃, UFP-Au catalysts did not exhibit high catalytic activity (Fig. 7). Bond *et al.* [31] have proposed that ultrafine Au particles supported on Al₂O₃ or SiO₂ become electron deficient by donating electrons to the supports and the catalytic properties of the electron-deficient Au particles then resemble somewhat those of Pt, which is the element to the left of Au in the Periodic Table.

It is well known that CO is adsorbed strongly on Pt in contrast to the weak adsorption on 'inherent' Au and that CO adsorbed on the Pt surface is too stabilized to be oxidized by O_1 at temperatures below 150 °C [23]. The adsorption energy of CO on Pt is 110 - 140 kJ/mole [23], while that on 'inherent' Au is about 40 kJ/mole [32]. If UFP-Au becomes electron deficient by donating electrons to the host oxide of α -Fe₂O₃-Ti⁴⁺ and then has intermediate CO adsorption between that of Pt and 'inherent' Au, UFP-Au can provide suitable sites for the CO adsorption, which would facilitate reaction with O₂ adsorbed on the α -Fe₂O₃ surface.

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Biographies

Tetsuhiko Kobayashi received B. E. and Doctor of Engineering degrees from Osaka University, in 1978 and 1983, respectively. From 1983 to 1984 he remained at Osaka University as a postdoctral fellow of the Japan Society for the Promotion of Science to continue his work on electrochemistry. Since joining GIRIO in 1984, his research has focused on the development of new gas sensors.

Masatake Haruta received a B. E. from Nagoya Institute of Technology in 1970 and a Doctor of Engineering from Kyoto University in 1976. Since joining GIRIO in 1976, he has worked on the preparation of combustion cately its and gas-sensing oxide materials.

Hiroshi Sano received a B. Sc. from Niigata University in 1955 and a Doctor of Engineering from Kyoto University in 1976. He has been the director of the Material Chemistry Department in GIRIO since 1986, and his current fields of interest are catalysts and their application to new energy technology and pollution control.

Masanori Nakane received a B. E. from Kumamoto Institute of Technology in 1948 and a Doctor of Engineering from Osaka University in 1971. He was the director of the Material Chemistry Department in GIRIO until 1986. His major activities are concerned with the development of new meterials.

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M. HARUTA1, H. KAGEYAMA1, N. KAMIJO1, T. KOBAYASHI1, and F. DELANNAY2 "Government Industrial Research Institute of Osaka, Midorigaoka 1, Ikeda, 563 (Japan)

FUniversité Catholique de Louvain. Department of Material Science and Processes, Place Sainte Barbe 2, 8-1348 Louvain-la-Neuve(Belgium)

ABSTRACT

Highly dispersed gold catalysts have been prepared by calcining the copricipitates obtained from an aqueous solution of MAUCI4 and the nitrate of Fe. Co. Ni. or Be. They are active for the oxidation of CO even at such a low Fe. Co. Ni. or Be. They are active for the oxidation of CO even at such a temperature as -70°C and become more active in the presence of moisture. The gold particles are hemispherical in shape and are nearly homodispersed with a mean diameter smaller than 10nm. Most of them are strongly attached at their flat planes to the metal oxide support exhibiting a specific crystal prientation. The chemical shifts in XPS showed that the ultrafine gold particles were electron deficien. The results of EXAFS studies suggested the possibility of the existence of gold atoms coordinated with iron atoms.

INTRODUCTION

Gold has attracted little attention as a catalyst because of it. inert character and the difficulties in preparing highly dispersed small particles. Schwank noticed unusual activities and selectivities of monometallic gold catalysyts in his review1). However, gold catalysts have been considered in many cases not to be competitive with other noble metal catalysts in terms of activity. This remains to be true even in the small supported gold particles?).

An interesting discovery made recently is that gold supported on borosilicate catalyzes the oxidation of many organic compounds by nitrogen dioxides3.4). The reactions are efficient and highly selective and are being applied to the detection of butylated hydroxytoluene(BHT). On the other hand, we have also found recently that coprecipitation enables is to prepare highly dispersed gold catalysts and that the combination with the oxides of group vill 3d transition metals5-7) and alkaline earth metals $^{(8)}$ makes yold so active that it can catalyze the caldation of CO. even at -70°C. The soove gold catalysts are now under way for the application to the removal of CO from air, the oxidation of CD in sealed CO2 laser, and selective CD pas sensors9).

The present paper deals with the characterization of the novel gold catalysts, specifically Au/a-FegO3 and Au/BeO, through XPS, EXAFS, and TEM. The mechanism is also discussed of the oxidation of CO on these _____id catalysts in ______ moistened air at a room temperature.

EXPERIMENTAL

The gold catalysts were prepared by coprecipitation with sodium carbonate from an aqueous solution of HAuClg and the nitrates of various metals. The coprecipitates were washed, vacuum dried, and calcined in air at temperatures from BO°C to 500°C. Catalytic activity measurements were carried out in a small fixed bed reactor, with 0.20g of catalysts that had passed through 70 and 120 mesh sieves. A standard gas consisting of 1.0 volZ H₂ or CO balanced with air to 1 atm. was passed through the catalyst bed at a flow rate of 66 ml/min.

The structures of the gold catalysts were observed using a Hitachi H-9000 electron microscope operated at 300 kV. X-ray photoelectron spectroscopic analyses were made using a Shimadzu ESCA 750 under a vacuum below 5×10^{-6} torr. The binding energies were calculated by reference to that of C1s carbon contamination peak assumed at 285.0 eV. Measurements of EXAFS and XANES were carried out at the beam line 108 of the 2.5 GeV storage ring of Photon Factory in the National Laboratory for High Energy Physics(KEK,Tsukuba). Data analysis was made following the method of Maeda et al.¹⁰).

RESULTS

Catalytic properties of gold catalysts prepared by coprecipitation

Table 1 shows the catalytic activities of the gold catalysis with different metal oxide supports for the oxidation of H₂ and CO. The activity is expressed in terms of the temperature corresponding to 20% conversion($T_{1/2}$). The particle sizes of gold were calculated from peak half-widths of XRD and in some cases from TEM photograghs.

There are two groups of metal oxide supports that can generate remarkably high catalytic activities for the oxidation of CO at -70°C. They are the oxides of group VIII 3d transition metals like Fe. Co. and Ni and the oxides of alkaline earth metals like Be and Mg. In contrast to the casalytic nature of pure gold that the oxidation of H2 occurs at lower temperatures than that of CO, the above gold catalysts can catalyze the oxidation of CO at much lower temperatures. On the contrary to the metal oxide supports alone which are seriously deactivated by moisture, the coprecipitated gold catalysts become more active in the presence of moisture. In the experiments carried out at 30°C using the large grain of Au/a-Fe2O3(7-9 meshes) calcined at 400°C under a space velocity of $4x10^4$ h⁻¹, the oxidation efficiency of CO was raised to 100% from 95% by the addition of water in a range from 0.6 to 4.0 vol%. These characteristic properties may permit us to distinguish the coprecipitated gold catalysts from the conventional ones which are prepared by impregnation on MgO,

A1203, and \$1021,2).

The oxides of transition metals other than Fe. Co. and Ni, namely ScgOg. LegOgs TiOg, ZrOg. CrgOg, etc., appeared to be far less effective as a support because they did not give rise to active catalysts even when gold particles smaller than 5 nm had been prepared. Semiconductive oxides like CuO, ZnO, IngOg. SnOg were relatively good supports to exhibit high catalytic activities for the oxidation of CO. All the above metal oxide supports changed the catelytic nature of gold and made it more active for CO oxidation than for Ng oxidation, however, as far as AlgOg and SiOg supports were concerned, gold retained its original nature.

The above results strongly support the indication by Schwank¹¹) that the

TABLE 1

Oxides	Au content (stom2)	Calc.Temp. (°C)	71/2[H2] (*C)	T1/2[CO] (*C)	Diameter of Au(nm)
BeOa) MgOa.b) CaOa.b)	520	200 200 200	«0 67	<-70 κ-70 5	41) (21)
Sc203 Le203e)	5	600 200	94 205	60 92	very small
T102 2+02	5 5	400 400	108 141	63 111	5 very small
Cr203	5	400	212	155	≥30
MnOg	10	400	152	100 get.	**
Fe203 Co304 N10	5 5 10	400 400 400	27 56 67	«70 «70 «70	4,1f) 6 f) 8 f)
CuO	5	400	154	24	***
2 n0 6 d0	5	400 200	60	*=700) 206	2 1 2 1
A1203 1n203	5	300 400	66 68	84 510)	5
\$102°) \$n02	5	300 400	184 54	204 0	20 3
Au Fe203 Co304 N10	100	120 400 400	103 314 139	295 190 81	c.a.20

Catalytic activities of gold catalysts prepared by coprecipitation for the exidation of H2 and CO, and mean clameters of cold particles.

a) hydrous oxides, b) deposition-reduction, c) deposition-precipitation d) 20% conversion at -70°C, e) 20% conversion at 1°C, f) by TEM

type of support might play cruicial role in determining the catalytic activity of gold in reactions of oxygen-bearing molecules. Especially, in the cases of gold catalysts prepared by coprecipitation the metal-support interactions could strinkingly change the nature of gold due to the small sizes of particles. Among a variety of supported gold catalysts, Au/α -Fe2O3 and Au/BeO, that were the most active, have been chosen as samples for the further investigation.

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Effect of calcination temperature on catalytic activity

Figure 1 shows the dependence of the conversion in H2 exidation at 30°C and . CO exidation at -70°C on the calcination temperature for the Au-Fe coprecipitate. Figure 2 shows the dependence of $T_{1/2}$ values for the exidation of H2 and CO on the calcination temperature for the Au-Be coprecipitate. The contents of Au in these catelysts were 5 atom2, which had been proved to be the optimum content.

In the Au-Fe coprecipitate, a maximum catalytic activity was obtained by calcination at 200°C for H₂ oxidation, but for CO oxidation it was obtained by calcination at 300°C. X-ray diffraction confirmed the sample calcined at 200°C as poorly crystalline hematite(α -Fe₂O₃) with no other apparent crystalline compounds. In the samples calcined at 300°C only hematite became more crystallized. At 400°C, broad diffraction peaks due to metallic gold appeared.







Fig. 2. Effect of calcination temperature on the catalytic activities of Au-Be coprecipitate for the oxidation of H2 and CO and on specific surface area. A, H2: O, CO; D, specific surface area. The arrows denote that T1/2 values are lower than indicated.

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On the other hand, calcination of the Au-Be coprecipitate at 200°C gave a maximum catalytic activity for both Hg and CD exidation. The XRD patterns showed that the sample calcined at 200°C contained very fine gold crystallites with no other apparent crystalline phase. Thermogravimetric and differential thermal analyses showed that the support was still in the phase of amorphous beryllium hydroxide at 200°C.

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X-ray photoelectron spectroscopy

In order to investigate the change of the correcipitates during calcination. XPS measurements were carried out for the samples calcined at different temperatures. Figures 3 (a) and (b) show the variation of the binding energies (BE) of Au4f7/2 and Au4d5/2 peak maxima with calcination temperature for Au/a-Fe2O3 and Au/BeD, respectively. The error bars correspond to the reproducibility of the BE measurements. The dashed lines in the figures indicate the BE values that were measured for pure gold powder.

The distinct shifts toward higher BE observed for the samples dried at 80°C are about 1 eV for Au4f7/2 and about 2 eV for Au4d5/2. Since shoulders were observed in the Au4f7/2 and 4f5/2 peaks at around 86 and 90 eV, respectively, which were very close to the BER of auric oxide(Au2O3), such oxidic species of



Fig. 3. Variation of BE of the maxima of Au4f7/2 and 4d5/2 lines as a function of calcination temperature for gold coprecipitates. (a), Au-Fe(1-19) coprecipitate; (b), Au-Be(1-19) coprecipitate.



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gold should be at least partly responsible for the large chemical shifts of the samples calcined at low temperatures. For the Au-Be coprecipitate dried at 80°C. a gradual shift of the position of the Audif peaks toward lower BE values was observed during analysis. This is the reason why two BE values for Audf7/2 are given in Fig. 3(b). This suggests that the coprecipitates dried at 80°C were less stable in Au-Be system than in Au-Fe system. The change of chemical shift with calcination temperature occurs in a nerrower temperature range(80°Cs Ts200°C) for Au-Be system than for Au-Fe system(80°CsTs400°C).

Both the Au-Fe coprecipitate calcined at 300-400°C and the Au-Be coprecipitate calcined at 200°C, that were the most active, exhibited the chemical shifts of about 0.5 eV toward higher BE values in Au 4f7/2 and 4d5/2 peak maxima. This indicates that such an amplitude of chemical shift is closely correlated with the catalytic properties of ultrafine gold particles.

EXAFS Analysis of the Au-Fo coprecipitate

Figures 4 and 5 show Fourier transforms of k^3 -weighted EXAFS oscillation of Au L3-edge of the Au-Fe and Au-Be coprecipitates calcined at different temperatures, respectively. In Fig. 4 the data for Au203 and Au foil measured at 300 K are also included for reference. The Au-Fe coprecipitate calcined at 200°C still contains oxidic species of gold as a main compound of Au. The decomposition of oxidic species into metallic gold occured at 300°C and was completed at 400°C. On the other hand, in the Au-Be coprecipitate calcined at 200°C the oxidic gold was already completely decomposed.

The coordination numbers, N. of a fu atom in the Au-Fe coprecipitate, which were obtained by curve fitting analysis of Au-Au peaks in the radial distribution function(RDF), are 4-8 for the sample calcined at 300°C, 7-10 for 400°C, 9-12 for 500 C. Since the ultrafine gold particles supported on α -Fe2O3 are hemispherical in shape as observed in TEM (Fig.7), the coordination number was calculated as a function of particle diameter. From the curve, the diameter of gold particles in the sample calcined at 300°C is estimated to be smaller than 2nm and that in the sample calcined at 400°C is 1.5-3 nm.

The mean particle diameter determined directly by measuring the size of more than 2,000 particles in TEM photographs was 4.1nm for the sample calcined at 400°C. The discrepancy seems to suggest a possibility that gold atoms are coordinated with other light atoms. In order to examine this possibility, the "difference" technique was applied for the Au L3 EXAFS signal of the sample calcined at 300°C to separate the signals of minor components12). As shown in Fig. 6, two peaks, one of which could be ascribed to Au-O coordination, were observed in the Fourier transform of the residual signal obtained by subtracting the Au-Au pair and noise signals from the measured signal. By

subtracting the Au-O pair signal from the residual signal and by Fourier transforming, the final RDF could be obtained. The curve fitting of the obtained peak on the assumption that it came from Au-Fe coordination output its distance of 2.48 Å. This value is very close to 2.51Å, which is the sum of metal bond radii of Au and Fe. Therefore, it is likely that there exist Au-Fe bonds, most possibly at the interface between gold crystallite: and hematite particles.

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Transmission electron microscopy

Figure 7 clearly shows the specific crystal prientation of gold crystallites; the Au(111) plane with a lattice spacing of 2.36A is in junction with the hematite(110) plane having a lattice spacing of 2.52A. This kind of epitaxy-like junction was very often observed not only in Au/α -Fe2O3 but also in Au/Co3O4 and Au/NiO^{6,7}). The reduction and oxidation treatment of Au/ α -Fe2O3 to transform the host oxide(α -Fe2O3 \rightarrow Fe3O4 \rightarrow y-Fe2O3) did not cause any tiny change of the sizes of gold particles. These facts support that there should be a strong bonding between Au and Fe at the interface.

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Fig. 5. Fourier transforms of k3-weighted EXAFS oscillation about Au L3-edge measured at 300 K for Au-Be coprecipitate calcined at various temperatures.



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Fig. 6. The first difference Fourier transform of the Au L3 difference signal obtained by subtracting the Au-Au pair signal from the measured signal.

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Fig. 7. TEM photograph of Ti doped Au-Fe (1-19) coprecipitate colcined at 400 °C.

DISCUSSION

In order to make highly dispersed gold catalysts, well mixed, most preferably homogeneously mixed coprecipitates, should be prepared. It is assumed that such precursors could not be prepared by coprecipitation of Au with Cr and Cd, and by deposition-precipitation of auric hydroxide on SiO2 sols, because the sizes of gold particles in the calcined samples were larger than 20 nm.

The stability of oxidic yold species including auric oxide in the well mixed coprecipitates depends on the kinds of counter metals. The oxidic species of gold in the Au-Be coprecipitate were stabilized only little and were completely decomposed into metallic gold at 200°C even though no appreciable thermal change occurred in the matrix of beryllium hydroxide. On the other hand, in the Au-Fe coprecipita.. The oxidic species of hold were much more stabilized and were not completely decomposed to +> 0°C. The above difference can be ' jecies for metal hydroxide explained by the affinities r - c-prec pitate, gold might be strongly matrices. In the case of embedded into the matrix of nydrous ferric oxide. During calcination, simultaneously with the crystallization of nematite, the oxidic gold species incorporated were decomposed into very tiny gold crystallites to come outside to the surface of hematite particles searching for favorable planes to the strong fixation. The epitaxy-like joining of gold crystallites with hematite particles could prevent gold crystallites from further coagulation,

1. 61.

It should be noted from the comparison between $Au/a-FegO_3$ and Au/BeO that strikingly high catalytic activities for the oxidation of CO were generated only when metallic gold was present while the oxidation of H₂ did not necessarily need the metallic gold. This might be due to the difference in the reaction mechanism. The clidation of H₂ may take place between gaseous hydrogen ar weakly adsorbed oxygen²), whereas the oxidation of CO can be assumed to occur between adsorbed CO and adsorbed oxygen. It is plausible that not oxidic but metallic gold surface is necessary for the adsorption of CO.

The results shown in Table 1 lead us to consider that the catalytic activities of monometallic gold catalysts for H2 oxidation are largely dependent on the size of gold particles, namely the exposed surface area of metallic gold, and almost independent of the kinds of metal oxide supports. In contrast to this, the markedly high activities for the oxidation of CO were obtained only when the sizes of gold particles smaller than 10 nm were supported on the selected groups of metal oxides. The effective oxides of the first group are a-Fe2O3, Co3O4, and NiO, which can adsorb oxygen to a certain extent and are themselves active for the oxidation of CO. The oxides of the second group are the hydroxides of Be and M9, where only little of oxygen but a large amount of water is adsorbed on the surface. Taking into consideration of

the enhancing effect of water in Au/a-fep03 as well as the surface propert of hydroxides of Be and Mg. it could be concluded that adsorbed water as well as adsorbed oxygen on the surface of support oxides is involved in the exidation of CO.

The XPS studies showed that in the active gold catalysts ultrafine gold particles were electron deficient. Hence, they become electronically similar to the metal to the left in the Periodic Table, namely Pt13), on which CO molecules are strongly chemisorbed. Therefore, the nature of ultrafine gold particles immobilized on the selected metal oxides might possibly be altered so that the sufficient chemisorption of CO occurs at low temperatures although bulk gold surface does not chemisorb CO1).

Based on the above considerations, the following mechanism can be speculated for the oxidation of CO in moistened air at a room temperature. Dwing to the transmutation of catalytic properties of ultrafine gold particles immobilized on a-Feg03, carbon monoxide and oxygen can be chemisorbed on the surface of gold. The adsorbed CO molecules react with HgO molecules and OH groups on the surface of metal oxides to form formic acid or formate ions. The adsorbed oxygen then reacts with the formate intermediates to produce carbon dioxido.

Without ultrafine gold particles, metal oxide catalysts like a-FegO3. Co3Oa are seriously deactivated by moisture because water prevents CO molecules from adsorption on the metal sites. Even with hydrous BeO and MgO where little oxygen species are a sirbed, high activities were obtained. This fact 'may indicate that oxygen is activated on the gold surface and is spilled over te the metal oxide surface.

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EXTENDED ABSTRACTS

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SELECTIVE CO SENSING WITH SEMICONDUCTIVE METAL OXIDES SUPPORTING ULTRAFINE GOLD PARTICLES

T. Kobayashi, M. Haruta, and H. Sano Government Industrial Research Institute of Osaka, Midorigaoka 1, Ikeda 563, Japan

There has been strong need for the selective and highly sensitive detection of CO, since CO is toxic even at a concentration of 100 ppm. The detection of CO with conventional Pd- or Pt-sensitized SnO2 semiconductor sensors is appreciably interfered by the presence of H₂. We have recently developed several supported gold catalysts which are extremely active for CO oxidation [1-3]. One of the advantageous features of these gold catalysts is that the oxidation of CO is accelerated with water and proceeds preferentially with respect to the oxidation of H₂. This property has led us to the investigation of the selective detection of CO with semiconductive Fe₂O₃ or SnO₂ supporting ultrafine gold particles (UFP-Au) [4,5].

Semiconducting materials, UFP-Au/Fe₂O₃ doped with Ti⁴⁺ (Au:Fe:Ti=5:97:3 in atomic ratio) and UFP-Au/SnO₂ doped with Na⁺ and Sb⁵⁺ (Au:Sn:Na:Sb= 1:99:5:1), were prepared by coprecipitation [4,5] from an aqueous solution of metal nitrate or chloride and chloroauric acid with an alkaline solution (Na₂CO₃ or NH₃). The coprecipitates were washed with distilled water, vacuum-dried, and then calcined at 400°C. The thick



Fig. 1 The sensitivities of UFP-Au/Fe₂O₃·Ti⁴⁺ sensor operated at 70°C as a function of gas concentration.

(○);CO, (●);CO with 100 ppm H₂, (△);H₂, (▲);H₂ with 10 ppm CO. film gas sensor devices were fabricated [4] from the above semiconducting powder, and they were finally sintered at 550 - 650°C. Gas sensitivity is expressed by the ratio of the electrical resistance of a device measured in fresh air with a relative humidity of 65 % to that measured in the presence of CO or H₂, R-air/R-gas.

Figure 1 shows the sensitivities of the UFP-Au/Fe2O3·Ti⁴⁺ sensor operated at 75°C as a function of the concentration of CO and H₂. The sensitivity to 30 ppm of CO is still higher than that to 1000 ppm of H₂. Furthermore, the sensitivity to 10 ppm of CO is hardly affected by the co-presence of 100 ppm of H₂. The sensitivity of this sensor tended to decline gradually in a continuous





Fig. 2 The sensitivities of UFP-Au/SnO₂·Na⁺·Sb⁵⁺ sensor operated at 200°C as a function of gas concentration.

(©);CO, (•);CO with 100 ppm H2. (△);H2, (▲);H2 with 10 ppm CO. operation for several hours, the reason of which might be the accumulation of water moleculer adsorbed onto the sensor surface. Such a decline is a temporary one and the original sensitivity can be restored by heat-flashing at temperatures above 200°C.

Figure 2 shows a similar relationship to that observed in Fig. 1, for the UFP-Au/SnO2·Na⁺·Sb⁵⁺ sensor operated at 200°C. While the selectivity for CO is a little inferior to that in the UFP-Au/Fe₂O₃·Ti⁴⁺ sensor, the sensitivity to CO is still appreciably larger than the one to H₂ and it is hardly affected by the presence of H₂. The UFP-Au/SnO₂·Na⁺·Sb⁵⁺ sensor can be free from the accumulation of contaminants like ammonia and water molecules adsorbed on the surface,

owing to its high operation temperature as 200° C, and is advantageous over the UFP-Au/Fe₂O₃·Ti⁴⁺ sensor in that it does not need the heat-flashing in the continuous operation.

Through the investigation of catalytic properties and microstructures, the following conclusions have been obtained concerning the ruling factors which determine CO sensitivity;

- The gas sensing properties of semiconducting metal oxides are closely related with their catalytic activities for oxidation reactions.
- (2) Smaller particle size of Au supported on the semiconductors gives higher sensitivity and higher selectivity towards CO.
- (3) Surface basicity of the oxides might play an important role in both CO sensing and CO oxidation, especially under the condition of humid atmosphere.

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35 Fellowships through STA

GLASS AND CERAMICS FOR THE FUTURE



(Growth of Superconducting Whisker, Bi-System, x1100)

GLASS AND CERAMIC MATERIAL DEPARTMENT GOVERNMENT INDUSTRIAL RESEARCH INSTITUTE, OSAKA

BRIEF HISTORY AND BACKGROUND

Government Industrial Research Institute, Osaka, (GIRIO) was founded in 1918 as a national research laboratory to develop and promote the chemical industry in the western area of Japan. It now belongs to the Ministry of International Trade and Industry. Glass and Ceramic Material department is one of five research departments. There is also a general administrative department within GIRIO.

Since the foundation we have pursued a wide range of studies on glass. Much of the work has borne fruits and has found application within the Japanese glass industry. For example, great advances in the camera industry have been made using the result of our research on optical glass. The department began ceramic research after the second world war.

Through comprehensive studies on the effects of constituents on the optical properties of glass, we succeeded in developing glasses with high refractive index and low dispersion, such as LaK and LaF. The Japanese camera industry has benefited greatly from this development. Our interest in optical glass led us to produce a large telescope mirror disc with a low thermal expansion coefficient. We succeeded in casting high quality discs 2m in diameter. A bobbin for a precise electrical inductance coil has made from glass with an extremely low expansion coefficient.

Borosilicate glass is well known for its durability. In the course of research on this glass system, we have developed the technique of preparing porous glass with high silica content by phase and subsequent leaching. Comprehensive developmental research is continuing on porous glasses.

Glass has the capacity to incorporate many elements. For this reason, it may be used for the disposal of high level radioactive waste from nuclear processes. We have conducted research and development in this field for more than a decade and have established the glass compositions and melting technology necessary for solidification of nuclear waste. The chemical and technical properties of the solidified wastes were investigated.

We have performed excellent basic researches on, for example, phase separation: nucleation and crystallization; volatilization; diffusion; electrical conductivity; mixed alkali effect. As will be explained after, We are now exploring so-called new glasses.

New ceramic materials have great potential and are currently finding

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industrial applications. We developed cutting tools made of alumina which show excellent performance compared with carbide tools. In the course of our research on MHD materials, we have succeeded in developing a new type of heating element, made of lanthanum chromite, which is commercially produced by a private company.

We have taken part in an R/D project on high efficiency gas turbines and are currently participating in fine ceramic project (one of the JISEDAI projects). In these projects, our attention has been mainly focused on composite ceramics, composed of a non-oxide ceramic matrix reinforced with ceramic fibers or whiskers to improve the reliability.

Ceramic joining is an important industrial problem. We have developed many joining methods, most of which could easily applied to large scale production.

Our research interests also include functional ceramics. We have worked on gas sensor materials and conductive perovskite-type oxides. We are currently working on superconducting oxides which are synthesized by melting process. So far the results look very promising.

ION-CONDUCTING GLASSES

Lithium-ion-conducting glasses

Lithium ion conductors are used as solid electrolytes in batteries, ECD devices, sensors, etc. Glassy solid electrolytes have certain merits compared with crystalline electrolytes. These are as follows:

higher ionic conductivity,

*wide composition region,

*isotropy, no grain boundaries,

·lower electronic conductivity.

Figure 1 shows the ionic conductivity of the $Li_2O-B_2O_3-Li_2SO_4$ system at room temperature. In this system, the conductivity exhibits a maximum at $0.3Li_2O \cdot 0.3B_2O_3 \cdot 0.4Li_2SO_4$, and decreases with increasing Li content. This result shows that the introduction of lithium salt enhances the conductivity, and that there is an optimum structure and Li content for high ionic conductivity.

Preparation of glassy thin films

Thin films formed by PVD or CVD show low resistance and are therefore suitable for device applications. We are investigating sulfide glassy thin films which are expected to have high ionic conductivity of the order of 10⁻³Scm⁻¹ at room temperature. Glass samples in the system Li₂S-GeS₂ are prepared by sputtering. Li₂S-containing films are very hygroscopic, hence they must not come into contact with oxygen or moisture. Therefore we handle both sputtering target and films in an argon-filled glove box. The film composition is analyzed by ESCA and the sample is isolated from air when placing it into the apparatus.

Mechanism of ionic conduction

Ionic conductivity is dominated by the concentration and mobility of the mobile ion. At low concentrations, the conductivity increases as the ionic concentration increases. However, at high concentrations, the conductivity becomes saturated. In order to obtain high conductivity, therefore, not only optimum ionic concentration but also the composition and structure which give high mobility are needed. Therefore we are currently investigating the structures and properties of ion-conducting glass by means of Raman spectroscopy; IMA; radio-isotope tracer methods; dielectric measurements; and pulse NMR. We are examining the relationship between these

properties and the ionic conductivity.

Raman spectroscopy is used to estimate the structural units of glass. In lithium borate glasses, we found an unknown structural unit by Raman spectroscopy as shown in Fig.2. Since lithium has no radio-isotope, the self-diffusion coefficient is measured by IMA using the isotope exchange method and by pulse NMR. The value is Li_2SO_4

used to calculate the Haven ratio. We are also examining the "mixed alkali effect" which may be related to the mechanism of ionic conduction.

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Fig.2. Raman spectra for lithium borate glasses.

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HIGH LEVEL RADIOACTIVE WASTE GLASS

High-level liquid radioactive wastes generated from the reprocessing of spent nuclear fuel must be converted to a durable solid form in order to immobilize the radio-nuclides before they can be permanently stored. As part of a government directive we have for the past two decades been carrying out a research program on the fundamental problems of the vitrification of spent nuclear fuel. The work is also in cooperation with PNC (the Power Reactor and Nuclear Fuel Development Corporation). The following topics have been studied by our group:

 We have searched for the optimum glass composition for nuclear waste containment. Phase separations such as crystallization and 'yellow phase' separation must be avoided in the glass formation. Screening of the glasses was done by a Soxhlet leach test.

2) Other solidification processes have been investigated. These include hot pressing and batch preparation of glass by the sol-gel method. The wet gels were first dried and then melted by microwave heating.

3) We have performed corrosion tests on refractory and electrode materials and have developed new corrosion resistant materials for use in ceramic melters. PNC have applied our results to the vitrification of nuclear waste program.

Atomic transport and surface analysis in waste glass

In order to assess the long term stability of the glass, we have investigated the material transport properties such as ionic diffusion. electrical conductivity of the melt, thermal conductivity, volatilization, crystallization, phase separation and corrosion in aqueous solutions.

An understanding of ionic diffusion in glasses is essential to elucidate the leaching mechanism. The self-diffusion coefficients of alkali ions and protons have been measured using radio isotope tracer techniques. The self-diffusion coefficients decrease in the order Na⁺>proton>Cs⁺. Protons play a key role in the corrosion of nuclear waste glass.

Corroded surface layers caused by water attack have been analyzed by ESCA and FTIR, which provide information about deposits in the layers and their dissolution in water. Figure 1 shows the ESCA depth profile of each element in a hydrated surface layer of nuclear waste glass which has been corroded by a Soxhlet leaching treatment. Alkali metals, alkaline earth elements, silicon and aluminum are depleted in the surface layer, whilst



the concentrations of rare earth elements, zirconium and iron are increased. The behavior is governed by thermodynamics, whether an element leaches out or deposits depends on the solubility product of its hydrate.

Comparison of the leachability of nuclear waste glasses with natural glasses

This project, 'Research on natural analogues for leaching behavior of nuclear waste glass forms', seeks to compare the long-term stability of nuclear waste glasses with that of natural glasses which have undergone weathering. However, the corrosion rate may be different owing to differences in composition. Basaltic glasses found in nature have a similar composition to nuclear waste glasses, although they contain no borate and comparatively few alkali oxide ions. We have therefore compared properties such as the Soxhlet leaching ability and the diffusion coefficients of Nu* and protons. Figure 2 shows an example of a Soxhlet leaching test.







POROUS GLASSES

Porous glasses, the products of leaching of heat-treated alkali borosilicate glasses in a certain composition range, have been known for more than forty years. The numerous studies on the structure and properties of the porous glasses have made a great contribution to our knowledge of chemically inhomogeneous glasses. The fundamental studies on the porous glasses having finished, we are currently examining the possible application in various branches of science and engineering.

Preparation

Borosilicate glasses of appropriate compositions can be separated by heat treatment into two phases. One phase is rich in silica, the other is rich in alkali borate. The alkali borate-rich phase is leached out by an acidic solution leaving behind a porous glass consisting of the silica-rich phase. Figures 1 and 2 show the pre-aration process and an SEM photograph of the porous glass, respectively. In order to obtain high-quality porous glass, we must have an initial glass satisfying the following requirements: (1) both silica-rich and alkali borate-rich phases must be interconnected; (2) the silica-rich phase must contain as much silica as possible; (3) the alkali borate-rich phase must be readily soluble in an acidic solution, and last but not least; (4) the stress developing during leaching at the interface between bulk glass and leached layer must not exceed a level determine^{AI} by the mechanical strength of the glasses.

Applications

We are currently researching the following applications of the porous glass:

1. Porous glasses have been successfully for biochemical catalysis as carriers of enzymes. Enzymes thus immobilized by absorption on the porous glasses maintain a high activity for a long period and may be used repeatedly.

2. Microporous glasses may be used as semipermeable membranes for separating liquid mixtures by reverse osmosis. The salt rejection rate is over 90 % using 0.5 wt% NaCl solution[1].

3. Alkali resistant glasses are useful as separating media or membranes and filling materials of gel permeation chromatography. These glasses may also be applicable for the phase separation method in a manufacture. We

have succeeded in making highly alkali resistant porous glasses which consist of a silica-rich phase with a certain amount of $ZrO_2[2]$.

4. Purous glasses may be used as membranes for separating gas mixture. The permeation rate increases dramatically with increasing the pore volume. We have produced a high speed membrane with a permeation rate 500 times higher than that of ordinary porous glasses[3].



Manufacturing process for porous glass

Fig. 1 Preparation process of the porous glasses.



Fig. 2 SEM photograph of the porous glass ×20000).

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HALIDE GLASSES

Our halide glass team is currently working on glasses based on chlorides, bromides and iodides, but not fluorides. The present research project of non-fluoride halide glasses started in 1986 following a two-year preliminary study.

Much attention on ZrF_4 -based glasses in glass science and technology in the past decade has stimulated research activities on the non-fluoride halide glasses. This is because the use of halides other than fluorides theoretically enable us to obtain glasses with ultralow optical loss, which is superior to fluoride glasses. Another more important possibility is the application as a glass fiber transmitting CO_2 laser energy. However, several problems arise which must be overcome before halide glasses can be successfully used e.g. they are highly hygroscopic and have low thermal stability. Every new glass forming system is being explored in order to solve these problems.

The final goal of our research is to establish the technology of glass preparation, purification, and fabrication for the halide glasses. We are also interested in the structure and fundamental properties of halide glasses as they are intermediate between oxide glasses with more covalent bonds and fluoride glasses with more ionic bonds.

The hygroscopic nature of halide glasses prevents them from being prepared by the usual method of melting in air. They are therefore prepared in a glove box containing dry N₂. Figure 1 shows the glass forming region of the ZnBr₂-BaBr₂-KBr ternary system[1]. Glasses with glass transition temperatures higher than 90°C have been obtained in this system. The infrared transmission of several different types of glass is shown in Fig. 2. This indicates that the ZnBr₂-based glass is transparent over a wider band of wavelengths than the other glass systems.

The crystallization behavior of the $ZnBr_2$ -based glasses has been examined by DSC. We are carrying out structural analyses of ZnX_2 -based and CdX_2 -based glasses using vibrational spectroscopy[2] and XAFS.

The practical application of halide glasses in the near future seems to be difficult. However, we believe that these materials will eventually play an important role in the region of far-infrared optics.



Fig. 1 The glass forming region in the system ZrBr2-BaBr2-KBr. Isothermal curves of the glass transition point are also depicted.



Fig. 2 Infrared transmission spectra of the ZnBr2-based glass compared with other types of glass.

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GLASS SCIENCE UNDER MICROGRAVITY CONDITION

New and pure glassy materials are required for highly functional elements in opto-electronics technology. It is, however, difficult to prepare such glasses on the earth, since glass cannot be produced without a erucible, which contaminates the glass. Nucleation of crystals also occurs at the container wall and the container material restricts the upper limit of the glass melting temperature. These difficulties can be solved if the melting process is carried out without the crucible. In outer space only very weak force is needed to control the movements of objects experiencing microgravity. Glasses may therefore be produced by the containerless melting process. The objective of this study is to develop the technology of glass production under microgravity and to prepare new glasses with superior transmittance in the non-visible region, particularly in the infrared region.

Development of Containerless Melting Process

The first stage of the research was to develop a suitable furnace for containerless melting. In cooperation with NASDA (National Space Development Agency of Japan) and Ishikawajima-Harima Heavy Industry(Fig.1) we developed an image furnace equipped with acoustic levitation apparatus which floats and holds the specimen at the center of the furnace under microgravity. Glass melting tests have been conducted using this furnace at gravitational forces of 1/10-1/100 G for 20-22 seconds produced by the ballistic flight of an aircraft. We succeeded in containing the material in the center of the furnace. On melting the floating materials a sphere of molten glass was obtained(Fig.2)[1,2].

Properties of Glasses Prepared under Microgravity Conditions

In 1991, we shall conduct a glass melting experiment in the NASA space shuttle. We plan to prepare glasses in the system $CaO-Ga_2O_3-GeO_2$, which have good near-infrared transmittance[3,4]. The glass-forming region, crystallization rate and glass transition temperature have all been investigated at a force of IG, in order to select the most suitable glass composition and operating conditions of the furnace[4]. Having developed the furnace and its operating conditions we then went on to examine the problems of hot working of glass production under microgravity.



Figure 1. Schematic representation of the acoustic levitation furnace.



Figure 2. Photograph of specimen floating in the furnace by acoustic power under microgravity.

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XPS and UPS measurement



Measurement of elect conductivity of ionicconductive glass





Preparation of halide glass in dry atmosphere



Probe for superconductive properties

Hot-pressing of ceramics in inert or varoum atmosphere



HIP apparatus

GLASSES FOR NON-LINEAR OPTICAL DEVICES

Materials which have non-linear optical properties have attracted much attention in the fields of information processing and optical communications. Switching devices, optical bistable devices, amplifiers and modulators all makes use of non-linear optical effects. Glass dispersed with semiconductors is one candidate material, since various optical properties of semiconductors are more easily controlled in a glass matrix. In this department, we are developing the basic technology for producing glasses dispersed with a semiconductor which have well controlled characteristics. Control of the composition and of the dispersion conditions is important as are the particle size and shape of the dispersed semiconductor. New processes for producing glasses with a dispersed semiconductor are also beir," investigated.

ACCURATE DETERMINATION OF REFRACTIVE INDEX OF SILICA GLASSES

Refractive index (n) and its temperature coefficient (dn/dT) have been determined accurately for various silica glasses. In the ultra-violet (UV) region, photo-lithography is of great importance for fabrication of semiconductor LSI circuits. Silica glass is an ideal material for transmitting UV light since it does not degrade on irradiation. It is commonly used as a lens material in UV light projectors. Lenses to be used at 248nm require n to be known to an accuracy of $\pm 1 \times 10^{-5}$ or better. The temperature coefficient of n is about $1 \times 10^{-5}/^{\circ}$ C for most silica glasses at room temperature so the ambient temperature of the lens must be controlled to within 1°C. The value of n is found to depend on the fabrication method and thermal history of the glass. We are currently studying the way in which n is related to specific gravity, which directly reflects the thermal history of the glass and is easier to measure than the refractive index.





LOW DISPERSION UV TRANSMITTING FLUORIDE GLASSES

Glass which can transmit UV light with low dispersion is requited for making achromatic projector lenses for use in UV lithography. Fluoride glasses transmit UV light well and are readily formed into the requisite lens shape without surface cleavage which occurs in may optical crystals.

We are currently studying fluoride glasses which have a wide band gap. Our interests include the melting technology, the transmittance, refractive index, dispersion and coloration of the glasses under UV lights.





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CERAMIC COMPOSITE

Brittleness is the weakest point of ceramics when they are used as structural components for example in a diesel engine or a gas turbine. One of the ways to overcome this problems is "toughening" of the ceramic through reinforcement with fibers or particles. In such CMC (Ceramic Matrix Composite), fibers or particles improve toughness by dissipating or absorbing fracture energy at the top of crack front. Research on CMC's with various combinations of matrix and fibers/particles is one of the main themes of ceramic research in our department. Current efforts in this area are as follows:

Whisker reinforced composite

As whiskers of SiC or Si_3N_4 have very high strength and refractivity, they are suitable for ϵ the reinforcing fiber in CMC's. CMC's with various matrices such ϵ 4, SiC, and Al₂O₃ are being investigated.

The SiC(whisker)/Si₃N₄ composite[1], which was first successfully fabricated in our department, exhibits an strength increase at temperatures over 1100°C. This is attributed to the fact that the whisker which prevent grains from sliding at high temperatures. Both fracture toughness and wear-resistivity are also enhanced[2]. Whisker incorporation has a remarkable effect on the electric resistivity, which decreased markedly from that of Si₃N₄ which is an insulator to about 10 ohm cm or less. Such a low resistivity means the composite can be subjected to electric discharging machining (see Fig.1).

One of current topics of interest is to tailor the texture of whisker-reinforced CMC's. SiC or Si $_3N_4$ whisker are mixed with organosilicon polymer, which forms a matrix of SiC or Si $_3N_4$ upon pyrolysis. The interface between the whisker and matrix is also being investigated.

Other research is aimed at clarifying the fundamental phenomena in the composite forming process, in order to develop highly reliable ceramics for gas turbine components. The effects of fibers or sintering aids on toughness and on densification behavior of CMC are being studied.

Long fiber-reinforced composite

CMC's reinforced with a continuous fiber such as SiC fiber or carbon fiber attracts much attention as they have extraordinary toughness similar to that of bamboo or other wooden material. Such composites are expect-





ed to be the next generation material of choice for structural application eg. in the aerospace industry. We are currently looking at ways of improving fabrication of the composite without damaging the fiber in the process.

Particle reinforced composite

Ceramics have medical applications as they can be used to make artificial bone or tooth. The texture of apatite is similar to that of human bone, but it suffers from low strength. The strength and toughness may be improved by dispersing ZrO₂ particles in it.

Several combinations of different matrices and particles are currently being studied. ZrC particle / B_4C matrix composite exhibits improved strength and toughness. A roller bearing made of Si₃N₄ dispersed with TiC particles hed a longer lifetime than one made from Si₃N₄ alone.



Fig.1 SiC(whisker)/Si3N4 composite pieces machined by means of electric discharging.

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TRIBOLOGY OF CERAMICS

Measurement of tribological properties

The efficiency and life time of gas turbines are expected to be greatly improved by using fine-ceramic materials instead of metals, since fine ceramics have superior mechanical strength, chemical durability and wear resistance at elevated temperatures.

A knowledge of the tribological properties of ceramics is indispensable for predicting to the friction and wear of mechanical parts. The ball-ondisk method is the most reliable way to measure friction and specific wear rate accurately (see Fig.1). Fundamental measurements are continuing in order to assess the reliability and reproducibility of this method. The next stage of our work is to measure the frictional resistance of ceramics to abrasive particles in the combustion gas running through gas turbine on frictional resistance of ceramics. The effect of solid lubricants on wear will also be investigated.

Development of ceramic bearings

ball and the ring under compressive loads.

The most interesting practical application of the superior tribological properties of ceramics is the ceramic bearing. It has many advantages over ordinary metal bearing, eg. it can be used at temperatures over 800°C in air without water cooling and oil lubrication. The chemical durability of a ceramic bearing enables it to be used in corrosive environment such as in a chemical reactor, where the presence of strong acid or alkalis means that a metal bearing will be rapidly damaged. Ceramics do not rust and can be used for long continuous period, so a ceramic bearing saves maintenance costs.

We are currently studying the performance of ceramic bearings made of Si_3N_4 , which is widely regarded as an excel ent material for this application because of its well-balanced properties such as strength, hardness and toughness. We have investigated the relationship between fatigue characteristics and the material properties. We found that fracture toughness predominantly determines the lifetime of the bearing. High hardness resulted in microfracture at the contact point between the

An application of ceramic composites to bearing material is also tried. TiC particle-reinforced Si_3N_4 composite, which has an improved toughness, exhibited longer life as a ball bearing material.

Development of ceramics containing solid lubricating agent

When ceramic bearings are used at higher temperatures, the ball holder (retainer) should withstand oxidation. An ability to lubricate is also desirable because ordinary oil is not stable at higher temperatures such as 500 or 600°C.

A new type of ceramic composite, containing graphite microcrystals in the sintered body, has been fabricated in our department(Fig.2)[1].

Composites with carbides such as SiC, TiC or B_4C as the matrix component shows a superior tribological properties when dispersed with graphite particles. A sliding bearing test showed that the graphite was effective in lowering the friction coefficient and specific wear rate and was therefore acting as a lubricating agent.



Fig.1 Friction and wear measurement by ball-on-disk method.

P applied load, x1 sliding distance, F1 friction force, intocerficient of friction, r1 radius of ball, d1 diameter of wear track, h1 wear depth, D1 diameter of sliding circle. S1 cross sectional area of sliding circle. W1 wear volume, Ws1 specific wear rate



Fig.2 Microstructure of Graphite/ TiC composite.

; 5 µm

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SUPERCONDUCTING CERAMICS

A Superconducting cuprate (La-Ba-Cu-O) was first reported by Bednorz and Muller in 1986, and a number of studies on this material have been undertaken all over the world. Our institute started to investigate the superconducting ceramics at the Glass and Ceramic Material department in 1987.

The effects of sulfur addition

The effects of sulfur addition on the superconductivity of Y-Ba-Cu-O system were studied[1]. We examined whether the substitution of oxygen with sulfur took place or not, and expected to improve the superconductivity by adding sulfur to the Y-Ba-Cu-O system. Sulfur-containing compounds with nominal compositions $YBa_2Cu_3O_xS_y$ were prepared by the solid state reaction method using CuS or Cu₂S instead of CuO. By this method, sulfur atoms were not incorporated into the crystal lattice of the 1:2:3 phase and made it multiphasic. Although T_c (critical temperature) was lowered with increasing sulfur content, J_c (critical current density) for y=0.03 was about twice as high as that for y=0.00, this is due to the increasing the bulk density by adding sulfur.

Superconducting glass ceramics

The new type superconducting oxide, Bi-Sr-Ca-Cu-O system, was found by Maeda et. al. in 1988. The characteristic nature upon processing is that the Bi-Sr-Ca-Cu-O compounds form a glassy state when it is rapidly quenched from a molten state. We have been researching into the crystallization of superconducting phases from a glassy melt-quenched solid. We found that the addition of Ga or Ge to the Bi-Sr-Ca-Cu-O system was effective for manufacturing when the glass was formed. The wide glass forming region is obtained by adding Ga or Ge. We made fibers and ribbons (10cm length) by the melt-quench method using a single roller with Ga or Ge additives. After heating at 820°C, the ribbons showed $T_c(end)$ at 77-20K. We determined the composition region where the superconductivity was maintained in spite of the improvement in manufacturing.

Superconducting whiskers

The superconducting whiskers of Bi(Pb)-Sr-Ca-Cu-O were prepared by heating a glassy melt-quenched plate in a stream of oxygen gas (Fig.1).

The dimensions of the whiskers are 2-10 μ m × 10-300 μ m × ~15mm. The whiskers have the 2212 structure (low-T_C phase) and show a zero-resistance state at 70K. Each whisker exhibits the structure where several platelike single crystals are stacked. The orientation of the longest c axis is perpendicular to the platelike crystal plane and the ab plane is parallel to the crystal plane. The whiskers are able to be bent to a radius of curvature(R) of 0.4mm without a decrease in the T_C value. The highest J_C is 67000A/cm² (63K, zero magnetic field) in a nonbending state. It surpasses 35000A/cm² even in a bending state of R≥1mm and finally decreases down to 3200A/cm² for R=0.4mm (Fig.2). Our further subjects for study related to the whiskers are the elucidation of the growth mechanism, preparation of high-T_C whiskers, lengthening the whiskers and evaluation of the mechanical properties[2,3].



Fig.1 The whiskers of Bi(Pb)-Sr-Ca-Cu-O grown by heating a glassy meltquenched plate in a stream of oxygen



Fig.2 The Tc and Jc (63K, zero magnetic field) of the whiskers as a function of the radius of curvature.

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CERAMIC COATING

Ceramic coating by the plasma spraying technique is being studied. METCO 9MB Low-Pressure-Plasma-Spraying (LPPS) system and TAFA 90HV Atmospheric-Plasma-Spraying (APS) system are used in this section. Plasma spraying is one of the coating processes in which molten raw material powders are accelerated and shot onto the substrate in the plasma flame. In principle, anything that can be melted is applicable as the sprayed material. Any material may be coated when it is adequately cooled.

Ceramic-ceramic composite coating and non-oxide ceramic coating are studied as part of the ordinary research and development "A study on the ceramic coating by low pressure plasma spraying". Coatings made of non-oxide ceramics, such as SiC, Si_3N_4 , etc., are expected to resist wear at high temperature. However, non-oxide ceramic coatings can not be made by plasma spraying in hir because of thermal decomposition. These coatings are achieved when the appropriate spraying aid is added and the spraying is carried out in a closed chamber with the a controlled atmosphere.



Fig.1 Low-Pressure-Plasma-Spraying system.



Fig.2 Plasma flame at 300 torr, 40kW.

FLUORINE RESISTIVE ENAMELS

This research aims at development of fluorine resistive coatings on metals to be used in the fluorine chemical and semiconductor industries.



In these fields, corrosion of tanks and pipes by fluoride gasses or fluoric acids causes severe safety problems. Traditional enamels composed of oxides are easily attacked by these fluorides leading to failure in a closed system.

Fluoride enamels, on the other hand, composed of metal fluorides such as MgF_2 , CaF_2 , SrF_2 , BaF_2 , NiF_2 , YF_3 , etc. are highly resistive to these atmospheres, even at moderately high temperatures. They are therefore promising materials for use in this field. We are surveying glass compositions which are durable to these gasses and acids, and are highly adhesive to metals without fracture at elevated temperatures under bending stress.

THEORETICAL STUDY OF CERAMIC INTERFACES

Various properties of ceramics depend on grain boundaries, mechanical and electrical properties and sintering ability. The joining between ceramics and metals is an essential schnique in order to realize the practical use of ceramics. It is essential to understand the fundamental properties of grain boundaries and interfaces in ceramics and of metal/ceramic interfaces from a macroscopic viewpoints. We are investigating theoretically the atomic structure and properties of grain boundaries in covalent ceramics such as SiC and of metal/ceramic interfaces such as alumina/transition metal In these systems, it is necessary to calculate the electronic interfaces. structure of the interface. With powerful modern computers, interfacial electronic structures, stable atomic configurations and interfacial energies can be calculated using the electronic theory of solids. The effects of impurities and the dependence on the kinds of ceramics and metals are also being examined. By comparing the calculated results with observations and experiments, such as HREM images and photoelectron spectroscopy, it is possible to understand the fundamental properties of grain boundaries and interfaces from a microscopic viewpoint.

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ORGANIZATION AND OTHER ACTIVITY

This department consists of 5 sections. Three of them deal with glass research and two with ceramics. The total research staff number 33. We have now more than 20 research projects both large and small. Some of them are inter-sectional and inter-departmental. We participate in national R&D projects, most of which are in cooperation with other national laboratories and private companies.

Organization and main research fields Director: Dr. M. Kinoshita

Glass technology section: Mr. J. Hayakawa

Glass melting under microgravity, optical properties of glasses.

Glass science section: Dr. H. Wakabayashi

lon conducting glasses, technology of waste management.

Advanced glass section: Dr. H. Tanaka

Halide glasses, porous glass, non-linear photonic materials.

Engineering ceramics section: Mr. S. Kose

Composite ceramics, ceramic coating, tribology, ceramic interfaces. Functional ceramics section: Dr. H. Yamashita

Superconducting oxides, fluorine resistive materials.

Other Activities

Most of the researchers in this department are members of the Ceramic Society of Japan. We also have close contact with other societies such as those of optical glass, applied physics, power metallurgy and fine ceramics. Ours is the only national laboration / dealing widely with glass.

We have therefore run a glass technology course for foreign researchers for many years.

There are several invitation schemes applicable to foreign researchers, whose numbers in our institute have been increasing in recent years.



The main building of the glass and ceramic material department



The annex of the department





GLASS AND CERAMIC MATERIAL DEPARTMENT GOVERNMENT INDUSTRIAL RESEARCH INSTITUTE, OSAKA

8-31 MIDORIGAOKA-1, IKEDA, OSAKA 563

.

PHONE: (0727) 51-8351 TELEFAX: (0727) 51-2156



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Gold Catalysts Prepared by Coprecipitation for Low-Temperature Oxidation of Hydrogen and of Carbon Monoxide

M. HARUTA.* N. YAMADA.† T. KOBAYASHI.* AND S. IJIMA‡1

*Government Industrial Research Institute of Osaka, Midorigaoka 1, Ikeda 363, Japan, †Kishida Chemicals Company, Ltd., Joshoji-machi, Kadoma 571, Japan; and ‡Research Development Corporation of Japan, Science Building, 5-2 Nagata-cho 2-chome, Tokyo 100, Japan

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Novel gold catalysts were prepared by coprecipitation from an aqueous solution of HAuCl₄ and the nitrates of various transition metals. Calcination of the coprecipitates in air at 400°C produced ultrafine gold particles smaller than 10 nm which were uniformly dispersed on the transition metal oxides. Among them, Au/α -Fe₃O₃, Au/Co₃O₄, and Au/NiO were highly active for H₃ and CO oxidiation, showing markedly enhanced catalytic activities due to the combined effect of gold and the transition metal oxides. For the oxidation of CO they were active even at a temperature as low as -70° C. e 1989 Academic Press. Inc

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INTRODUCTION

During the course of an investigation into new oxide catalysts useful for the low-temperature catalytic combustion of hydrogen (1-4), it became evident that the catalytic activities of transition metal oxides for hydrogen oxidation had a volcano-like relation with the heat of formation of oxides per gram-atom of oxygen (5). The volcano relation indicates that the formation of metaloxygen (M-O) bonds is rate determining for the oxides of Ag and Au, which are located on the left side, while the breaking of M-O bonds is the slow step for the other metal oxides located on the right side. Therefore, an attempt was made to develop composite oxides of Ag with the 3d transition metals, for which an enhancement in both catalytic activity and thermal stability was expected.

Our earlier paper (5) reported that an appreciable enhancement in catalytic activity was, in fact, achieved in some composite oxides of silver with 3*d* transition metals which were prepared by coprecipitation. Specifically, a mixed oxide composed of

¹ Present address: N² , Corp., Miyazaki 4, Mivamae, Kawasaki 213, Japan.

Co, Mn, and Ag (20:4:1 in atom ratio) was both thermally stable and highly active for the oxidation of H_2 and CO. The successful results obtained for these composite oxides of silver led us to expect that a significant enhancement in catalytic activity might also be exhibited by composites of gold and the other metal oxides. The present investigation into gold-based oxide catalysts was undertaken to test this hypothesis.

Previous work on gold catalysts has been reviewed by several authors (6-10). All the gold catalysts investigated so far are gold supported on inactive ceramic oxides, such as SiO₂ (11-17), Al₂O₃ (14-16, 18), MgO (15-17, 19), and TtO₂ (20), or unsupported gold filaments (21), powder (22, 23), sponges (24), filings (25), and gauze (26).

The chemical reactivity of gold catalysts has been studied for the oxidation by oxygen or nitrogen oxides of CO (11, 22, 24, 26) and H₂ (12, 15, 17, 21-23), selective oxidation of organic compounds by nitrogen dioxide (13), hydrogenation of alkenes (7), and so on. However, the conventional goid catalysts prepared by impregnation have been reported to be usually far less active than platinum-group metal catalysts, although they are superior in selectivity for only a few reactions such as the oxidation

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of 1-pentanol to 1-pentaldehyde by NO₂ (13) and the hydrogenation of 1-pentene to *n*-pentane (7). The present paper deals with a new type of gold catalyst prepared by coprecipitation instead of impregnation. In a previous short communication (27), we reported that combination with the oxides of Group VIII 3d transition metals makes gold so active that it catalyzes the oxidation of CO, even at -70° C.

EXPERIMENTAL

The gold catalysts were prepared by coprecipitation. An aqueous solution of chloroauric acid and a transition metal nitrate was poured into an aqueous solution of sodium carbonate under stirring. The precipitate was washed, vacuum dried, and calcined in air at 400°C for 4 h. Gold powder and the simple oxides of Fe, Co, and Ni were also prepared in this manner. Conventional gold catalysts were prepared by immersing support oxides in an aqueous solution of HAuCL with a small excess volume. After drying, the impregnated samples were reduced with hydrogen at 200 or 300°C. They were then washed in hot water several times and dried in air at 200°C. The support oxides used were α -Fe₂O₃ calcined at 400°C (SA = 42 m²/g) and γ -Al₂O₃ (anhydrous extra pure, Merck, $SA = 97 \text{ m}^2/\text{g}$). Another type of gold catalyst was prepared by the reduction of HAuCL with diammonium citrate in an aqueous dispersion of colloidal SiO₂ (STO, Nissan Chemicals Ind., Ltd.). This catalyst was finally calcined in air at 300°C after washing and drying.

All the starting materials used were reagent grade (Kishida Chemicals Co., Ltd.). Elemental analysis of chloroauric acid tetrahydrate (HAuCl₄ · 4H₂O) was conducted by means of atomic emission spectroscopy using an inductively coupled argon plasma source (Hitachi, Ltd., ICP Emission Analysis System 306) to determine the impurity

els of Pd, Pt, and Ir, because trace amounts of these Pt-group metals may sometimes account for part or almost all the

catalytic activity of the gold sample under study (10). The impurity levels were approximately 11, 2, and 7 ppm, for Pd. Pt, and Ir, respectively.

Catalytic activity measurements were carried out in a small fixed-bed reactor, with 0.20 g of catalyst that had passed through 70- and 120-mesh sieves. After the pretreatment of catalysts in a dried air stream at 200°C for 30 min, a standard gas consisting of 1.0 vol% H2 or CO balanced with air to 1 atm was passed through the catalyst bed at a flow rate of 66 ml/min. This reaction gas was dried by passing through columns of silica gel and P2O3 granules. In the experiments at temperatures below 0°C, the reaction gas was further passed through a silica gel column cooled to -77°C to prevent the accumulation of moisture in the catalyst bed. In order to investigate the influence of moisture on the catalytic activity, the reaction gas was passed into a constant moisture device (28) which contained saturated NH4Cl aqueous solution. The relative humidity of the reaction gas was kept constant at 76%, as confirmed by humidity measurements using a Humidector (Shinyei Co., Ltd.).

Activated carbon and molecular sieve 13X were used as column packing agents to analyze CO_2 and CO, respectively, in the effluent gas by a gas chromatograph. The conversion efficiencies, determined from the changes in concentrations of CO_2 and CO, were in good agreement with each other under all the conditions tested.

The activity of a given catalyst is usually expressed in terms of the temperature corresponding to 50% conversion $(T_{1/2})$, which can be obtained from the conversion vs temperature curves. A lower temperature for $T_{1/2}$ indicates correspondingly greater catalytic activity.

Specific surface areas were determined by the single-point method using a Quantasorb surface area analyzer. A mixture of 30 vol% N_2 in He gas was used, with N_2 as adsorbate at a temperature of -195.8°C. The reproducibility of the data was within

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Catalysts

±5%. X-ray diffraction experiments were performed after each thermal treatment in a Rigaku X-ray powder diffractometer with Ni-filtered CuKa radiation. Crystallite sizes of Au were calculated from peak halfwidths by using the Scherrer equation with corrections for instrumental line broadening.

The fine structure of the gold catal was observed using an Akashi EM-0 electron microscope operated at 120 X-ray photoelectron spectroscopic ar ses were made using a Shimazu ESCA under vacuum below 5 × 10⁻⁶ Torr. surface composition was determined by ing a sensitivity factor for each eler (29). The bulk composition was determ by X-ray fluorescence analysis.

RESULTS

Oxidation of Hydrogen

A variety of mixtures of metallic gold with single and binary oxides was prepared to seek appreciable enhancement in the catalytic oxidation of H2. The results are summarized in Table 1. Among the 3d transition metal oxides investigated, the oxides of cobalt, iron, and nickel, the Group VIII metal oxides, were found to exhibit appreciably enhanced catalytic activity in the presence of gold.

Figure 1 shows the catalytic activities ex-

	Au/MnO,	10	152	69
ysts	Au/MnO,	50	134	115
XO2A	Au/a-FerOs	10	73	73
4V	Au/ConO.	10	66	69
alu	Au/NIO	10	73	116
ialy-	Au/CuO	10	143	20
750	MnO ₂	0	244	32
The	a-FerOs	0	225	43
y us-	Co ₁ C ₄	0	128	47
nent	NIO	0	221	55
ined	CuO	0	> 300	6
incu	Ag ₂ O	0	89 ~ 103	
	PdO	0	1	-
	PtO ₂	0	38	-
	Auron	100	131	1
nold	5 M			

Au

(at.%)

TABLE 1

Catalytic Activities for H. Oxidation and Specific Surface Areas of Various Metal Oxides and Gold Catalysts

112

(°C)

pressed by $T_{1/2}$ and specific surface areas as
a function of gold content. Maximum activ-
ties were obtained at 5 at.% of gold for the
Fe-Au and Co-Au systems and at 10 at.%
for Ni-Au. As the T1/2 values for gold pow-
der and the host metal oxides individually
were above 130°C, it was clear that their
combination resulted in noticeably en-
hanced catalytic activities.



FIG. 1. Dependence of T12[H2] and specific surface area on gold content in the coprecipitates calcined at 400°C. (a) Au/α -Fe₂O₃; (b) Au/Co₃O₄; (c) Au/NiO.

303

Surface area

(m¹/g)

69







The changes of specific surface area with composition appeared to be bimodal for all three systems. The maximum catalytic activity was obtained at the minimum in specific surface area in the Fe-Au system, while it was observed at the second peak in the Co-Au and Ni-Au systems.

Oxidation of CO

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The coprecipitated gold catalysts were much more active for the oxidation of CO than for the oxidation of H2. The three typical gold catalysts, hereinafter denoted Au/ α -Fe₂O₃ (Au/Fe = 1/19), Au/Co₃O₄ (Au/Co = 1/19), and Au/NiO (Au/Ni = 1/9) (see characterization), were able to oxidize CO even at -70°C (27). They were also able to oxidize the CO completely at 30°C even under a relative humidity of 76%. While a Hopcalite catalyst (mixed oxides mainly composed of Mn and Cu) commercially used for CO safety gas masks had lost its activity after 20 min, the Au/a-Fe₂O₁ and Au/Co3O4 catalysts maintained their activity for at least 7 days.

Comparison of Catalytic Activity for CO Oxidation with That for H₂ Oxidation

It is useful to compare the new gold catalysts with the support metal oxides, gold powder, impregnated gold catalysts, and a supported Pd catalyst, for the oxidation of CO and H2. Figures 2 and 3 show such a comparison. The impregnated gold catalysts, Au/y-Al2O3 and Au/a-Fe2O3, were active only at temperatures above 100°C, although they were calcined at 200°C; which was lower by 200°C than the calcination temperature for the coprecipitated catalysts. The support oxides, namely Co3O4, NiO, and a-Fe2O3, and gold powder are active only at temperatures in the range 60 to 300°C under the same experimental conditions. These comparisons clarify that it is only the gold catalysts prepared by coprecipitation that exhibit marked enhancement in catalytic activity.

Figure 4 shows $T_{1/2}$ values for CO oxidation and H₂ oxidation as a function of mean diameter of Au crystallites determined from TEM and/or XRD. The correspirated gold catalysts exhibited the highest activities when they were calcinced at 300-400°C, while the impregnated and reduction-deposited catalysts exhibited lower catalytic activities with an increase in calcination or reduction temperature. For example, the gold catalyst supported on α -Fe₂O₃ by impregnation (No. 5 in Fig. 4) had higher catalytic activity, although much lower than those of coprecipitated catalysts, when calcined at 70°C than at 200 and 300°C.

GOLD CATALYSTS FOR H2 AND CO OXIDATION

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A general trend is that catalytic activity increases with the decreasing diameter of Au crystallites in the oxidation of both CO and H2. However, small crystallites of Au do not necessarily lead to catalytic activity at such an extremely low temperature as -70°C. In the Au/Al₂O₃ catalysts prepared by coprecipitation, the $T_{1/2}$ value for CO oxidation is much higher than those for Au/ α -Fe₂O₃, Au/Co₃O₄, and Au/NiO even though the crystallite size of Au is similar in all cases. On the other hand, the $T_{1/2}$ value for H2 oxidation is comparable for all four catalysts. Accordingly, the oxidation of CO at low temperatures seems to require both the control of Au crystallite size and the selection of appropriate support oxides.

Characterization

The X-ray diffraction patterns showed that the coprecipitates calcined at 400°C in air were composed of metallic crystallites of Au and the oxides, namely α -Fe₂O₃, Co₃O₄, and NiO. The crystallite sizes of Au were estimated to be 3.6 nm for Au/ α -Fe₂O₃ from the peak half-width of Au(111) at $2\theta =$ 38.2°, ca. 6.0 nm for Au/Co₃O₄ from Au(200) at $2\theta =$ 44.4°, and ca. 8.0 nm for Au/NiO from Au(220) at $2\theta =$ 64.6°. The crystallite sizes of Au in the impregnated catalysts were estimated to be 16, 20, and



FIG. 4. Catalytic activities for H₂ and CO oxidation as a function of mean particle diameter of Au. 1-4, coprecipitates calcined at 460°C; 1, Au/ α -Fe₂O₃ (Au/Fe = 1/19); 2, Au/Al₂O₃ (Au/Al = 1/19); 3, Au/ Co₃O₄ (Au/Co = 1/19); 4, Au/NiO (Au/Ni = 1/9); 5, Au/ α -Fe₂O₃ (5 wt%, impregnation, reduction at 200°C); 6, Au/SiO₂ (17 wt%, reduction, calcination at 300°C); 7, Au/ γ -Al₂O₃ (5 wt%, impregnation, reduction at 200°C).





23 nm for α -Fe₂O₃-, SiO₂-, and γ -Al₂O₃-supported catalysts, respectively.

Figure 5 shows a high-resolution TEM photograph of Au/α -Fe₂O₃ (Au/Fe = 1/19). Gold particles are uniformly dispersed on the hematite particles of size around 20–50 nm. The histogram in Fig. 6 shows that gold particles exist with a rather sharp size distribution. The mean diameter of 2131 particles of Au is 4.1 nm with a standard deviation of 1.4 nm (34%). This diameter agreed well with the value estimated from XRD data.

In the XPS spectra of coprecipitated Au/ α -Fe₂O₃ (Au/Fe = 1/19), the binding energy of Au 4f_{3/2} was 83.9 eV, a little larger but very close to that of metallic gold evaporated onto α -Fe₂O₃ and appreciably different from that of Au₂O₃, 86.3 eV. No detectable differences in the Fe 2p_{3/2} and O 1s



FIG. 6. Histogram of gold crystallites in Au/α -Fe₂O₃ (Au/Fe = 1/19) prepared by coprecipitation and calcination at 400°C.

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TABLE 2

Composition of Au/a-Fe₂O₃ Catalysts Prepared by Coprecipitation

Oxides	Fe	Au
Prepared (at.%)	95	5
Elem. anal. (at.%)	94.6	5.4
XPS (at.%)	93.4	6.6
XRD	Au particles and	a-Fe,O

spectra between Au/α -Fe₂O₃ (Au/Fe = 1/19) and α -Fe₂O₃ were observed.

Table 2 summarizes the results of analyses for the surface composition calculated from the peak area of Au $4f_{s2}$ and Fe $2p_{32}$ in XPS and the bulk composition obtained from X-ray fluorescence measurements. The compositions of the starting solution, surface layer, and bulk were nearly identical. This result shows that gold particles are deposited mainly on the surface of hematite particles with little, if any, being incorporated into the bulk of the hematite.

DISCUSSION

It has been demonstrated in the present study that gold becomes a very active catalyst for the oxidation of CO when small gold particles are prepared in the presence of 3d transition metal oxides. In the impregnation and reduction methods, which were used for the preparation of almost all the conventional gold catalysts, gold was obtained only as large particles, usually above 10 nm in diameter. This is because gold has a very low melting temperature, low sublimation energy, and very low Tammann temperature (10). These properties, in addition to the intrinsically poor reactivity of gold metal, make gold markedly different from Pt-group metals in catalytic behavior.

An attempt has recently been made by Zhang (10) to prepare small gold particles supported on MgO and Y-zeolite by incipient wetness impregnation of chloroauric acid and by ion exchange with gold diethylenediamine trichloride, respectively. Although gold particles smaller than 2 nm

were obtained by calcining at temperatures below 200°C, the reported catalytic activity for H₂ oxidation was not appreciably high. It could be assumed that chloride ions remained in the MgO carrier because the catalysts were not washed by hot water after reduction. In fact, the catalytic activity of Au/Al2O3 prepared in our laboratory by impregnation was appreciably improved by washing with hot water. In the coprecipitated gold catalysts, the coprecipitates were thoroughly washed before calcination and therefore were considered to be almost free from chloride ions. It could also be expected that Pt-group metal impurities contained in HAuCL might be mostly excluded during the coprecipitation of gold and 3d transition metals because of the extremely low concentrations of those impurities in the starting solution. The small gold particles held inside the supercages of zeolite Y containing Na* grew at temperatures above 100°C, indicating that they were not thermally stable when used as an oxidation catalyst.

From a comparison with Zhang's work, it is evident that coprecipitation is an effective method to prepare small gold particles with good thermal stability and possibly to avoid contamination from chloride ions and Pt group n...tals which are usually contained in the starting materials. During calcination, gold components were decomposed to form metallic gold crystallites which move out from the inner part of the coprecipitate particles toward the surface. This process might cause the gold particles to be strongly held by the support oxides thereby preventing their coagulation.

The parabolic changes in catalytic activity for H_2 oxidation shown in Fig. I may be closely related to the dispersion and exposed surface area of Au metal. The total surface area of exposed Au metal increases with an initial increase in Au content and may then decline with a further increase in Au content due to the coagulation of Au particles. This compensating effect of Au content gives rise to the maximum in the exposed surface area of Au and, accordingly, to the maximum catalytic activities at 5-10 at.% of Au. It seems to be a coincidence that a catalyst supporting smaller gold particles requires a smaller content of Au to reach the maximum catalytic activities: Au/ α -Fe₂O₃ (4.1 nm, 5-7 at.%) < Au/ Co₃O₄ (ca. 6 nm, 5-10 at.%) < Au/NiO (ca. 8 nm, 10-14 at.%).

The bimodal change of the specific surface area appears to be complex in comparison with the simple parabolic change observed for the composite oxides of Ag with Co and Mn (5). The initial increase in the specific surface area up to 2 at.% of Au might be due to the incorporation of gold into the precipitates of Fe, Co, and Ni, as in the case of Ag-Co and Ag-Mn oxides. Since the ionic radius of Au³⁺ is 1.37 Å, larger than 0.65-0.78 Å for divalent or trivalent ions of Fe, Co, and Ni, gold may retard the crystal growth of the coprecipitates leading to a larger specific surface area than that of the pure oxides of Fe. Co. and Ni.

In a sodium cart nate solution of pH around 8.5, gold tetrachloride anions were progressively transformed, before coprecipitation, to gold hydroxide anions thereby releasing free chloride ions. This reaction is not as fast as the precipitation of hydroxides of Fe, Co, and Ni. With an increase in Au content, the amount of chloride ions released appreciably increases during coprecipitation to change the precipitation conditions. Au at 5 at.% causes release of chloride ions of more than one-tenth the concentration of nitrate ions. The coexistence of the chloride ions in a concentration comparable to that of nitrate ions might possibly result in a change in the size of primary solid coprecipitates and their coagulation phenomena. The above phenomena can be considered to be closely related to the occurrence of the bimodal change in specific surface area.

The final decline toward the small surface area of gold powder for a gold content above 20 at.% could be ascribed to the sintering of gold and the decreased proportion

of metal oxides. In fact, the catalyst with 20 at.% Au had the same appearance as that of metallic gold rowder. It should be noted that the maxit metally cativity was obtained at the minimum in specific surface area for Au/α -Fe₂O₃, but at the second peak or plateau in Au/NiO or Au/Co₃O₄.

It is noteworthy that the catalytic activity of Au/Al₂O₃ was comparable to those of Au/a-Fe₂O₁, Au/Co₁O₄, and Au/NiO for H₂ oxidation but remarkably inferior for CO oxidation. As shown in Fig. 4, the catalytic activity for H2 oxidation seems to be solely dependent on the particle diameter, namely the exposed surface area of gold metal. On the other hand, the support oxides may also play an important role in the oxidation of CO. Even though gold was supported in the form of small particles of 5 nm in diameter, the catalytic activity is not as high with inactive Al2O3 as that obtained for Au/a-Fe2O3, Au/Co3O4, and Au/NiO. These results indicate that either a kind of metalsupport interaction occurs or oxidation may proceed through a bifunctional mechanism in which both gold particles and support oxides activate different steps of the CO oxidation.

Hydrogen is considered not to chemisorb on the gold surface while CO chemisorbs weakly on the gold surface (10). Therefore, it is probable that the contribution of support oxides may differ in H2 oxidation from that in CO oxidation. The interaction of gold crystallites with the support oxides, a-Fe2O1, Co1O4, and NiO, which are all semiconductors, migh' alter the surface properties of gold crystallites so as to favor CO adsorption. According to simple calculations, outer surface atoms compose about 40% of the total atom content for hemispherical gold particles of diameter 4.0 nm. This suggests that the electronic states of the outer surface gold atoms can be readily modified by the interaction with the support oxides.

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GOLD-SUPPORTING TIN OXIDE FOR SELECTIVE CO SENSING

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Tetsuhiko KOBAYASHI, * Masatake HARUTA, and Hiroshi SANO

Government Industrial Research Institute of Osaka, 1-8-31 Midorigaoka, Ikeda, Osaka 563

The sensitivity and selectivity of semiconducting SnO2 towards CO at 150 - 250°C was appreciably enhanced by supporting highly dispersed gold and by doping with ${\rm Mg}^{2+}$ ions. The addition of Mg^{2+} prevented the Au/SnO₂ solid from sintering and maintained it catalytically active for CO oxidation, even in a moist atmosphere.

In a recent paper we reported that Ti4+-doped Fe203 supporting ultrafine gold particles with a diameter of about 4nm (hereafter denoted as UFP-Au) exhibits excellent CO selectivity against ${\rm H_2}$ and ethanol at an operating temperature below 100°C (1). This sensing property originates from the high catalytic activity of UFP-Au/Fe203 for CO oxidation at low temperatures [2]. The gas sensors operating below 100°C, however, seem to have the disadvantage that they usually need periodic heat-flashing in longtime operation to avoid the accumulation of water and contaminants on the surface [3]. A commercial semiconductor CO sensor, which is fabricated from Pd-supporting SnO2, also uses periodic heatflashing to remove CO molecules which are adsorbed at around 90°C [4].

In order to obtain a CO sensor operated at a constant temperature without the need for heat-flashing, a new gas-sensing semiconductor must be developed that exhibits sufficient CO sensitivity and selectivity at temperatures above 150°C. Since CO oxidation over UFP-Au/SnO2 needs a temperature at least 100 degrees higher than UFP-Au/Fe2O3 [5], a relatively high sensing temperature of CO would be required for the former material. Therefore, we have made an attempt to prepare new CO selective semiconductors

composed of SnO2 and Au. Semiconducting SnO2 incorporating Au. ${\rm Mg}^{2+},$ and Sb5+

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(Sn:Au:Mg:Sb = 100.1:5:1, in atomic ratio) was prepared by coprecipitation from a mixed aqueous solution of SmC14, HAuCl4, and SbCl5 with an aqueous solution of NH3. Since the electrical conductivity of SnO2 markedly decreased with the addition of Mg²⁺, the doping of Sb⁵⁺, which scarcely affected the gas selectivity of the semiconductor, was necessary to maintain the resistance of a SnO2 thick film at a level of 300 K Ω at 200°C. The precipitate was washed with distilled water, vacuum dried, and then impregnated with an aqueous solution of Mg(NO3)2. This precursor was vacuum dried again and calcined in air at 400°C for 3h. For comparison, Au/SnO2 (Au:Sn = 1:100, in atomic ratio) and SnO2 were also prepared in a similar manner to that described above.

Thick-film gas sensor devices were fabricated from the above semiconductors using final calcination temperature of 600°C for stabilization [1]. Gas sensitivity is expressed by R-air/R-gas, this being the ratio of the electrical resistance of the device measured in air with a relative humidity of 65% to that measured in



the presence of 300ppm CO or 300ppm H₂. The catalytic activities of these materials for CO and H₂ oxidation were measured by using a small fixed bed reactor $\{2\}$.

Figure 1 (a) shows the sensitivities to CO and H₂ of the Au/SnO₂/Mg²⁺/Sb⁵⁺ sensor as a function of operating remperature. A high sensitivity with a fairly good selectivity to CO against H₂ was obtained at temperatures between 150° C and 250° C.

Figure 1 (b) and (c) stow gas sensitivities for Au/SnO₂ and SnO₂. The addition of Au into SnO₂ improved the sensitivity for CO more appreciably than for H₂, but such a high selectivity to CO as observed in Fig. 1 (a) could not be obtained without the addition of Mg^{2+} . Since the



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addition of Mg2+ into SnO2 without Au did not improve the original sensitivities of SnO2, selective CO detection obtained with Au/SnO2/Mg2+/Sb5+ might be due to a synergism between the Au particles and Mg2+.



The XRD patterns of sensor F18. 2 materials. a; Au/SnO2/Mg2+/Sb5+ (color; black), b: Au/SnO2 (color; deep purple). . : Au. V: SnO2,



Fig. 3 The efficiency of CO and H2 oxidation over sensor materials as a function of temperature: a: $Au/SnO_2/Mg^{2+}/Sb^{5+}$, b: Au/SnO_2 .

P2Os column),

1% CO + air (saturated @ ---- B ---- i with H2O at O°C), - 0 - - 0 - -: 1% H2 + air.

Space velocity; 20000h-1 w1 g-1

Figure 2 shows the XRD patterns of Au/Sn02/Mg2+/Sb5+ and Au/Sn02. of The XRD peaks Au/SnO2/Mg2+/Sb5+, which are broader than those of Au/SnO2, clearly indicate that Mg2+ suppresses the sintering of Au/SnO2 to give smaller gold particles. The Mg2+ doped sample show no appreciable XRD peaks due to metallic Au even after heat-treatment at 600°C. The black color of this material may indicate the presence of highly dispersed Au particles, which are smaller than those in Au/SnO2.

Direct information concerning the difference in the sensitivities of Au/SnO2 with and without Mg2+ was obtained by measuring the catalytic activities of the materials for CO exidation. Figure 3 shows that CO oxidation takes place in preference to H2 oxidation over the above two materials in the dried reaction gas. In the humidified reaction gas, water vapor does not affect the catalytic activities of Mg2+ doped Au'SnO2 but inhibits CO oxidation over Au/SnO2 without Mg2+ decreasing the preference to CO oxidation.

The difference in the size of Au particles may account for the difference in the sensing

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properties of the Au/SnO₂/Mg²⁺/Sb⁵⁺ and the Au/SnO₂ sensor. It has been found in the case of UFP-Au/Fe₂O₃-Ti⁴⁺ that the sensitivity as well as the selectivity towards CO increases with smaller Au particles [1]. The measurements of gas sensitivities were conducted under humid conditions, despite this Au/SnO₂/Mg²⁺/Sb⁵⁺ remains selective to CO due to the fact that it is unaffected by water in the catalytic oxidation of CO. Recently, UFP-Au supported on hydrous oxides of alkaline earth metals have been found to catalyze CO oxidation even at -70°C [5,6]. An increase in the basicity of SnO₂ by doping with Mg²⁺ might play an important role in the vater-resistant o ... n of CO.

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COを選択的に検知する金担持酸化スズ

小林哲彦、* 春田正穀、佐野 寛

工業技術院 大阪工業技術試驗所 平563 徳田市議ケ丘1-8-31

150~200°Cの中濃銀域における SnO2ガスセンサのCOに対する検知感度および選択性 は、高分散した金粒子およびMg²・イオンの添加により著しく増越された。 Mg²・の癒 加により Au/SnO2の検結が抑制され、選分を含む雰囲気中でもCO酸化の触媒活性 が雄神された。

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METHODOLOGY FOR MAKING R&D PROGRAMS OF CHEMIC . "NSORS

Masatake HARUTA, Kazuo HIIRO, Hideo TANIGAWA, Hiroyasu TAKENAKA, Susumu YOSHIKAWA, and Hiroshi SANO

Government Industrial Research Institute of Osaka Midorigaoka 1, Ikeda 563, Japan

ABSTRACT

The R&D programs of chemical sumsors should particularly be focused on the practical needs. Sensing materials and signal transducers should be exploited efficiently and should be combined synergetically by taking into consideration of the sensitivities, selectivities, and stybilities required. The methodology for program making is presented by showing the examples of Government Industrial Research Institute of Csaka. The practical needs, technological bottle necks, prospects of development, and fields of application were disclosed through the styvey by distributing questionnaires to the people engaged in chemical sensors. The results of needs analyses together with seeds exploration substantially helped the establishment of the needsorientated and motivation-enforced R&D programs. The activities and achievements of research carried out in the framework of the programs at GIRIO are also described.

1. INTRODUCTION

Chemical sensors are a sort of artificial eyes to recognize chemical species. They measure the presence or connentration of gares, ions in solutions, and organic compounds. They are widely used in microwave ovens, gas safery alarms, air-fuel ratio control systems fc. : 'omobiles, monitoring of water pollution, medical drug delivery devices for diabetes, process control of fermentation and so forth. They are really becoming indispensable for many industries and for our daily lives as well.

There are various kinds of chemical sensors: multifarious sensors are existing and will be needed even for a single chemical compound depending on the conditions of operation, sensing materials, transducers, and detecting principles. Especially, sensing materials used are diverse from organic metal complexes, conductive polymers,



biological materials, metals, and metal exides. This situation makes it difficult for users to choose suitable chemical sensors without a vast amount of knowledge and for engineers and makers to find the efficient approach to the development of a sensor specifically desired.

From the above points of view, it is very important to establish the methodology for making R&D programs of chemical sensors. The R&D programs of Government Industrial Research Institute of Osaka(GIRIO) were made based on the results of questionnaire survey of social needs to chemical sensors and on the results of literature and patent survey of the past ten years. Several distinguished results were obtained in the framework of the R&D programs. These activities at GIRIO for the past five years are presented to show one of the examples of the methodology for program making and its validity.

2. THE STRUCTURES AND PERFORMANCES OF CHEMICAL SENSORS

Chemical sensors can be classified into four groups: humidity, gas. ion. and bio-sensors. Structually, they are divided into two types[1] as shown in Fig. 1. The first type is called "unstructured sensors", where sensing of chemical species and transducing the resulting physico-chemical changes to an electrical signal is performed in a single phase. Humidity and gas sensors using sintered ceramics are typical examples of this type. The adsorption and reaction of water or flammable gases on the surface results in a change of the electrical conductivity of the sintered body ceramics, thus leading to an increase in electrical current under a constant applied voltage.

The second type is "structured sensors", b-cause the part of receptor for chemical species and the part of transducer are clearly separated into two phases. Conventional ion selective electrodes are a typical example of this type. They are combined with reference electrodes to measure the potential difference between the two electrodes. Almost all biosensors are also included in this category. For example, glucose oxidase recognizes glucose and then catalyzes the reaction to produce H_2O_2 , the concentration of which is transduced with an oxygen electrode to an electrical signal.



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11. STRUCTURED SENSORS

Fig. 1. Structure of chemical sensors,

Many advanced sensors have been developed in the structured type by using FET(Field Effect Transistor)[2], SAW(Surface Acoustic Wave) devices[3,4], piezoelectric crystals[5,6,7], fluorescence[8,9], and fiber optics[10] as a transducer. One of the advantageous features of this type is that it allows us to choose and prepare sensing materials without taking into consideration of their transducing properties. The typical examples of sensing membranes are cellulose acetate butyrate[11] for humidity, Pa[1] and SnO2[13] for gases, Si3N4[14], NAS glass[15], valinomycin[16] for ions, and urease and glucose oxidase[17] for bio-related i spounds.

The R&D work of chemical sensors consists of exploiting sensing materials and transducing devices, fabricating whem into sensing

devices, and testing sensor performances. There are three "S"s in the performances of chemical sensors. The first is gensitivity which includes range of detectable concentration, resolution with respect to a certain concentration change, and response time, namely, three "r"s. The second S is gelectivity. The interference by other coexisting species is one of the serious problems of the present chemical sensors. It should be examined beforehand which kinds of species coexist in the atmosphere where the target sensors are used. The third S is galability which includes reproducibility of signal output during the repetition of measurements and durability in the long term use.

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3. SURVEY AND ANALYSES OF NEEDS TO CHEMICAL SENSORS

In order to establish a self-consistent and effective strategy for the R&D programs of chemical sensors, a comprehensive survey of social needs was made by GIRIO in 1983[18]. Questionnaire sheets were sent to 341 people who were engaged in chemical sensors at industries(256), universities(65), national and public research laboratories(14), and technical consultants(6). Forty eight per cent of them sent us back their answers, from which the needs to chemical sensors were analyzed as described below.

3.1 Overail Chemical Sensors

Figure 2 shows the number of answers to the question "Please choose three chemical sensors which you think will grow remarkably in the future.". The area of each circle is proportional to the number of answers. In the category of gas sensors, oxygen sensors gained the largest number. It seemed to be reasonable because oxygen sensors were being commercially used for the control of air-fuel ratio in automobiles. The second is CO sensors. This is probably because there were no reliable CO sensors to monitor whether combustion of fuel in household appliances is safely taking place. The third is H2O, namely, humidity sensors. This might be due to the very humid climate in Japan. Humidity sensors are in a wide-spread use for air conditioners, video tape recorders, automatic cooking oven ranges and





Fig. 2. Sensors to be expected to grow. Figures in the circles show the number of answers.

so forth. It should also be noted that nitrogen oxide sensors obtained more points than hydrocarbon fuel sensors which had been commercially used for gas leak alarms.

In the category of ion and bio-sensors, the expectation to sensors for bio-related compounds is far the largest. Sensors for organic compounds gained answers as many as oxygen or CO gas sensors did. The broad definition of biosensors includes not only the sensors with biological materials such as enzyme, immuno-assay, micro organisms for sensing but also other types of sensors which simply measure bio-related species such as Li⁺ concerned with psychology and E⁺ in body fluid. These fields of biosensors are expected to make the greatest progress in the future. As for ion sensors, heavy metal ions and nitrogen compounds such as NO₃⁺ are ranked at the top and at the second, respectively.

The opinions concerning chemical sensors described freely by answerers are schematically summarized in Fig. 3. The area corresponds to the number of opinions. In the category of gas sensors, PH3 and AsH3, doping gases used in semiconductor industries, CO, and humidity are attracting large concerns of answerers. Relatively great demands are seen to the high temperature use of gas sensors and to some extent



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Fig. 3. Schematic summary of the opinions written freely by correspondents.

to the low temperature use. In the category of ion and biosensors, smell and taste sensors are pointed out as a future target of research. In sensors for rare earth metal ions, NO3⁺⁺, organic acids are also a big concern. Among the requirements for the performances of chemical sensors, selectivity and stability are of the greatest.

It is important for making R&D programs to grasp properly the technological problems. Figure 4 shows that the most serious problems are degradation, reproducibility, stability of signal output, and contamination by oil and dust. These all concerns the stability of sensors in a wider sense of meanings.



Fig. 4. Problems of chemical sensors.



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Fig. 5. Annual growth in the number of papers and patents concerning chemical sensors.

A literature and patent survey was done for the past ten years to know the state of the art of chemical sensors mostly from the point of seeds. Figure 5 shows that there is a big contrast in the number of scientific papers and patents between ion sensors and the other chemical sensors. In the field of ion sensors, scientific papers are much more than patents. probably because R&D work is mainly done in university laboratories. especially as the subjects of analytical chemistry. On the other hand, in the fields of humidity and ges sensors the number of patents largely exceeds that of scientific

papers indicating that R&D work has been carried out mainly in industries.

3.2 Humidity Sensors

As shown in Fig. 6, few people think that the present humidity sensors have no serious problems and that their tost are too high. The biggest problems are reproducibility and precision, secondly, stability of signal output, thirdly, durability and life and then the range and conditions of measurements. The former three constraints are more or less concerned with the stability properties of sensors in a short and long term use. Therefore, an improvement in stability is considered to be the most important task of R&D on humidity sensors. Figure 7 shows the fields of application of humidity sensors. They are expected to be the most frequently used for air conditioning in residences and offices, probably because the humid hot summer and dry cold winter in Japan. The second largest field of application is drying processes, where high temperature performance and durability is particularly required for humidity sensors.



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Fig. 6. Problems of humidity sensors.



Fig. 7. Fields of application of

humidity sensors.

3.3 Gas Sensors

Figure 8 shows several problems of gas sensors with their



Fig. 8. Problems of gas sensors.

s sensors with their respective percent fractions in the number of answers. Selectivity occupies the largest fraction. Then, limitation to the applicable conditions, life, and stability occupy almost equal fractions to each other.





Table I shows more detailed data how differently producers, sellers, users, and researchers recognized the problems of each gas sensor. Conterning CO gas sensors, although producers and sellers do not think there are any serious problems, users and researchers complain of selectivity, stability, and applicable conditions. As for fuel gas sensors which had been already commercialized in Japan, selectivity and stability are considered similarly by the four groups to be very serious problems. Therefore, it may be suggested that the types of CO gas sensors under consideration are different between the group of producers and sellers and the group of users and researchers; the former is actually dealing with electrochemical sensors and the latter is developing oxide semice, uctor or other types of compact sensors.

In the cree of oxygen sensors, durability is the largest constraint and the second largest ones are convenience and conditions for use. There also observed a big discrepancy in answers between the group of producers and sellers and the group of users and researchers. This may again suggest that users and researchers were seeking for stabilized-zironia based oxygen sensors or other ones of the new type which were different from conventional elect. Amical sensors, Concerning hydrogen sulfide and sulfur dioxide gas Lensors, for which electrochemical cells have long been used, no serious problems are pointed out for the present.

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Durability			-	-	-	+ 44	100	**				-	*		-	-	15
alar+++++++	-		-	-	(4.44	-	-	-	*		-	-			-	-	93
Conditions		-	-			+	-		+	-		-	-		-	14	*3
Stability	**	+	-	-			****	-	+	-		*	+	*		-	68
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Table I. Constraints of producers, sellers, users, and researchers on the respective gas sensors.





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Fig. 9. The number of people engaged in gas sensors.

Figure 9 shows the number of answerers who were engaged in respective gas sensors at the time of questionnaire survey. The number was by far the largest for those who were engaged in O_2 sensors. Sensors for H_2 , CO, CH₄, and hydrocarbons had almost the same number of people engaged, however, the fraction of R&D was relatively large in the case of CO gas sensors.

The comparison of the present situation shown in Fig. 9 with the future situation estimated from the number of answers to the question "In the future which gas sensors do you wish to develop, use, sell, or produce?" can tell us the growth potentials of respective gas sensors. The ratios of the numbers of the future involvement to those of present one, namely, "future indices", are shown in Table II. The production and selling of CO sensors will appreciably grow. Nitrogen oxide sensors are exceptional that they will grow appreciably in every sector. Users will increase in the sensors for alcohols, NO_X, halogens, and SO₂. Research & development activities will be strengthened in the sensors for SO_X, halogens, H₂S, NO_X, and H₂, Strong interest in MOSFET sensors might be responsible for the large index of H₂.

Among the fields of application of gas sensors, prevention of gas explosion and of accidental death by oxygen shortage is considered to be the most important(Fig. 10). Industrial applications to process control in chemical facturies and energy savings are regarded as the second most important.

Table II. Future indices for various gas sensors.

	1.11	Fatara.							
Present		P	1	ų	ġ.				
81	100	122	108	- 11	146				
00	300	200	142	108	104				
CH4	100	15	118	-61	81				
нć	100	.61	6.9	78	139				
alconols	100	161	. 58	217	80				
01	100	10.6	13	6.3	1.13				
00.	100	125	TIS	70	103				
halogens	100	50	43	122	161				
NRS	100	34	80	6.2	61				
H ₁ 5	100	. 30	36	114	14				
501	150	60	44	120	201				
U.S.	100	580	1118	162	1.143				

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3.4 Ion Sensors

The general problems of ion sensors that were pointed out by correspondents are first life, secondly selectivity, and thirdly inconvenience in using sensors(Fig. ll(a)). What appear to be characteristics of ion sensors are that relatively large fraction of people feel no problems and that many people regard them as inconvenient to use. The former is due to the long history of commercial use of ion selective electrodes, as typically represented by pH electrodes. The latter is due to the fact that they need a reference electrode to measure potential difference.



The problems of several respective ion sensors are also shown in Fig.11. Concerning proton ion sensors, more than 70% of correspondents answered "there is no problem". In the following three ion sensors which were expected to grow markedly in the future, no one answer "no problem". The heavy metal ion sensors need more improvements in fundamental properties such as life, selectivity, and applicable conditions than in stability of operation, reproducibility, and precision of measurements. Similarly, for organic compounds, life, selectivity, and applicable conditions are the major constraints. In the case of bio-related compounds, the problems concerning stability and reproducibility are as serious as the problems of life, selectivi-

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ty, and applicable conditions. This might be because living biological materials like enzyme, microorganism are used as sensing materials.

Figure 12(a) shows the number of answerers who are concerned with respective ion sensors. Those who are concerned with proton occupy the largest number, halide ions the second, and alkaline and cyanide ions the third and then bio-related compounds. It should be noted that the fractions of R&D on organic compounds and on bio-related compounds are very large while that of users is more than half in proton sensors.

When the future involvement shown in Fig. 12(b) are compared with the present one, a marked increase is observed in organic and bio-related compounds whereas an appreciable decrease is observed in proton. It is worthy of noting that the fractions of R&D are relatively large in both present and future involvement in heavy metal ions and nitrogen compounds.





As can bee seen from Fig. 13, ion sensors are expected to be very useful for the automatic chemical analyses and speedy and simple clinical analyses in medical treatment. The second largest fields of application are automation. quality control in factories, and monitoring of pollution by poisonous heavy metal ions.

Fig. 13. Fields of application of ion sensors.

3-5 Biosensors

Table III shows one of the typical data obtained from an intensive investigation of biosensors which was made by Osaka Science & Technology Center in 1986[19]. Glucose and Na⁺ and K⁺ ions are most frequently measured in clinical analyses. Ions such as Cl⁺ and H⁺, as well as O_2 and CO_2 gases, are also very often measured.

There are four types of chemical sensors which use biological materials for molecular recognition, that is, enzymes, microbes, immunity, and the others like organera. Figure 14 shows the number of Japanese patents concerning biosensors published in the period of 1980-1984. Enzyme sensors occupy by far the largest portion. More than 90 species can be detected by using different types of enzyme sensors, eleven of which are already commercially available. In this sense, among biosensors enzyme-based ones are the most popular and successful.





Table III. Frequency of measurements with biosensors in clinical analyses.

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liucove	1.3	3	1	1.8
.20110 -0610	2	1.	3	
/r:1.acid	3	1.1	1	5
red	1	1	1.1	- 1
creatinine	2	2	0	4
aev) dsa	1	3	0 1	2
N 4 *	1.14	1	1	19
К "	1 1\$	1	1 2	20
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¢1*	1 12	1	1	6
NR:*	1.1		4	6
0,	1.13	5	1 3	1.8
ć 0,	10		4	6
pH	1 12	8	- 3	21



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(32 answerers)

4. R&D PROGRAMS OF CHEMICAL SENSORS IN GIRIO

Based on the results of needs and seeds analyses, a few targets were set up for humidity, gas, ion, and bio-sensors. The first target for humidity sensors, the operating temperature above 200°C, is mainly for the application to combustion exhaust gases. The second target for linearity of signal output is for the simple signal processing in monitoring process gases and air conditioning. As for gas sensors, selectivity to CO is not only one of the largest practical requests but also the most rewarding fundamental research subject. Low temperature operation is also attempted for the future combination with silicon devices. In ion sensors, extension of measurable ions to heavy metal ions and organic acids and miniaturization through ISFET are two main targets. Lastly for biosensors, proteins in blood and K⁺, Na⁺, Ca²⁺, Cl⁻ ions in body fluid are targeted together with multi-enzyme electrodes.

4.1 Humidity Sensors

A glassy material composed of $V_2O_5 - TeO_2 - Li_2O + Ag_2O - K_2O(2O)$ was exploited for a humidity sensor which could be operated at above 200°C. The main component $V_2O_5 - TeO_2$ is one of electrically conductive glasses, to which Li_2O was added to provide moisture sensitivity. The addition of Ag_2O was to reduce the electrical resistivity. The role of K_2O was to enhance the servitivity to trace amount of moisture. The above compounds are sixed and ground well. To the fine powder urea was added and then it was pressed and calcined at 300-350°C. The addition of urea made the sintered elements macro-porous to facilitate the diffusion of water molecules into the inside of the elements. 16

The material without K_2O can be used as a dew point sensor for VTR heads and the rear windows of automobiles. Figure 15 shows that the electrical resistance decreases only slightly until it sharply drops at a relative humidity around 90%. This sharp drop can be operated as an on-off switch which works at around a dew point.

On the other hand, if K_2O is added, the material turns to be sensitive to the whole range of humidity as shown in Fig. 16. The resistance change of two order of magnitude can be obtained for a



relative humidity variation from 10 to 70 %. The response was also sufficiently rapid that the steady-state signal values toward increased humidity could be reached within 30 seconds and toward decreasing humidity within 60 seconds. It was also confirmed that this sintered glassy material could respond to a small change of voisture from 16 ppm to 31 ppm at least up to 230°C. Another type of humidity sensitive material mainly composed of P205 and potassium titenate whisker was also developed[21].

In the second line of approach to the development of humidity sensors with linear response, the techniques of solid polymer electrolyte(SPE) water electrolysis[22] have been applied. Figure 17 snows the structure of SPE amperometric humidity sensor. Noble metals of 2+4mg/cm² are chemically deposited on both sides[13]. When a potential over 3V is applied between the two electrodes, the observed



Fig. 17. Structure of pPE amperometric humidity sensor 1. Solid polymer electrolyte. 2.3 Electrone. * SPE-electrope composite. 5.4 Leto



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A. Rt/Nafion120/Rh (0.1cm²), O.Rh/Nafion112/Rh (0.1cm²), Rh/Nafion15/Rh (0.25cm²) Rt/Nafion125/Ir (0.25cm²), Rh/Nafion125/Pt T0.1cm²). current is proportional to the amount of water decomposed, which is again in proportion to the concentration of water in a gas stream. This amperometric sensors exhibited perfectly linear responses in the whole range of relative humidity as shown in Fig. 18. As monometallic electrode catalysts Rh, Ir, and Pt were found to be preferable owing to their corrosion resistivity. 18

4.2 Gas Sensors

There are two types of CO sensors commercially available. Semiconductive SnOg catalyzed by Pd needs periodical heat flashing to sweep out CO adsorbed at a low temperature[24]. Electrochemical CO sensors are not compact and are troublesome in maintenance[25]. Therefore, new sensing materials were exploited for the development of highly selective CO gas sensors which could hopefully be operated at low temperatures for the future combination with silicon devices.

The material newly developed was a novel gold catalyst which exhibited an extremely high activity for the oxidation of CO at low temperatures (26-30). Figure 19 shows TEM photographs of the gold catalyst prepared by calcination of Au-Fe coprecipitate in air at 400°C. The gold loading is about 13wt%, which corresponds to an atomic ratio of Au/Fe = 1/19. Very fine gold particles are homogeneously dispersed with a mean diameter of 4.1nm and a standard deviation of 36%. These gold particles are not spherical but hemimic spherical in shape and are contacted at their flat planes with α -Fe₂O₃ (hematite) exhibiting a specific crystal orientation of Au (111) plane toward the (110) plane of hematite. This epitaxial-like growth of gold particles makes them very stable against heating and reduction-oxidation treatments.

The catalytic proverties of the Au/a-Fe₂O₃ are very unique compared with those of single oxides and gold powder itself (Fig. 20). Over gold powder, CD oxidation takes place at much higher temperature than hydrogen oxidation. The same nature can be seen for Pd supported on Al₂O₃. On the other hand, on almost all the metal oxides, the oxidation of CD occurs at lower temperatures as typically shown by a-Fe₂O₃. The coprecipitated gold catalyst is so active that it can catalyze the oxidation of CD even at -70° C and has similar nature to





Fig. 19. TEM photographs of Au/q-FesOs catalyst calcined at 400°C.





- Ha exidation ---- : CO exidation
- 1: Au/a-FeiOs(Au/Fe=1/19. coprecipitation, 400°C)
- 2: 0.5 wt% Pd/AlrOs (impregnation, 300°C)
- 3 : Au fine powder
- 4 : CosOs (carbonate, 400°C)
- S: NIO (hydrate, 400°C).
- 6 : a-FeiOs (hydrate, 400%).
- Reaction conditions : catalysts 42-70 meshes. Hs or CO 1 vol % in air. space velocity = 2×10° h" milg



those of transition metal oxides. In addition, the catalytic activity for the oxidation of CO is not depressed but is enhanced by moisture.

In order to make hematite really n-type semiconducting to reduce electrical resistance, Ti^{4+} ion was doped at a concentration of 3 atom% (Ti/Ti+Fe)[31]. The thick film sensor devices were prepared by painting the paste of semiconducting materials calcined at 400°C on an alumina substrate with comb-shaped gold electrodes and then by calcining in air at 600°C. The electrical resistivity(R) was measured in a synthesized air moistened at a relative humidity of 65%.

Figure 21 shows the sensitivities expressed by the ratios of Rair to Rgas to CO, H₂, and EtOH as a function of device temperature. Without gold, Ti⁴⁺ doped hematite exhibited only small sensitivities to CO and H₂ even at temperatures above 150°C. Only EtOH was detected with high sensitivity, which was one of the common features of metal oxide semiconductor gas sensors. With finely dispersed gold, temperatures at maximum sensitivities shift by about 100°C toward lower temperatures for H₂ and EtOH. For CO, much larger shift was observed and an appreciably high sensitivity over 30 could be obtained at a device temperature below 50°C.

The logarithmic plots of both sensitivities and concentrations yield straight lines. Figure 22 shows that at 40°C 20 ppm CO can be detected with the same sensitivity as that to 1000ppm EtOH and with much larger sensitivity than that to 1000 ppm H₂. Furthermore, the slope for CO is larger than the other two showing higher resolution for CO.

In the next step, an attempt was made to develop an optical gas sensor with a structure schematically represented. Fig. 23[32]. The fiber optic temperature sensor under a thin 1. a oxidation catalyst detects a temperature increase due to the oxidation of CO over the catalyst surface. Since highly dispersed gold catalyst will possibly have high absorbability of infrared light and be readily warmed as a commercial infrared light sensor using evaporated gold thin film does, the surface of the gold catalyst can be kept clean from moisture and other adsorptive gases.

Optical chemical sensors are advantageous over the conventional electricity-based sensors in that they are free from the interference by electromagnetic noise and the danger of inducing gas explosion and





that they are capable of the control at remote location and of the direct linkage with an optical information system.

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A film of 0.5µm thick composed of a-FejO3 with highly dispersed Au was prepared by simultaneous sputtering. The pillar-structured thin film obtained was still less active than the thick film prepared from the paste of coprecipitated powder, however, it was sufficiently active at temperatures around 150°C. The preparation of thin film gold catalysts in connection with their optical properties is under study.

4.3 Ion Sensors

The first line of approach to the development of new ion sensors was the utilization of a natural lacquer. Urushi as a matrix material of ion sensitive membranes. Urushi is an oriental natural lacquer used extensively for Japanese lacquer-wares[33] and has excellent durability and mechanical strength. Many ion-selective electrodes for iodide[34], perchlorate[35], nitrate[36] and thiodya - 37] ions have already been developed using Urushi at GIRIO. The second line of approach was the application of FIT to the fabrication of miniaturized ion sensors.

Selenocyanate ion selective electrodes were prepared from tri-noctylmethylammonium selenocyanate ion+exchanger and Urushi(38). Figure 24 shows that linear potential-concentration curves with a slope of 60mV per decade were obtained within the concentration range of 10^{-1} - 10^{-3} mol/1 selenocyanate for the membranes composed of ionexchanger at 45+50wt%. The stellic response time was less than 30 seconds. The electrode exhibited constant potential within the pH range 1-10 and good selectivity except for a few cases. Figure 25 shows selectivity coefficients, Kij in the Nernstian equation $E = E^{\circ} \pm [(2.303RT)/zF]log(a_1+K_1+a_3).$

As the first step for the miniaturization, a coated wire electrode was linked to the gate lead-wire of a conventional field effect transistor [39]. This coated lead-wire ion sensitive FET(CLISFET) was prepared by using even weight of selenocyanate ionexchanger and Urushi. A linear response was obtained in the range of 10^{-2} to 10^{-5} mol/l selenocyanate.







Fig.24., Furential-concentration curves of idenocyanate identities relectrodes. with of ionic inanger: (7) 50 (2) 45 (7) 40 (4) 25

içit ⊒. Pig,25, Selectivity coefficients.



Fig. 26. Measuring circuit of Urushi ISFET.

A selenocyanate ion sensitive FET (ISFET) was fabricated by coating a mixture of selenocyanate ionexchanger(45%) and Urushi(55%) directly on the gate of a FET device with a size of 0.5mm x 6.5mm[40]. The FET device was water resistive owing to SigN4 coating. Figure 26 shows the measuring circuit of Urushi ISFET. A linear response was also obtained in the range of $10^{-1}-10^{-5}$ mol/1. The selectivity coefficients

and other characteristics of the above two FET sensors were almost the same as those of the ion selective electrode.

A chemically modified CLISFET for silver ion was prepared using silver-7.7.8.8-tetracyanoquinodimethane anion radical(Ag+TCNQ τ) film formed by dipping the wire in the TCNQ dry acetonitrile solution[41]. A CLISFET for copper(II)ion was also developed by using Cu=TCNQ τ film[42]. These sensors showed linear responses to the concentration of their respective ions in the range of 10⁻¹ to 10⁻⁵ mol/1 with Nernstian slopes.

New ISFET sensors using silica on sapphire(SOS) wafer are under development for the detection of copper(II) ion and gluconic acid.

4.4 Biosensors

As a typical ion sensor useful for clinical monitoring a K* CLISFET was fabricated by using valinomycin as an ion-sensing material



Response of CLISFET with 10-1 mol dm-1 interfering ion. Partov. Parts and Urushi as a membrane matrix[43]. Figure 27 shows i linear response in the range of 1 to 10^{+5} mol/1 of potassium ion. Sodium ion interferes slightly. However, rubidium ion terribly interferes by yielding a drainsource current as large as potassium ion does at the same concentration, which is also a constraint on commercial potassium ion selective energy.

Among the biosensors using biological materials for sensing, enzyme-based ones are the most successfully developed and becoming popular. More than 90 analytes can be detected with

various enzyme electrodes, eleven of which were commercially available until 1985. Multi-data processing has become very important for the beasurement of multi-component systems. An auto-talibration system with a multiplexer was developed for detecting simultaneously more than two components in an attempt to apply to freshness and taste sensors in the future[44].

Compared with the enzyme-based biosensors, immunosensors still have the problem how to transduce efficiently the antigen-antibody binding into a processable signal, because antibodies merely attach to their analytes.

An attempt was made at GIRIO to develop a direct amperometric immunosensor[45]. In body fluid, the antigen-antibody bindings are transduced into the enzymatic reactions of complements, which consist of the cascadic enzymatic systems of serin proteases and some regulator proteins. Terminal reaction of this cascade 's the

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formation of membrane attack complex, which is a channel of 11 nm in diameter, in the cell membranes or artificial lipid bilayers. Lipid membranes containing some hapten ligand, which were complete insulators, were formed in the filter and attached to the end of electrode. The membranes become conductive in proportion to the degree of the immunological reactions induced by the addition of antiserum sample. The conductivity change is measured by an usual amplification apparatus. 25

This detecting system, an unlabeled method named by us "Complement-mediated Amperometric Immunosensor(CAIS)", is highly sensitive up to sub-nanomolar concentration, easy-handled, and free from background noise caused by non-specific protein adsorption. This is applicable to the detection of both immuno-assay and complement assay.

5. CONCLUSION

Research on chemical sensors at GIRIO has been initiated as a practical subject of surface science and at the same time as one of the attempts to explore new functional surface materials. The social needs to chemical sensors were gathered through the questionnaire survey. The analyses of the answers led to the establishment of selfconsistent and efficient strategy for the R&D of humidity, gas, ion, and bio-sensors. Through the exploitation of new sensing materials and the application of material technology already accumulated in different lines of research, several new chemical sensors were successfully developed.

The above experiences concerning the methodology for making R&D programs of chemical sensors led us to the following conclusions.

- Needs-orientated programs are particularly important for the R&D of chemical sensors. It has t en often the case that necessity is the mother of invention of chemical sensors.
- Needs to chemical sensors should be properly translated into the required properties represented by the three Ss. namely, gensitivities, gelectivities, and gtabilities.
- 3) Seeds of chemical sensors, namely, sensing materials and transducing devices should be synergetically combined so as to

fulfill the requirements to chemical sensors.

4) Liaison between universities, national laboratories, and industries will be the key for the efficient program making and successful accomplishment. Scientists in university laboratories and engineers in industries are, expected to be good sensors to needs and seeds, respectively. Those who work for national laboratories like GIRIO will have to be good transducers for seeds to needs, and vice versa. 26

Ackowledgment. We thank Dr. Nakane, former director of material chemistry department, GIRIO for his guidance and support throughout the research. We also thank Mr. S. Wakida and Dr. T. Kobayashi for their significant contribution to the R&D work on ion sensors and gas sensors, respectively.

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•NEW_____Vol.17 No.2 May, 1989 TECHNOLOGY JAPAN

INNOVATIVE PRODUCTION NOW

A complise Lighting Manufacturing Plant

TOPICS

Shimone Prefecture's Exchange Groups between Different Trades Development of 1 - 5-Sea Submersible Shinkai 6500

HIGH-TECH INFORMATION

S. com for Efficient Treatment of Evalsticing Oil Wastercater
Tachnology for County Ceramic Materials on Motols
Water Alcohol Separation with Molos Fiber Membrane Module
Factorization of Policer Transistor in IC Form
Prodownia Saplings Cultured by Entechnological Method

NATIONAL R&D PROJECTS

V. Water Treatment System Actual Renaissance 90 Project —From MITT's Large-Scale Project—

FLASH

Special-Purpose Calculator for Dietetic Treatment Automatic Finger Sphygmananometer





NEW TECHNOLOGY & PRODUCTS

contact with coolant, losing their cooling effect, nowever. Hitachi Metais has conducted intensive research to develop a high rust-resistant stainlessteel for molds. If lowering of the cooling effect due to rust can be prevented, plastic products will continue to harden promptly as they do while the molds are new and no rust is generated.

This enhances productivity, and plastic molding manufacturers will have less downtime for descaling

* Mitachi Metals, Lid Public Relations President's Office 1+2 Marunouchi 2-chome Chicagasku Takro Tel: 03-284-4552 Telex, HITAMET J 24494

89-5-002-381 Inorganic Microcapsule

The General Industrial Research Institute, Usaka, has developed an inorganic microcapsule manufactured by the interface reaction process and it is being produced commercially by Suzah, Yoshi Industrial Co., Ltd

This microchsule consists of a powder and the a diameter of from several to several dozen micrometers Uses of conventional microcapsules are limited because their wall material is an organic substance having little resistance to heat whiter, and solvents in the manufacturing process, they are mutually agglomerated or adhere to production machinery which significantly reducts production yield.

The wall naterial of the new microcapsule is inorgatic such as calcium carbonate or iron oxide, so it has excellent resistance to we er, heat and chemicals. Being harmless, it eliminates the toxicity problem of convencional chemicals for forming capsule wills. As a result, it can be used as a base material for manufacturing cosmitics and pharmaceuticals.

In the interface reaction process, an evidesion of oil and an aqueous solution is used when precipitating the powder particles and for forming hollow spherical particles. Dispersing the substance to be encapsulated in th aqueous solution for a while causes it to be encapsulated in the microcapsules. Since the capsule wall is porous, the psulated substance is released ally so the capsule can be used

surposes other than those for conventional microcapsules. Research is in progress by the private sector to use the microcapsule for controlledrelease pharmaceuticals, such as powder deodorant and organic phosphorous medicines over a period of time, as a filler for functional sheets and films, and as a base material for cosmetics.

 Government Industrial Research Institute Osoka, the Agency of Industrial Science and Technology 8-31. Midoeigaoka Ischome Ikeaa Circ.

Osaka Tel: 0727:31:8351

89-5-002-382 Garments Madr of New Endothermic, Heat-Storage Fiber

Unitika. Ltd and Descente. Ltd have jointly diveloped a new endothermic, heat-storage fiber, called Solar α and using this fiber. Unitika has begun producing a new line of garments including winter work cloth inp and healthcare apparel for serile besons Solar α was used for producing skiwear for use in the Cargary Olympic Games. These sold ver well, so the company has trebled production of sportawear, autumn and winter apparel, including winter work clothing; and healthcare apparel for senile persons.

Solar α is a double-layer solar fiber consisting of a core and a sheath, with airconium carbide incorporated in the core layer. Zirconium carbide absorbs visible light rays and reflects infrated rays, so shortwave energy of less than 2 μ m, which comprises 95% of sunlight is efficiently absorbed for conversion into heat and storage in the fiber Since the fiber reflects longwave heat equivalent to infrared rays of over 2 μ m, it increases heat retention ω preventing the infrared rays (roughly 10 μ m) generated by the human body from being dissipated outside.

Initially, the companies tried costing zirconium carbide on the fiber, but the substance's tremendous hardness prevented spinning of the treated fiber. Subsequent research led to the devel-



Solar a Skiwear

opment of a core-sheath fiber whose endothermic and heat storage effects were fully proved at the Calgary Olympic Games, leading to brisk sales of skiwear



Based on these results. Unlika began mass production of various subtain and winter apparel and ketatstic (foot warmer) coverlets.

* Uninko Lid. Public Relations Sec. in 4.4 Nihonbashi Muromachi 3-c 1998 Chuo-ku Tokso Tel 03-246-7336 Telex J23303

82-5-002-383 New High-Performance Polystyrene

Asahi Chemical Industry Colluct has developed two new types of tighperformance polystyrene (PS) having superfative strength and fuster

One of these, general-purpose polystyrene (GPPS), has a high molecular reight and can be formed into arge objects. The other, high-impact polystyrene (HIPS), is mixed with rubber and has a remarkable laster and superlative impact strength. Both are characterized by performances comparable to those of conventional highperformance resins such as actionitrile styrene (AS) and action the butadiene styrene (ABS). The com-



通商産業省 工業技術院

大阪工業技術試験所

GOVERNMENT INDUSTRIAL RESEARCH INSTITUTE, OSAKA AGENCY OF INDUSTRIAL SCIENCE AND TECHNOLOGY MINISTRY OF INTERNATIONAL TRADE AND INDUSTRY

研究概要

当所は新材料の開発とこれに関連する 計測技術の確立を目標に研究を行ってい る。研究は各研究者の創意により、基礎 的な問題を取扱う経常研究とこれを基盤 として組立てられる特別研究に大別され る。更に国民経済上重要かつ緊急に必要 とされ昭和41年度から発足した大型工業 技術研究開発(大型プロジェクト)、エネ ルキーの長期的な安定供給を図るために 昭和49年度から発足した新エネルギー技 術研究開発(サンシャイン計画)、昭和53 年度から発足した省エネルギー技術研究 開発(ムーンライト計画)、また昭和56年 度に発足した次世代産業基盤技術研究開 発制度、昭和63年度から医療福祉機器技 術研究開発、平成元年度から重要地域技 術研究開発 更に平成2年度から地球環 **境技術研究計算**に参加して利々の技術の 研究開発を行っている。

Outline of Researches

Ordinary Research and Development

Special Research and Development

R & D Program

Research and Development on New Energy Technology Research and Development on Energy Conservation Technology

Research and Development Project of Basic Technologies for Future Industries National R & D Programs for Medical and Welfare Apparatus Specific Regional Technology Development System Global Environment Project

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圖平成2年度予算額(単位:百万円)

Budget for Fiscal Year 1990

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試験研究費 Research expenses	1.029
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合 Total	2.570

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・指定職を含む Including Director+General

關試験研究費内訳

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Research Expenses



平成2年度試験研究費内訳

Breakdown of Research Expenses in 1990

平成2年度研究課題 !

List of Research Project in FY 1990

特别研究

翻計測·標準控載 嘉柏厳欄会材料の強度斜価技術に関する研究 1. 機械的特性値評価法の研究 加速教会試験法の研究 日 新 は お お 利 福 ウィスカー複合セラミックスの構造を駆い関する研究 1. ウィスカー権合セラミックスの作製 権合セラミックスの撤損造と期性 超折案変化型記録材料に関する研究 1. 単一波長光感応型材料の研究 二波長光感に型材料の研究 複合化機能性セラミックスの界面伝導特性制固に関する 研究 1. 微細構造制卵の研究 2. 海索機構の研究 ハロゲン化物ガラスの光穿特灯制御に関する研究 1. 光学特性の研究 2. 成形性の研究 センシング能をもつ分子権合系に関する研究 1. 分子認識型バイオセンサーの創製技術の研究 イオン売郷性ガラス募扱の鋳製技術に関する研究 イオン伝導性ガラス薄膜の観髪技術の研究 ガラス薄膜の特性評価技術の研究 鹿性能適明セラミック兼顧製造技術に繋する研究 1. 精光制御薄膜製造技術の開発 意記建層層間化合物の合成に関する研究 1. 合成法の研究 2. 権能開発の研究 **潮反応,分離技術** 機切子・舞闘を利用した触家設計技術に関ける研究 1. 微粒子設計の研究 - 権限技術による国体表面の構造なび機能設計の研究 触媒表面のキャラクタリゼーション 關床線基繁確立技術 X線光学園子の裏担化に関する研究 職康子力平和利用技術 対点次変化体の漫出複動に関するナチュラレ・アナログ 92143 1. ガラス固化体の化学的发让性の研究 2. 天然ガラス等の実験半環境における反変の研究 翻公審防止技術 商業推出物処理材の開発に関うる研究 1. 無機質マイクロカブセル化級人はの開発 難分解性有害有機化学物類の触媒放化分解に関する研究 1. 固体輕鬆湿式酸化用輕磁の開発 翻国際特定共同研究事業 オプティカルマイクロガスセンサに関する研究 徽官民道那共同研究 定素及び定業系ハイプリッド確念材料の対価 向上接街に開する研究

SPECIAL RESEARCH

Reaction and Separation Technology

Technology for Establishment of Industrial Foundation Technology for Peaceful Use of Atomic Energy

Anti-pollution Technology

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次世代産藥基盤技術研究開発

個 召成ガス化用セラミックスタービンの要素評価等開発 製品特性評価(効果耗性)

- 製造プロセス技術(転性強化技術) 翻塞電流・寒磁界和電源材料評価
- 教会プロセス技術
- 職非離形光電子材料の研究制発
- 発現機構の解明及び評価技術構発

R & D PROJECT OF BASIC TECHNOLOGIES

Ceramic Turbine for Coal Gasification

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新エネルギー技術研究開発 (サンシャイン計画)

水業エネルギー技術研究開発 關水業の製造技術の研究 裏黒水藻気電解活の研究

四体務分子電解調水電解法の任职 日本課の醸造・貯蔵技術の研究

金鵬水業化物による水素的概括肉の研究 腰水漏利斯指衛の研究

水敷吸蔵含金を用いるエネルギー窒弊技術の研究 水県吸蔵合金電池の解析・評価

R & D ON NEW ENERGY TECHNOLOGY (THE SUNSHINE PROJECT)

Hydrogen Energy

Hydrogen Production Technology

Transportation and Storage of Hydrogen

Utilization of Hydrogen

省エネルギー技術研究開発 (ムーンライト計画)

新型電池電力貯蔵システム 顧新型電池の研究開発

新型電池の適用性拡大のための研究

新型電池の性能評価

鰓科電池の研究開発 羅藩融炭融進型堅料電池の研究開発

新規構成材料の研究 材料試驗與価 先導的・基盤的省エネルギー技術

翻塞リチウムイオン講覧性固体電解質の研究 鹽鯛触線帯を用いた省エネルギー的有機合成の研究

R & D ON ENERGY CONSERVATION TECHNOLOGY (THE MOONLIGHT PROJECT)

Advanced Battery Electric Power Storage System New Types of Batteries

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Fuel Cell Power Generation Technology Molten Carbonate Fuel Cells

Research on new component material Evaluation and lesting methods for componental

Leading and Basic Technology for Energy Conservation Research on High Lithium Ion Conductive Solid Electrolytes

Research on Energy Saving Organic Synthesis Using Cooper Catalyst

大型工業技術研究開発 (大型プロジェクト)

水総合再生利用システムの研究開発 (アクアルネッサンス'90計画) 顧査生物の研究開発 1. 国宝化散生物内高活性化 醫療業材の研究解発 1、 ガラス複合種の研究開発 2. 無機硬素料の評価技術 超先端加エシステムの研究開発 職イオンビームプロセス技術 発電施設用部材高だ加工装置の要要技術評価 聪品级 雛郎材の要局御船特性評価 高機能化学製品等製造法(海洋生物活用)の研究 開発(海洋バイオ) 题海洋生物活用基础技術 解紛嚢類の骨層 医胞技術(含ま・ベクター系の解発) 翻有用生体種種の提集、解明幕に係る技術 報生物の皆する天然有機化合物姿势機能の探索・解明 離音送輸送機用推進システムの研究制発 職トータルシステムの研究開発 新教四十年十二个技术办开京

NATIONAL R & D PROGRAM (LARGE-SCALE PROJECT)

New Water Treatment System (Aquil-renaissance: 90 project) Micro-organisms

Membrane Materials

Advanced Material Processing and Machining System Ion Beam Processing Technology.

Evaluation of Essential Technology in Advanced Material Processing System for Power Plants

Characterization of Micro areaed Surface ' ayer.

Fine Chemicals from Murine Organism

Fundamental Tech in for Utilizar on of Marine Organisms

Basic Technology for Research and Utilization of Useful Bio-

logical Functions Research on offensis in formal sources and environments and a second organic subfraction

Super/Hyper Sonic Transport Propulsion System riermia ibarrier boarris team







1. レーザー光による新敏硬化粉よの研究開発

翻動展内レーザー手術装置

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重要地域技術研究開発

職業対加工における高度表面絶理技売

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2. 城税利法のための素材またと、

SPECIFIC REGION TECHNOLOGY DEVELOPMENT SYSTEM

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データベースマネジ ジント システム等の開発

職者エネルギーは料開発支援システムの研究開発

DATA BASE MANAGEMENT SYSTEMS

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科学技術振興調整**螢蘆**点 基礎研究

剤プラズマ村和の合成と経歴。 ち労ル離極発道のキャラクタリゼーションと機能発現に関 する研究 毎化用繊維のミクロ構造と強度発現機構の整要的研究

行子権自有における光電子取出業員程の研究

金田格効能を利用した触客及び15学などが用新規材料の書。 を、1991年2月か

技術・情報サービス

Technology and Information Services

●技術指導・相談

当所では技術交流推載センターを設け、各種要望ご答えるた たに技術相談を否いは技術指導を行っている。

•受託研究

研究成果の普及、実用化のために、現行の研究内容を若干紙 長するとか、現行設備そのまま力若干の手直しをすることで直 接有効な資料を提供しうる場合に取り、民間企業力らの委託を 受けて、受託研究を実施している。「工業技術規受託研究規程」 (例和40年9月1日改正)

●共同研究

新技術戦争における研究活動の効率的な運用を図るために、 其間研究者ガイ份な技術的能力及び経理の基礎を有することを 要件として「工業技術院共同研究規程」(平成元年7月1日改正) に導づき民間企業等と共同研究を実施している。

●見 学

当所の設備や機器などを見学される場合はあらかじめ聞い合 わせの上、文**島**であ申し込み下さい。





刊行物·図書

の主な刊行物

大阪工業技術試験手報告

大阪工業技術試験計學報

大工业ニュース

大阪工業技術試験所詳衡

大阪工業技術試験研要賞

Publications and Library

• Z *

当所の認顧室には臨内外の専門認識、文献、雑誌などを常備 している。当認識証所載の認識は、所替の紹介で覚護すること だできる。なお、外面特認、国内維持の母々について、工業技 内院試験研究機関認識達超会議編「1988年版、外面雑誌総合告 録。、1988年版 国内結誌地合目録。を希知下さい。

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8t Total	17,800	42,300	60,100



鄹 内

大阪工業技術試験所

〒563 大阪 6池田市緑丘1-8-31 TEL 0727-51-8351(代表) FAX 0727-51-2156(総務部)

Government Industrial Research Institute. Osaka

Midongaoka 1-8-31, Ikeda. Osaka. 563 Japan



案内図 Guide Map

Chemistry Express, Vol.3, No.3, pp.159-162 (1968)

KINKI CHEMICAL SOCIETY. JAPANO

PREPARATION AND CATALYTIC PROPERTIES OF GOLD FINELY DISPERSED ON BERYLLIUM OXIDE

Masatake HARUTA, Kenji SAIKA, Tetsuhiko KOBAYASHI, Susumu TSUBOTA, and Youhiko NAKAHARA

Government Industrial Resea ch Institute of Osaka, Midorigaoka 1-8-31, Ikeda 563 † Osaka Institute of Technology Omiya 5-16-1, Asahi-ku, Osaka 535

Ultrafine gold particles supported on beryllium oxide have been prepared by calcination in air of the coprecipitates obtained from an equeous solution of $HAuCl_4$ and $Be(NO_3)_2$. The coprecipitate with an atomic ratio of Au/Be=1/19 calcined at 200°C was the most active and could catalyze the oxidation of H_2 at 0°C and CD at -70°C.

Experimental data presented in the previous papers[1,2,3] have demonstrated that coprecipitation, instead of impregnation, enabled us to prepare gold finely dispersed on metal oxides and that the oxide of Fe, Co, or Ni as a support could yield very active catalysts for the exidation of carbon monoxide at 1-w temperatures. During the course of an invertigation undertaken to see if other metal oxides could also be used as a support suitable for markedly enhancing the catalytic activity of gold, we have recently found that some alkaline earth metal oxides act as an effective support. The present investigation deals with the preparation of gold finely dispersed on beryllium oxide and its catalytic activities for the oxidation of hydrogen and carbon monoxide.

The new gold catalysts were prepared by coprecipitation with sodium carbonate from an aqueous solution of HAuCl_A and Be(NO₃)₂. The coprecipitates were washed, vacuum-dried, and calcined in air at different temperatures for 3 - 18h. Catalytic activity measurements were carried out in a small fixed bed reactor, with 0.10g of catalysts that had passed through 70 and 120 mesh sieves. A standard gas consisting of 1.0 vol² H₂ or CO balanced with air to 1 atm. was passed through the catalyst bed at a flow rate of 33 ml/min. The reaction gas was dried by passing through a silica gel column cooled to -77°C to prevent the accumulation of moisture in the catalyst layer. The analytical techniques were

Chemistry Express

similar to those described elsewhere[1,3].





Fig.1. Catalytic activities for the oxidation of H, and CO as a function of Au content. A, H: at 30°C; O, CO at -70°C.



Fig. 2. Effect of calcination temperature on the catalytic activities for the oxidation of of Ha and CO and on specific surface area. Till expresses the temperature where 50% conversion is attained. A , H: at 30°C; O, CO at -70°C; D, specific surface area. The arrows denote that Tils values are lower than indicated.

Figure 1 shows exidation efficiencies for H2 at 30°C and CO at -70°C as a function of Au content in the Au-Be coprecipitates calcined at 200°C. The oxidation of CO over the Au-Be catalysts can take place at much lower temperatures than the oxidation of H_2 . This feature presents a striking contrast to the catalytic behavior of the conventional SiO₂ or Al₂O₃ supported Au catalysts, where H2 can be oxidized at lower temperatures than CO. Maximum catalytic activities were obtained in the range of S-10 atom%(Au/Be=1/19-1/9) for both H₂ and CO oxidation. This range of Au content coincided with the optimum Au contents for Au/Fe203, Au/CogO4, and Au/NiO.

> Figure 2 shows the effect of calcination temperature on the catalytic activities of the Au-Be(1/19) coprecipitate. Maximum catalytic activities were obtained for both H_2 and CO oxidation when the coprecipitate was calcined at 200°C, where a maximum value was also attained for specific surface area. This may indicate that the larger surface area was preferable in obtaining higher catalytic activities in Au-Be system. On the other hand, higher calcination temperatures ranging from 300°C to 400°C were required for the Au-Fe coprecipitate to generate markedly enhanced catalytic activity for CO oxidation[1,3] This is probably because strong interaction at the interface between Au and Fe₂O₃ which might be created by calcination at

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relatively higher temperatures was more important than the large surface area of Fe_2O_3 . The activity of the Au-Be(1/19) catalyst was so stable even at -70°C that 100% efficiency of CO oxidation was maintained in a continuous run fr. 42 h, while the efficiency declined to about 50% in a few hours over Au/Fe₂U₃[1]. In the I-ray diffraction patterns shown in Fig. 3, a broad plateaux

In the 1-ray diffraction patterns when a sample calcined at 150 °C. The assigned to metallic Au was first observed in a sample calcined at 150 °C. The peak became more apparent at 200 °C and from the peak half-width the mean particle diameter of gold was estimated to be 4.3 nm. Since DTA data indicated that the formation of unhydrous BeO starts at 280 °C, the most active catalysts were considered to be composed of ultrafine Au particles and hydrous BeO. The particle diameter of Au increased only slightly with an increase in calcination temperature; it was 4.9 and 5.1 nm for the samples calcined at 400 and 500 °C, respectively. Therefore, it is likely that not unhydrous but hydrous BeO played an important role in enhancing the catalytic activity.

Figure 4 shows the binding energies for the maxima of Au $4f_{7/2}$ and $4d_{5/2}$ IPS peaks for the Au-Be coprecipitate. The binding energies for the sample cricined at 200°C are larger by 0.5eV than those for Au bulk metal, whereas the sample calcined at 400 and 500 °C exhibited no such chemical shifts. Since the Au-Fe coprecipitate calcined at 300-400 °C, which exhibited catalytic activities close to that of Au-Be calcined at 200°C, showed chemical shifts of almost the same degree, it enpears that the electron defficiency that gives the chemical shift of about +0.5 eV leads to the most suitable surface state for the low temperature oxidation of CO. Accordingly, on ultrafine Au particles supported on hydrous BeO the adsorption of CO might turn to occur moderately, neither too weakly as on pure gold surface nor too strongly as on platinum surface[A].









Calcination Temperature / *C

Fig.4. Variation of the binding energies for the maxima of Au $4f_{3/3}$ and $4d_{3/3}$ XPS peaks with calcination temperature.

The reason why only Au/hydrous BeO could maintain high catalytic activity for CO oxidation at -70°C is not clear at present. It might be, at least pertly, due to the difference that carbonate species are hardly formed and unstable on the surface of BeO while they are readily formed and relatively stable on the surface of the oxides of Fe, Co, and Ni.

The authors thank Professor S. Minami of Osaka Instants of Technology for his continued encouragement throughout the work.

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愈を高分数担待した酸化ペリリウムの調製と触媒特性

春田正驗,* 西家離司,* 小林哲娜、 坪田 年、 中原佳子

大阪工業技術試験所 〒563 地田市緑ケ丘 1・8・31 +大阪工業大学 〒535 大阪市旭区大宮 5・16・1

塩化金酸と硝酸ペリリウムの混合水溶液から得られた共沈物を空気中で焼成することに より、金を超微粒子状にして酸化ペリリウム上に分散担持した。 200℃で焼成したAu/Be= 1/19の組成をもつ共沈物が豊も触媒活性が高く、H2及びCOをそれぞれ0℃、・70℃でも酸化 できる。

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Paper No 181

PREPARATION OF HIGHLY DISPERSED GOLD ON TITANIUM AND MAGNESIUM OXIDE

Susumu TSUBOTA, Masatake BARUTA, Tetsuhiko KOBAYASHI, Atsushi UEDA, and Yoshiko NAKAHARA Government Industrial Research Institute of Osaka Midorigaoka 1, IKEDA 563, Japan

ABSTRACT

Gold could be highly dispersed on titanium oxide and magnesium oxide in their aqueous dispersion containing Mg citrate. The mean diameters of gold particles are smaller than These gold catalysts are active for the oxidation of CO Snm. even at a temperature below 0°C. The magnesia support, Mg citrate acts not as a reducing agent as a sticking agent which blocks the coagulation of gold particles. On titania support dispersed in neutral solution Mg2+ ions instead of citrate ions are mainly suporbed. It is likely that Mg2+ ion suppresses the transformation of amorphous titania to anatase during calcination and prevent gold particles from coagulation caused by earthquake effect.

INTRODUCTION

Gold has been regarded as catalytically far less active than platinum-group metals. This is because of its chemically inert character and of low dispersion in supported catalysts. We have recently reported that through coprecipitation gold particles smaller than 10 nm can be highly dispersed on Co304, d-Fe203, NiO1-3), and Be(Ch)24). These gold catalysts are active in the oxidation of CO at a temperature as low as -70°C. However, coprecipitation is valid only for a selected group of metal oxides as mentioned above, because the precipitation rates of support metal hydroxide and gold hydroxides and their affinity might determine in the dispersion of gold.

This paper deals with the methods for supporting gold in a highly dispersed state on pre-formed TiO2 and MgO powder, on which ultrafine gold particles have been difficult to be supported by the conventional methods.

EXPERIMENTAL

Preparation of gold catalysts

The following materials were used for catalyst supports; magnesia (Ube Industries, Ltd.; crystalline small particles prepared by vapor method; BET=14Om2/g), titania-A (Idemitty Optimized Co.; amorphoue dried at 120°C; BET=110m2/g), and titania-B (JRC-TIO4; anstase; BET=40m2/g). Each of these supports was dispersed in an aqueous solution of HAuCl4. The pH of titania dispersion was adjusted to 7.0 with Na₂CO₃, while the pH for magnesia dispersion, which was not intentionally adjusted, was naturally settled at around 9.6. The aqueous dispersions were stirred for 2 hrs after the addition of a variety of reagents (citrates of Mg, Na, or NH4, or HCHO; 2.5mol/Au for magnesia, and 5.0mol/Au for titania). These precursors were washed with distilled water and then filtered. The cake was vacuum dried and calcined in sir for 5 hrs at 400°C and 250°C for TiO₂ and MgO, respectively. The gold content of these catalysts thus obtained were latomX (Au/Ti) in Au/titania and 2atomX (Au/Mg) in Au/magnesia.

Catalytic Activity measurements

The activities of the gold catalysts were measured in the oxidation of CO or H₂. Experiments were carried out in a small fixed bed reactor with 0.10g of catalysts that had passed through 70 and 120 mesh sieves. A standard gas of 1.0 vol.% H₂ or CO balanced with air to 1 atm was passed through the catalyst bed at a flow rate of 33m1/min. The conversion of CO and H₂ was determined through gas chromatographic analyses (G-2800, Yanagimoto Co. Ltd.) of effluent from the reactor.

Characterization of Catalysts

The structures of the fold catalysis were observed using a Hitachi H-9000 electron microscope operated at 300 kV. X-ray diffraction (XRD) analysis was made by using a Rad-B system (Rigaku Denki Co.Ltd.). Infrared spectra were taken with a Nicolet 20-SXC spectrometer. For the IR analysis, each sample was mixed with KBr(2wt.% for magnesia; 10 wt.% for titania), and pressed into a thin wafer. Differential thermal analysis (DTA) was made by using a 3SC-5200 thermal analyzer (Seiko Denshi Kogyo Co.Ltd.). X-ray photoelectron spectroscopy (XPS) was measured with a SSX-100 spectrometer (Surface Science Laboratories, Inc.).

RESULTS

Gold supported on magnesia

Table 1 shows the catalytic activities of Au/magnesia prepared with different additives. It was found that catalytic

activities were enhanced by the addition of Mg citrate. When Mg; citrate was added into the suspension before the addition of HAuCl4, the activity enhancement could not be observed. The use of Na citrate or HCHO caused lower catalytic activity. The pH of the suspension during the preparation, usually 9.6, was increased to 11 when Na citrate was added. The addition of HCHO to the suspension produced a purple color, which indicated the reduction of Au³⁺ to colloidal gold.

Figure 1 shows the XRD patterns of Au/magnesia catalysts, where the presence of Mg(OH)2, not MgO, are evidenced. The starting material, MgO, changed to Mg(OH)2 by hydration in the aqueous suspension. From the width of the XRD peak of Au(200), the particle size of gold is calculated as about 14nm for Au/MgO prepared without additives, and this value is in good agreement with 10 nm determined by TEM observations. On the other hand, in the catalyst prepared with the addition of Mg citrate, gold particles smaller than 3 nm are observed by TEM. Although such a very small particles of gold did not show the diffraction peak in XRD, the presence of metallic gold were confirmed by the binding energy of 84.2 eV for the <u>XPS peak</u> of Au4f5/2. The catalyst prepared with the addition of HCHO contained only large gold particles (more than 20nm, by TEM observation).

Figure 2 shows the IR spectra of the precursor of Au/magnesia before calcination. Without Mg citrate, the IR absorption of surface H₂O and MgCO₃ are observed at 1638cm⁻¹ and 1449cm⁻¹, respectively. In the case of the precursor prepared with Mg citrate, other absorptions are detected at 1595cm⁻¹, 1423cm⁻¹, 1263cm⁻¹, 1083cm⁻¹, and 1061cm⁻¹. These absorption peaks coincide with those obtained for pure Mg citrate powder.

TABLE 1

Additives	co	Catalytic conv.,%	activity $T_{1/2}[H_2]$, °C
none		10	>200
Mg ct.		100	67
Na ct.		5	>200
HCHO		0	>200

Catalytic activity of Au/magnesia prepared with various additives.

CO conv.:CO conversion at -70°C

 $T_1/2$:temperature for 50% conversion

ct.:citrate













WULL SUPPORTED ON TRIANA

In Table 2, the effect of the addition of Mg citrate on the catalytic activity is compared on the two different types of TiO2 supports. While the catalytic activity of Au/titania-A (amorphous) is enhanced by use of Mg citrate, Au/titania-B (anatase) shows a high catalytic activity regardless of the addition of Mg citrate.

Figure 3 shows TEM photographs of Au/titania catalysts. When Mg citrate is added in the dispersion, the gold particles are highly dispersed on titania-A (the average particle size of gold is about 4nm), and gold particles become larger without Mg citrate. In the case of titania-B, however, the small gold particles are highly dispersed even when Mg citrate was not used.

The IR absorption spectra of the precursor of Au/titania prepared with Mg citrate were shown in Fig. 4. The adsorption peak at $1400cm^{-1}$ on titania-A might correspond to the adsorbed citrate species. Compared with the case of Au/Mg(OH)₂, however, the amount of citrate species is much less on the titania support.

TABLE 2

Catalytic activity of Au/titania prepared with and without Mg citrate, (Comparison of two different titania supports).

Titania Support	Addition of $T_1/2[CO]$	Catalytic ac Mg citrate T1/2[H2] ,°C	.T1/2[CO]	T _{1/2[H2]}
Titania-A(amorphous)	<0	25	35	139
Titania-B(anatase)	<0	35	<0	34

Ty/2:temperature for 50% conversion

TABLE 3

Catalytic activity of Au/titania~A prepared with additives.

Additives	Catalytic T _{1/2} [CO],°C	activity T1/2[H_],°C
none	35	139
HCHO	83	165
Na ct.	23	93
NH4 ct.	5	85
Mg ct.	<0	25
Mg (NO3)2	<0	35

ct.:citrate

T1/2: temperature for 50% conversion








Fig. 5. XRD patterns of Au/titania-A prepared with various additives. (a)without any reagent(blank); (b)with Mg citrate; (c)with Na citrate, (d)with $Mg(NO_3)_2$.

智

ger"

Table 3 shows the catalytic activities of Au/titania-A prepared with a variety of additives. The appreciable enhancement of the catalytic activity is observed not only through the addition of Mg citrate but also through Mg(NO₃)₂ addition. Other citrates bring about a slight increase in activity. Similarly to the case of the magnesia support, HCHO causes the reduction of Au³⁺ in the suspension of titania-A giving a poor catalytic activity.

Figure 5 shows the XRD patterns of four kinds of Au/titania. The amorphous titania-A is transformed into anatase by calcination, and Mg^{2+} seems to suppress this crystallization. The catalytic activity tends to become low with an increase in crystallity of the support.

Figure 6 shows DTA curves for the precursors of Au/titania-A before calcination. There is an exothermic peak at around 460°C in each signal. These peaks corresponds to the transformation from amorphous titania to anatase. It is clear that the addition of Mg^{2+} shifts the temperature for the crystallization toward higher temperature.

DISCUSSION

It has been demonstrated that Mg citrate plays an important role in the preparation of highly dispersed gold catalysts with $Mg(OH)_2$ and TiO_2 as supports.



Fig. 6. DTA curves for A /t/~ania-A before calcination. (a)without any reagent(blank); (b)with Na citrate; (c)with Mg(NO₃)₂, (d)with Mg citrate. (heating rate : 5°C/min in air).







However, it has appeared that the role of Mg citrate is different between TiO_2 and $Mg(OH)_2$.

Gold supported on magnesia

The pH of the suspension of magnesia is 9.6 during the preparation. At such a pH region, AuCl4⁻⁻ should be sufficiently hydrolyzed into Au(OH)3 judging from the stability of gold species in aqueous solutions⁵). Then the hydroxide of Au might be deposited on the surface of Mp'OH)2 before the additives are introduced into the suspension. Since the point of zero charge (PZC) of Mg(OH)2 appears at pH = 126), the positively charged surface is suitable for the adsorption of anions such as citrate ion, as observed in the IR spectrum.

The reduction of Au^{3+} by HCHO in the suspension made the Au particles large and lowered the catalytic activity (Tables 1 and 3), and therefore, the reducing power of citrate ion seems not to be related to the high activity. The adsorbed citrate ion is considered to act as a sticking reagent which can block the coagulation of gold species in the suspension and/or during calcination. A speculated behavior of citrate ion is illustrated in Fig. 7.

The pH of the suspension containing Na citrate was 11 and close to the PZC of $Mg(OH)_2$. Since the effective adsorption of citrate ion is not expected at such a pH region, the enhancement of the catalytic activity might not be observed in $Au/Mg(OH)_2$ prepared with Na citrate.





Cold supported on titenis

Since the pH of the suspension of titania is adjusted to 7.0, Au species to be deposited on the support might be Au(OE)3 as in the case of $Au/Mg(OH)_2$. However, since the surface of titania in the suspension was proved to be negatively charged, as was reasonable from the PZC of TiO_2 at pH = 4 -66), cations such as Mg^{2+} should be more easily adsorbed on titania than citrate anion.

Figures 4 and 5 show that the addition of Mg citrate or $Mg(NO_3)_2$ enhances the catalytic activity of Au/titania-A and suppress the crystal growth of anatase. On the other hand, when anatase support, titania-B, was used as a starting support, there was observed no appreciable effect of Mg citrate addition. The liability of the surface of the carrier will accelerate the coagulation of supported species due to the so called "earthquake effect". The presence of Mg2+ prevents amorphous titania-A from crystallization and therefore from the "earthquake effect". On the other hand, since anatase support has already crystallized structure, there is no appreciable effect of Mg citrate

addition on the dispersion of gold.

A small arount of citrate species are detectable on titania-A in Fig. 6. The citrates other than Mg salt might have also a certain effect on an increase in catalytic activity of Au/titania-A. It is probable that citrate ions in Au/titania-A play similar role to that in Au/Mg(OH)2.

Acknowledgment

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ULTRAFINE GOLD PARTICLES IMMOBILIZED WITH OXIDES OF Fe. Co. or Ni FOR THE CATALYTIC OXIDATION OF CARBON MONOXIDE *T -70°C

M. HARUTA". T. KOBAYASHI", S. IIJIMA"", and F. DELANNAY"**

* Government Industrial Research Institute of Oaaka Midorigaoka 1, Ikeda, 563 Japan

** Research Development Corporation of Japan, c/o Department of Physics, Yagoto-urayama, Tenpaku, Negoya 468, Japan.

*** Department of Material Science. Université Catholique de Louvein, Place Sainte Barbe, B-1348, LLN, Belgium

ABSTRACT

Gold particles smaller than 10 nm in diameter, immobilized with a-Fe203, Co304, or NiO, were prepared by coprecipitation from an aqueous solution of HAuCl4 and the nitrate of Fe, Co, or N1 and by calcination of the coprecipitates in air at 300-400°C. The ultrafine gold particles were hemispherical in shape and strongly held by the host oxides. In most cases, hemispherical gold crystallites were deposited directing their flat (111) planes toward a-Fe203 (110). Co304 (111), and NiO (111) planes. I-ray photoelectron spectra showed that the gold particles with a mean diameter of 4.1 nm immobilized on a-Fe203 were more electron deficient than evaporated gold particles of the same size. The ultrafine gold particles immobilized with 3d transition metal oxides were active for the oxidation of CO even at such a low temperature as -70°C.

INTRODUCTION

It has generally been understood that gold is catalytically far less active in most of the reactions than platinum metal catalysts. Since gold particles were supported with a size larger than 10 nm and mostly ranging from 50 to 100 nm in the conventional gold catalystal.2), the difficulty to obtain highly dispersed gold particles might be another additional reason for the poor catalytic activities of gold catalysts. Although an attempt has recently been made to prepare highly dispersed gold catalysts by impregnation, they are thermally unstable at temperatures above 200°C and are poorly active for the oxidation of H2 and CO3).

On the other hand, our previous letter has reported that gold catalysts prepared by coprecipitation with group VIII 3d transition metal oxides exhibit extremely high activities for the oxidation of CO even at -70°C4). The present paper will deal with the characterization of the coprecipitated gold catalysts by TEM and MPS and will discuss the origin of the catalytic

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ULTRAFINE COLD PARTICLES SOR CO OXIDATION

carbonate from an aqueous solution of HAuCl4 and the nitrat. . : various transition metals. The coprecipitates were washed, vacuum dried, and calcined in air at temperatures from 80°C to 500°C. Catalytic activity measurements were carried out in a small fixed-bed reactor, with 0.20g of catelysts that had passed through 70 and 120 mesh sieves. A standard gas consisting of 1.0 volt of CO or H2 balanced with air to 1 atm. was passed through the catalyst bed at a flow rate of 66 ml/min.

The fine structures of the gold catalysts were observed using an Akashi EM-002A electron microscope operated at 120 kV and a Hitachi H-9000 electron microscope operated at 300 kV. X-ray photoelectron spectroscopic analyses were made using a Shimazu ESCA 750 under vacuum below 5x10-6 torr.

RESULTS

Gold immobilized with a-Fe203

Figure 1 shows the dependence of the oxidation efficiencies of H2 and CO on calcination temperature of Au-Fe coprecipitate containing 5 atom2 Au. As is clearly understood from the reaction temperature, the gold catalysts prepared by coprecipitation are highly active for the oxidation of both H2 and CO. In addition, they are much more active in the oxidation of CO than in the oxidation of H2. This is one of the characteristic features of coprecipitated gold catalysts because the gold powder and the conventional gold catalysts supported on innert metal oxides such as MgO, A1203, SiO2 are less active in the oxidation of CO than in the oxidation of H2. There was also observed a pronounced contrast between H2 oxidation and CO oxidation in the effect of calcination temperature on the catalytic activity. While calcination at 200°C gave a maximum catalytic activity for H2 oxidation, it did not create the catalytic activity for CO oxidation. The catalytic activity for the oxidation of CO appreciably increased only when the coprecipitate was calcined at temperatures above 300°C.



Fig. 1. Efficiencies for CO and Ha oxidation as a function of calcination temperature of Au-Fe(1/19) coprecipitate.

In order to clarify the change of the coprecipitate occurring during calcination, XPS measurements were carried out. Figure 2 shows comparison of the Au 4f5/2+7/2. Au 4d5/2. and 0 is lines for the four samples calcined at different temperatures. The most conspicuous result is the shift of the binding energy (BE) of Au 4f and 4d lines toward lower energy values with increasing calcination temperature. The magnitude of the shift between 80°C and 400°C is about 1.0eV for Au 4f7/2 and about 2 eV for Au 4d5/2. The width of the peaks also decreases with increasing calcination temperature. For Au 4f lines, this manifests itself by a decrease of the depth of the "valley" between the two lines. For the samples calcined at 80°C and 200°C, the Au 415/2 line exhibits a shoulder on the

high BE side. As the BEs for Au203 were 86.4eV and 90.1eV for Au 417/2 and 2. 2.2 . . .

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ULTRAFINE GOLD PARTICLES FOR CO OXIDATION

coprecipitate calcined at temperatures below 200°C. No significant difference of the BE value or of the shape of Fe 2p3/2 and Fel3VV lines were recorded for samples calcined at 200°C and 300°C, and for pure Feg03 calcined at 400°C.



Fig. 2. Change of XPS spectra of Au-Fe(1/19) coprecipitate with calcination temperature,

The presence of metallic gold could be detected by IRD as a broad peak of Au(111) at 20-38.2° only for samples calcined at 400°C. It was also only for samples calcined at 400°C that sharp as 1 intensified diffraction peaks corresponding to a-Fe203 were observed. How ver, the diffraction pattern for the samples calcined at 300°C showed the onset of crystallization toward o-Feg03. The above results obtained by XPS and XRD indicate that coprecipitates calcined at temperatures above 300°C consist of metallic gold particles and a-Fe203, while coprecipitates calcined at lower temperatures consist of metallic and oxidized gold, and amorphous precursor of a-F#203.

Figures 3 and 4 shows TEM photographs for Au-Fe coprecipitate (Au/Fe=1/19) calcined at 400°C. Very small gold particles are homogeneously dispersed on the surface of a-Fe203 particles. The mean diameter of the gold particles measured for 2131 particles was 4.1nm with a standard deviation of 1.4. The TEM observation both in the bright and dark fields incidated that the gold particles were not twinned or poly-crystals but single crystals. The enlarged view of the interface between ultra-fine go'd particles and hematite shows that the gold particles are not spherical 's hemispherical in shape and are strongly bound at their flat planes with the host cride. It was most commonly observed that the crystal of gold particles grew with a specific crystel direction: Au(111) plane with a lattice spacing of 7.36# was in junction with o-Fe203(110) plane having a lattice spacing of 2.52%. The difference of the lattice spacings was within 71.

The strength of interaction operating between ultrafine gold particles and hematite appeared to be very high because the reduction and oxidation treatment of Au/a-Fe203 to transform the host oxide (a-Fe203 - Fe304 - Y-Fe203) did not cause any liny change of the mean particle diameter of

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Fig.). Fine structure of Au-Fe(1/19) coprecipitate calcined at 400°C.

Fig. 4. Ultrafine gold particles immobilized with o-Fe.O..





Fig. 5. Fine structure of Au-Co(1/19) coprecipitate calcined at 400 °C.

Fig. 6. Ultra-fine gold particles ismobilized with Cos0.

Au immobilized with Co304

Figures 5 and 6 show TEM photographs for Au-Co coprecipitate (Au/Co-1/19) calcined at 400°C. Ultrafine gold particles of 6-7 nm in diameter were dispersed on Co304 crystallites of about 15 nm in diameter. Comparison with Au/a-Fe203 suggests that the size of gold particles is larger while the size of host oxide is smaller. This might be because Au-Co coprecipitate was obtained as small primary particles and during calcination the gold crystallites had difficulty in finding suitable crystal planes of CO304 to be strongly bound, thus being more or less exposed to coagulation. In Au/Co304, typically one gold particle is attached to one Co304 (111) is 4.67 Å, about double of that for Au(111) with only 1.1% deviation.

Gold immobilized with NiO

Figure 7 shows a TEM photograph of Au-Ni coprecipitate (Au/Ni=1/5) calcined at 400°C. Since nickel oxide particles were very small with a diameter of around 10 nm and were comparable in size with gold particles, this catalyst looked like a sixture of gold and NiO particles. However, the interface between the two particles also showed that gold particles had grown facing Au(11) plane to NiO(111). The lattice spacing of NiO(111) is 2.41%, slightly larger than the lattice spacing of Au(111) by 2.1%. As this epitaxial-like growth of gold on NiO has to find the specific crystal plane for stabilization during calcination, host oxide with smaller size might cause larger probability for the coagulation of gold.



Fig. 7. Fine structure of Au-N1(1/9) reprecipitive calcined at 400 °C.







IN TRAFINE COLD PARTICLES FOR CO OXIDATION

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Comparison of catalytic activities between various gold catalysts in the oxidation of CO and H2

Figure 8 shows catalytic activities for the oxidation of CO and H₂ as a function of mean particle diameter of gold. The catalytic activities are expressed by the temperature for 50% conversion and the crystallite sizes of gold were determined by TEM and/or peak-half widths of XRD. The mean diameter of gold exceeded 15 nm in gold catalysts prepared by impregnation and reduction. On the other hand, the mean particle diameter of gold in the coprecipitated catalysts was smaller than 10 nm even though they were calcined at a temperature (400°C) comparably higher than the final heat-treatment temperatures (200°C or 300°C) for impregnated and reduced catalysts.

A general trend is that catalytic activity increases with decreasing diameter of Au crystallites in the oxidation of both CO and H₂. However, small crystallites of Au do not necessarily lead to catalytic activity at -70 °C. In the Au/Al2O3 catalysts prepared by coprecipitation, the Ti/2 value for CO oxidation is much higher than those for Au/a-Fe2O3. Au/Co3O4, and Au/NiO even though the crystallite size of Au is similar in all cases. On the other hand, the Ti/2 value for H₂ oxidation is comparable for all four catalysts. Accordingly, the oxidation of CO at low temperatures seems to require both the costrol of Au crystallite size and the selection of appropriate support oxides.



Fig. 8. Catalytic activities for CO and H_a oxidation as a function of mean particle diameter of gold.

- 1 Au/a-FerOs (Au/Fe=1/19, coprecipitation, 400°)
- Au/Al₂O₂ (Au/Al=1/19, coprect₂ vation, 400 °C)
- Au/CosO. (Au/Co = 1/19, coprecipi stion, 400 *C1
- 4: Au/NiO (Au/Ni=1/19, coprecipitation, 400 °C)
- 5: Au/a-FerOs (Au 5 wt %, impregnation, 200 °C)
- 6: Au/SiOs (Au 17 wt%, reduction by citrate, 300%)
- 7: Au/u-AliOs (As 5 wt%, impregnation, 200 °C)

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ULTRAFINE GOLD PARTICLES FOR CO OXIDATION

DISCUSSION

The characteristic features of gold catalysts immobilized through coprecipitation with a-Fe203, Co304, NiO can be summarized as follows.

- Gold particles are hemispherical in shape and stabilized with host oxides through epitaxial-like crystal growth.
- 2) The mean diameter of gold decreases with an increase in the size of host oxides, namely $Au(4, 1nm)/\alpha$ -Fe2O3<Au(6-7nm)/Co3O4<Au(7-8nm)/NiO and much smaller than the conventional gold catalysts prepared by impregnation or reduction.
- 3) The ultraline gold particles immobilized with α -Fe203. Co304 and NiO exhibit appreciably large chemical shift toward higher binding energies in XPS than the simple, separated ultrafine gold particles.

The above features strongly suggest that there should be a kind of metal-support interaction between ultra-fine gold particles and group VIII 3d transition metal oxides. In comparison with the data reported by Oberli⁶) that the chemical shift for Au particles of 2-6 nm in diameter is only 0.2eV, the chemical shift, 0.5eV, observed for the gold particles of the same size which are immobilized with α -Fe₂O₃ is appreciably large. This result provides a strong evidence for the metal-support interaction, however, it is not clear at the present stage whether the interaction is physical or chemical nature.

If the interaction is physical electron transfer to the host oxides from hemispherical-spherical gold particles, their junction with flat interface might be advantageous for the enhancement of electron transfer. Usually, the electron transfer causes little effect on the metal side of the junction because conduction electrons are abundant. However, it is possible when gold particles are very small that the change created on the particles by such an effect becomes sufficient to affect the values of the binding energy of the core electrons. According to this model, the progressive shift of the PE of gold 4f7/2+5/2 and 4d5/2 with decreasing calcination temperature would be due to the decreasing average size of Au particles.

It is also probable that Au and Fe can form intermetallic compound, or, at least, that Fe has some solubility in Au. It is reported that iron is soluble in Au upto about 10 atomZ at $300^{\circ}C-400^{\circ}C^{7}$. The formation of such compounds or "alid solution should affect the BE of Au lines. For example, literature8, indicates that the BE of Au6f7/2 is higher by about 1 eV for AlAu2 in comparison with Au metal. It may be thought that the preparation method induces such a type of alloying of Fe into Au during calcining the coprecipitate. Such an alloy formation may require relatively high temperature as $300^{\circ}C$. The abrupt increase in the catalytic activity of Au/a-Fe203 for CO oxidation at a calcination temperature of $300^{\circ}C$ might probably be ascribed to the formation of intermetallic compounds. As the junction interface between gold particles and hematite was flat and was not broken by the reduction-oxidation treatment, it could be considered that at least at the interface gold and iron forms intermetallic compounds.

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PROCEEDINGS OF THE

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OPTICAL DETECTION OF CO IN AIR THROUGH CATALYTIC CHROMISM OF METAL-OXIDE THIN FILMS

Tetsuhiko KOBAYASHI, Masatake HARUTA and Hiroshi SANO, Material Chemistry Department, Government Industrial Research Institute of Osaka, Midorigaoka 1, Ikeda, Osaka 563, Japan Bernard DELMON Groupe de Physico-Chimie Minérale et de Catalyse, Université Catholique de Louvain,

Place Croix du Sud 1, B-1348 Louvain-la-Neuve, Belgium

Abstract

Transparent thin films of transition-metal oxides (Cr. Mn. Fe. Co. Ni. or Cu-oxide) were prepared by pyrolysis of organic metal salts on glass substrates. At 250°C - 350°C, the thin films of Mn_3O_4 , Co_3O_4 and NiO showed detectable decreases of optical absorption in visible region due to the presence of CO in air, which can be regarded as <u>catalytic chromism</u>. Since the optical response of these oxides toward CO reversibly occurred with the change in the concentration of CO (0.5 - 10 vol.% in air), the catalytic chromism is applicable to the optical detection of CO.

Introduction

The optical detection of gases has been attracting a growing interest owing to the following advantages; (i) less danger against ignition of explosion. (ii) resistance to electro-magnetic noise, and (iii) operation through optical fibers without electricity. We have recently reported that combustible gases can be optically detected by a combination of an optical thermometer and the oxidation catalyst through exothermic catalytic oxidation of the gases¹.

In the present study, the absorption spectra of transition-metal oxides have been investigated in order to develop a novel optical gas sensor which can directly output the optical signal responsive to the presence of gases. The thin oxide films were adopted as a specimen because of the high transmission efficiency of light. The high ratio of "surface" to "bulk" of the thin films enable us to expect a large optical response induced by the interaction of gases with the surface.

Experimental

Transparent thin films of Cr_2O_3 , Mn_3O_4 , Fe_2O_3 , Co_3O_4 , NiO, and CuO were prepared by pyrolysis of naphthenates or 2-methylhexanoates of corresponding transition metals². The butanol solution or the toluene solution of the organic salts of metals (2.5 - 6 wt.% of metal) was deposited on the one side of glass substrate (18 x 18 x 0.1 mm) by use of a spin-coater at a rotating rate of 5000 rpm. After dried in the ambient atmosphere and temperature for 1 h, the organic salts deposited on the glass subtrate were pyrolyzed in air

for 2 h at appropriate temperatures where the pyrolysis of the organic salts could be readily completed. The temperatures confirmed by TG and DTA beforehand were 500°C for Cu-oxide and 400°C for other oxides. Thickness of the films were in a range of 300 - 700 nm, determined by SEM observation.

Absorption spectra of the oxide films were analyzed with a transmitted visible light by use of a spectrometer which has a multi-channel photodetector and a halogen lamp (150W) as a light source. In order to control the atmosphere and the temperature of the films during the optical measurements, the thin film specimen was set in a quartz cell (200 ml) with an electric heating coil and a pair of parallel flat windows for the transmission of the light. The atmospheric gases were passed through the cell at a flow-rate of 100 ml·min⁻¹.

Table 1 Change in the absorbance (△Abs) of the thin oxide films induced by 1 vol.% CO in air.

	Absorban	ce in air	Abs	by CO
Oxide	at 250'C	at 350°C	at 250°C	at 350'C
Cr203	1.4.4	0.039	4.4.4	nd
Mn304	0.107	0.108	-3.7 %	-4.6 %
Fe203		0.129	4.4.4	nd
C0304	0.170	0.174	-2.9 %	-1.7 %
NIO	0.042	0.054	-2.3 %	-21.1 %
CuO	* * *	0.248		nd







Catalytic activity of the oxide films for the oxidation of CO was measured calorimetrically by a DSC equipment in a stream of air containing 1 vol.% CO.

Results and discussion

Effects of CO on optical absorption of the thin oxide films were measured at 250°C 350°C, and and the results at the wavelength of 700 nm were summarized in Table 1. It was found that the thin films of Mn304, Co30d, and NIO showed detectable decreases in the absorbance when 1 vol.% of CO was introduced to air in the quartz cell. The optical response to CO was most remarkable on the NiO film at 350° C.

The absorption spectra of the NiO film in the visible region at 350°C are shown in Fig. 1. The decrease in the absorbance by CO was not specific to a certain wavelength but was observed in the whole visible region.

Figure 2 shows the time-response of the optical change of the NiO film to CO. Since 99.9% of the atmosphere in the cell is replaceable within 14 min after the flowing gas was changed, the optical change of the film seems to follow rapidly the change in the composition of the atmosphere.



Fig. 2 Time-response of the change in the absorbance of a NiO thin film at 350°C. a: in air, b; in 1 vol.% CO + air. Wavelength = 700 nm.





Fig. 3 Decrease in the absorbance of a NiO thin film with the concentration of CO at 350°C. Wavelength = 700 nm.

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Figure 3 shows the dependency of the absorbance on the concentration of CO. The decrease in the absorbance is proportional to the logarithm of CO concentration in the range of 0.5 to 10 vol.%. It should be noticed that the absorbance change is observed due to a small amount of CO even in the co-presence of a large excess of oxygen, and therefore, a simple reduction of the oxide film by CO can not be regarded as an origin of the phenomenon.

Similar results shown in Figs. 1 - 3 were also obtained in the cases of Co_3O_4 film³ at 250°C and Mn₃O₄ film at 350°C.

Catalytic oxidation of CO over a NiO film could be calorimetrically detected at temperatures above 250°C and the rate at 350°C was evaluated as ca. 8 x 10^{-9} mol·s⁻¹·cm⁻². This strongly suggests that the change in the absorbance of the NiO film is caused by one or a combination of the elementary steps in the catalytic CO oxidation (<u>Catalytic Chromism</u>); adsorption of CO, elimination of adsorbed oxygen species, or formation of carbonates.

The decrease in the absorbance of the film was also observed when the atmosphere was changed from air to N₂ at 350°C. According to the reported results⁴ on the TPD of oxygen over NiO, O₂⁻ species seem to be adsorbed and desorbed depending upon the concentration of O₂ at around 300°C. Since NiO shows p-type semiconductivity⁵, the adsorption of O₂⁻ produces the positive holes, p⁺, in the valence band. Catalytic CO oxidation reduces the concentration of adsorbed O₂⁻ and hence that of p⁺.

 $O_2 \longrightarrow O_2^- ad + P^+ vb$ (1)

 $2 \operatorname{CO} + \operatorname{O_2^*ad} + \operatorname{p^*vb} \longrightarrow 2 \operatorname{CO_2}$ (2)

If it is assumed that the electron-excitation in the valence band from the filled states toward p^* causes optical absorption of wide wavelength, the decrease in the absorbance is explainable as the decrease in the concentration of p^* due to the catalytic oxidation of CO.

The reversibility with a fast response and the concentration-dependency observed in Figs. 2 and 3 have proved that the <u>catalytic chromism</u> is applicable to an optical detection of CO contained in air.

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RESEARCH ON HLW MANAGEMENT IN GOVERNMENT INDUSTRIAL RESEARCH INSTITUTE - OSAKA

Our laboratory has been contributing to the development of solidification processes of high-level nuclear waste, HLW, from the fundamental aspects. The studies are carried out at cold and laboratory scale, in co-operation with the Power Reactor and Nuclear Fuel Development Corporation, PNC. Testing methods for the characterization of solidified products are concerned with.

Present Subjects

1. Natural analogues for leaching behavior of nuclear waste glass forms

- Chemical stability of glass forms
- Corrosion of natural glass in laboratory environments
- Evaluation technique for long term stability in waste glasses
 - (in co-operation with PNC)

Finished Subject

1. Vitrification process

(1) Phase separation of molybdates from melt

- Solubility of MoO3 in glass melt
- Change of physical properties of borosilicate glass by MoOg-phase separation

(2)Volatilization

- Volatilization of HLW containing borosilicate glass at elevated temperature
 - Effect of water vapor on the rate of volatilization from B2O3-containing glass
- Volatilization mechanism of cesium from molten borosilicate glass (3)Corrosion of materials for glass melting furnace
 - Corrosion of heat-resisting alloys by melts contained HLW at high temperatures
 - Corrosion test of various refractories for glass melting furnace
 - Corrosion mechanism of INCONEL electrode by glass melts

(4)Measurement of physical properties of HLW-containing borosilicate glasses at elevated temperatures

- Viscosity
- Electrical conductivity

2.Improvement of melter elements

(1)Development of long-life refractory materials for ceramic melter

- Synthesis of ZnCr2O4 spinel by sintering and the corrosion test
- (2)Developmer. of long-life electrode materials for ceramic melter
 - Synthesis of Cr-metal and ZnCr2O4 spinel composites by hot-pressing and the corrosion test

3. Characterization of solidified products

(1)Thermal conductivity

- Measurement of thermal conductivity of borosilicate glasses by laserflash method
- Thermal conductivity of mixed alkali glasses
- Thermal conductivity of glass-copper composites
- Calculation of thermal conductivity in glass-copper system by sphere packing models



(2)Leachability

- Chemical durability of HLW containing borosilicate glasses

- Effect of surface roughness of glass on the leachability

- Effect of pH of leachants on the leachability

- Leach models of alkali from HLW containing borosilicate glasses (3)Crystallization

- Effects of crystallization on thermal properties and chemical durabil-

ity of HLW containing glasses

(4)Characterization of glassy solid form

- Depth-profiling of leached glass surface by ESCA

- Characterization of the simulated waste glass produced in France (COGEMA); Collaboration work with CRIEPI,

Density, thermal expansion coefficient, transition temperature, thermal conductivity, elastic constants, leachability.

4. Alternative solidification processes

(I)Alcoholated gelation of HLW and melting by micro-wave furnace - Preparation of alcoholated gel from HLW

- Drying, calcination and melting of alcoholated gel by micro-wave irradiation

(2)Pressure sintering process

- Pressure sintering of simulated HLW with glass powders

- Pressure sintering of copper-glass composites

- Development of continuous pressure sintering process with HLWcontaining glass and copper metal powders

(3)Normal sintering of HLW powders

- Normal sintering of HLW powders with porous high-silica glass powders

5. Other

(1)Possibility of application of solid form containing HLW as heat generator

TEC. NOLOGIES DISCUSSED WITH MITTE

- Rapid Measurement Technique of Pollutants in Groundwater
- Treatment of Wastewater Chemicals by Super Critical Fluid
- Enhancement of Microbial Activity in Membrane Bioreactor
- Evaluation of Organic Membrane Materials for Bioreactor
- Microhabitat of Microbes Degrading Hazardous Compounds in Activated Sludge Floc
- Decomposition of Organic Compounds by Oxidation Methods
- Oxidation of Organic Compounds with Biomimetic Catalysts
- Trace Metal Species in Inland Sea Bottom Water
- Interferometer Type Sensors Using Optical Fibers
- New Waste Water Treatment System Using High-Concentrated Bioreactor and Separation membrane, for Water Reuse and Energy Recovery
- Remote Fiber Sensing of Some Organic Contaminants in Water by Laser Spectrometry

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"Budget, Staff and Scale Information", MITI, 7 pages.

National Research and Development Program

AQUA RENAISSANCE '90 PROJECT

R&D on new wastewater treatment system, using high-concentrated bioreactor and separation membrane, for water reuse and energy recovery







Aqua Renaissance '90 Project

To cope with short supply of water in the near future and water pollution, industrial wastewater, sewage, and other aqueous effluents must be processed to get new water resource. But the prevailing water treatment process require large treatment facility sites or large land as sludge-disposal dumps and much energy consumption.

Japan depends heavily on imported petroluem for its energy supply and its energy supply structure is fragile compared with other advanced countries. For this reason, it is necessary to establish novel ways of obtaining new, inexpensive energy in order to make the nation less dependent on petroleum.

The Ministry of International Trade and Industry (MITI) has been conducting research and development on new wastewater treatment systems permitting water reuse and energy recovery under the name of "the Aqua Renaissance '90 Project" This undertaking is part of the National Research and Development Program "large-scale project". The total budget of the project is about 12 billion yen during six years (1985 – 1990).

Development on biotechnology and separation-membrane technology in the new water treatment system.

The principal research and development items are the following

1. R & D on Microorganism	(Selection and activation of valuable microorganisms)
2. R & D on Membrane	(R & D on efficient membrane for separation which is resistant to the deterio- rating effects of sewage and microorganisms)
3. R & D on Final Treatment Reactor	(R & D on bioreactor for denitrification, and R & D on production of oil from the sludge)
4. R & D on Membrane Module	(R & D on compact membrane module which can effectively separate the micro- organisms and organic materials)
5. R & D on Bioreactor	(R & D on high-efficiency bioreactor for methane production)
6. R & D on Control System	(R & D on optimum control and sensor system consist of instrumentation for monitoring flow rates, methane fermentation, etc.)
7. R & D on New Watertreatment System	(Completion of new system for water reproduction and energy recovery)

Research & Development Schedule

Fiscal year	1985	1986	1987	19	88		1989	1990	
Items of R & D	and the second sec				-		5	CONSISTER OF	-
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R&D on membrane		Basic research	Fundamental exper	iments			the second second		
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R&D on control system		Enduar		Detail	ation		The state		
R&D new watertreatment system	Overall planning -	related technology	Schematic design	and constr	Uct	ion	Pilot plant o	* • - • • • • • • • • • • • • • • • • • •	
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Epoch-making water treatment and methane production technology.

Advantage of the new wastewater treatment

- 1) compact facility
- 2) large reduction in the amount of sludge generated
- 3) little energy cosumption
- 4) easy to control and maintain

Comparison between prevailing water treatments and the Aqua Renaissance '90 Project



The Aqua Renaissance '90 Project will realize the followings





Extensive Spin-off of Aqua Renaissance '90 Project



Members of Association

5

1

Ishikawajima Harima Heavy Industries Co., Ltd. Ebara Corporation Japan Organo Co., Ltd. Kawasaki Heavy Industries, Ltd. Kubota, Ltd. Kurita Water Industries Ltd. Kobe Steel, Ltd. Sanki Engineering Co., Ltd. Shimizu Construction Co., Ltd. Water Re-use Promotion Center Chivoda Chemical Engineering & Construction Co., Ltd. Director of DIC Degremont Co., Ltd. General Affairs Toshiba Corporation Toto Ltd.

Organization chart of the associaton



Nishihara Environmental Sanitation Research Corporation Nitto Electric Industrial Co., Ltd. NGK Insulators, Ltd. Nippon Petroleum Refining Co., Ltd. The Japanese Association of Industrial Fermentation Hitachi Plant Engineering & Construction Co., Ltd. Mitsubishi Electric Corporation

Mitsubishi Rayon Engineering Co., Ltd.

Aqua Renaissance Research Association

Aqua Renaissance Research Association

Address:

Toranomon Takagi Bidg. 2F 7-2, Nishi-Shimbashi 1-Chome Minato-Ku, Tokyo 105 JAPAN

Bidg. 2F Tel : (03) 503-2131 hi 1-Chome 105

0-2131 Fax: (03) 503-2139

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公害資源研究所 平成元年度 予算, 人員及び規模 Budget. Staff and Scale, FY1989

李 算 (算位百万円) Budget Last Million Yen



Scale

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数地面積	262.497m	10.197m	20,7691	80,268m°
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Call More Labor, Sciences

平成元年度研究課題

List of Research Project in FY1989

特别研究

安全·保安技術

「創田保安技術に関する研究(火火対策と過ぎシステム) 「液化石油ガス保安対策技術に関する研究 「不連続性岩盤の安全性評価に関する研究

豊康・エネルギー技術

○天然ガスからの液体燃料の製造に関する研究
○注素鉱物からの高付加価値シリコン系素材の製造に関する研究
○湿式法によるレアメタルの濃縮積製技術に関する研究
○応崗岩の資源化技術に関する研究
○高温ビートバイブに関する研究

新材料技術

○気相熱分解法による炭素系材料の製造に関する研究 ○レアアース系ゼロ次元複合機能性材料の創製に関する研究

產業基盤確立技術

○超幾粉砕技術に関する研究 ○レーザーによる炭高複合クラスターの創製に関する研究

戴山保安技術研究

二坑内大災衛火技術に関する研究 二醇電気着大防止技術に関する研究

官民瀛寨共陶研究

○地下空間におけるAE、地圧、ご盤特性の積密評価技術に関する 研究

题子力平和利用技新

こ放射性魔魔物感分空洞の岩盤力学特性に関する研究

公審防止技術

- ○不活性化学物質の不助一系光反応による変換、分解に関する研究 ②物理化学的方法による窒素、リンの高度処理技術に関する研究
- ○正場騒音の複数伝搬モデルに関する研究
- ○活性汚泥の生態学的創催による有害化学物質の高効率処理手法に 関する研究
- ○工業都市域における浮遊粒子状物質の大気環境激度子欄手法に関する研究
- ○化学工場周辺の場然における製処理性合成有機化合物の差動に関する研究
- ○地下水汚染防止に係る水中汚濁成分の迅速分析技術に関する研究 ○海外炭の利用拡大に対応する大気汚染防止技術に関する研究
- ○ディーゼル重量素NO(低減対策に関する研究
- (酸性物質生成に係る環境中反応性化学種の高密度簡易要定に関す も研究
- 回録海工場地先の停滞海域における貧酸素水塊構造の解明に関する 初立

Special Research & Development

Mine and Industrial Safety

- Mine Safety Technology (Fire prevention and emergency escape system)
- Salety technology on the Liquefied Petroleum Gas
- Safety in Discontinuous Rocks

Resources and Energy Technology

- Production of Liquid Fuel from Natural Gas
- Production of Silicon Materials from Low-grade SiO: Ores by Chlorination
- Refining of Rare Metals
- Utilization of Granite for industrial Use
- High Temperature Heal Pipes

New Materials

- Formation of Carbon Materials by CVD
- C Formation of Functional Suspension Colloids of Ultrafine Powder Containing Rare-Earth Compounds

Basic Technology for Future Industry

- OUltra-fine Grinding Process
- Production of carbon Clusters and Carbon Metal Complexes Using Laser

Coal Mine Salety

OMine Fire Extinguish Techniques OCountermeasures against Electrostatic Charges

Gavemment & Private Organization Joint Research

Precise Measurement of AE, Earth Pressure and Deformation of Rockmass around the Cavern

Nuclear Research

Rock Mechanics about Storage Caverns for Radioactive Wastes

Pollution Control

- Transformation and Decomposition of Less Active Chemical Substances in the Troposphere
- Physicol Chemical Wastewater Treatment for Nitrogen and Phosphorus Removal
- Complex Propagation Model for Factory Noise
- Ecological Control of Activated Sludge Process for Hazardous Compounds
- Simulation Models of Suspended Particulate Concentration from industrial Emission
- © Behavior of Synthetic Organic Compounds in Coastal Environment / Rapid Measurement Technique of Pollutants in Groundwater
- Preparation Technology of Ultra Clean and Combustion Control Technique for Low Volatile Coal
- CReduction of NO, emitted from Heavy-Duty Diesel-Powered Vehicles
- Measurements of Various Reactive Species Related to Environmental Acidification by Developing Sensitive Analytical Methods

Anoxic Water Mass Formed in the Stagnant Waters of Inland Sea

- ニキノール発酵域,長の進化管理技術に関する研究。
- 逐漸環境の長期的変遷過程の解明による最適で地の予測技術に関 する研究
- ご治院海域における深遊生物組成変化の予測と日朝計測技術に関す る研究
- (有代有機化)物況に反産業廃棄物の処理技術に関する研究 、先属技術産業に任る環境方面物資の応数方面とモニタリング手法 の開発に関する研究
-)挿発性有機、ウケン化つ物の人気数比別面技術に関する研究 人気中則子との相関を考慮した発生液ダストの評価と測定方法に 物件を研究
- 「超臨県流体による将水中の化学物質の処理技術に関する研究 、群小発生薬からのNO、低減化のための触線機械技術に関する研究
- 「フロン代替物質の対流圏内における分解過程に関する研究

- Liquid Fuel Production from Ethanol Fermentation Stillage by Thermochemical Conversion
 - Long Range Prediction Model for the Change of Shallow Water Environment for Optimum Industrial Development

Structure of Lower Trophic Ecosystem in an Europhicated Bay-Automated Measurement and Prediction of Plankton Organisus Succession

- . Treatment of Industrial Wastes Containing Mixed Hazardous Organic Compounds
- Diffusion Processes and Monitoring Method of Pollutants Related to High Technology Industries
- Emission Control of Volatile Organic Halides by Adsorption
- . Evaluation and Measuring Method for Source Dust in Consideration with SPM
- K Treatment of Wastewater Chemicals by Super Critical Fluid
 - Catalytic Combustion Technique for Reduction of NO, from Small Scale Stationary Sources
 - . Removal Processes of Alternatives to the Legislated Chlorofluorocarbons in the Troposphere

她球環境

ご確業活動起源の温暖化関連物質の単動解明に関する研究

國聯產重技術研究事業

○中国レアメタルの分離積裂技術に関する研究 (執帯勉強における大気汚染でセスメント手法に関する研究)

Global Environment

Environmental Behavior of Green House Gases of Industrial Origin

- International Research Cooperation
- Separation and Refining of Chinese Rare Metal Ores
- Air Pollution Assessment in a Tropical Area. India

公審防止事前調查	Assessment Technology for Industries
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②大気汚染シミュレーションモデル開発 ②水質汚染予測手法開発

Assessment Models of Air Poliution

Assessment Models of Water Pollution in Coastal Area

指定研究

Designated Research & Development

-----大型工業技術研究開発 (大型プロジェクト) ------

マンガン団境探転システムの研究開発

ーマンカン団場の採用性に関する研究 -マンカン日場の採動特性に関する研究 -マンカン日場の採動に伴う環境保全に関する研究

水総合再生利用システムの研究開発

大深度地下空間開発技術に関する研究

酒精度地下構造評価技術 人深度地下空間構築技術 大変度地下空間環境制度,防火技術。

-----新エネルギー技術研究開発(サンシャイン計画)---

石炭エネルギー

「石炭ガス化の裏効率化に関する研究」

Large-Scale Project

Manganese Nodule Mining System

Manganese Nodule Collecting Technology

Hydraulic Lifting Technology of Deep Sea Manganese Nodules Marine Environmental Assessment Studies for Ocean Mining of Manganese Nodules

New Water Treatment System

Enhancement of Microbial Activity in Membrane Bioreactor & Evaluation of Organic Membrane Materials for Bioreactor

Underground Space Development Technology

- Geological Survey and Evaluation Technology
- Dome Construction Technology
- Environment Conditioning and Hazard Preventing Technology

----- New Energy Development Project

Coal Energy

improvement of Gasification Efficiency of Coal

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増勝工ネルギー 二高量的体の統紛料借出技術に関する研究 二総無共御前技術等に関する研究 高量的体料抽出システムの解析評価

新工業ルギー技術の基礎的研究開発 木質系パイオマスから液体性料の製造に関する研究

先編的基礎的省エネルギー技術 単使技術の高度化に関する研究

(産業起源の二酸化炭素数出低減技術分析モデルの構築) 二酸化炭素還元反応の研究

Geothermal Energy

- Fracturing and heat Extraction Technology of Hot Dry Rock
- Geothermal Well Drilling Technology
- Data Analysis and Evaluation on the Heat Extraction System of Hot Dry Rock

Basic Study on New Energy Developmont Liquid Fuel Production from Woody Biomass by Thermochemical Conversion

-----Energy Saving Project

Leading and Basic Technology for Energy Conservation Advanced Combustion Technology

Global Environment Project

- System Studies for Reduction of Industrial Carbon Dioxide Emission
- Research for Reduction of Carbon Dioxide

科学技術庁重点基礎研究

二酸化炭素の再資源化技術の基礎的研究 二地下環境制備技術の基礎的研究 二光機能時性を有する含塑素化合物の基礎的研究 二基盤的研究分野における知識ペース構築手法の基礎的研究

経常研究

石炭鬱

②石炭組織成分の分光学的手法による構造研究 ②石炭の夜化反応機構の研究 ②炭素一鉱物系多孔質への微生物固定化の研究 ③熱プラズマ法によるダイヤモンド合成の基礎研究 ③高分子炭化物の吸着分離能向上の研究 ③高分子炭化物の吸着分離能向上の研究 ③芳香族系ポリマーからの新炭素体の合成

燃料資源部

出现技術部

シーキー料果」「の基礎研究 た助起ビロール諸体の料性及び利用の研究 育機スラーン油化の基礎的研究 医能化水素の報分離技術の研究 重要算分解油から総合多環方香族分離技術の研究 全値化合物を含有するカーホンブラックの研究 シュール油の高度利用のための基礎的研究 ディタレン類及びナフトール類の酸化反応の研究

Accelerated Basic Research

- Preparation of Chemicals from Carbon Dioxide
- Control Technology of the Underground Space
- Synthesis of Nitrogen Compounds with Photoinduced Functions
- Knowledge Base Systems in the Fields of Pollution and Resources

Ordinary Research

Coal and Carbon Department

- Structural Study on Petrographic Components of Coal by Spectroscopic Methods
- Reaction Mechanism of Coal Liquefaction
- Porous Materials from Coal for Microbes
- Synthesis of Diamond using a Plasma Jet
- Improvements of Adsorptive Ability of Carbonized Products
- Novel Carbon Materials from Aromatic Polymers

Fuel Department

- Fundamental Interactions between Hydrocarbon Molecules and Laser Photons
- Characterization and Utilization of Aromatic Hydrocarbons-Pyrroles Exciplex
- Thermochemical Treatment of Urganic Sludges
- Membrane Separation for Hydrocarbons
- Separation for Aromatic Hyrl ocarbons from Heavy Cracked Oils
- Carbonblack Intercalated with Metal Compounds
- Distinctive Utilization Method of Shale Oil
- Oxidation of Naphthalenes and Naphthois

Combustion Engineering Department

- Combustion Diagnostics by CARS
- Combustion Technology by Oxygen Rich Air
- Structure of Particle-Laden Turbulent Jet
- Diamond-Like-Carbon Formation during Compustion



アルコール無機の研究 無量測定技術の研究 臨界域の輸送現象の研究 (高約荷水イラーの研究)

お新業源語

- ●酸化鉱力進行起業の基礎的研究 希主類元素の分析法の研究 日実衛星度分の運到分離の研究 レマイキを範疇の選組法の高度化の研究 高純度セラミークス原料形束の生成の研究 コンルトンラストの精錬性術の研究 マンルトンラストの精錬性術の研究 モリマギンの高純度化位者の研究 非水溶液を利用するレアメタルの精製微粉体化の研究 レアメタル多環合素材の氏局合成の研究 乙濃酸十分成における熱ブラズマの特性の基礎的研究 「新た体を填持性の基礎的研究」 「新た体を填持性の基礎的研究」 「新た体を填持性の基礎的研究」

實源開発部

- ○海底現状鉱床の開発技術の基礎研究 ○原位素地行計(例法の研究 ○連任年における対撃評価の研究 ○南後大陸鉱物資源開発の調査研究
- 「セラリークスピートによる岩石庵削の研究
- 「超高圧未續洗高度利用の基礎的研究」

燈業保安部

-] 閉鎖系空間における人人、爆発性状の研究
- 「高濃度領域料」との約一分数技術の研究
- 「地下構要料のセーフティアセスメントの研究
- 」含水体集の動加圧下の様ごう特性の研究
- 「カスカンサーの基礎的研究」
- 「不均益物体内の、変動物性値分布のリモートプルービング技術の研究
- 市に大ス設備の説明爆食等の研究
- 北内通民量計判値評価法の研究 シューアンエラー発生機構とその分析手法の研究

立戦環境部

- 二法物理通程を含む汚染質の上空に散の研究
- 二斉周連昇モデルの基礎的研究
- 三組ム後え他の米系環境内での筆動に関する基礎的研究
- 」遺法情報による海洋衛生物群業の機能の評価手法の研究
- (1) キートセンシングによる大陽光散乱粒子の広域水平匹散現象の 研究
- 、山口海流の環境調査手法の基礎的研究

大繁環境保全部

「低濃度早素酸化物の詳去機能材料開発の研究

- Combustion of Alcahola
- Precise Combustion Calorimetry and Electrical Calibration for Bomb Calorimeter
- Transport Properties near Critical Region
- Study of Direct-Contact Type Boiler

Materials Processing Department

- Selective Flocculation of Chide Ores
- Determination of Rare Earth Minerals
- Separation and Concentration of Valuable Trace Elements in Coal by Advanced Coal Cleaning Processes
- Advanced Separation for Rare Metal Minerals
- Preparation of Ceramic Powder
- Metallurgy of Cobalt-rich Ferromanganese Crusts
- Relining of Molybdenum
- Refining of Rare Metals with Non-aqueous Solutions
- Preparation of Highly Controlled Ultra-fine Particles for Rare Metal Composites
- Ceramic Processing for Siliceous Materials
- Diagnostics and Modeling of Therma! Plasmas in Connection with Ultra-fine Particles Formation
- Improvement of Fine Grinding Using Non-aqueous Solutions
- Packing Characteristics of Powders
- Characterization and Utilization of Fine Particulate Slurry

Mining and Geotechnology Department

- Massive Mineral Deposits on the Ocean Floor
- Measurements of In-situ Earth Stress
- Rockmass Behaviors under Heavy Earth Pressure
- Research for the Development of Antarctic Mineral Resources
- Rock Drilling by Ceramic Bit
- Fundamental Studies on the Application of Ultra High Pressure Water Jet

Industrial Safety Department

- Characteristics of Confined Fires and Explosions
- Homogeneous Dust Dispersion Technique at Higher Dust Cloud Concentration
- Safety Assessment of Underground Construction
- Detonability of Water-gel Explosives under Dynamic Pressure
- Development of Gas Sensors
- Remotion proving of Wave Properties through Heterogeneous Bodies
- Crevice Corrosion of the High Pressure Facilities
- Measuring System of Mine Airflow
- Outbreak Mechanism and Analysis Technique of Human Erroru

Environmental Assessments Department

- Upward Diffusion Process Accompanying with Cloud Formation
- C Fundamental Model of "Aoshio" Water Upwelling
- Ecological Modelling of Engineered Bacteria in Natural Coastal Environment
- © Ecological Characterization of Marine Microbial Communities Basad or Genetic Information
- Hcrizontal Diffusion Detected by Remote Sensing Techniques
- New Approach Methods of Coastal Environmental Study

Atmospheric Environment Protection Department

New Materials Removing Low Concentration Nitrogen Oaides from Air



○環境中粒子状物質の反応活性発現機構の研究 シマロン、トリタレン等の脱ハロゲン化反応の研究 宇宙豪廉開発利用に係る要素技術の調査研究 ホスト・ゲスト相互作用の応用による有害化学物質の補促の研究 作業用ディーゼル機関から排出される準確低減の研究 超微粒子の補業性能評価の研究 粒子は物質への土壤成分高与についての研究 有機内染物質の多年の、十夕解析力法の研究

水讚遺保全部

- ○化学分類手法による混合像生物集団解析法の基礎的研究 ○活性汚泥の微細構造と後化活性の研究
- 「相気凝集法の研究
- ご各種種化法の組合せによる有機物の分解の研究
- 二生体間運触媒による有機化合物の酸化の研究
- 一種谷化分析技術による水中微量汚濁成分の分析法の研究
- 2.治岸城風層水中の金属の単動の解明

北海道石屋鉱山技術試験センター

□ 坑内作業用移動機構の基礎的研究 □本質安全化超音波監視センサーの基礎的研究

○光ファイバー下渉型センサーの基礎的研究 ○連続容量回路の本質安全防爆化の基礎的研究

九州石炭鉱山技術試験センター

○ガス度動記数の研究 ○プロバレー空気混合気の点人限界の 所充 三樽重検定試験法の小型化に関する研究

- Development Mechanisms of Catalytic Action of Particulate Mattars in the Environment
- Catalytic Dehalogenation of Halocarbons
- Exploitation of Extraterrestrial Resources
- Capture of Hazardous Chemicals Applying Host-Guest Interaction
 Reduction of Smoke Emitted from Diesel-Powered Vehicles in Working Environment
- Estimation of Collection Efficiency for Ultra-fine Particles
- Estimation of Soll Source Contribution to Metal Components in Suspended Particulate Matters
- Multi-compone. a Analysis for Organic Vapor Pollutants

Water Pollution Control Department

- Characterization of Microbial Community Structures by the Cellular Matericia
- Microhabitat of Microbes Degrading Hazardous Compounds in Activated Sludge Floc
- Magnetic Coaguiation in Effluent and Water Treatment
- Decomposition of Organic Compounds by Oxidation Methods
- A Dividation of Organic Compounds with Biomimetic Catalysts
- Trace Pollutants in Water by Hybrid Analytical Metho Trace Metal Species in Inland Sea Bottom Water

Coal Mine Safety Research Center, Hokkaido

- CLocomotive Mechanism for Underground Development
- Cintrinsic Sale Littraweet Sanage for Integrated Monitoring System
- Cinterlerometer Type Sensors using Optical Fibers
 - Cintrinsic Salety for Continuation Circuits with Capacity

Coal Mine Safety Research Center, Kyushu

- Fluidity and Diffusion of Methane Gas
- Ignition Limits of Propane-Air Mixture
- Miniat rization of Testing Methods for Permissible Explosives



TECHNOLOGIES DISCUSSED WITH AIST:

- Superconductivity
- High Performance Ceramics
- Synthetic Membranes for New Separation Technology
- Synthetic Metals
- High Performance Plastics
- High Performance Materials for Severe Environments
- Photoactive Materials
- Non-linear Photonics Materials
- Biotechnology (Utilization of Recombinant DNA)
- Molecular Assemblies for Functional Protein Systems
- New Electron Devices (Superlattices Devices)
- Three Dimensional ICs
- Bio-Electronic Devices
- New Models for Software Architecture
- Manganese Module Mining System
- New Water Treatment System
- Interoperable Database System
- Advanced Materials Processing & Machining System
- Fine Chemicals from Marine Organisms
- Super/Hyper-Sonic Transport & Propulsion System
- Underground Space Development Technology
- Advanced Chemical Processing Technology
- Human Sensory Measurement Application Technology
- High Performance and Low Cost Solar-Photovoltaic Conv
- Solar-Thermal Applications Systems for Industrial Process
- Coal Liquidification Technology
- Coal-Based Hydrogen Production Technology
- Integrated Coal Gasification Combined Cycle Power Gene
- Geothermal Energy
- Technologies on Hydrogen Production, Storage, Transpor
- New Energy technologies (Wind, Ocean, Bio)
- High-efficiency Membrane Complex Methane Production
- Advanced Battery Electric Power Storage System
- Fuel Cell Power Concentration Technology
- Super Heat Pump
- Superconducting Technology for Electric Power Apparate
- Ceramic Gas Turbine Project
- Synthetic Technology of Artificial Clay for HP Ceramics
- Re-utilization System Technology of Composite Materials

BIBLIOGRAPHY OF LITERATURE RECEIVED FRO



LABORATORIES AND INSTITUTES

Technology is a repository of great hope in today's world. At the research laboratories of AIST, work is carried on in developing the leading and basic technologies that will form the groundwork for future technological innovations.

R&D

Research carried out at AIST laboratories and institutes includes the following characteristics.

- Research and development of leading technologies to form a base for future technological innovation.
- As national institutes, AIST facilities conduct research needed for the propagation of technical standards required for government administration, the establishment, maintenance and supply of standards, and the creation of sophisticated experimental methods.
- Research addressing Social needs in earthquake prediction and environmental protection.
- Government support makes possible fundamental and comprehensive experimental research which would be beyond the resources of the private sector.

New R&D projects aim at finding solutions to energy shortages. Use depletion of the world's natural resources, environmental pollution and other pressing problems.

Research projects are classified into two broad categories: ordinary fundamental research, and special research. Research institutes under AIST have over 600 ordinary research themes and more than 150 special ones. These are further classified into 17 fields, such as electronics, earthquake prediction and biotechnology.

Besides these, the designated research is executed. This project research is aimed at industrialization, and includes the national R&D large-scale project. R&D project on basic technology for future industries. R&D on new energy technology called the "Sunshine Project" and R&D on energy conservation technology called the "Moonlight Project". AIST institutes are taking charges of fundamental fields represented by the above mentioned project.

THE TSUKUBA RESEARCH CENTER

In fiscal year 1979, nine iesearch laboratories under AIST moved to Tsukuba Academic City to form the Research Center of the Agency of Industrial Science and Technology. Having previously beet, scattered over the Tokyo metropolitan area, consolidation of these nine institutions - National Research Laboratory of Metrology, Mechanical Engineering Laboratory, National Chemical Laboratory for bidustry (formerly the Industrial Laboratory of Tokyo), Fermentation Research Institute, Research Institute for Polymers and Textiles, Geological Survey of Japan, Electrotechnical Laboratory, Industrial Products Research Institute, and National Research Institute for Pollution and Resources, helped the Center forge closer relations among AIST institutions and supported the efficient development of advanced research acuvities.

1. Project for Expanding Infrastructure for Research and Information

To support research and development and permit more effective use of research and technological information and more advanced computerization and processing. AIST is developing system for promoting laboratory automation, constructing and expanding data bases on research and technology, and expanding networks, aided by the Research Information Processing System (RIPS), installed for joint use by AIST institutions at the time of their move to Tsukuba. 2. Project for Promoting Research Cooperation

AIST is taking a variety of steps to promote interaction between AIST institutes in Tsukuba, while stepping up private/public international technical exchanges and more effective use of research and technical information. This includes holding comprehensive symposiums and other forums at Tsukuba, accepting researchers from foreign countries and receiving technical trainees from local public entities and other organizations. These strangements are aimed at strengthening research projects and encouraging studies in Japan.

3. Activities of RIPS (Research Information Processing System) At the end of fiscal 1987 a large-scale, general purpose computer system (FACOM M-780/20) and a super computer system (CRAY supercomputer XMP/216 and its IBM front-end processor 3090/18E) were installed as the third stage to meet increasing demands for high-speed calculations and large-scale memory capability. Furthermore, software for structural analysis, image processing, models of simulation and scientific calculations were installed in this system. In addition to the present one, a high-speed charmel (EATHERNET) enhanced the network among the laboratories. Now, RIPS is aimed at supporting the advanced, efficient research activities demanded by the AIST laboratories.



NTRODUCTION TO INDIVIDUAL LABORATORIES AND NSTITUTES

The National Research Laboratory of Metrology (NRLM) is e national representative institute for standards of length, time, ass, temperature and related quantities in Japan, and takes the lead unifying units of various physical and engineering quantities and nproving standards for science and technology. The research orks cover broad fields for the development and the improvement I standerds. The NRLM is responsible for establishing of working andards and calibration of measuring instruments in compliance ith the Measurement Law. The technical consultancies are also eing carried out. Another important responsibility is to promote uernational cooperation for metrological unification, in pursuance ith the Metric Convention. The NRLM keeps close contact with he International Bureau of Weights and Measures, the International ureau of Legal Metrology and the research institutes for standards a many countries. The major research projects of the institute are as ollows

Standards and Metrology (1) Basic standards of length, time, temperature and mass (2) Industrial standards of density, force, pressure, flow rate, vibration, shock acceleration, surface roughness, microparticles and viscosity.

Applied Precision Metrology (1) Precision measurement of laser frequency (2) Precision nonlinear spectroscopy (3) Precision long distance measurement (4) Nanometrology (5) High temperature thermophysical properties (6) Thermal and mechanical properties of solids (7) Thermophysical properties of fluids (8) Precision dimensional metrology (9) Measurement system and evaluation (10) Reliability of measurement apparatus (11) Measurement for high temperature superconductivity.



New Cestum Time and Frequency Standard Adopts the "Optical Pumping Technique"

and the second se	A DESCRIPTION OF A DESC	and a second which is not if the post of the second s	the little starting of a starting of the start
Hational Research Laboratory of Methology	Tsukubs Gakuen 0298 (54) 4118	Total personnel	218
1-4. Umezono 1-chome, Tsukuba-shi, Ibaraki, 305	Senior Officer for Research Planning	Total budget	2,243 (million yen)

The Mechanical Engineering Laboratory (MEL) was stablished in 1937 with the objective of promoting advancement of he Japan's machine industry. Today, still maintaining its traditional ole, MEL has changed and expanded its role for the development if new engineering technologies through association of mechanical ingineering with other technical fields, development of traditional echnologies in mechanical engineering towards their limits, and nitellectualization of machines and systems. The major R&D fields are shown below:

- Basic Researches in Mechanical Engineering (1) Optics. Instrumentation and control. (2) Micro machines. (3) Tribology. (4) Control of noise and vibration.
- Materials and Manufacturing Engineering (1) Synthesis and evaluation of novel materials for machines. (2) Advarced metal forming and high-precision machining/grinding. (3) Energy processing and binding. (4) Intelligent manufacturing system.
- Energy Saving and New Esergies (1) Wind power conversion system. (2) Advanced combustion engines. (3) Highperformance heat transfer/exchange. (4) Environment control.
- Robotics and Intellectualization (1) Locomotion and manipulation. (2) Motion control. (3) Computer-aided analysis and design for manufacturing
- Bioengineering (1) Engineering for medical diagnosis. (2) Biocompauble materials. (3) Elucidation of biological functions.

Micro Gripper Prototype-I for Miniature Operation

provide the state of the state	a second design of the second s	E	27	334
Mechanical Engineering Laboratory	Tel 81-298-54-2521	Fax. 81-298-04-2010	Total personnel	#70
1-2. Namiki, Tsukuba shi, Ibaraki, 305	(Research Planning Office)		Total budget	3,320 (million yen)


DIFFUSION OF TECHNOLOGICAL ACCOMPLY MENTS

The Agency of Industrial Science and Technology registers as industrial properties both at home and abroad the technological developments of its 16 research laboratories and several projects under outside contract, and works to ensure their effective utilization and diffusion.

Patents and other industrial properties (collectively referred to as "patents")* within the jurisdiction of AIST can be licensed to both domestic and ferign companies under certain conditions They are (1) a license fee is paid. (2) the licensee is capable of using the patents and (3) the licenses is non-exclusive.

Since October, 1985, NEDO conducts management and propagation of accomplishment of development contacted to NEDO.

Industrial properties under AIST's jurisdiction (registered or pending as of March 31, 1990)

	Domestic	Foreigt
Laboratories	7,702	1,70
Commissioned research and development	6.236	42
Total	15,938	2,12
	Laboratories Commissioned research and developmens Total	Domestic Laboratories 7,702 Commissioned research and development 6,236 Total 15,938

- Patents under AIST's Justich, now and "Weit Loopening. The present status of patents in the table below. The Agency is in charge of about 15,900 patents in Japan and about 2,100 abroad Of the total, 762 patents are licensed to private and semiprivate enterprises. The revenue from licensed patents totaled \$330 million in fiscal 1989
- System of Disseminizing Technological Accomplishments Permission to use patents under the jurisdiction of AIST, with exception of some patents, is granted to abroad segment of Japanese and foreign businesses by the Japan Industrial Technology Association (JITA).

JITA is a nonprofit foundation intended primarily to diffuse the technological achievements of ALST. The Association offers (1) mediation by specialized consultant engineer. (2) conclusion and mediation of state-owned statemis for conclusion of license agreement and agreement management, and (3) briefings and publishings of information on state-owned patents likely to be exercised in the near future, in order to ensure effective dissemination of state-owned patents.

In a tie-up arrangement with the Research Development Corporation of Japan, JITA calls on it to promote application of unused state-owned patents while presenting mutual characteristics.



INDUSTRIAL TECHNOLOGY COUNCIL

1. Overview

The Industrial Technology Council was established on July 25, 1973, as an affiliated institution of the Ministry of International Trade and Industry. ITC officials investigate and deliberate on important matters related to scientific technology in the mining and manufacturing industries in response to inquines from the Minister of International Trade and Industry.

Conditions affecting Japanese technological development have changed in recent years. The time has come for Japan to develop original technologies in a way that can give full play to national ingenuity and creativity. Moreover, Japan is pinning high hopes on technological development as a means of enhancing the quality of national life, upgrading the domestic industrial structure and contributing to international society.

Under this situation. ITC is working on a broad range of issues related to technological development from a standpoint of MITI as a whole.

2 Activities (Recommendations and Reports Since FY 1982)

* Industrial technology development policies

(Report of the Planning Subcommittee, Coordinate Committee, November 27, 1984)

- · Future system of the Second Round
- (Report of the Planning Subcommittee, Future Technology Development Committee, June 24, 1988)
- Industrial Science and Technology Policies for 1990s (Report of the Technology Innovation Subcommittee for 1990s, Coordination Committee, May 11, 1990)
- New Evolution of the Sunshine Project (Interum Report of the New Energy Technology Development Committee, June 15, 1990)

Organization Chart of the Industrial Technology Council (as of August 30, 1990)





RECENT TRENDS INVOLVING AIST

The situation surrounding the science and technology is changing substantially. For example, the so-called approaching/ tesonance phenomenon of the science and technology is going on and the necessity of harmony between the science/technology and nature/society is increasing. Japan is now expected to contribute positively to the international society in terms of science and technology. In this background, it is essential to promote technology. In balance established between the science and technology. In particular, the global point view as well as to proceed with R & D with balance established between the science and technology. In particular, the global environmental problems are common subjects of human being, and their solution requires technological break-through. The "Vision for 1990s" recently publicized by the Industrial Structure Council and Japan Industrial Technology Association as well as the Recommendation No 17 of the Science and Technology Conference stress this concept.

In the field of standardization administration, active discussions are under way concerning the future industrial technology, such as proposition made on the basic concept of long-term plan for the future J15 policy.

Vision for 1990s

The Technological Policy for 1990s Subcommittee was erected under the Coordination Committee of the Industrial Technology Council in the course of review on "Trade and Industry Policies for 1990s" of MITT. The industrial technology policies in 1990s were studied in its joint meeting with the Industrial Technology Policy Subcommittee of the 1990s Policy Committee of the Industrial Structure Conference, and the report publicized in May 16, 1990.

The science and technology is one of development infrastructures of a country and the world and at the same time a powerful means to solve subjects common to all human beings. In this recognition, the importance or expectation on the science and technology is growing rapidly. In this situation, Japan is requested to make international contribution worthy of its power in the field of science and technology. On the other hand, the present state of science and technology of Japan indicates relative lagging in the field c. fundamental researches and heavy shortage of talents for research and development. Moreover, this report sets forth following four vital columns of industrial technology policies for 1990 in view of changes in tides surrounding of the science and technology, such as progress of approach and resonance phenomenon between science and technology, harmony between science/lechnology and nature/society, or necessity of achieving the comfortable and affluent national living. These columns are promotion of techno-globalism from a global point of view. promotion of R & D with balance established between science and technology, promotion of R & D to realize the comfortable and affluent national life, and improvement and expansion of infrastructures for development of the science and technology.

Recommendation and report of the Science and Technology Conference

(1) Report for Inquiry No.7

"R and D basic plan on environmental protection technology" Concerning global environmental protection technology, the guideline of science and technology policies determined by the cabinet in March. 1986 set forth planning of the research and development basic plan as a priority field of promotion. With an aum placed on deepening of comprehensive understanding on the earth and full exploitation of the result to contribute to prosperity of humankind, with due consideration on international contribution, the prime minister made inquiry on the R & D basic plan of global science and technology in March, 1989 to the 42nd Science and Technology Conference. In response, the discussions were made in the global environmental protection technology committee and its subcommittees and the report made in June, 1990.

(2) No.18 mouiry, "Comprehensive basic policies for future"

In compliance with the guideline for science and technology policies determined in the cabinet in March, 1986, the science and technology promotion policies have been pushed forward toward creation of more affluent society and national life while centering around the creative and affluent science and technology. However, the environment surrounding the science and technology has undergone substantial change. In order for Japan to play the role as a member of the international society and to make effort for enhancement of the national life and continuous prosperity, the prime minister made inquiry on planning of comprehensive science and technology policies for coming 10 years with insight into the coming new century.





INDUSTRIAL STANDARDIZATION

1. Outline

10

y.

Promoting industrial standardization is one of AIST's most important tasks. By law, deliberations on JIS (Japanese Industrial Standards) are the responsibility of an AIST subsidiary organization, JISC (the Japanese Industrial Standards Committee).

Industrial standardization has contributed to the building of Japan's industrial infrastructure and helped rationalize production u. its industries. JIS operates through delibrations by some 9,000 experts from industry and academia, as well as consumers. As all JIS are voluntary standards, it is essential that they reflect the opinions of all concerned.

2 JIS and JIS Marking System

Like many other countries, the purpose of JIS in to promote (i) improved quality and rationalized production, (ii) smooth and fair trade, and (iii) rational consumption through appropriate and rational "standards." Some 8,400 JIS are established at the end of FY 1990.

The JIS marking system is used to encourage standardization (see (igures below). Under the JIS Marking System, following government inspection regarding quality control and other factors, authorized manufacturers are permitted to attach the JIS mark to products which belong to categories designated by the relavant Minister(s) as worthy of the mark, thus helping users and consumers to judge the quality and performance of the product. So far, some 1,100 items bear the JIS mark, and some 16,300 permits, including about 150 approvals for overseas factories, have been granted.

There are two possible ways of obtaining JIS marking approval for foreign factories: procedures A and B, as explained in the note below.

Fig. A



Fig. A. The mark to be affixed by the manufacturer of the designated commodities provided in Article 19, of the Industrial Standardization Law, to the designated commodities. their packages, containers or involces when the manufacturer has obtained permutation or approval

Fig. B. The mark to the affixed by the manufacturer using the designated processing techniques provided in Anucles 25, of the Industrial Standardization Law, to the processed commodities, their packages, containers or invoices when the manufacturer has obtained permission or approval.

Fig B

the sig to character

3. International Standardization (1) Participation in ISO and IEC

A large number of international standards have been established by the International Organization for Standardization (ISO) and the International Electrotechnical Commission (IEC). Both organizations are involved in a wide range of activities. There were about 7,400 ISO standards and about 2,100 IEC standards at the end of 1989. The Japanese Industrial Standards Committee (JISC) is a member of ISO and IEC and has participated actively in ISO's work since 1952 and in IEC's since 1953. Whenever possible. JISC has taken on important duties in the secretariats of the organizations' technical committees and

sub-committees and has been active in promotin international cooperation and exchanges of tectmology an information

At the 13th General Assembly of 190 was held a Tokyo t 1985, Mr. Isamu Yamashita, vice-president of Kesdattren, wa elected the 14th ISO president. The first Japanese presider of ISO, he served until 1988.

(2) Technical Cooperation with Developing Countries

The Standards Department carries out technical cooperatio in the field of Industrial standardization and Quality Control These cooperation are designed to help developing countriprogress in industrial standardization and Quality Control ; collaboration with the Japan International Cooperatio Agency (JICA) and the Japanese Standards Association (JSA)

Three group training courses are conducted each year: a month course on implementation of TQC and standardizatio activity, a 2-month course on certification and inspectic system and a 1-month course on a senior seminer o industrial standardization and quality control. Participants : these courses are Governmental officials it charge c standardization in government agencies or national standard bodies in developing countries.

Aside from holding training courses, the Standard Department sends experts and survey learns on industri-Standardization and quality control to developing countrie upon official request. A nd in order to transfer technolog from JA ian to a developing countries by means of training engineers and the governmental officials in charge (standardization and quality control in developing countries. center for standardization is scheduled to be built, whe Japanese experts will work for technical cooperation. Th center will be a base to establish and promote standardizatio in the country. In 1989, five year cooperation plan started the Industrial Standardization and Certification Test Center Thailand.



A foreign factory seeking JIS marking approval may proces



according to scheme A or B.

TECHNOLOGY RESEARCH AND INFORMATION

1. Technological Surveys

In order for Japan to make sound economic progress and contribute to global welfare, it must deal successfully with a wide variety of issues, including trade friction, resources and energy supply, and employment. The creative and independent development of technology in a comprehensive and efficient manner will be indispensable to achieve these goals. In the planning and preparation of a reasonable and effective industrial technology policy, there is a need to come to grips with research and development both at home and abroad and to analyze industry and its problems while checking these findings against actual research and development in Japan

To this end, AIST surveys trends in research and development, technology trade, and patents in Japan and overseas: the technology policies and development status of other countries, and more comprehensively, the important and urgent questions involved in the pursuit of creative and independent technological development.

Domestic surveys

AIST conducts surveys in Japan to understand trends in technological development, research and development activities of enterprises, problems surrounding technology, and technological policies.

(Themes for 1989)

- · Survey on technology transfer
- Survey on reorganization and location trends of research agency of enterprises in Japan
- Survey on the present state of industrial technology level of NIEs in Asia and the research and development strategy of our enterprises for NIEs of Asia

Overseas surveys

The office conducts collection and analyses of various overseas technological publications, reports, and literature as well as surveys and analyses of technological development and technological policies in advanced countries such as the USA and Europe in cooperation if JETRO and other organization while utilizing the fund for scientific technological promotion adjustment.

[Themes for 1989, related to the fund for scientific technological promotion adjustment]

Survey on the method of advanced scientific technological promotion adjustment fund

The Committee for Science and Technology Policy (CSTP) of the OECD conducts various programmes including the exchange of experiences and information on science and technology policies of member countries in order to promote the international cooperation in the field of science and technology. AIST makes positive contribution to the activities of CSTP, participating in various conferences.

In May, 1988, OECD started the 3-year programme named TEP (Technology/Economy Programme) for the purpose of analyzing the inter-relationship between science, technology, economy and society and also developing useful indices for the governments of member countries to plan, execute, and evaluate the technologyrelated policy.

In this regard, the Government of Japan hosted an international conference for TEP in Tokyo in March, 1990.

2. Propagation Activities

To enhance awareness of mining and manufacturing technologies and the industrial technology policies of ALST, the Agency publicizes its policies and the technological achievements made at its 16 research laboratories in its bulletin "ISAT (Industrial Science and Technology)". In addition, ALST issues "An Introduction to ALST", distributed domestically and overseas in both Japanese and English, in which on-going projects are described. ALST also publishes details of its work in the "ALST Annual Report". Newspapers, television, and radio are also used to report on the progress of ALST projects under way.





a) Julin Research and Whet Advanced Countries

	Name of Research Project	Japanese Research	Counterpart Research in Developed Country	Counserport is statuse in Developing Country	Durnaum (Fiscal year)
1	Research on exploration techniques for rare mineral rasources	Geological Survey of Japan	United States Geological Survey, USA	Bureau of Mines and Geo-Sciences. Philippines	1987-1991
2	(Kare mineral resources) Chernical assessment in Nile River	National Research Institute for Pollution and Resources	United States Environmental Protection Agency, USA	Mansoura University. Egypt	1990-1992

5) Joint Researches for Glob dEnvironmental Technology

Name of Research Project	AIST Research Institute	Counterpart Research Institute	Country	Duration (Fiscal year)
 Studies on afforestation with functional soil improving materials (Functional soil) 	National Chemical Laboratory f~ Industry Government Industrial Development Laboratory, Hokkaido	Central Arid Zone Research Institute Industrial Technology Development Institute	India Philippines	1990-1993
 Prevenuion of expanded pollution in the tropical zone with the metal mining development (Pollution with metal mining) 	National Research Institute for Pollution and Resources	National Department of Mineral Production	Brazil	1990-1992

(2) Promotion Program of Research and Development Cooperation

Name of Research Project	Japanese Research Institute	Counterpart Research Insutute	Country	Durkion (Fiscal year)
Research and Development Project of Machine Translation System with Japan's Neighboring Countries	Electrotechnical Laboratory	China Sofiware Technique Corporation	China	1987-1992
	Center of the International cooperation for Computer-	National Electronics and Computer Technology Center	Thailand	
	relation	Language Research Division	Malaysia	1.1.1.1
		Agency for the Assessment and Application of Technology	Indonesia	
International Research Cooperation on Recovery of Valuable Resources in Brine	Governmental Industrial Research Institute, Shikoku	Comision de Fomento Minero The Institute of Saline, Academia Sinica	Mexico China	1989-1994

(3) Bilateral and Multilateral Cooperation

AIST Promotes bilateral cooperation with China. Korea, etc. through science and technology agreements and multilateral cooperation with ASEAN countries

3 NEDO International Research Exchange Center NEDO has established the International Research Exchange Center in 1989 to conduct the following four programs:

THE JAPAN KEY TECHNOLOGY CENTER

The Japan Key Technology Center, established in response to a proposal by the private sector, conducts activities directed at the overall improvement of the environment for private research and development in fundamental technologies.

(a) Capital Invesument

The Center provides capital investment for researches carried out by companies established for joint research purposes. ('90FY ¥21.7 billion)

(b) Loan Service

The Center provides conditional interest-free loans to aid in reducing R&D related risks and costs. ('90FY' ¥6.3 billion)

(c) Mediation in Arranging Joint Research

Mediation is performed for private companies wishing to conduct joint research with national research institutions. (1) International joint research program

- (2) International research exchange program to invite foreign researchers in the long-term and help them do research and line in Japan smoothly.
- (3) On the Research Training Program to re-educate researchers in and out of Japan.
- (4) International Joint Research Grant Program to contribute the advancement of international exchange.
- (d) Execution of Consigned Research
- The Center brings together experts from government industry and academia to conduct research consigned to The Center by private companies.
- (e) Japan Trust International Research Cooperation Service The Center has established a charitable trust called the Jarun Trust Fund. The operating profits from this fund will be used to invite foreign researchers in key technologies to Japan.
 (f) Research Information Service
- The Center collects and sorts a wide variety of important research literature which is kept on file at national research institutes and other organizations.

(g) Surveys Service The Center conducts various kinds of surveys to aid privatesector research in key technologies.



ROMOTION OF TECHNOLOGICAL DEVELOPMENT IN THE RIVATE SECTOR

To encourage R&D by the private sector, tax incentives are ered for technological development as well as financing for the velopment of industrial technology (through the Japan velopment Bank) and conditional loans for P&D projects, and at operates a research association is operated for promoting ning and industrial technology.

Tax incentives for Technological Development

- (1) The following tax incentives have been instituted (effective until March 31, 1993) for facilitating research and development in fundamental technologies. These cover a 15 percent maximum deductible for corporate or income taxes or 10 percent in (a) below.
 - (a) Tax method for Deducting Additional Research Expenses These are deductible from corporate or income taxes, and are equal to 20 percent of the excess of current qualified R&D expenditures over the highest amount of the previous R&D expenditure.
 - (b)Tex Incentives for Promoting R&D in Fundamental Technologies

Also deductible from corporate or income taxes is seven percent of the cost of acquiring facilities for conducting R&D in fundamental technologies. Categories of facilities are stipulated in the Ministry of Finance Notifications -No.47 dated March 30, 1985, No.60 dated March 31, 1986, No.126 dated September 29, 1987, No.52 dated March 31, 1988, No.58 dated March 31, 1989, and No.55 dated March 31, 1990.

(c) Tax Incentives for Promoting R&D by small and medium enterprises

Six percent of the cost of R&D by small and medium enterprises during the business year, applied selectively with (a) above are deductible from corporate or income taxes.

- (2) Tax Incentives for Mining and Industrial Technological Research Associations
 - A. Special depreciation allowances are given to members of research absociations for acquiring fixed assets used in experimental research in promoting mining and industrial technology.

- B. Condense recording, of down to one yen, of charges imposed by cooperatives for the equisition of fixed assets required for the study of mining and industrial technology. C. Tax reductions are given on fixed assets used for research.
- (3) Special depreciation allowances are permitted for assets used in subject research.
- (4) Donations to Research Corporations, by special permissions, may be calculated as losses.
- 2. Promotion of International Joint Research In order to develop future industrial technology and improve both domestic and international cooperation on researches and studies, grant is given to international joint research teams in material fields. This program is administered by the New Energy and Industrial technology development Organization.
- 3. Conditional Loans for R&D Projects

Other conditional loans are available for R&D development of energy-saving technology and alternative energy technologies for petroleum and new power generation techniques.

 Financing for the Promotion of Industrial technological Development (Development of new technology) (Japan Development Bank)

Funds are provided at attractive interest rates for the commercialization of important industrial technologies and the construction of special structures for advanced basic research which will make a significant contribution to the advancement of industrial technology and play a key role in upgrading the industrial structure.

 Research Association System for the Development of Minung and Industrial Technology

This system, taking into account the efficiency and importance of joint research by enterprises, gives legal status to cooperative research organizations producing technology related to industry and mining. It was started in May, 1961 and 53 associations are currently active.

utline of Finance System for the Promotion of Industry and Technology: Budget for FY1990

1.11. A re-residence of the second state of the local distribution of the local second s	In present to be an other provide and the second science of the second state of the se	And the second	
	12.5.6.4 二气花浸渍	Development of new technology	
	Improvement of research facilities	Development for commercialization	Commercialization of new technology
Construction costs ligible for financing	Cost entailed in acquiring special buildings and structures for basic and applied research	 Construction of demonstration plants Trial manufacture of machinery and equipment 	 Production line construction Development of heavy machinery
Ratio of financing	Approximate	y 50% of construction costs of works elig	ible for financing
Financing period		15 years or less (in principle)	
Redeemable period		Two to three years (in principle)	
Budget for FY 1990		750,000 (million yen)	
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(4) Multilateral Cooperation

1) International Energy Agency (IEA) Energy R&D Projects MITT participants in cooperative research and information exchange under the Committee of Research and Development of IEA.

		1990. 9
Working Parties	MITT's Joining Implementing Agreements	Start of MITT's Participation
End Use Technology	Advanced Heai Pump Alcohol and Alcohol Blends as Motor Fuels Energy Conservation in Combustion Advanced Fuel Cells CADDET June, 1990 Assessing the impact of High temperature Superconductivity June, 1990	April, 1979 February, 1986 April, 1984 (April, 1990)
Renewable Energy	Bioenergy Wind Energy Conversion Systems Solar Heating and Cooling Systems Production of Hydrogen from water	May, 1987 April, 1978 October, 1977 October, 1977
Fossid Energy	Cost Technology Information Service Cost/Otil Mixtures Enhanced Oil Recovery Aunospheric Fluidized bed combustion	March, 1977 March, 1981 May, 1979 February, 1980
Fusion Power	· Reversed Field Punches	(May, 1990)
Other	Energy Technology System Analysis Energy Technology Data Exchange	September, 1981 January, 1987

2) International Cooperation Projects proposed by the Working Group on Technology, Growth and Employment (Summit). These projects will be carried out on an independent basis, separated from the Summit framework.

Projects promoted by AJST	Participants (Observers)
Photovoltaic Solar Energy	Italy, France, Germany, UK, EC, (US)
Advanced Robotics	France, US, UK, Germany, Canada, Italy, (EC)
VAMAS (Versailles Project on Advanced Materials and Standards)	UK, US, Canada, EC, France, Germany, Italy

3) Organization of Economic Cooperation and Development AIST takes part in the Committee of Science and Technology Policy of OECD. In 1990, Technology/

Economy Program (TEP) Symposium will be held in Tokyo under the auspice of CSTP.

2. Cooperation with Developing Countries

(1) Institute of Transfer of Industrial Technology (ITTT project) 1) Joint Research for New Technology

Fiscal year: April 1 - March 31

-	Name of Research Project	AIST Research Institute	Counterpart Research Institute	Country	Duration (Fiscal year)
1	Research on sensing technology for cutting process information (Sensing for cutting process information)	Mechanical Engineering Laboratory	Korea Institute of Machinery and Metals	Korea	1987-1991
2	Research on geology and muneral resources of the collision zone in Pakistan (Geology of the collision zone)	Geological survey of Japan	Geological Survey of Pakistan	Pakistan	1987-1990
3	Research for industrialization of thermomechanical pulping of oil palm by-products (Thermomechanical pulping of oil palm)	Government Industrial Research Institute, Shikoku	Palm Oil Research Institute of Malasyia	Malaysia	1987-1990
4	Measurement of three dimensional object and nondestructive testing (Measurement of three dimensional object)	Mechanical Engineering Laboratory	Research and Development Center for Calibration, Instrumentation and Metrology	Indonesia	1988-1990
-	Hydrogenation of palm oil components (Hydrogenation of palm oil)	National Chemical Laboratory for Industry	Palm Oil Research Institute of Malaysia	Malaysia	1988-1990
	Enhancement of reference sections of sedimentary basing)	Geological Survey of Japan	Mines and Geosciences Burea	Philippines	1988-1991
	Separation and refining of rare metal cores from China (Rare metal ores from China)	National Research Institute for pollution and Resources Government Industrial Research Institute, Tohoku	Guanzhou Research Insutute of Non-ferrous Metals	China	1988-1993



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Name of Research Project	AIST Research Institute	Counterpart Research Institute"	Country	(Fiscal year)
Research on the new coal combustion technology by fluid and bad	Government Industrial Development Laboratory, Hokkaido	Institute of Coal Chemistry, Academia Siruca	China	1988-1990
Study on the effective utilization of neglected hydrocarbons (Utilization of neglected hydrocarbons)	National Chemical Laboratory for Industry	Korea Advanced Institute of Science and Technology	Korea	1989-1991
Air polluuon assessment at tropic area.	National Research Institute for Pollution and Resources	Indian Institute of Technology	India	1989-1991
(Air politice assessment) Research on utilization of natural	Government Industrial Research Institute, Tohoku	Mineral Technology Development Center	Indonesia	1989-1992
Development of AL/Li Alloy	Mechanical Engineering Laboratories	Nayang Technological Institute	Singapore	1990-1993
Research on rare-metal resources in weathering crusts of granitoid.	Geological Survey of Japan	Department of Mineral Resources	Thailand	1990-1992
Study on countermeasures for the ignition source of gas and coal dust explosions in coal mine (lamit on source in coal mine)	Nauonal Research Institute for Pollution and Resources	Central Coal Mining Research Institute	China	1990-1993
Effective activation treatment of lignite and peat materials (Lignite and peat materials)	Government Industrial Development Laboratory, Hokkaido	Thailand Institute of Scientific and Technological Research	Thailand	1990-1992
 Study on utilization of sepiolitic Mg bearing day from Turkey (Sepiclitic Mg bearing day) 	Government Industrial Research Institute, Nagoya	Mineral Research and Exploration Institute of Turkey	Turkey	1990-1993
 Study on evaluation and utilization of Brazilian Quartz (Brazilian Quartz) 	Government Industrial Research Institute, Nagoya	h State University of Campinas	Brazil	1990-199

2) Joint Research for transfer of Technology

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	Name of Research Project	AIST Research	Counterpart Research In-titute	Country	Duration (Fiscal year)
1.	Research on evaluation for standard of large force (Standard of	National Research Laboratory of Metrology	National Institute of Metrology	China	1987-1990
2	Research on reliability of volume measuring instruments in the tropics (Wolume measuring instruments)	National Research Laboratory of Metrology	Directorate of Metrology National Institute of Science and Technology	Indonesia Philippines	1988-1990
3	Evaluation for material selection and corrosion prevention of materials in the tropical environments (Materials telection and corrosion prevention)	Government Industrial Research Institute, Chugoku	Research and Development Center for Metallurgy	Indonesia	1990-1992
4	Development of durable concrete based on the utilization of indigeous resources (Durable concrete)	Government Industrial Research Institute, Kyusu	Institute of Human Settlements	Indonesia	1990-1992

3) General Research

Name of Research Project	AIST Research Institute	Duration (FY)
 Research on plasticization of tropical and subtropical plant materials (Plasticization of plant materials) 	Industrial Products Research Institute	1987-1990

INTERNATIONAL COOPERATION IN RESEARCH AND DEVELOPMENT

International research and development cooperation advances Japan's own R&D while contributing to the formation of harmonious economic ties with other nations.

AIST is therefore an active research partner with developed and developing countries alike.

AIST conducts joint research programs in the area of advanced technology with developed countries and invites foreign researchers. Besides, under the Institutes of Transfer of Industrial Tachnology (TTTT), AIST conducts joint research and exchange of researchers with developing countries.

Further, NEDO help foreign researchers work and live in Japan smoothly.

1 Cooperation with Developed Countries

(1) Invitation of foreign researchers

- 1) AIST has established on program in FY1988 to provide foreign researchers with an opportunity to conduct research for a certain period of time with researchers at the institutes of the Agency of Industrial Science and Technology (AIST) in order to advance scientific and technological knowledge in their respective fields and to promote creative research and development in the open environment of the institutes.
- (2) Joint Research Project
 - 1) Specific International Joint Research Projects

- a Qualifications
- Generally, a researcher under the age of 35, holding a doctorate in science or engineering
- b Number and period of invited researchers Approximately 13 persons for a period of one year c. Host insulutes
- Sixieen research institutes belonging to AIST d Compensation
- Round trip aufare, living expense, housing allowance. family allowance and relocation allowance
- e. Japanese language course
- A Japanese language course is given as a general rule at the beginning of the researcher's stay.

Besides Foreign Researchers can be invited to AIST laboratories by

- 3) AIST accepts researchers in EC countries through Japan-EC Industrial Cooperation Center
- 4) Foreign Researchers are invited by a charitable trust called the Japan Trust Fund which is administered by the Japan Key Technology Center.
- 5) AIST has made a memorandum of understanding with National Science Foundation to accept up to thirty US researchers a year to AIST laboratories.
- (Research conducted jointly by AIST research institutes and research institutions in developed countries)

Name of Project	Japanese Research Institute	Counterpart Research Institute	Country	Duration (Fiscal year)	
Research on optical microgassensors	Government Industrial Research Institute, Osaka	University Catholic Luban Belgium		1987-1990	
Research on precision evaluation of new superconductors and development of precision measurement devices	Electrotechnical Laboratory	National Institute of Standard	USA	1988-1992	
Research on the synthesis of fluorine- containing heterocyclic compounds and	Government Industrial Research Institute, Nagoya	National Institute of Health	USA	1989-1992	
Research on generation and utilization	National Chemical Laboratory for Industry	Technische Universität München	Germany	1990-1993	
Research on the thermobiology. emphasizing on water structure in	Fermentation Research Institute	National Institute for Medical Research	United Kingdom	1990-1992	
Research on mechanisms for release of methane into the atmosphere	Fermentation Research Institute Geological Survey of Japan	Ohio State University Tubingen Universität United State Geological Survey	USA Germany	1990-1993	
Research on Acid rain Mechanism by the advanced observation and modeling	National Research Institute for Pollution and Resources Government Industrial Research Institute, Nagoya	National Center for Atomospheric Research Iowa State University	USA	1990-1993	

2) Research Grants to International Joint Research Team In order to develop the future industrial technology and to contribute to the improvement of both domestic and international cooperation on researches and studies, the system is to promote international joint research team carrying out original research related material fields. This program is administered by the New Energy and Industrial Technology Development Organization.

(3) Bilateral Cooperation

AIST cooperates with developed countries such as United States, West Germany, France, Italy, United Kingdom. through science and technology cooperation agreements. industrial cooperation talks and so on concerning joint research, exchange of researchers, and information.



Country	Frame Work	Yew of Invusion	Field of Cooperation (Mutually Selected Areas)
USA	US-Japan Conference on Natural Resources (U.J.N.R.)	1964	Fire Research and Safety, Marine Mining, Marine Instrumentation and Communications, Marine Geology, Others
	Agreement between the Government of Japan and the Government of the United States of America on Cooperation in Research and Development in Energy and Related Fields	1979	Fusion, Coal Energy, Solar Energy, High-energy Physics, Other energy and energy-related research and development areas, as may be mutually selected.
	Agreement between the Government of Japan and the Government of the United States of America on Cooperation in the field of Environmental Protection	1975	Stationary Source Pollution Control Technology, Management of Bottom Sediment Containing Toxic Substances, Air Pollution-related Meteorology. Others
	Agreement between the Government of Japan and the Government of the United States of America on Cooperation in research and Development in Science and Technology	1988	Life sciences, including biotechnology; Information science and technology; Manufacturing technology; Automation and process control; Global geoscience and environment; Joint database development; and Advanced materials, including superconductors.
United Kingdom	MITI-DTI Talks.	1988	Superconductivity, Marine Technology Biotechnology, Others,
France	Agreement between the Government of Japan and the Government of the French Republic on Cooperation in the field of Science and Technology	1974	Marine Science and Vectoology, Biological and Medica, science and Technology, New Energy Technology, Energy Conservation, Others
West Germany	Agreement between the Government of Japan and the Government of Federal Republic of Germany on Cooperation in the Field of Science and Technology	1974	Marine Science and Technology, Biological and Medica Science and Technology, Environmental Protection Technology, New Energy Resources, Transport Technology, New Materiala, Data processing, Information and Documentation, Mechanical Engineering, Others
Australia	Agreement between the Government of Japan and the Government of Australia on Cooperation in Research and Development in Science and Technology	1980	Experimental Petrology, Lower Atmosphere Physics, Fluidized Bed combustion Technology, Vision Technology for Robots, Others
	Cooperation between Japan and Australia in Energy Research and Development and Related Areas	1978	Coal Technology, Solar Energy Utilization, Energy Conservation, Others
Canada	Agreement between the Government of Japan and the Government of Canada on Cooperation in Science and technology	1986	Environment Technology, Energy Technology, Space and Communications, Computers and Robotics, Others
5weden	Japan (AIST) — Sweden (STU) Research and Development Cooperation	1981	Medical and Welfare Technology, Biotechnology, Materials (Polymer and Composite, Ceramics, Lignan), Others
lialy ,	Agreement between the Government of Japan and the Government of Italy on cooperation in Science and Technology	1988	New materials, Biotechnology, Volcanology and Seismology, Physics Environment, Articial Intelligence Energy





R&D on earth environment technologies for 1990 (Designated researches)

(Unit million ver

Name of project	Budget for 1990	Outline of project
Study on cartion dioxide fixation technology by artificial shotosynthesis etc.	165	R&D to transform carbon dioxide (which is the major cause of greenhouse effect) into other useful materials by using energies (solar rays, etc.) currently not used.
Study on carbon dioxide fixation by algae		R&D on the counteraction ability of the nature against increase in carbon dioxide and countermeasures using such counteraction through quantitative definition of photosynthesis of algae by conditions and study on its mechanism. Technical development to enable easier and more correct measurements of the concentration of carbon dioxide in seawater with less sample.
Study on coral mers as sinks of atmospheric carbon droxide		R&D on fixation of carbon dioxide to calcium carbonate by clarifying the past concentration level of carbon dioxide and time stone generation state, and actual state of present coral rol and factors affecting growth of coral
Analysis and evaluation of technologies for reducing anthro- pogenic carbon dioxide emissions		R&D to establish an engineering model to analize and evaluate of technologies for reducing anthropogenic carbon dioxide emissions and to clarify the effectiveness of reducing technology and composition of an energy system with less emissions
Study on recovery technology of carbon diatide		R&D on effective separation and concentration of carbon dioxide
Development of biodegradable plastics		R&D on materials compatible with environment, such as plastics degradable in nature

International Joint researches on global environmental technology for 1990

Classification	Name of project	Counterpart	Period	Outline of project
Specific international joint research projects (global environmeni)	Research on mechanisms for release of methane into the atmosphere	USA, West Germany	90-93	Biological and geometrical researches to clarify the release mechanism of methane gas (whose green house effect is about 20 to 40 times larger than carbon dioxic i) into atmosphere.
	Research on Acid rain Mechanism by the advanced observation and modeling	USA	90-93	Precision analysis of the acid rain components and creation of a simulation model based on in-situ observation of generation, long-distance transpon, and transformation process of acid rain components.
Joint researches for grobal envirormenial technology	Studies on afforestation with functional soil improving materials (Functional soil)	India. Philippine	90-93	Development of functional soil recovering agent with water and ferulizer retaining capacity and soil organisms compatibility in order to recover soils suffering degradation due to expansion of desert, excessive pasturing
	Prevention of expanded pollution in the uppical zone with the metal minung development (Pollution with metal minung)	Brazi	90-92	Development of wide area pollution survey technology and prevention technology for mining production of tropical forests which is presumed to affect animals and plants over a wide area.

CONSOLIDATING RESEARCH AND DEVELOPMENT SYSTEMS RELATING TO INDUSTRIAL TECHNOLOGY

Outline of Policy

If Japan, now holding a key position in the world economy, is to play a role worthy of its position toward well-balanced development of the world economy while achieving further development toward the 21st century, it is essential for Japan to promote positively the cre-tive technical development and to contribute to the world community through technical development. What is required for this purpose is to improve and expand large-scale research and development infrastructures, to put into full swing the research and development in basic and advanced fields, and to establish a system s t consolidated augmentation of international research cooperation.

Under the "Law for Consolidating Research and Development System Relating to Industrial Technology" enacted in May, 1988, the New Energy Development Organization (NEDO) was expanded in October to the New Energy and Industrial Technology Consolidated Development Organization. This organization is given responsibility for undertaking industrial technology.

- 1 Research and Development Program
 - (1) Description
 - te addition in concentional P (D Program on Basic

Development Program (Large-scale Program), and Research and Development Program on Medical and Welfare Equipment Technology, NEDO has started newly the Research and Development Program on Earth Environme-Industrial Technology, NEDO will undertake comprehensiv and mobile R&D for these programs while carrying theout in closer cooperation with government, and academ. sector and private industry as well as foreign institutes. In 1991, NEDO will attempt efficient and steady execution of programs continued from the previous year. At the sam time. NEDO will set upon new six themes, including a nesoftware structuring model for the Future Industrie Program, advanced function creation machining technolog and application technology of human senses to measuremer for the Large-scale Program, and the condition moniton: system for the aged, digital hearing aid, and continuou blood sugar measurement system for the Medical an Welfare Program.





In the field of earth environment, NEDO will undertake general planning and research, technical development for carbon dioxide fixation and effective utilization. development of materials with less environmental load, and technical development for production process well matched to the environment.

(2) Budget

General accounts of ¥5.7 billion will be earmarked as fund for NEDO, while special accounts of ¥13.7 billion as grant to NEDO.

2 Research Facility Development Program

(1) Description

Large and high-level research facilities, which are indispensable for promotion of creative R&D in the advanced field, will be improved and expanded, and opened wide for domestic and foreign researchers.

Improvement and operation of these facilities are performed by the third sector erected for each facility. The third sector will finance one half of the initial investment by the capital and another half by the load. Two-thirds of the capital will be provided by NEDO while remaining one third by private and local government agencies. On the other had, 70% of the loan can be obtained interest-free from Japan Development Bank and Hokkaido and Tohoku Development Corporation, and Industrial Infrastructure Improvement Fund will provide guarantee of obligation for the commercial bank loan.

In cases where it is particularly difficult to establish basic research facilities due to the high level of research and development required, NEDO will also undertake to develop the needed equipment and support systems.

- (2) Outline of projects
 - 1) Ion Engineering Center

(Location: Kansai Culture and Science City, Hirakata, Osaka)

A facility to study the technology of applying ion beam to industries will be founded for general utilization. Incorporated in November, 1988 and partially opened in July, 1990

2) Research Center for the Industrial Utilization of Marine Organisms

(Location: Kamaishi, Iwate: Shimizu, Shizuoka)

A facility to study the technology of utilizing marine organisms in mining and industries for general service. Incorporated in January, 1989 and opened in April, 1990

3) Japan Microgravity Center

(Location: Kami Sunagawa Town, Hokkaido) A vertical drop facility which enables various nongravity tests for about 10 seconds using existing vertical pits of old mines will be improved and expanded for general service. Incorporated in March, 1989 and opened at the end of 1990 Fy

4) Applied Laser Engineering Center (Location: Nagaoka, Niigata) A facility to study the technology of applying laser to industries will be erected for general service. Incorporated in March, 1990

5) Advanced Material Research Center (Location: Ube, Yamaguchi; Tajimi, Gifu)

A facility to study and evaluate material physical properties and functions in super-high temperature environment will be improved and expanded for general service. Incorporated in March, 1990

(3) Budget

Industrial investment account capital of ¥2.2 billion will be earmarked as a component of investment of NEDO to the third sector and general accounts of ¥200 million as a fund necessary for improvement of facilities by NEDO itself.

3. International research per

(1) Background

Today, not only overseas demand on research and technical cooperations is increasing quantitatively, but also the content of demands is growing more and mr ; diversified. It is difficult to meet these demands by NEDO alons. On the other hand, foreign researchers accepted by NEDO is increasing steadily in number and apected to increasing further in fature. Norwithstanding II's situation, a system of providing lodgings and various benefits to researchers accepted is not yet completed, and it is urgently requested to improve and expand the comprehensive acceptance system. Moreover, domestic and foreign enterprises are showing an attitude to sugment further fostering and retraining of researchers to cope smoothly with worldwide structural adjustment of industries.

In view of above present state, NEDO will execute following four programs under integrated control.

- (2) Outline of programs
 - 1) International research cooperation

Joint research with foreign research institutes will be made on subjects requested from foreign countries under cooperation of the private sector while making the best of talents and knowhow of National Laboratories.

In 1990, two themes started in 1989 will be put into full gear and simultaneously the new "Research and Development concerning Improvement of Lowtemperature Starting Performance of Methanol-fueled Automobiles" will be started jointly with the USA.

2) International research fellowship

In addition to invitation of foreign researchers (mainly young researchers) for the long period, various benefits. such as training of Japanese, consulting on daily life. leasing and facilitzing of lodgings, etc. will be offered to accepted foreign researchers to enable them to spend comfortable research life in Japan. R&D information necessary to expedite international research exchange will be supplied to foreign countries.

Researcher training

The researcher program including lectures and on theresearch-training will be executed in an attempt to foster and mutual exchan, . of researchers of domestic and foreign enterprises.

4) International joint research support program

The grant will be provided to cover expenses incurred by the research activity to international joint research teams, including foreign researchers, which engage in R&D in the basic field (physical properties, etc.) expected to become fruitful seeds of industrial technologies toward the 21st century.

New four programs will be added to continuous 12 programs for 1990.

(3) Budget

1) International research cooperation

Special accounts of ¥197 million will be earmarked as grant to NEDO and ODA of ¥59 million as fund entrusted to NEDO.

- 2) International research fellowship General accounts of ¥54 million will be earmarked as grant to NEDO and general accounts of ¥161 million as fund entrusted to NEDO.
- 3) International joint research support program General accounts of ¥516 million will be earmarked as financial resources of grant.





MEASURES FOR REGIONAL LECHNOLOGY DEVELOPMENT

- Specific Regional Technology Development System (Regional Large-Scale Projects; see table below)
 - This system was inaugurated in 1982 to promote regional technological development, and executes 7 projects in 7 areas in 1990.
- System of Advanced General Regional Technology Development This system promotes joint research for development to introduce advanced technology into local areas.
- Regional Technical Cooperation Promotion Projects Major projects under this heading include:
- Expansion of AIST's "Research Information Processing System (RIPS)" network to include regional research institutes, wid, the aim of closing "research information gaps" between different loc dities;
- (2) Preparation of a high technology application manual containing research information and know-how on advanced industrial technology, gathered by regional research institutes of AIST;
- (3) Promotion of joint research and cooperation with local societies, as well as dispatch of researchers for technical instruction, seminars for technical learning for local researchers, and technical symposiums.

(Unit milkm yen)

Projeci Name	Period (FY)	Tokal Expenditure	Budget for FY1990	Outline of Project	R&D Results
Inteligent snow removing technology for cold regions (Hokksido region)	1987 1991	500	30	Development of an advanced snow remover equipped with newest sensing and automatic control systems to promote safety increase efficiency and decrease opersuing loads.	Experiments on the obstacle detector
Visual recognition and identification for flexible manufacturing systems (Chugoku region)	1987 1990	400	28	Development of a system to promote the efficient production of mechanical and electrical components in small lots of way many varieties through visual recognition of the components position, shape, and surface properties, and a database consisting of the knowledge of skilled workers.	The apparatus detecting surface defects on mechanical and electrical parts has been developed together with a classifying technique. The image recognition system has been also developed, which can identify complicated objects correctly using the knowledge database.
Synthetic technology of artificial clay for high performance ceramica (Chubu region)	1988 1992	600	40	Development of economical synthetic technology of highly pure and plastic kaolinitic clay for plasticizer and raw material of high performance ceramics.	The effects of the addition of several anions and ostions on the hydrothermal symbols was studied. Usefulness of seed crystals was clarified and has been applied to the tkg/hach scale synthesis. The relations between synthesic conditions and plasticity of produced kaolinus were studied in detail. These results indicated the necessity of break down of cosgulated particles.
R&D on re-utilization system technology of composite materials (Shukoku region)	1988 1992	350	28	For manufacturing valuable products from large FRP wastes, i.e., fishery boats and bath tube, this project is aimed at developing automatic cutting and crushing apparatuses for wastes and constructing new re-utilization system. from the crushed and separated composite.	For development of efficient automate cultury machine, automatic supplying equipment was made on trial. In composite of the recised FR powder with plaster, products had higher bending strength that of cement concrete. In thermal decomposition of FRP waste under steam, phthalic acid and styrene were easily recovered. The reinforced glass fiber were investigated to endow porosity.
Advanced utilization of time and time-based compounds for materials development (kyushu region)	1988	500	33	Development of technology for advanced utization of time and time based compounds such as calcium carbonate, calcium siticate hydrates, apatites etc. for pigment and fuler for paper, substitute for arbestos, fuler medium, adsorbent, fixed bed for bioreactor, etc.	Plate type particles of basic calcium carbonal with high dispersibility were synthesized in th larger scale, and stocchiometric hydrotvianus with high purity were obtained. Synthesizing conditions for incalcium silicate hydrate (TSI of longer crystals than 100sum were cardied
Advanced surface modification in material processing "Kinki mgion)	1989	600	35	Development of technology on advanced surface modification for matenals such as metals, plasues and ceramics for mechanical, electrical, magnetical, and optical surface functions.	Improvement of adhesion strength between thin films and the substrates of various kind to ton beam imadiation was carried out. The energies of ions were selected in the followin ranges, 0.5-2keV, 10-40keV and some MeV
Advanced Internal Inspection Technology for Composite Substances (Tohoku region)	1990	500	11	Development of the internal inspection system by using ultrasonic imaging. X-ray imaging and image analysis to ceruly the reliability of an internal sinucture and of a bonded interface in the composite subtrances such as cenamic-metal hond, micro-electronic device and carbonfiber reinforced plastic.	(Project launched in "Y 1990)





THE HUMAN FRONTIER SCIENCE PROGRAM

Since the Industrial Revolution, technology has long been recognized as a method of conquering, managing, and controlling nature. Technological developments have helped us to move into scientific domains of higher temperature, higher pressure, higher speed, and greater magnitude. As a result, our knowledge base, as well as the range of human activities, has greatly expanded. In the meantime, a variety of serious problems have emerged which include increases in resource and energy consumption and the heavier load imposed on the environment as well as more intense man-machine conflicts. In order to ensure more humanity and greater prosperity in the 21 at century, it is necessary to create a new system of scientific technology which will foster harmonious relationships with society and nature.

Based on these observations, it is believed that basic research on the precise mechanisms of organisms has the potential to become a driving force in developing various research areas and could become a treasure chest of scientific and technological weeds as it is expected to exploit the frontier of scientific technology for the 21st century.

Living organisms possess superior functional characteristics which have become extremely sophisticated and precise over a billion years of biological evolution. In contrast, only several hundred years have passed since the Industrial Revolution.

Today's most advanced scientific technology is no match for the superior mechanisms of living organisms. If these superior biological functions were to be clearly elucidated and properly utilized, it would help develop a new system of scientific technology characterized by "anti-pollution" and "energy saving". thus bringing humans as unlimited number of benefits. 1. Human Frontier Science Program

The Human Frontier Science Program is an international joint project in which basic research to elucidate superior functions of living organisms will be conducted in an attempt to utilize its results for the benefit of all human beings.

Voices which call for Japan to contribute more in the field of basic research are growing stronger. In response in the international area to this, the Japanese government proposed the Program at the Venice Summit in 1987 in an effort to exploit the scientific frontier of the 21st century. After the proposal was adopted at the Venice Summit, an international feasibility study was conducted by scientists in 1987 and the successful results were reported at the Toronto Summit in June, 1988.

The organization for the implementation of the HFSP was established in Strasbourg. France in October, 1989, starting the undertakings of research grants to international joint research teams, fellowships for travel and accommodation charges for research outside the country, and holding and sponsoring of workshops for information exchange and discussions. The first awardees have been designated in March, 1950.

 Research and Development Program for the Elucidation of Biological Functions.

While promoting the Human Frontier Science Program, the Research and Development Program for the Elucidation of Biological Functions was newly established in 1988 and research is currently in progress at research facilities of the Agency of Industrial Science and Technology in an attempt to elucidate biological functions under investigation.

RESEARCH AND DEVELOPMENT OF TECHNOLOGIES RELATED TO GLOBAL ENVIRONMENT

Global environmental problems such as global warming (greenhouse effect) by carbon dioxide, depletion of the ozone layer may exert substantial effects on the industrial society and human life. The solution of such problems is urgently demanded as a subject to be overcome commonly by human beings. In this respect, the designated research frame in the global environment field was created in 1989 to start promotion of advanced R&D in an attempt to promote researches on fixation of carbon dioxide by artificial photosynthesis etc., while exploiting technical knowhow and resources of laboratories and institutes of the Agency of Industrial Science and Technology.

In 1990, the international joint researches on global environment technologies by the AIST and foreign research institutes were started. The R&D program of global environment industrial technology was added to research and development activities of NEDO to proceed with improvement and expansion of the integrated R&D organization participated by the industry, government, and college. R&D projects of global environment industrial technologies for 1990

	Name of project
(1) Devi	elopment of environmentally benign production processes
1) R	& D on advaned bioreactor system
(2) Devi	elopment of environmentally benign substances
1) D la	evelopment of CPC substitutes without global warming or ozone yer depleting charactenistics
2) D	evelopment of biodegradable plastics
(3) Devi	elopmient of full carbon cycle technology
1) R	& D on fixation and reutilization of carbon dioxide by hotosynthesic microorganisms and algae
2) R	& D on fixation and reutilization of carbon dioxide by catalytic . ydrogenation
3) 5	tudy of carbon cycle mechanism in the ocean

R&D ON ENERGY CONSERVATION TECHNOLOGY -THE MOONLIGHT PROJECT-

Launched in 1978, the Moonlight Project is a comprehensive progaram of R&D for energy conservation under which work is

carried out cooperatively by national laboratories, industries and universities.

(Unit million yen)

Project Name	Period (FY)	Budgei for FY 1990	Outline of Project	R&D Result
idvanced Battery lectric Power Storage system	1980 - 1991	2.273 26 2,247]	Development of an electric energy-storage system including high-efficiency, large scale advanced batteries. The system is expected to have a load leveling function, allowing electric energy to be stored during off-peak hours and discharged during peak hours.	Developed a type Advanced Batteries with capacity of 50k W, and with efficiencies of 77%. Developed 1000k W class power storage, system using improved lead-acid batteries, and with efficiency of 70%.
uel Cell Power Ceneration Technology	1081 - 1995	3,180 (3,144)	Development of design concepts for systems adaptable to both dispersed and centralized power stations, using fuel cell power generating devices whose potential efficiency can reach as much as all to 60%. Natural gas, methanol and coal- der red gas are used as fuels.	(Phosphoric acid fuel cell) Developed two 1000k W plants and two on-site 200k W systems which were insulted in Tokashiki Island and Osaka (Molten carbonate fuel cell) Developed 1k W 10k W and 25k W class cell stacks (Solid oxide fuel cell) Developed 500W class cell stacks (Alkaline fuel cell) Developed 1kW class cell stacks and tested more than 2000 hours continuously
Super Heat Pump Energy Accumulation	1984 - 1992 -	(1.784 (1.786)	Development of several new systems, each of which consists of a high-performance electric driven heat pump system and a chemical heat storage system. These are expected to be used for air conditioning for large buildings, for distinct heating and cooling, or as process heat sources. The systems are to be operated so as to store energy at night and to discharge the stored energy in the daytume in order to contribute to a leveling of electric power demand.	Applied for 81 patents as a result of studies on working fluids, materials, elemental apparatuses, systematization etc. Developed bench plant.
Supercounducting Technology for Disatric Power Apparatuses	1988 1995	2.610 [2.52 ³]	Development of a more efficient and stable electric power system using supercounducing power apparatuses, among which generators are the closest tanget. The system will assist in overcoming problems such as power loss and lack of suitable sites for transmission lines which occur as power stations become bigger and more removely situated.	Developed high stability and high current density type 10KA class NbT) conductors for field windings Designed 70MW class model machine Designed compressor unit of reingenation system.
Ceramic Gas Turbine Project	1988 1996	L 1.132 L 1.067	Development of ceramic gas turbines applicable to co-generation and electric power generation systems. These turbines, which use non-petroleum, fuels such as natural gas and methanol, offer thermal efficiency which may be increased to 42% by raising the turbine inlet temperature to 1350°C	(Projeci launched in FY 1988)
Other		[⁶⁴⁴ [²⁹¹ 353	Leading and Basic Technology for Energy Conservation, International Cooperation in R&D. Technology Assessment on Energy Conservation: Conditional Loans for Energy Conservation Promotion, Promotion of Energy Conservation through Standardization, etc.	

Upper columns in parentheses represent general accounts and lower ones special accounts. Those not in parentheses represent general accounts only NOTE Completed R&E Projects on Energy Conservation Technology (Unit: billion yen)

Waste Heat Utilization Technology System (1976-1981, 4)
 Magneto-Hydro Dynamic (MHD) Power Generation Technology (1966-1983, 11.4)
 Advanced Gas Turbine (1978-1987, 26)

4 Surling Engine for Wide Use (1982-1987, 8)





R&D ON MEDICAL AND WELFARE EOD HENDREY 70 0%

Japan is putting much effort into raising the standard of its medical and welfare services, and there is an urgent need for more tolvanced equipment in this field. Often however, the development of technology for medical and welfare apparatus is hampered by large risks. Since 1976 fiscal year, ALST has addressed this problem by carrying out R&D aimed at the tapid development and marketing of reasonably priced, high-performance apparatus in this "high-risk" category. Research work is conducted at ALST's national research laboratories or on a consignment basis at the Technology Research Association of Medical and Welfare Apparatus (administered jointly by MITT and the Ministry of Health and Welfare). By the end of fiscal 1969, R&D had been completed on realise types of equipment for medical care and thirtees for Nursiago "handicap" eleves these are already in use.

Development of non-invasive continuous blood glucose monitoring system, digital hearing kids and health monitoring system for the elderly started this year by the New Energy and Industrial Technology Development Organization.

Development of four types of equipment for medical care and three for nursing "handicap" continued into 1990 (fiscal year) from the previous year.

(Unit: million yen)

Project Name	Period (FY)	Budget for FY1990	Outline of Project
Mode call equipment technology	and the statement of the statement of the		
1. Automatic HLA typing system	1987-1990	40	Device to automatically classify leukocyte forms to treatment of knukemia
2. Laser angloplasty system	1988-1991	221	Device to remove thrombus into averus with laser beam
 Three dimensional imaging system for medical diagnosis 	1988-1991	114	System providing three-dimensional imaging for medical diagnomia
4. Laser offeniority system	1989-1992	74	System to perform osteonomy accurately with exima laser beam
5. Non-Invasive Continuous blood glucose Monitoring system	1990-1993	8	System to measure the value of blood glucose non-invarively and continuously with two-sensor
Nursing "Handicap" equipment technology			
1 Anu-decubitus mechanical mattress	1987-1990	24	A bed to provule prolonged prevention of decubitus ulcers for those who are
2. Evacuation care system	1989-1993	68	System to cruth and remove the solidified feces standing in the rectum with
 Threadimensional tactile display terminal for visually handicapped 	1989-1992	54	supersonic vibration. System to form tactile solid body out of pin display of high density for visually hadicapped
4. Digital hearing aids	1990-1994	8	Hearing slids to be able to adjust correct indent to each hearing characteristic and to change the sound speed
II Support Equipment for participation of the elderty an society			
1. Health monitoring system for the elderly	1990-1992	57	System to be able to participate and detect early in case of emergency of the elderly

Completed R&D on Medical and Welfare Equipment (Unit million yen)

Medical equipment technology

- Muluchannel automated biochemical analyzer (1976-1978, 251)
- Automated differential blood cell analyzes (1976-1978, 269)
- (3) Anuficial heart for clinical use (1976-1979, 480)
- (4) Ponable antificial kidney (1976-1980, 617)
- (5) Laser scalpel (1978-1981, 533)
- (6) Positron computer technology (1979-1982, 470)
- (7) Laver function support device (1979-1984, 653)
- (8) Diagnossi and therapy support system for neural disorders (1981-1986, 600)
- (9) Block imaiment system for minuno-related diseases (1983-1947, 362)
- (10) Photochemical reaction system for diagnosis and therapy of cancer (1984-1987, 306).
- (11) Immunological cancer diagnosis system (1985-1988, 306).
- (12) Hyperthermia system for cancer therapy (1986-1989, 388)

- 2. Welfare equipment technology
- (1) Modular type motorized wheelchair (1976-1978, 226)
 - (2) Braile duplicating system (1976-1975, 143)
 - (3) Gost pattern analyzer for the handicspped (1976-1978, 161).
 - (4) Multifunctional ted for the severely handicapped
 - (1976-1978, 101)
 - (5) Middle ear implant (1978-1982, 429)
 - (6) Guidance device for the blind (1979-1983, 366)
 - (7) Vocal and speech training device (1979-1983, 346)
 - (8) Power driver antificial wheel leg (1980-1985, 427)
 - (9) Chair capable of 3-dimensional movement (1981-1985, 297)
- (10) Book reader for the bland (1982-1988, 585)
- (11) Transfer supporting system for the handicapped
- (1983-1988, 522)
- (12) Automated body temperature adjuster (1984-1988, 272)
- (13) System for processing prosideus sockets (1986-1989, 209)

THE NATIONAL RESEARCH AND DEVELOPMENT PROGRAM -THE LARGE-SCALE PROJECT-

Under the National Research and Development Program (popularly known as the Large Scale Project), AIST conducts K&D projects on technologies which are of particular importance and urgent need to the nation. Government funds are given by contract to participating private enterprises, which work closely with national lateratories and academic organizations

A total of 27 projects have been undertaken since 1966. Sixteen of these have already been completed, with various technical

results, including large-scale integrated circuits, high performance electric car battery technology and the practical use of desatination equipment. The results of such efforts are all available to the public, and have attracted worldwide attention. In 1990, ALST will continue to work on eight R&D projects currently in progress, and will also start two new projects. "Advanced Chemical Processing Technology" and "Human Sensory Mesurement Application Technology"

(Unit million ver.)

Project Name	Period (FY)	Budget for FY14CD	Outline of Project	R&D Results
Manganese Module 1981. 978 Mining System 1994		978	Rd: D on an efficient and reliable hydraulic mining system in which manganese modules are obligened by a towed vehicle for commercial scale mining to help ensure a stable supply of non-ferrous mineral resources.	The midlerm assessment on detailed design and experiment development of fundamental componenta. Manufacturing of underwater particip, underwater cable, air compressor and collector
Automated Sewing System	luiamated Sewing 1982- 894 lystem 1990		R&D on an automated industrial sewing system, involving processes such as preparation, making - up and finishing, to cope with rapid changes in the diametric apparet market.	Fundamental technologies essential to the automated sewing system were developed, and experimental machines were manufactured and opersited. Software and hardware of the demonstration system were designed.
Advanced Robox [983 2.483 Technology [990 [2.266]		2.483 [2.205]	R&D on advanced tobes technology for systems to support people working under difficult or dangerous conditions.	Basic key technologies such as locomotion, merupulation, and sensor technologies have been developed for three types of robots. For nuclear power plants, for undersa, and for oil plant fure for oil refuseries. Using these technologies, robusts for feasibility study were designed.
New Water Treatment System	Vew Water Treatment 1985- 1.519 System 1990 [1,289]		R&D on a new wastewater treatment system using a high-concentration bioreactor and separation membrane, for water reuse and energy recovery (e.g. mechane gas from anaerobic bioprocess).	Bench-scale experiments made it possible to design a bioreactor-process for high-rate methane fermentation and a membrane-module with a persistent flux.
Interoperable Database System	1985- 1991	1.708 (1,255)	R&D on technology for interoperable information systems with such features as distributed databases and multi-media technology, to form an infrastructure for the "information-orients" boost v	Some of the Implementation Specifications required for the Infrastructure to assure the uneroperability among networked heterogeneous computers have been developed in conformity with OSL.
Advanced M. Inal 1986 2.9 Processing and 1993 [2.3 Machining System		2.935 [2.766]	R&D on advanced surface processing using excimer laser beam and/or ion beam, and on ultraprecision mechanical processing, for advanced industries such as emergy, precision machining and electronics.	Elementary techniques for the high power, high repetition ratio, long life excimer laser and the ultra fine ion beam and the deep ion injection were developed
Fine Chemicals from 19 Marine Organisms 19		1.186 [396]	R&D on biotechnological production of fine chemicals such as pigmenta, dyestaffs, mosturizing materials, and coucies materials for underweater sinuctures.	Preliminary investigations on useful materials from marine organisms and on utilization technology of biofunction
Super Hyper Source Transport Propulsion System	1989	1.62 191 1,430	R&D on a combined-cycle engine which will combine the "rampet" and "high performance turboret", and provide high reliability and efficiency at both the subsonic and the hyper- sonic level.	Conceptual study for combined cycle engine and its components for SST/HST
Linderground Space Development Technology	1989.	[371]	R&D on underground space development technology are as follows: (1) geological survey and Evaluation technology (2) dome construction technology (3) environment conditioning and hazard prevenuion technology (4) pilot dome construction	Preliminary investigations on key technologies such as geo-tomography, constructing machines and fire hazard prevention.
Advanced Chemical Processing Technolog	y 1990-	29	R&D on advanced chemical processing technology for producing the functional materials such as functionally gradient materials, pure metals, polymers with fine augment of molecules.	(Project launched in FY1990).
Human Sensory 1990- Measurement Application Technology		50	R&D on technologies for measurements of psychological and physiological effects, method of quanutarize analysis and evaluation of computated human sensation, and sensory evaluation simulator	(Project launched in FY1990)
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Upper columns in parentheses represent general accounts and lower thes special accounts. Those not in parentheses represent general accounts only. NOTE

Completed National Research and Development Projects (Unit: million yen)

- Super High Performance Electronic Computer (1966-1981, 10,100)
- New Method of Producing Olefin (1967-1972, 1,200)
- Sea water Desalination and By-product Recovery (1969-1976, 6,700)
- Comprehe-isive Automobile Control Technology (1973-1978, 7,300)
- 9 Direct Steelmaking Process using High-temperature Reducing Oas (1973-1980, 13,700)
- 10. Olefin Production from Heavy Oil (1975-1981, 13,800)
- Resource Recovery Technology (1973-1982, 12.600)
 Subsea Chi Production System (1978-1984, 18,200)
- 16 C, Chemical Technology (1980-1986, 10,500)
- 18 High Speed Computing System for Scientific and Technological Uses (1981-1989, 17,500)
- 2. Desulfunzation process (1966-1971, 2,700) Remote controlled Undersea Oil Drilling Rig (1970-1975, 4,500)
- 6. Electric Car (1971-1976, 5,700)
- 8. Pattern Information Processing System (1971-1980, 21 900)
- 11. Jei Aircraft Engines (1971-1981, 19,700)
- 13 Flexible Manufacturing System Complex using Laser (1977-1984 13.50)
- 15 Opucal Measurement and Control System (1979-1985, 15,700)
- 17. Observation System for Earlt Resources Satellite 1 (1984-1988, 10.900).

R&D ON NEW ENERGY TECHNOLOGYA A 3298

-THE SUNSHNE PROJECT-

The Sunshine Project was started in July 1974 to secure a stable energy supply for Japan which has a vubnerable energy structure. High priorities are given to the development of the following five projects.

Budger

The Agency promoting the Sunshine Project is also active in international cooperation through IEA and other international organizations.

(1) Solar Energy

(2) Geothermal Energy

(3) Coal Energy

(4) Hydrogen Energy

(5) Comprehensive Research

(Unit million yes)

	Project Name	FY1990	Outline of Project	R&D Results
1	Solar Energy	7,324 190 (7,134)	 Research and development of high- performance and low-cost solar-photovoltaic- conversion technology which we hope will be widely used by early 21st century. Development of the application of solar- thermal-application systems for industrial processes which require sophisticated thermal controls. 	 The price of solar cell has been reduced from 20,000-30,000 yen/W peak to 720 yen/W peak. The cost of solar photovoltaic-conversion system has been reduced from 2,000 yen /k Wh to 200 yen/k Wh. Technology of fixed-temperature-stock-room system (-5°C) driven by solar-thermal energy has been achieved.
2	Geothermal Energy	5,378 [133 5,245]	 Nation-wide geothermal exploration survey assessing geothermal potential in Japan. Research to confirm the effectiveness of exploration techniques for deep geothermal resources. Development of binary cycle generation plant and hot dry rock generation system. 	 Completion of geothermal potential map in Japan. Geothermal potential assessment and development of optimum exploration methods for high potential areas. Development of high-precision magneto-telluric method. Design and test operation studies of Downhole Pump for binary cycle power generation. Production of geothermal researvoirs by fracturing techniques developed for hot dry rock power generation system.
3	Coal Energy	24,901 (266 24,635)	 Coal Liquefaction Technology Development of original liquefaction processes for both bituminous and brown coal. Coal-based Hydrogen Production Technology Development of mass-production technology for low-cost clean hydrogen energy. Integrated Coal Gasification Combined Cycle Power Generation Technology (IGCC; Sponsored by ANRE) Development of the technology of IGCC which is more efficient and have less environmental impact than conventional coal-fired power generation. 	 Design of 150 Vd pilot plant for liquefaction of bituminous coal. Operation of 50 Vd pilot plant for liquefaction of brown coal. Construction of 20 Vd pilot plant for coal-based hydrogen production. Construction of 200 Vd IGCC pilot plant with entrained bed reactor.
4	Hydrogen Energy	108	Development of technologies on hydrogen production, storage and transportation, use and safety Development of technologies on producing highly efficiently, and on transportation and storage using metal hydrides Development of hydrogen batteries and hydrogen- fueled engines	The pilot plant of alkaline-water electrolyzer with a 20N m ³ /h capacity was successfully operated for a long period at the highest efficiency in the world.
W.	Comprehensive Research	1.784 [145 1,639]	 Basic studies of other new energy technologies, such as wind energy, ocean energy, bio energy, but excluding four areas (solar, geothermal, coal and hydrogen) are proceeding. Development of a high-officiency membrane complex methane production unit. 	The pilot plant of 100kW-class wind turbine generator system was successfully operated for a long period.
6	International Cooperation	61	 International cooperation through IEA. Bilateral cooperation with Australia, etc. 	



NOTE: Upper columns in parentheses represent general accounts and lower ones special accounts. Those not in parentheses represent general accounts only.

FY1990 BUDGET AND PERSONNEL IN AIST

1 Budget



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Note: Ordinary R&D : Personnel expenditures Designated R&D : Research expenditures and Moonlight Plojec Technologies for futur Special R&D : Expenditures

Personnel expenditures and ordinary research expenditures of AIST laboratories. Research expenditures incurred by research laboratories through work connected with the Large-Scale Project, the Sunshine and Moonlight P ojects, the R&D Project on Medical and Welfare Equipment Technology, the R&D Project on Basic Technologies for future Industries, and the Regional Large-Scale Project.

Technologies for future Industries, and the Regional Large-Scale Project. Expenditures incurred through Special Research, Expansion of Laboratory Facilities, Operation of Geological Research Vessel, Nuclear Research, R&D Promotion for Small Industries, Research Related to Prevention of Environmental Pollution Expenditures in operating joint facilities at Tsukuba.

Tsuk oba-related expenditure Budget for individual projects

The total budget for the large-Scale Project, Sunshine and Moonlight Projects and R&D Project on Basic "schoology for Future Industries, minus the budget for Designated R&D. (Designated R&D is also omitted from the Total Budget for AIST)

2 Budget and Personnel for Government Laboratories

	Budget	Personnel	Researchers	Administrators
	(million yen)	315	1	314
Assessment of Industrial Science and Technology (Headquarters)	75,799	010	128	90
Agency of Dictational Deviatory of Metrology	2,186	210	217	59
Automat Research Laboratory	3,289	2/0	276	74
Mechanical Engineering Energineering	3,936	350		18
National Chemical Laboratory for Diducely	1,122	90	100	33
Fermentation Research Institute	1,532	125	10.	+
Research Institute for Polymers and Textiles	4.574	353	237	110
Geological Survey of Japan	9.217	686	554	132
Electrotechnical Laboratory	1 413	125	102	23
Industrial Products Research Institute	1010	319	245	74
National Research Institute for Pollution and Resources	1.224	96	73	23
Government Industrial Development Laboratory, Hokkaido	650	53	39	14
Covernment Industrial Research Institute, Tohoku	239	241	186	55
Government Industrial Rescaron Institute, Nagoya	2.500	210	168	51
Covernment Industrial Research Institute, Osaka	2.048	52	40	12
Government Industrial Research Institute, Chugoku	720	45	35	10
Covernment Industrial Research Institute, Shikoku	513	01	71	20
Covening Industrial Research Institute, Kvushu	969			and the survey of the survey o
Government moust at reason and the	41,185			
Common Expenditures	5.525			1.100
Other Laboratories	112,509	3,654	2,546	1.100

FYTOM BUS STREETSIAMORAITING

RESEARCH AND DEVELOPMENT PROJECT ON BASIC TECHNOLOGIES FOR FUTURE INDUSTRIES

This project is aimed at the development of innovative basic technologies essential for eat-blishing new industries

The five fields covers , are superconductivity, new materials, biotechnology, new eir-gonics devices and software. The following fourieen special stegories, all of which have theoretically

or experimentally shown potential for application in new industrial technologies, have been selected. Research and development in these categories are conducted until the materials involved are ready for practical application.

(Unit: million yes)

Project Name	Period (FY)	Budges for FY1990	Outline of Project	R&D Results
 Superconductivity Super conducting Materials and Devices 	1968-	2.347 910 [1,437]	Development of new superconducting materials, processing technologies for applying superconducting materials to electric power equipments, e.g. magnets and wirds, and technologies for fabricating superconducting electronic devices.	Effective persning centers were introdued in oxide superconductors. New method to durectly observe magnetic fus distribution in oxide superconductors was developed. Y-type proximity effect superconductor device was developed.
2) New Materials 1) High-Performance	1981- 1992	1313	Development of high - trength certainics at extremely high temperatures to be used as materials for gas turbure components.	Si, N_a and SiC commiss with high reliability which can mand high temperature (~1200°C) were developed as materials for gas turbure components.
2) Synthetic Membranes for New Separation	1981- 1990	192	Development of synthesic membranes for new separation technology, to separate and refine freely mixed games or liquid restures by utilizing differences in physical properties.	Symbetic membranes which can efficiently separate ethanol/water mixed solutions, CO/N ₂ mixed gases, and optical isomers of amino acid were developed.
3) Synthesic Mesais	1981- 1990	162	Development of synthetic metals and polymerse materials with electroconductive properties of metals.	Layered symbotic graphite with the highest electroconductivity of 92103 S/cm was found. The first example of soluble conclucting polymer was demon strated by the ustrockaution of a long alkyl chain on polythiophene rangs.
4) High-Performance Plastics	1981- 1990	126	Development of high-performance plastics and polymenc materials with mechanical properties of metal.	Polymers with high flexarel moduli, and molding- techniques for obtaining high performance liquid-crystalline polymers and molecular composites have been developed.
5) High-Performance Materials for Severe Environments	1989- 1996	[1,001 [153 838]	Development of carbon/carbon composities, antermetallic compounds, and fiber reinforced intermetallic compounds which can be used to develop a space plane and SST/MST.	SiC fibes, modified by the electron beam method, was-developed to stand high temperature (~1500 ^M C).
6) Photoscuve Maurials	1985- 1993	471 [76 396]	Development of photoactive materials, which characteristically exhibit a revensible change is the structure or arrangement of spokecules in response to a light stimulus.	Photochromic LB films for multiplex recording and photochemical hole burning materials which work at liquid Ng temperature have been developed.
7) Non-linear Photonics Materials	1989- 1998	540	Development of photonics materials which exhibit high nonlinear optical susceptibilities and short response times for application of optical information systems.	Consugned polymers and CuCl dispersed glasses having the highest susceptibility reported so far have been developed.
(3) Biolechnology 1) Uulization of Recombinant DNA	1981- 1990	152	Investigation into the use of recombanant DNA technology for the development of the w microorganisms for practical use in industry.	Noble host-vector systems of yeasts were developed. Various enzymes and bioactive substances have been efficiently produced by the improvement of host-vector systems of industrial microsorganisms.
2) Molecular Assemblies for Functional Protein System	1980. 1998	321	Development of molecular assemblies of functional proteins for reactions with sophisticated functions roch as production and conversion of complexed biomatenals coupled with selective uninport and recognition.	Photosynthesis protein complexes were extracted from photosynthesis bacteris and absorbed to the surface membrane with omentation of the molecule. Phycobaliprotein was isolated from thermophilic cyanobacteris for the first time and evaluated.
 (4) New Electron Devices 1) Superlattices Devices 	4) New Electron Devices 1981- 1) Superitatices 1990 Devices		Development of superlattices electron devices with extremely fine atructure tailored to atomic scale for utilizing new electronic effects.	Multifunctional high speed devices which utilize resonant tunneling, high mobility, and ballistics of electrons in superlatices were fabricated by molecular beam epilaxy (MBE) and metal originic chemical vapor deposition methods.
2) Three Dimensional	1981-1990	. 30	Developmens of ICs characterized by a three- dimensional arrangemons of active elements made of semiconductor layers.	Prototype of 3D devices with optical detector, A/D logic and memory circuits were fabricated by SOI formation are through-hole wiring technologies.
3) Bio-electronic Devices	1986	28	Development of bio-electronic devices for future computer elements by using biological information processing functions based on plasticity, molecular recognition and self organization in a cell level.	A new optical method for detecting neural activities in the brain was developed. A photo-electro device which had wery high quantus efficiency was also developed by paling up organic membrane.
(5) software 1) New Models for Software	(5) software 1990- 1) New Models for 53 1997 Development of annovative models for flexible software architecture so that software can func- according to the surrounding mulation.		3 Development of innovative models for flexible software architecture so that software can function according to the surrounding situation.	(Project lautocheel in FY1990)

NOTE: Upper columns in parentheses represent general accounts and lower ones special accounts. Those not in parentheses represent general accounts only. Completed Research and Development Projects on Basic Technologies for Future Industries (Unit: million yen) Fortifield ICs for Extreme Conditions (1981-1985, 1,315) Advanced Alloys with Controlled Crystalline (1981-1988, 3,903) Advanced composite Materials (1981-1988, 4,649) Bioreactor (1981-1988, 2,978) Large Scale Cell Cultivation (1981-1989, 3,362)

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FY1990 APPROPRIATIONS RELATED TO INDUSTRIAL TECHNOLOGY IN MITI

(Utat: one hundred million wa

ltern	Fiscal 1989 Appro- priacions	Fiscal 1990 Appro- priati ais	Increase over the Previous Year	Noke
Request for R&D-related appropriations General accounts Special accounts Industrial investment accounts	2,336 694 1,360 282	2,498 685 1,531 282	162 59 171 0	Growth rate against previous year 6.9% Δ1.3% 12.6%
Major Projecta [Positive contribution to international society] * Promotion of Human Frontier Science Program (HPSP) * Promotion of International R&D Center * Development of global environment technology	9	13	4	International HFSP Organization (Tentative name for the fund) To be located in NEDO.
 R&D cooperation with developing countries Support to international joint R&D 	1 16 4	2 21 4	1 5 0	Specific R&D at national experimental research laboratories
[Further promotion of technological development projects] (ALST-related) * Preparation project of research foundation by Integrated of New Energy & Industrial Technology Development	22	22	0	Participation from industrial investment accounts
 Organization R&D expenses of national experimental research institutes (Special R&D, Government-private sector joint R&D, important area technology R&D, operation expenses of laboratorea atc.) 	147	145	2	including development of global environment technology
* R&D project on basic technologies for future industries	68 (21)	75 (40)	6 (19)	Applied technologies for new models for soft architecture
 Large scale project Development of medical and welfare equipment 	139 (92) 7	(102) 7	2 (10) Δ0	Advanced chemical processing, Human sensory measurement application Non-invasive continuous blood gluease monitoring system, digital hearing aids, health
* R&D on new energy technology	(259)	275	4 (6)	monitoring system for the elderly.
R&D on energy conservation technology	107 (101)	(110)	8 (9)	
 Interna Anal joint research on aircrafts (YXX, V2500) Unmanned space experiment system (Free flier) 	42 45 (40)	39 53 (48)	(5)	
(Data processing-related) * R&D for Sun generation computer	65 (28)	70	5	
(Technological development relating to superconductivity)	44 (29)	52 (43)	8 (14)	Moonlight, etc.
* Promotion of development for new industrialized housings	(1)	10 (9)	9 (8)	Development of new industrialized housings to realize the needs of those living
The Service of the Japan Key Technology Center Promotion of standardization	260 (260) 9	260 (260) 10	(0) 1	Parucipation of financing from industrial investment accounts

Note Figures in () belong to special accounts, which are part of the upper figures

FY1990 Appropriations Related to Science and Technology in Japanese Government (Summary)

(Unit multic) yer						(run withes hev)
Agency/Mosistry	General accounts	Growth rate against previous year (%)	Special accounts	Growth rate against previous year (%)	Total	Growth rate against previous year (%)
Ministry of Education	204,580	4.0	689.721	49	894 301	4.7
Science and Technology agency	369,838	4.]	124.937	12.4	494 775	6.0
Ministry of International Trade and Industry	68,492	Δ1.3	181,340	10.4	249 832	6.0
Defense Agency	104.268	12.0	the same of the state of the st	NAME OF TAXABLE PARTY AND ADDRESS OF TAXABLE PARTY.	104 268	12.0
Mitustry of Agriculture, Forestry and Fishery	66,707	3.2	3.300	12.4	70.007	2.9
Minustry of Health and Welfare	40,150	7.0	11.092	23	51 242	5.9
Minustry of Posts and Telecommunications	4.657	47	26 543	0.5	11100	11
Ministry of Transport	16.371	6.4	1.039	13.3	17 410	6.8
Environment Affairs	9.217	16.9	and a state of the	and the second	9.217	16.9
Minustry of Foreign Affairs	7.095	10.7	and a second second second second	A REAL PROPERTY AND ADDRESS OF TAXABLE PARTY.	7.095	10.7
Others	12.006	5.3	4.250	A10	16.257	10
Total	903 381	4 7	1.016.222	5.6	1919 600	5 7





WINGAMIZALION OF AISI

Ministry of International Trade and Industry (MITT)

- Industrial Technology Council

Agency of Industrial Science and Technology (AIST)

Japanese Industrial Standard Committee

Headquarters

General Coordination Department

Standards Department

- · General Coordination Division
- · Personnel Affairs Division · Sudget and Accounts Division
- · Deputy Director-General for
- Technology Affairs (Research and International Affairs)
- · Research Admanistration Division
- · Director for Global Environment
- Research System Planning Division
 Director for Planning of Regional
- Technology International R&D Cooperation
- Division
- Deputy Director-General for Technology Affairs (Policy Planning)
- Technology Policy Planning Division
 Director for Life science Application
- Technology
- Technology Promotion Division
 Technology Research and Information
- Division International Technology Research and laformation Office
- · Deputy Director-General for Technology Affairs

- (Technology Development) Director for Planning of Basic Technology for Future Industries Director for Development of Basic
- Technology for Future Industries Director for General Coordination of
- Development Programs (Large-scale industrial technology
- development) · Director for Development Program
- (Large-scale industrial technology development)
- * Director for General Coordination of Development of Programa
- (New energy technology development) Director for Development Program
- (New energy technology development) Director for General Coordination of
- Development Programs (Energy conservation technology development)
- · Director for Development Program (Energy conservation technology development)
- . Tsukube Administration Office

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The Human Frontier Science Program
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- · Standards Division
- · Material Standards Division
- Textile and Chemical Standards. Division
- · Machinery Standards Division
- · Electrical Standards Division
- . Information Standards Division
- · Director of International
- Standard lation Affairs

Research Laboratories and Institutes

· National Research Laboraotry of Metrology

- · Mechanical Engineering Laboratory
- National Chemical Laboratory for Industry
- · Fermentation Research Institute
- Research Institute for Polymers and Textiles
- · Geological Survey of Japan
- * Electrotechnical Laboratory
- · Industrial Products Research Institute
- · National Research Institute for Pollution and Resources
- · Government Industrial Development Laboratory, Hokkaido
- · Government Industrial Research Institute, Tohoky
- · Government Industrial Research Institute, Nagoya
- · Government Industrial Research Institute, Osaka
- · Government Industrial Research Institute, Chugoku
- · Government Industrial Research Institute, Shikoku
- · Government Industrial Research Institute, Kyushu

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HISTORY OF AIST

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OVERVIEW OF THE AGENCY OF INDUSTRIAL SCIENCE AND TECHNOLOGY

R&D in advanced fields of electronics, new materials, biotechnology, and so forth is remarkable contributing to the international society through technical development while taking into account dom surrounding science and technology. Japan will promote Techno-Globalism to stimulate scientific well as distribution and transfer of the results of such activities. Domestically, Japan, which is now o technological powers in the world, will maintain and enhance the bases for building up an affluent. This will enable us to triablish a long-and med⁴ uniterm basis for the economic development of positively to the international society through R&D it is essential for Japan to play a role in promotin areas which will benefit all humankind.

For 1991, the Agency of Industrial Science and Technology will promote R&D in such areas as n and pollution control to tackle environmental problems while making efforts to expand and argume research organizations in developing global environmental technologies, promoting interns environmental technologies, and establishing R&D organizations to study industrial technology rel We also intend to develop overall industrial technology policies, in particular the following.

First of all, for this year, international research cooperation will promote the Human Frontier explain the superior functions of the living body and to search for possible applications. Specifically improve and expand international research exchange by:

- R&D in specific areas through task sharing among the Agency of Industrial Science and Tech advanced countries.
- (2) assistance to R&D development by international joint study teams in explaining physical prope
- (3) invitation of foreign researchers to the research institutes of the Agency of Industrial Science a
- (4) incorporation of the International Exchange Center established in NEDO in 1990 into the Int Research Exchange Center.

Secondly, regarding R&D in basic and advanced fields, the following projects will be further exp Basic Technologies for Future Industry to promote advanced technology development (new mat Scale Project of Industrial Technologies vital in terms of the national economy (effective utilization

R&D for of Medical and Welfare Equipment Technology to contribute to the welfare of society thro Continued promoti it a also planned for the Sunshine Project aimed at developing clean net

energies, etc.) and the N Y n-light Project for generating energy conservation technology to achiev of Japan's energy security.

Technological R&D for "tal Regions will be actively promoted to stimulate economic growth of In line with these R&D focusing on basic areas will continue in 16 research laboratories under and Technology, while furthering mutual exchanges through joint research among industries, gover

Thirdly, in order to encourage R&D in the private sector, R&D founding will be provided Promotion Center, utilizing various tax incentives (Added R&D Expenditure tax deductions in infrastructure technology for R&D). The latter refers to improving the research infrastructure in R&D.

Finally, regarding the industrial standardization system (IIS) which has supported our developm international cooperation through making positive contributions to international standardizati technical cooperation, promotion of standardization in advanced technology fields (information biotechnology).

We hope this pamphlet will help in understanding the policies of the Agency of Industrial Science



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masaru

Dr. Masaru Sugiun Director-General. Agency of Industri Science and Techn

Mandana Anguna I be Government Industrial Roman established in 1967 as a research facility for dewakoping saming and industrial technology for the Toboku region. Initially, is conducted a major project on automics processing for Kuroko ore which is abundant in the Tohoku district and its vicinity. Since then, it has made a names of research contributions ranging from utilization of regional resources to advanced technology areas, all sterming from the desire to develop industries in this region as well as share the responsibilities for national projects. In addition, since 1975, it has taken part in international joint research and development projects with various countries such as Thailand, Indonesia and China.

At the present time, Research is focused on the following three fields:

1. Researches on regional resources and energies

- (1) Separation and refining technique for finechemicals from low utilized biomass (2) Technique for extracting lipids (3) Developing materials for geothermal power plants (4) Recovery of useful metals from geothermal hot water (5) Geothermal data analysis associated with well-stime ation by hydro-fracturing (6) Development of groundwater velocimetry.
- 2. Development and evaluation of new materials (1) Development of nano-composites by intercalation (2) Development of mesoporous materials from synthetic silicate-bearing smectites (3) Functionally gradient material by a seif-propagating high temperature synthesis process (4) Internal inspection system for composite substances (5) Mechanical properties of austempered ducule iron.

Government Industrial Research Institute, Tohoku 4-2-1, Nigatake, Miyagino-ku, Sendai-shi, Miyagi, 983

The Government Industrial Research Institute, Nagoya (GIRIN) was established in 1952 as a national research center to contribute to R&D on advanced regional technology as well as national program. Research activities are conducted in six departments, namely, Mechanical Engineering, Metallurgical Engineering, Chemistry, Radiation Research, Ceramics Science, and Ceramics Technology. It has 241 staffs and 2.6 billion-yen budget in total in fiscal year 1990. One division in Ceramics Technology is located at Seto-city, which is well known for her largest production of pottery and porcelain wares in Japan, to contribute to her regional technology.

Since its establishment, GIRIN has played an important role for R&D on such industrial science and technology as liquid bulge forming, casting and foundry technologies, synthesis of organic fluorine compounds, radiation graft polymerization, solar energy utilization, functional and engineering ceramics, and pottery and porcelain production technologies.

GIRIN has recently focused her research activities more on inorganic material (ceramics) research aiming to Energy (including hard energy technologies for high temperature gas turbine, unclear fission, and large power transportation, soft energy technologies such as passive solar device, and energy conservation technologies) and Space & Aircraft technologies. GIRIN is executing 42 research projects in conjunction with national R&D projects such as Basic Technologies for Future Industries (Engineering Ceramics, High Temperature Ceramic Super Conductor, Metal Based Composite Materials and Inter Metallic Compounds), Sunshine (Passive Solar Device, Photo Catalysis), Moonlight, Large Scale Project (Surface modification of ceramics by beam technology), Special Programs for Mines & Technologies (Bio Ceramics and other Functional

Government Industrial Research Institute, Nagoya 1-1. Hirale-cho, Kita-ku, Nagoya-shi, Aichi, 462

refining or rare metals produced in China (a) Research utilization of natural zeolite in Indonesia.



High Temperature Electrochemical Measurement Apparatus

Sendai 022 (237) 5211	Total personnel Total budget	53 575 (million yen)
	LOTHER CHANDER.	their A sharehouse and a printer or movies should be strengthen a work of

ceramics, Functional Organic Fluorine Compounds) and so forth. It is also conducting 50 basic research programs. Among those, new metals and casting technologies, environmental protection technologies, radiation physics and chemistry, biotechnologies and, pottery and percelain technologies are included.

GIRIN has actively joining to the bilateral international cooperative research programs under AIST scheme in the field of fluorine bionics organic compound, acid rain project, and utilization of indigenous materials in developing countries as well as to multi national cooperative programs under IEA in the fields of ceramics and solar materials.



Ultra-High Pressure Cold Isostatic Press for Ceramics Forming

Nagoya 052 (911) 2111	Total personnel Total budget	241 2,600 (million yen)
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The Government Industrial Research Institute, Osaka was established in 1918. Since then it has produced a number of outstanding research achievements in the exploration and development of new materials including carbon fibers and electrically conductive transparent thin films. Our research institute consists of five research departments and is giving priority to the following research fields:

- Energy-related Materials materials for energy conversion such as for batteries, fuel cells, electrolysis processes, hydrogen energy, and high-temperature ceramics for gas turbines, etc...
- Optical Materials: glasses and thin films for optics, nonlinear optical materials, optical chemical sensors, etc..
- 3) Functional Surface Materials: heterogeneous catalysts, biocompatible materials, atomic-scale designing of graphite intercalation compounds, new functions and theoretical analyses of the interconnection in composite materials, surface modification with ion implantation, etc..

In addition to the above fields of materials research, the following lines of approach are also being put forward:

- Intensification of innovative, fundamental studies to create highly advanced functions of materials.
- Initiation of unexplored approaches to materials through human sensation and feelings.
- Fermentation of basic science concerning the creation and analyses of novel materials through atomic scale techniques and computer calculation and graphics.

Government Industrial Research Institute, Osaka 8-31, Midorigaoka 1-chome, Ikeda-shi, Osaka, 563 Promotion of R&D programs for global environment technology.

Through the above research activities, our institute encourages contacts and cooperation between industrial and university research on a regional, national, and international level.



Ion Implantation Machine

a la	Dkeda 0727 (51) 8351	Total pers	sonnel	219	
563		Total bud	iger	2,560 (mill	ion yen>

The Government Industrial Research Institute, Chugoku was established in 1971 to conduct pollution control studies in the Seto In'and Sea and engineering studies to develop new industrial technology in the Chugoku district. Since then, there have been many noteworthy achievements by the Institute results, including research on dissolution of pollutant out of the bottom sediment in the Seto Inland Sea, development of a fresh surface characterizing microscope using exo-electron, and materials evaluation in severe environments.

Two research department, the Marine Science and Technology Department and the Industrial System Department, are affiliated with the Institute. The first has four research divisions, and studies ocean engineering using the largest hydraulic model of the Seto inland Sea in the world, shown in the picture, as well as physical, chemical and biological oceanographies. The second has three research divisions, and studies machining technology, the surface science of new materials and computer science in factory automation.

The institute also conducts marine biology studies under a major national R&D program, a materials study of hydrogen energy and ocean thermal energy conversion (OTEC) under national R&D projects focusing on new energy, computer image processing studies for developing specific regional technology, and international joint research with Indonesia in the area of corrosion.



Large-scale Hydraulic Model of the Seto Inland Sea

Government Industrial Research Institute, Chugoku	Kure 0823 (72) 1111	Total personnel	52
2-2. Hirosuehiro 2-chome, Kure-shi, Hiroshima, 737-01		Total budget	726 (million yen)





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The Government Industrial Research Institute, Shikoku was established in 1967 as a R&D center for developing mining and industries in the Shikoku region, taking advantage of its mild climate and location near the sea and rich forest resources. Its R&D centers upon pulp and paper technology and in developing marine resources, it is primarily concerned with extraction and uses of minor elements dissolved in sea water, and underwater welding and cutting.

Balancing its regional and national interests, the institute has emphasized research in marine resources, functional resources and mechatronics. Furthermore, our institute is the leader in the Shikoku region for research and technologies.

The major research area of the insutute are as follows:

1. Marine resources

 Manufacturing process of high-functional chemicals from sealife.
 Developing excellent absorbents for uranium and lithium.
 Developing functional sheets from acidic polysaccharides, the main constituents of seaweeds.
 Manufacturing degradation-controlled sheet.
 Development of magnesium pyroborate whisker.

2. Local technologies

(1) R&D on re-utilization system technology of composite materials. (2) Control of a flexible long arm and development of a small active mass damper. (3) Swing and vibration control of a grane. (4) Sophisticated surface processing with laser beam and ion beam.

International R&D cooperation projects and fundamental research

(1) Research on industrization of thermomechanical pulping of oil paim by-products. (2) International research cooperation on recovery of valuable brine resources. (3) Molecular mechanism of interactive recognization between cell surface and polysacchandes.



Bio-degradeble Polymer Film

Government Industrial Research Insutute, Shikoku	Takamatsu 0878 (67) 3511	Total personnel	45
3.3. Hananomiya-cho 2 chome. Takamatsu-shi. Kagawa, 761		Total budget	513 (million yen)

The Government Industrial Research Institute, Kyusyu was established in 1964 to contribute to developing mining and industries in Kyusyu.

The institute has conducted 17 special research and 35 general research projects in the following major fields:

- Production and processing technologies for new materials: (1) Study of ductile ceramics at high temperatures, (2) Development of intelligent ceramic composite materials, (3) Research on fabrication and thermal characterization of heat resistant carbon ceramic composite materials. (4) Machining technology in ceramics and cutting tool application. (5) Development of ceramic cutting tools for steel. (6) Development of highperformance engineering carbons. (7) Development of advanced carbon/carbon composites with oxidation resistance. (8) Technology for improving properties of materials by the powder forming method, (9) Development of particle dispersed composite metals by high pressure solidification method. (10) Research on metallic materials using quantitative stereology. (11) Study of production of multifunctional microspheres. (12) Study of spectroscopic characterization of ceramics.
- Advanced technology for utilizing natural resources; (1) R&D utilizing lime and lime-based compounds in advanced materials, (2) Processing and evaluation of inorganic polymer having layer structure. (3) Production of porous ceramic materials from rice husks, (4) Advanced utilization of volcanic glass, (5) Production and utilization of molecular sieves from coal. (6) Research on concentrating a trace amount of gallium. (7) Refining process of fine parts of weathered granite.

3. Energy and pollution control technologies: (1) Study of the behavior of coal and solvent mixtures in the initial stages of coal liquefaction, (2) Hot-gas corrosion of ceramics for gas turbin blades. (3) Development of new muffler adaptable with controlled resonators. (4) Research on the advanced biologica treatment of organic waste water. (5) Liquefaction of coal arx extraction of liquefied products under the condition of superentical state.



Friction and Wear Testing Apparatus

Government Industrial Research Institute, Kyusyu Syuku-machi, Tosu-shi, Saga, 841 Total personnel Total budget

91 946 (million yen)

The National Chemical Laboratory for Industry (NCLD) was established in 1900 for promoting the chemical industry in Japan. The laboratory conducts numerous research projects in four sreas: 1. development of new substances and highly functional materiels. 2 bio and biomimetic chemistry; 3. conversion and conservation (echnologies for energy and resources; and 4. standardization and safety technologies. Many strategies aiming the first area are introduced in chemical reactions and processes, for examples, ultrahigh temperature plasma, ultra-high pressure, laser beams, computer-aided molecular design systems. For the elucidation and application of biological functions, the laboratory has developed icchnologies for genetic engineering, cell membranes, artificial and super enzymes, and artificial photosynthesis. In effective utilization of energy and natural resources, extensive studies are being made on catalysis technology, coal liquefaction, heat storage using chemical reactions, fuel cells, superconductors, membrane technology, and biomass utilization. Regarding the final area, intensive studies are also made on standardization of chemicals, as well as a means of controlling environmental pollution, eliminating industrial hazards, and preventing explosions of gases and explosives. The main research projects of the laboratory are as follows:

(1) Ultra-high temperature: generation, measurement and utilization (1) Ultra-high temperature: generation, measurement and utilization (2) Solid state polymerization under ultra-high pressure (3) Laser regulated chemical reactions (4) Development of organo-silicon compounds (5) Research and development of superconducting materials and devices (6) Research on the analytical and evaluational technology for high-quality functional materials by beam technology (7) Advanced material processing and machiness systems (8) Synthetic membranes for new separation technology (9) and Research for morphogenesis and expression of genetic information (10) Development of proteins with new function (11) Liquefaction of ceal (12) Super heat pump energy accumulation system (13) Reference materials for calibration of analytical instruments (14) Estimation and prevention of explosion hazards of special material gasses.



Application of Excimer Lasers to Chemical Syntheses

and the second se		Total personnel	349
in the second Laboratory for Industry	Tsukuba Gakuen 0298 (54) 4451	TOTAL PROTOCOLOGICA	4 200 (million ven)
National Chemical Laboratory for Distance	Personal Planning Office	Total budget	a, 300 (nuturon yen)
Higashi Lichome, Tsukuba-shi, Ibaraki, 305	Kesearen Flandung Gritter	free lands and provide an a second second as and	

The Fermentation Research Institute (FRI) was established in 1940 with the objective of contributing to the development of industries involved with microorganisms. The Institute conducts a broad range of activities including the development of a variety of enzymes, techniques for biologically treating industrial waste water, and improved industrial processes related to microorganisms. Recent years have brought advances in such areas as recombinant DNA technology, bioreactors using immobilized enzymes and coenzymes, cell growth and gene expression control in cultured animal and plant cells, hydrogen producing microorganisms, the production of substances regulating cell function, the development of new enzymes and the utilization of thus far unused resources. As the authorized depository for patent microorganisms in Japan, the Institute also handles the deposition and distribution of domestic and foreign strauns of microorganisms.

Major research performed at the Institute centers on the areas below

Designated Research: Protein Molecular Assembly Technology. R&D on a New Water Treatment System, Fundamental Technology for Utilization of Marine Organisms, Basic Technology for Utilization of Useful Biological Function, Research on Energy Conversion by Photosynthetic Microorganisms, Algal Potential for Carbon Dioxide Fixation, Molecular and Cellular Biological Study of Morphogenesis in vitro, Regulation of Plant Gene Expression, Molecular Mechanisms for Regulatory Protein Functions, Research on Photosynthetic Molecular Assembly, Molecular Recognition and Response of Smooth Muscle Cells, Expression Regulation of an Intestine Contractor Peptide Gene, and Theoretical and Experimental Investigation on Ribozymes.

Special Research: New Transferring Enzymes and their Functions, Construction of Bioreactor, Biocatalyst for Oxidation in Micro aqueous System, Application of Recombinant DNA Technology to Hydrocarbon Utilizing Soil Pseudomonads, Development of Plant Genetic Engineering, Development of Substances Regulating Cell Function, Regulatory Mechanisms of Cell Proliferation on Eukaryotic Microorganisms, Studies on the Thermobiology, Mechanisms for Release of Methane into the Atmosphere by Microorganisms, Treatment of Offensive Odors Using Microorganisms, Flocculant Produced by Microorganisms, and Studies on the Role of Calcium Ion in Signal Transduction in Animal Cells.

Special Coordination Funds for Promoting Science and Technology: Basic Study on Safety of Genetical Engineering Techniques in Open System, Structure Function Relationship of RNA Molecules and Application of Synthetic Ribozymes, Development of Experimental System for the Analysis of the Response Mechanism of Plant Cells, and Studies on Angiogenic Factors.



Various Kinds of Biodegradable Plastics



The Reserved EastWhete for Perysters and Textiles was glassby established in 1918 as the SED Laboratory and in 1937 came the Textile Research Institute covering the entire area of tile technology. After the War, polymer science and technology is integrated into this research field, and the present name was opted in 1969. The present organization was founded in 1988 mprised of 4 research departments; Polymer Chemistry Dept., o-engineering Dept., Material Physics Dept. and Material Design i Engineering Dept. In recent years, research activity has been used on upgrading polymer materials, the synthesis of new inctional polymers, bio-function utilizing technology and tovative technology for textile industries. And, future emphasis to be placed on the development of functional materials based on ilecular-level science. Main research items are as follows:

Synthesis and structuring of polymeric materials (1) Synthesis of ordered polymers (2) Organization of polymer surfaces (3) Synthesis of biologically functional polymers

Functional Molecules (1) Photo-reactive polymers (2) Energy transforming polymers (3) Conductive polymers (4) Polymeric materials for specific separation

Bio-functional materials (1) Structure analysis and molecular design of biopolymers (2) Elocompatible materials and pharmacologically active polymers (3) Biosensor and biodevice (4) Biomimetic materials

Composite and high performance materials (1) Highly durable materials under extreme conditions (2) Light-weight strong polymeric materials (3) Polymer alloys

- 5 Innovative areasing to be an an an analysic of the sector of the se
- Measurements and analysis of materials (1) Characterization and evaluation of polymers (2) Nondestructive evaluation sechniques for polymeric meterials (3) Weathering of polymeric materials.



Plasma Reactor

	the second se		to an end of the second s
esearch Institute for Polymers and Textiles	Tsukuba Gakuen 0298 (54) 6229	Total personnel	124 1.607 (million mm)
-4, Higashi 1-chome, Tsukuba-shi, Ibaraki, 305	Senior Officer for Research Flamming	Total bubger	1,007 (Renton yes)

Established in 1882, the Geological Survey of Japan is the ily national research institute in the country concerned with the stematic investigation of geology and mineral resources. It is sponsible for geological sheet mapping and for research on ology and various kinds of resources (metallic and non-metallic inerals, fuel, geothermal energy and ground water) in the Japanese chipelago and adjoining offshore areas. Its work has contributed bstantially to environmental conservation and to mitigating mage from geological hazards such as earthquakes, volcanic uptions and landslides. The Survey also takes an active part in forts. Experts on geology and mineral resources are sent overseas d foreign trainees are admitted to training course in the Survey. addition, the Survey provides technical guidance to other encies, local governments and the general public. The results of work are published in the form of various scales of geological id thematic maps, bulletins and special publications. Major search programs in each field are as follows:

Field of geothermal resources (1) Confirmation study of the effectiveness of prospecting techniques for deep geothermal resources (2) Basic study on nationwide and regional geothermal assessment.

Field of utilization and development of resources (1) Study on metal concentration mechanism in the hydrothermal system. (2) Three-dimentional modeling for fuel resources assessment, etc. Field of disaster prediction and environmental research (1) The geological study of earthquakes. (2) Geological, geochemical and geophysical study of active volcances (3) Long range prediction model for changes in the shallow water environment to enable optimum industrial development use.

- Field of geological study for atomic energy utilization (1) Geological study of deep underground disposal of high-level radioactive waste (2) Geochronological study on fault activity, etc.
- Field of international cooperation (1) Mechanism of methane discharge into atmosphere, etc.



Taking Samples of Sea Bottom Sediment by the Geological Survey Vessel "Hakurei-Maru"

Geological Survey of Japan	Research Planning Office 0298 (54) 3572	Total personnel	349
1.3. Higashi 1-chome, Tsykuba-shi, Ibaraki, 305		Total budget	4,300 (million yen)





The Electrotechnical Patient IN MES AN OI as a mating laboratory for electrical hamalantical Communications. After several restor organizations charges since then, including the separation of what is now the Electrical Communication Laboratories, NTT, in 1948, the ETL now stands as the largest national research sustitute in Japan. For promoting future industrial science and technology, the ETL is responsible for conducting advanced research and development in electronics. standards and measurements, energy, and information and computer technologies. A list of ETL's notable schievements begins with the wireless telegraph, developed as early as 1896, and includes, just to name a few, Japan's first transistorized computer --- the Mark IV (1959); the Kondo effect (1964), which later earned the London Award for Dr. Kondo, ETL Advisory Fellow; the first genuinely data-driven computer SIGMA-1 (1987); the discovery of a new type of oxide superconductor and the development of Josephson Computer ETL-JC1 (1989, see picture); and the record high-power eacimer laser ASHURA (1989).

The ETL consists of 14 research divisions located in Tsukuba Science City and one research center in the Osaka area. Within 58 sections some 550 researchers, including approximately 250 Ph.D's are now actively working in the vast new frontiers of science and technology. The major research topics are: (1) Electronics fundamentals: physical studies on superconductivity, dynamics of elementary excitations, etc., development of new superconductors, opto-electronic materials, and amorphous semiconductors. VLSI technologies based on superlattice and three dimensional structures, advanced microfabrication technologies, supermolecular technology utilizing organic molecular assemblies, and biochemical and physiological studies on information processing in living organisms: (2) Standards and measurements: establishment and supply of national standards of electricity, photometry, acousties, and ionizing Tadiation and factoration, and an an anti-factor in the set of the

The ETL, keenly aware of the increasing importance of technical exchanges both with the private sector and academia, is also actively participating in a wide range of cooperative research efforts.

"Josephson Computer ETL-JCI"



	Teukuba Cakuen 0298 (54) 5006	Total personne!	686
Electrotechnical Laboratory	Description Office	Total budget	9,280 (million yen)
1-4. Umezono 1-chome, Tsukuba-shi, Ibaraki, 305	Research Flanning Office		Contract of the state of the second state of t

The Industrial Products Research Institute (IPRI), established in 1928, specializes in the fields of improving the quality of life. Fundamental researches in this field are (1) biomimetic chemistry; (2) materials evaluation technology; (3) biometrics and sensor technology; (4) psychometrics and cognitive science. Those researches have been applied to development of materials and apparatuses for medical use, equipment related to human health and welfare, to design and evaluation of housing systems, and also to evaluation of consumer goods, by combining material and human engineering. Since we have various specialists in physics, chemistry, mechanical engineering, electrical engineering and electronics, information science, psychology. physiology, forestry, industrial design, and so on, IPRI is able to synthetically and systematically solve interdisplinary problems. which maight be difficult for an institute engaged in one specific field to solve. The current research topics are listed below.

 Research related to materials and apparatuses for medical use, human health and welfare; (1) Biometric transduction of sensory information, (2) Cell compatible biomaterials, (3) Three dimensional display for the blind, (4) Non-invasive measurement of functional decreases in humans, and etc.

Servi .

- Research related to housing systems; (1) Fundamental system technology for emergency in living space, (2) Psychological and physiological measurement of the influence of low frequency noise on the body, and etc.
- Research related to consumer goods; (1) Measurement of human fuzzy information processing, (2) Research on the ergonomic design of visual display terminals, (3) Modelling of thinking process in conceptual design of products, and etc.

4. Special research projects; (1) Fundamental research on organiliquid and gas separation by membranes, (2) Research andevelopment of advanced composite materials. (3) Application of measuring human sense to product design. (4) Design o synthetic receptor molecules, and etc.



Three-dimensional Optometer

physical second leave to be a second to be an example of the second s	Taukuba Gakuen 0298 (54) 6610	Total personnel	125	
Industrial Products Research Institute	Dense Blancing Officer	Total budget	1,441 (million yen)	
1-4, Higashi 1-chome, Tsukuba-shi, Ibaraki, 305	Kesearch Planning Ottesa	Revenue of the Property of States of the Sta		

Was willistent Wate are's hestitate for Publishes and Ransonsouth edfinited in 1920, is concerned with a wide range of resuscrip ds related to apploitation, processing and utilization of minoral surces and natural energy resources, mining and industrial safety, environmental protection. Research on safety maintenance in I mimes is also conducted at the institute's Coal Mine Safety search Centers in Hokkaide and Kyuryu (including the Usui erimental coal mine). At the Institute extensive research efforts focused on the following fields.

Mineral Resource Development and Utilization

- · Exploitation and development of marine mineral resources off shore or in deep seabeds, such as manganese nodules. hydrothermal deposits and cobalt-rich manganese crusts.
- · Advanced construction technology for underground space utilization.
- · Production of new materials, such as functional silicon materials and ultrafine powder.
- · Processing and refining technology for low quality ore and unexploited resources, especially rare metals.

Energy Development and Utilization

- · Comprehensive utilization technology for oil-alternative fuel resources such as coal, natural gas, oil sand, oil shale and biomass, including organic material technology.
- · Advanced combustion technology utilizing various low-grade fuels and energy-saving technology.
- · Geothermal energy exploitation and heat extraction technology.
- Environmental Protection
- · Comprehensive industrial pollution control technology for emission abatement, pollutant measurement and environmental assessment.

- or on regional scales A
- · Global environmental studie s on chimatic changes. formation and transformation of chaspicals in the troposphere. Mining and Industrial Safety
- . Coal mine safety technology, such as gas and coal-dust explosions, mine fire and gas outbursts to support the domestic coal mining industry.
- · Safety assessment for utilization of underground space.
- . Demolition of old constructions using explosives and its safety asse ismeni.



Environmental Pollution and the Circulation of Substances

Intional Research Institute for Pollution and Resources	Tsukuba Gakuen 0298 (54) 3026	Total Personnel	319
6.3. Onogawa, Tsukuba-shi, Ibaraki, 305	Research Planning Office	Total budget	4,016 (million yen)

The Government Industrial Development Laboratory, okkaldo (GIDLH) was established in 1960 as an institute for veloping industries and mining in Hokkaido. The GIDLH nsists of three research departments. The Resources and Energy ngineering Department is engaged in a wide range of basic and plied researchers in the field of energy and natural resources. The pplied Chemistry Department covers analytical chemistry, mihetic chemistry, and the chemistry field including life sciences. he Material Science and Technology Department is carrying out ne R&D in new and functional materials, the research on valuation techniques for these materials and advanced atilization of rese materials for cold regions.

In recent years, the GIDLH has been working on the following & D projects:

Energy technology R&D; (1) Research in coal liquefaction. gasification and combustion, (2) Development of heat pump technology for cold regions.

Environmental protection R&D; (1) Evaluation of new snow tire having low dust pollution, (2) Development of combustion catalyst for reducing NOx, (3) Development of technology processing wastes from advanced industry.

New materials R&D; (1) Development of fine ceramics from silica in rice husks, (2) Development of inorganic fibers and non-crystal materials with a high functional ability, (3) Development of a new preparation method for ultrafine particles, (4) Evaluation of functional single crystal produced in high-reessure hybrid system.

Biomass and biotechnology R&D; (1) Synthesis of optically active substances by enzymatic reactions, (2) Construction of unique strains of yeast with hydroxylation ability on

polyaromatic compounds by genetic engineering, (3) Advanced pyrolysis of biomass resources.

- 5. Regional technology R&D; (1) Intelligent snow removing technology for cold regions, (2) Research on medical diagnostic remote system.
- International cooperation in the R&D with developing countries; (1) Research on new coal combustion technology by fluidized bed. (2) Research on afforestation with functional soil improving materials, (3) Effective activation treatment of lignite and peat materials.

Non-frost type heat exchanger





Sapporo 011 (851) 0151

Total personnel 1.228 (million yen) Total Budget

TECHNOLOGIES DISCUSSED WITH KOBE STEEL

- KSL Technology of Microwave Melter
- Management of Alpha-Contaminated Wastes
- Incineration and Ash Melting for Plutonium-Contaminated Combustible Wastes
- Microwave Solidification Treatment of Incinerated Ash Contaminated by Radioactive Materials
- Crud Slurry Solidification System
- Copper Alloy for High-Cycle Plastic Molding
- Cryopump for Producing a Good Quality Vacuum
- Twin-Head Arc Welding Robot- GT-5000

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Reprint from

"MANAGEMENT OF ALPHA-CONTAMINATED WASTES"

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INTERNATIONAL ATOMIC ENERGY AGENCY VIENNA, 1981

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DEVELOPMENT OF A NEW SOLIDIFICATION METHOD FOR WASTES CONTAMINATED BY PLUTONIUM OXIDES

Utilization of microwave power

F. KOMATSU, Y. SAWADA Mechanical Engineering Research Laboratory, Kobe Steel Ltd, Kobe

K. OHTSUKA, J. OHUCHI Power Reactor and Nuclear Fuel Development Corporation, Tokai, Japan

Abstract

DEVELOPMENT OF A NEW SOLIDIFICATION METHOD FOR WASTES CONTAMINATED BY PLUTONIUM OXIDES: UTILIZATION OF MICROWAVE POWER.

Non-combustible wastes such as incineration ash contaminated by radioactive materials were immobilized with cement or birumen, and were also vitrified with low melting glass powder. These methods are in practical use at present. For permanent storage or disposal these treatments are excellent because they offer stabilization. As a result much reagent or flux is used, but there is one disadvantage: the volume of solid material is increased. In 1978, at the Tokai Works of PNC, research and development work was started on the treatment facilities of non-combustible wastes containing plutonium oxides, such as incineration ash, residues of inorganic acid digestion, components of the HEPA filter and glass or ceramic pieces. A new treatment method using microwave power is being investigated in this project. These wastes are solidified directly in the cylindrical metal crucible of a microwave melter, and are then continuously converted to ceramic-like solid materials. As a result of this research, fundamental data such as the physico-chemical properties of these wastes and the physico-chemical and elution properties of solid materials were obtained. Furthermore, having produced an experimental microwave melter of 5 kW output at 2450 MHz, the melting conditions and engineering data for developing the practical equipment for an output of 100 kW at 915 MHz are now being investigated. In this paper, the treatment system for these wastes, the fundamental data of these wastes and solid materials, and the microwave melter are described.

INTRODUCTION

Heating by microwaves, as is represented by the microwave oven, has various advantages compared with other heating methods - facility of remote control, heating speed and low occurrence rate of deteriorated or damaged parts.











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Products for Zud treatment [Products rate [kg/Day]] * [kg/Yess] Pretreatment for the microwave metting

Final treatment for

reduction and stabilization



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FIG.1. Outline flow sheet of microwave treatment for the photonium contaminated wastes.

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The principle of microwave heating is the dielectric heating method intended to oscillate directly by electric waves the molecules which compose the substance. Hence, it could be one of the most excellent methods for melting non-combustible wastes composed of inorgenic dxide substances such as Al_2O_5 , SiO₂ and CaO.

The Power Reactor and Nuclear Fuel Development Corporation (PNC) is currently investigating the solidification treatment method by microwaves for non-combustible wastes, other than metals, of the plutonium fuel facilities at Tokai Works. This PNC plan has set a timetable, with completion in 1985, of treatment facilities including the secondary treatment process for reduction.

1. PROPERTIES AND OUTPUT OF WASTES

At present plutonium-contaminated HEPA filters, neoprene gloves and combustible wastes are stored temporarily in the storage yard. When the treating facilities are completed they will be removed from the storage yard and treated.

The kinds of waste to be treated, the secondary treatment method for reduction and the output of secondary wastes, are shown in Fig.1. The major waste outputs are HEPA filter elements, residues of acid digestion and incinerated ash, which occur at the rate of 1:0.28:0.19.

The physico-chemical properties of the wastes for solidification treatment are presented in Table I.

Samples possessing the same analytical values as given in this Table were also used in the microwave solidification experiment. Since the actual wastes containing plutonium cannot be used for the experiment, HfO_2 , whose physical properties with respect to density, vapour pressure and free energy of oxides are similar to those of PuO_2 , was added by 0.1 wt% in the solidification experiment, as the dummy substance for PuO_2 .

1.1. Incinerated ash

In the plan, combustible matter such as tissue paper, cardboard boxes, swabs, latex gloves and plywood frames of the HEPA filter, after dismantling, are burnt in the incinerator installed at the facility. The ash output is assumed to be about 7 kg/d.

The chemical composition of incinerated ash varies with the kind and quantity of the materials incinerated. The ash of the chemical composition presented in Table I is the result of incineration in an ordinary incinerator of tissue paper, 85%; cardboard boxes, 5%; latex gloves + gummed tape, 1.5%; and LPDE sheets, 1.5%.

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TABLE I.	COMPARISON OF	CHEM	ICAL C	OMPOSITI	ON,
	MINERAL PHASES	S AND	BULK	DENSITY	OF
	WASTE MATERIA	LS			

Waste material		HEPA	filter elements	High temp	Acid digenion	Incinerated	
i te mis		Glass fitter	Asbestos separator	filter	residues	Ash	
	sio,	56.60	25.48	48.88	16.08	30.72	
	AI. 0,	5.37	1.23	0.36	13.60	24.69	
	Fe, 0,	013	4 69 N	45.18	2.71	5.53	
	CaO	3.15	1.03	1.29	0.03	8.43	
	MgO	< 0.005	35.89	5.62	0.22	5.38	
	TKO,	0.045	0.032	0.057	0.55	1.23	
	Na, O	5.26	0.026	0.022	0.05	5.29	
120	K,0	1.57	0.057	0.12	0.19	0.69	
-	Ç2, 0,	1.57	0.11	< 0.005	0.10	0.03	
cal Computatio	CuO	1.57	0.11	< 0.005	< 0.01	0.07	
	210	3.49	0.37	< 0.006	4.85	0.81	
	960	3.49	0.37	< 0.005	1.49	0.05	
Juens	MnQ	< 0.005	0.03	< 0.45	0.02	0.13	
Ĭ	NiQ	< 0.005	0.16	< 0.005		< 0.01	
	c	1.38	5.31	< 0.27		< 0.1	
	P.Q.	1.38	5.31	< 0.005	0.026	0.19	
	CI	1.38	2.10	< 0.27	0.14	0.32	
	50,			< 0.005	49.68	2.85	
	NO,			< 0.01	< 0.01	eneter	
	8,0,				-	0.48	
	¥.,		autor			1.11	
	19 10%8 R	4.34	22.11	< 0.1		10.44	
Mineral Phases		Amergneus	Cinochrystatile	Gruneitt, Rankinite	A(; (\$0,);	a-Ounz Anatase Calcula Anatase	
Bulk Density (g/cm*)			0.11	0.062 -	0.4 ~ 0.53	0.293	

* ig = ignition



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The samples used in this case were based on the data of the kinds of combustible at present used in the facilities and the results of investigations into the percentages of such materials.

It was found that the volume of combustible waste was reduced to about 1/100 of the original by incineration.

1.2. HEPA filter elements

HEPA filters used in the facilities consist of cases made of plywood frames and elements made of glass fiber filter and asbestos separators. HEPA filters are subjected to secondary treatment (dismantled) for reduction [1]. The output of waste elements is assumed to be about 37 kg/d.

In the analysis and experiment, new HEPA filters were used after dismantling.

1.3. Residues of acid digestion

Chloride compounds such as PVC and neoprene gloves are treated by the acid-digestion method. The residues are white powder or lumps. The residues presented in Table 1 were obtained when neoprene gloves were treated in the test equipment.

The bulk density ranges from 0.4 to 0.53, and these values are the highest owing to the effects of oxides of heavy metals such as ZnO and PbO. The output is estimated about 10 kg/d.

The chemical components are sulphate compounds such as aluminium sulphate anhydrate. Residues of ac.d digestion contain sulphate ions by about 50 wt%. At present, a digestion technique to reduce this content is being investigated.

1.4. High-temperature filters

This is a wool-type filter which is used as a dust trap in incinerators. The estimated output is as low as 50 kg/a. The bulk density is the lowest of all, about 0.062 to 0.071.

1.5. Glass and ceramic pieces

These pieces include experimental implements such as pyrex beakers, measuring flasks, porcelair dishes, etc. The output is as low as 100 kg/a.

2. EQUIPMENT OF MICROWAVE SOLIDIFICATION

When materials composed of inorganic oxide matter such as Al_2O_3 , SiO_2 , and CaO (which are generally called dielectrics) are placed in a high-frequency



(A) Composition of microwave melter



(B) Cross-section of microwave melter

FIG.2. Merowave solidification test unit.



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FIG.3. Photograph of the microwave solidification test unit.

electric field (microwave), the molecules, which are in an electrically neutral state, are turned into molecules with electric dipoles by the strong electric field.

Molecules violently change polarity owing to the frequencies and they generate frictional heat. As a result, the substance reaches the molten state.

By using microwave energy the solidifiable temperature is determined by the material of the crucible and its melting point. When a metal crucible manufactured from low carbon steel is used, the maximum applicable temperature is 1500°C. The microwave melter shown in Figs 2 and 3 was newly developed for the purpose of treating waste containing radioactive substances.

Microwaves from a microwave power generator are radiated into the microwave melter through a wave guide. The melter is the so-called cavity type, which can concentrate the microwave energy on to the sample by vertically adjusting the head tuner.

The sample is continuously fed into the metal crucible – which is also used as a canister – from the hopper by means of a screw feeder and is continuously solidified. During solidification the metal crucible is rotated at a speed of 2 to 5 rpm to radiate the microwaves uniformly.





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TABLE II. COMPARISON OF CHEMICAL COMPOSITION, MELTING POINT, MINERAL PHASES AND DENSITY OF SOLID MATERIAL

		Single	Meit Treat	ment	ħ	Aixec Mai	(Treatment	
liems		Incinerated Arth (1,A)	HEPA Litter (H.F)	Acid digen tion residuate (A.D.A.)	A.D.A.+1.A (1.4711)	H.F + 1.A	H.F.+ A.D. B (3.57:1)	N.F + A.D.F + LA 11 0 28 0 19
	510, (wth)	42.45	45.82	32.82	32.72	45.64	43.88	42.96
	A1, D	22.82	3.98	25.77	31.30	8.70	9.24	13.26
	Fe, D,	3.75	3.27	6.73	2.73	4.70	4.37	5.05
8	040	5.78	1.90	0.28	5.20	2.60	0.48	1.50
2	Mg0	7.46	28.07	1.45	3.48	28.58	28.54	27.52
ateria	TiO, -	4.16	0.07	0.18	1.11	0.30	0.38	0.42
2	N& 0	1.01	0.30	0.27	3.17	3.07	1.82	1.62
1 Sol	K,0 ~ 0	0.27	0.05	1.39	1.08	0.87	0.75	0.48
e vo	0101 -	0.00	0.10	0.47	0.25	0.14	0.16	0.15
Ossile	Cub -	D.11	< 0.01	0.03	0.06	0.02	0.01	0.02
Com	200	0.38	C.36	11.10	6.37	0.81	2.45	1.58
3	PbO	0.08	<0.01	4.90	2.60	< 0.01	0.85	0.24
Cliem	MnO -	0.04	0.09	8.0.0	0.13	0.09	0.07	0.09
	P. D	0.50	0.07	0.372	0.308	0.16	0.09	0.19
	SC: ····	0.27	0.16	0.14	0.247	0.02	0.01	0.01
	C1	< 0.1	< 0.005	< 0.01	< 0.01	< 0.01	<0.01	<0.01
Melting point (° C)		1250	1450	1440	1310	1345	1325	1330
Mineral phases		Angrihita Augriz	^g arstwrite	Melownite Gennite	Ampritaita R impurpod i R Gatorica	Restarite	Forsterine	Forsterile
	Density	2.85	2.97	3 57	3.16	3.01	3.04	3 08

The crucible is cooled with nitrogen gas on the outside in order to prevent oxidation and meltdown due to high temperatures. Once a crucible is filled with solidified matter it is replaced with a new one. Since the produced solidified matter is slowly cooled in the crucible it is converted to a crystalline structure with the most stable physico-chemical properties.





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FIG.4. Cross-section of solidified solid material in the metal crucible.

The off-gas released during solidilying is discharged from the exhaust pipe and is treated in a scrubber. The off-gus temperature is about 60 to 70°C, which is lower than that of other melters such as an electric furnace.

The equipment crucible in this paper measures 100 mm in diameter and is 130 mm high. In practical facilities, where the treating capacity will increase, the crucible size will be much larger, about 300 mm in diameter and 300 mm high.

One crucible can solidify about 50 kg material and it is planned to treat 15 to 20 kg/h at 60 kW microwave output power. As a result, the microwave power generator to be used should have a maximum output of 100 kW at 915 MHz.

3. PRETREATMENT FOR SOLIDIFICATION

Satisfactory solidification by microwaves largely depends on the pretreatment to increase the microwave absorption and insulation effects. For this purpose the sample should be in powder or granular form.

<u>Residues of acid digestion</u> and HEPA filter elements are heated for 15 minutes at 800°C in an electric furnace. As a result, SO_3 contained in residues of acid digestion is almost completely eliminated so that the furning phenomenon by sulphate mist may be avoided. Also, HEPA filter elements become fragile and can be crushed very easily.



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TABLE III. COMPARISON OF ELUTION TEST ON SOLID MATERIALS AND WINDOW GLASS

Solid	Solub		Element (mg/1)										
material	(ert %)	T.Fe	Ca	Mg	ті	AI	Si	Na	к	PD	Zn	s	н
H.F	0.48	<0.05	1.0	0.9	з	<1	24	10.5	1.32	<0.1	<0.05	<1	< 3.7 x 10 ⁻³
H.F+A.D.R+I.A	0.26	<0.05	0.9	1.3	э	2	9	3.45	0.45	<0.1	<0.05	<1	< 3.0
Window glass'	1.75	<0.05	10.2	1		0.5	65	187	10.1	-	-		-

*Chemical composition (w1%): \$10, 69.52, A1203 1.04, Fe203 0.61, Mg0 0.01, Na20 14.15, K20 1.06, Ba0 0.17, Po0 < 0.01, B203 0.40

By this pretreatment the residues of acid digestion and the HEPA filter elements are reduced by about 50% and 17% respectively by weight.

High-temperature fillors and glass and ceramic pieces are crushed for easy transport and melting.

4. RESULTS OF TREATMENT BY MICROWAVES

As shown in the flow-sheet in Fig.1, wastes occurring at highest rates were HEPA filter elements, residues of acid digestion and incinerated ash. Although these wastes are treated by daily operation, in actual treatment it is difficult to melt according to the determined mixing ratio and sequence.

. Therefore, in this experiment single treatment and mixed treatment were individually assumed. The mixing ratio was based on the product rate of these wastes.

The physico-chemical properties of the solid material obtained by melting are presented in Table II. The sample, which was continuously solidified in a metal crucible by using microwaves, is shown in Fig.4. The amount of solid material melted in the crucible was 2.6 to 3 kg. For instance, the HEPA filter element with the highest melting point (1450° C) was solidified at the rate of 960 g/h by using a microwave energy of 4.7 kW (55-W/cm²). The components of the off-gas in this case were dust concentration, 500 to 600 mg/m³; SO₁, 600 to 1000 ppm; HCl, 1.5 to 2.0 ppm; NO_x, 70 ppm. Production of SO₃ gas might be caused by the decomposition of SO₃ in the residues of acid digestion and CaSO₄ in the incinerated ash.

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Waste material	Volume reduction rate
Incinerated ash	1/10
HEPA filter elements	1/27
Residues of acid digestion	1/8

TABLE IV. VOLUME REDUCTION RATE OF WASTE MATERIALS FOR MICROWAVE MELTING

The results of an elution test on the sample shown in Fig.4 are presented in Table III. The elution properties of the solid material and window glass were compared under identical conditions. The 5-g sample, crushed to under 200-mesh size, was immersed in 200 ml boiling (100°C) distilled water for an hour.

After the test, the sample was filtrated by a 0.45-µm membrane filter, and the solubility and soluble ions were analysed. The results showed that the solid material was more resistant to elution than the window glass.

5. CONCLUSION

In this paper, the microwave solidification technique for plutoniumcontaminated non-combustible wastes, the treatmant process, treating conditions, and properties of the obtained solidified matter are outlined.

The treatment process is illustrated in Fig.5. By applying the solidifacion treatment technique, as shown in Table IV, a reduction effect of, for instance, more than 1/10, is expected in the case of ash, and this method may be an excellent technique when compared with the cement or bitumen immobilized method.

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DISCUSSION

C. SOMBRET: The experimental furnace which you described has a low output (5 kW). The corresponding throughput which you quoted is less than 1 kg per hour, which is understandable. Work on the processing of other types of waste in microwave furnaces has been carried out in France and also, I believe, in Harwell. Mr. Grover could, I'm sure, provide a few details on this point. In France, the studies were discontinued owing to the difficulty of finding a highoutput industrial generator, which is essential in order to process any form of waste from an industrial installation. Do you know of any manufacturer in Japan or elsewhere in the world who could supply a suitable generator, let us say, with an output 100 kW?

F. KOMATSU: The magnetron tube of 100 kW output (915 MHz) is commercially available in Japan.

J.R. GROVER: Microwave heating is being studied in the United Kingdom for both the evaporation stage and melting stage of a possible vitrification process for high-level wastes to produce a borosilicate glass product.

C. BAUER: Mr. Komatsu, did you investigate the structure and composition of the phase or phases present in the crucible after the melting? And did you consider the possibilitity of accumulation of Pu in one of the phases?

F. KOMATSU: For the experiment we used HfO_3 whose physical properties are similar to those of PuO_3 ; it was added up to 0.1 wt%. We investigated the structure and composition of the solidified matter using the electron probe microanalyser. The Hf element was found to be distributed uniformly over the matrix. We have performed no experiments with plutonium so far.



INCINERATION AND ASH MELTING FOR PLUTONIUM-CONTAMINATED CONBUSTIBLE WASTES

K.MIYATA*, J.OHUCHI, E.INADA and N.TSUNODA Power Reactor and Nuclear Fuel Development Corporation (PNC) Tokai-mura, Ibaraki, Japan 319-11

1. Introduction

Plutonium-contaminated solid wastes have been generated during NOX fuel fabrication in PNC. These wastes are classified into combustibles, non-combustibles and chlorine-containing organic materials such as PVC and chloroprene, and packed in 2001 drums or 1.7m² cubic containers at the NOX fuel facilities.

These wastes have been treated for the volume reduction and conditioned in the Plutonium-contaminated Waste Treatment Facility (PWTF) since late 1987. The combustible wastes have been treated in the conventional incinerator. The incinerated ashes are melted without any additives to be a 20~30kg ceramics-like block in the stainless steel made canister by powering microwave. The products are packed in the 2001 drums and stored temporarily in the Plutonium-contaminated Waste Storage Facility (PWSF).

2. Process Description

2.1 Waste Feed

10mm

The combustible wastes are introduced into the glove box from drums after being assayed Tarcontents. The wastes of paper and rags are sealed in a paper big by 2kg automatically and wood-framed HEPA filters are cut into s veral pieces using circular saw. These wastes are sent to the incine ator by the belt conveying system.

2.2 Incineration

The conventional incineration unit consists of feeder, incinerator, ash transfer unit and off-gas treatment unit and has the throughput of 50kg/h in one-shift operation. The incinerator is composed of primary and secondary combustion chamber, and covered with stainless steel casing to keep alpha-tightness and prevent the leakage of radionuclides. The pressure and temperature in the incinerator are regulated automatically by controlling the flow rates of exhaust gas and kerosene.

Paper bagged combustible wastes and cut pieces of HEPA filters are fed to the primary combustion chamber from the top of the incinerator through the airlock room with adiabatic shutter. These wastes are incinerated on the inconel made fire grades by two kerosene burners. The combustion off-gas with coarse dusts from primary combustion chamber is transferred to the secondary chamber and high-temperature filter (HTF) for post combustion and filtration. Secondary chamber is filled with silicon carbide lumps. HTF is mainly composed of many vertical cylindrical tubes that are coated with asbestos fiber.

Figure 1 and 2 show the artistic view of the conventional incineration unit and the cross section of the incinerator.





Fig. 2 Cross Section of the Incinerator

3.



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2.3 Off-gas Treatment

The off-gas treatment don't there is of dilutor, pre-filter, HEPA filter, scrubber, mist the off blowers. The off-gas from high-temperature filter is miss that is and cooled below 80°C by the dilutor. Radionuclides are the from the off-gas by the HEPA filters, and environmental pellow such as SOX and NOX gas are removed by the scrubber prior to discnarge to the atomosphere.

2.4 Ash Handling

The ash handling unit consists of hopper, crusher, vibrating sieve, and belt conveyers. The ash is crushed and separated from nails after being removed from the bottom of the primary chamber. The ash is sent to the microwave melting process by flight conveyer.

2.5 Microwave Melting

The microwave melting unit consists of a microwave generator, wave guide and melter. The melter is equipped with tuner, power monitor, isolator, discharge detector and ITV monitor, installed in a glove box. Microwave from microwave generator is radiated to the ash in the cavity-type melter through a wave guide. Microwave energy is focused on the ash in the canister by adjusting the tuner vertically. The microwave generator has the output of lOkW with 2450NHz frequency. The melter has the throughput of 5kg/h.

The ash is melted in the canister (130mm^bX 770mm^x) at 1200 - 1400°C, and converted into ceramics-like blocks, which are bagged out for packing in 200£ drum. The drum with eight ceramicslike blocks is stored in PWSF. The major mineral phases of the products are forsterite, anorthite and augite. The off-gas released during melting is sent to the off-gas treatment unit of the conventional incinerator.

Figure 3 and Table 1 show the microwave melting process flow and the operating data of incineration and melting, respectively.



Fig. 3 Microwave Melting Process Flow

Table 1 Operating Data

	Incineration	Meltir
Feed Rate	2.0 kg/3 min. (Paper Bag) 5.0 kg/5 min. (Cut HEPA Filter)	4 kg/
Temperatura	Primary Chamber 800~ 500°C Secondary Chamber 500~1000°C HTF 500~ 500°C	1200-14
Operating Pressure (mmH ₂ O)	- 30 40	- 45~-
Exhaust Gas (Nm ³ /h)	7000-9000	.5~
Density (g/cm ¹)	0.1~0.3	3

2.6 Cost

The construction cost of the PWTF is about on million US dollars. The incineration unit and micro share 14% and 4% in the construction cost of the PW These equipments need the glove boxes, alpha-tightn control system and licensing/inspection to treat th contaminated wastes, which are not necessitated for wastes.

3. Operational Results

As shown in Table 2, approximately 12 tons of wastes such as paper, rags and HEPA filters have be for the first one-year-operation in PWTF, and appro of ashes have been generated. The weight reduction 1/13. HEPA filter composed of glass fiber and asbes cause of small weight reduction ratio. The volume r combustible waste is about 1/130 by incineration an

Figure 4 shows a temperature and pressure chamincineration of paper, rags and HEPA filters. Durin the temperature of primary chamber and secondary chamintained at 800 to 900°C and 900 to 1000°C, resperence at -40 mmH20.

Treatment Method	Incineration	Melting
Treated Waste Weight	12 ton	760 kg
Product	900 kg (Ash) ·	34 blocks (800 £)
Weight Reduction Ratio	1/13	, 1
Volume Reduction Ratio	1/1:	30

Table 2 Weight and Volume Reduction Ratio of the Incineration and Melting



Fig. 4 Temperature and Pressure Change of the Incinerator

.

4. Conclusions

- Incineration and microwave melting for plutoniumcontaminated combustible waste has been demonstrated successfully for the volume reduction and immobilization.
- (2) The alpha-tightness of the conventional incinerator have
- been maintained during the incineration up to date.
- (3) The volume reduction ratio of 1/130 for the combustible wastes have been attained by the combinated process of conventional incineration and microwave melting.
- (4) The incinerated ash have been conditioned stably without any additives by powering microwave.
- (5) Treatment for plutonium-contaminated waste from MOX facilities will be continued to reduce the stored waste volume and the characterization of the conditioned waste will be also progressed.

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U.S. Patent May 18, 1982 Sheet 1 of 3 4,330,698





FIG.1B













FIG.4B



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		21 04 79 JP P54-49599				Sawada. Sanada, K	Yoshihisa: Komatsu, Fumiaki, Nishinomiya, JP; Kazuo, Kobe, JP, Sakaki, Yorihisa, Akashi, JP	
	0	Patentinhaber:			8	Enigegen	haltungen	
		Kope bieel, Lig., Kope, Myp	90. 4			US	40 39 797	
	3	Vertieter.				US	25 86 754	
-		Tiedtke, H., DiplIng.; Böhli DiplIng.; Grupe, P., DiplIr PatAnw., 8000 Mönchen	ng. G	., Dipl. Chem ; Kinne, R., elimenn, H., Dipl. ing.,				

B Mikrowellenofen



Patentansprüche:

1. Mikrowellenolen zum Erwärmen eines Gutes. der aus einem das Gul aufnehmenden und abnehmbaren Unterteil und einem Oberteil besteht. das über einen Mikrowellenleiter mit einem Mikrowellenoszillator verbunden ist. dadurch gekennzeichnet, daß zum Verflüssigen eines schmelbaren Gutes (M) im Unterteil (2) ein Schmelzliegel (8) vorgesehen ist, der über eine im ju Oberteil (1) vorgeschene Fullollnung (5) beschickbar ist, wobei die Leistung des Mikrowellenofens (1, 2) über eine auf dem Oberieil (1) angeordneie Abstimmvorrichtung (4) steuerbar ist, die von einem kohlen, zylindrischen Metallkörper gebildet ist, der 15verschiebbar im Oberteil (1) geführt ist, und daß die Mikrowellenzuleitung (3) gegen im Olenraum entstehende Gase abgedichtet ist.

2. Mikrowellenofen nach Anspruch I. dadurch gekennzeichnet, daß der Schmelztiegel (8) in einem 20 Behälter (12) aufgenommen ist, der drehbar innerhalb des Unterteils (2) montiert ist.

3. Mikrowellenofen nach Anspruch 1 oder 2, dadurch gekennzeichnet, daß das Unterteil (2) mit einem Inerigasanschluß (14) versehen ist, durch den 25 in einen Spalt (G) zwischen der Innenwand des Unterteils (2) und der Außenwand des Behälters (12) ein Inerigas einleitbar ist, welches einen Druck bositzt, der über dem Innendruck des Mikrowellenofensliegt. 30

4. Mikrowellenofen nach Anspruch 2 oder 3. dadurch gekennzeichnet, daß der Behälter (12) eine Antriebswello (13) trägt, die aus dem Unterteil (2) herausführt und über eine Antriebsvorrichtung angetrieben ist.

5. Mikrowellenofen nach Anspruch 1 bis 4. dadurch gekennzeichnet, daß die Mikrowellenzuleitung (2) ein Hohlleiter ist, in dem zueinander im Abstand stehend mehrere luftdichte Zwischenwande (Si: Si) eingesetzt sind und somit zwischen sich und 40 der Innenwand des Hohlleiters (3) einen Innenraum begrenzen, dessen Druck höher gehalten ist als der Innendruck das Mikrowellenofens.

6. Mikrowellenolen nach Anspruch I bis 5. doppelwandiger Zylinder ist, dessen Zentralhohlraum (15) auf der dem Schmelztiegel (8) zugewandten Seite durch ein Netz (16) aus leitfahigem Material und auf der anderen Seite durch ein Sichtfenster (17) abgeschlossen und mit einem 50 Inerigas-Anschluß (18) versehen ist und dessen Wandungs-Hohlraum (19) einen Anschluß für eine Mantelkühlung besitzt

Die Erfindung bezieht sich auf einen Mikrowellenofen gemaß dem Oberbegriff des Patentanspruchs 1.

verschiedener Materialien durch Einwirkung von Mikrowellenstrahlen hat im Vergleich zu anderen Anwarm-und Schinelzverfahren eine Reihe von Vorteilen, zu denen die Vergleichmäßigung des Erwärm- und Schmelzprozesses und die Möglichkeit der genaueren 63 Teil des Patentanspruchs 1 angegebenen Merkmale -Steuerung der Geschwindigkeit des Schmelzvorgangs durch Einstellen der aufgebrachten Mikrowellenleistung gehören.

Das Prinzip der Mikrowellen Erwärmung kann auf zahlreichen verschiedenen Gebieten und zu verschiedensien Zwecken angewendet werden. Beispielsweise konn das Volumen von Abfallschlämmen, die bei diversen industriellen Prozessen anfallen, durch Trocknen oder Schmelzen (mit nachfolgender Erstarrung) mit Hille von Mikrowellen verringert werden, om die weitere Behandlung zu erleichtern. Die Schmelz- und Erstarrungsbehandlung zur Volumenverringerung durch Mikrowellenbestrahlung kann auch bei radioaktiven Abfallen angewendet werden, die in kerniechnischen Anlagen anfallen, gesommelt werden und während langer Zeit an abgeschirmten Orten gelagert werden, um Behalter und Raum für die Lagerung einzusparen und dadurch die Lagerkoppoliat zu erhöhen, wobei pullerdem der Arbeitspulwand zur Handhabung der Abfalle verringert wird.

Aus der US-PS 2586754 ist ein Mikrowellenofen gemäß dem Oberbegriff des Patentanspruchs 1 bekannt. bei dem die Energieübertragung vom Mikrowellenoszillator in den Mikrowellenolen durch eine Koasialkabel-Einrichtung erfolgt, deren Aufbau auf eine vorbestimmte Mikrowellenlänge hin so optimiert ist daß die Mikrowellen Übertragungsverluste vom Mikrowellenoszillator zum Mikrowellenofen möglichst klein gehalten werden. Die Mikrowellenleistung bzw. die auf das zu schmelzende Gut zu übertragende Mikrowellenleistung ist bei diesem bekannten Mikrowellenolen allein über den Mikrowellenoszillator beeinflußbar.

Der Wirkungsgrad eines Mikrowellenofens hängt allerdings im wesentlichen davon ab, wie exakt die Mikrowellenfrequenz auf den Mikrowellen-Resonanzkörper abgestimmt ist. Wenn das zu erwärmende Out während des Erwärmungsvorgangs sein Volumen andert, wird der Mikrowellen-Resonanzkörper dadurch ebenfalls beeinflußt, so daß diese Abstimmung nicht mehr exakt vorliegt. Der aus der US-PS 2386754 bekannte Mikrowellenofen eignet sich deshalb nur für die wirtschaftliche Erwärmung eines Gutes, das im Verhältnis zur Ofenkonstruktion bzw. zum Erwärmungsbehälter oder Schmelzbehälter so klein ist daß es den sich aus Mikrowellenofengehäuse, Mikrowellenleiler und zu erwärmendem Gut zusammenseizenden »Mikrowellen-Resonanzkörper« nur unwesentlich dadurch gekennzeichnet, daß der Hohlzylinder ein 45 prägt. Das Resonanzverhalten des aus den oben angegebenen Komponenten bestehenden Mikrowellen-Resonanzsystems ist dabei weitestgehend unabhängig von dem zu erwärmenden Gut: mit einem Mikrowellenolen, wie er aus der US-PS 25 86 754 bekannt ist, und der ein derartiges Resonanzsystem besitzt können. somit nur Güter erwärmt werden, deren Größe und Form beim Erwärmungsvorgang gleich bleibt. Für die wirtschaftliche Aufschmelzung von anderen Oütern, wie sie im industriellen Anwendungsgebiet beispielsweise in 55 sich stellig andernden Mengen anfallen, kann dieser

bekannte Mikrowellenofen nicht mehr wirtschaftlich eingesetzt werden.

Der Erfindung liegt die Aufgabe zugrunde, einen Mikrowellenolen gemäß dem Oberbegrill des Patentan-Das Verfahren zum Erwärmen und Schmelzen 60 spruchs 1 zu schallen, mit dem selbst größere Mengen eines im Rahmen eines industriellen Verlahrens. anfallenden Gutes in einem Schmelztiegel wirtschaftlich aufgeschmolzen werden können.

> Diese Aufgabe wird durch die im kennzeichnenden gelöst

> Die sich beim Aufschmelzen und durch die kontinuierliche Zuführ des zu schmelzenden Gutes stetig

verändernden Volumina des aufzuschmeizenden Gutes im Mikrowellenolen bestimmen wesentlich das Resonanzverhalten des oben beschriebenen Mikrowellen-"Resonanzkorpersystems« aus Mikrowellenleiter. Ofenwandung und Gut, so daß sich bei herkommlicher Konstruktion der Wirkungsgrad des Mikrowellenofens mit dem montentanen Zustand des Gutes andert. Durch die erfindungsgemaße Maßhahme, die Abstimmvorrichung justierbar im Oberieil des Olens au lühren, kann der Wirkungsgrad des Ofens den jeweiligen Beiriebsbedingungen des Aufschmelzvorgangs optimal angepabi werden. Diese zur Wirkungsgrad-Optimierung führende Abstimmung erfolgt zudem mit geringstem Aufwand. indem durch einlaches Verschieben des Abstimmkörpers die Resonanzliäche und damit die Resonanzlähigkeit bzw. die Resonanzbedingungen des Mikrowelleno-(en-Resonanzsystems exakt justient und abgestimmt werden. Unter Losiosung von der herkommlichen Lehre, die Mikrowellenfrequenz auf den Mikrowellen-Resonanzkörper im Hinblick auf eine wirtschaftliche Erwärmung starr abzustimmen, eröffnen die erfindungsgemäßen Maßnahmen erstmals die Möglichkeit, auch variable Volumina bzw. sperrige Guter wirtschaftlich zu erwärmen, deren Ausdehnung sich im Schmelzraum beim Erwärmen stark andert. Dadurch kann auch kontinuierlich aufzuschnelzendes Out zugeführt werden, ohne Wirkungsgradverluste in Kauf nehmen zu müssen.

Vorteilhafte Weiterbildungen hinsichtlich der zusätzlichen Anhebung des Wirkungsgrades des Mikrowellenofens sind Gegenstand der Unteransprüche.

Nachstehend werden anhand schematischer Zeichnungen mehrore Ausführungsbeispiele der Erfindung näher erläutert. Es zeigt

Fig. 1(!) eine Draufsicht auf einen Mikrowellenofen. der als Schmelzvorrichtung dient.

Fig. 1(11) eine Seitenansicht des Mikrowellenofens

Fig. 2 einen senkrechten Schnitt, der einen Schmelztiegel zeigt, der sich in einem Unterteil des Mikrowellenofens befindet.

Fig. 3 eine Schnittdarstellung der Mikrowellenzuleitung des Mikrowellenofens.

Fig. 4(1) eine Schnittdarstellung einer einstellbaren Abstimmvorrichtung des Mikrowellenofens, und

Fig. 4(11) Ansichien von Ausführungsbeispielen der im Mikrowellenofen benutzten Netze.

Im folgenden wird zunächst auf Fig. 1 eingegangen. Die darin dargestellte Mikrowellen-Schmelzvorrichtung umfaßt einen Mikrowellen-Schmelzofen, der aus einem Oberieil 1 und einem Unterteil 2 besteht. Um den Schmelzofen herum ist eine nicht dargestellte Kühleinrichtung angeordnet, die normalerweise aus Rohren besteht, durch die ein Kühlmittel umgewälzt wird. Das Oberteil bzw. die obere Ofenhälfte 1 ist mit einem Hohlleiter bzw. Wellenleiter 3 für Mikrowellen, einer einsteilbaren Abstimmvorrichtung 4 und einer Zuführleitung 5 für Schmelzgut, d.h. für zu schmelzendes Material, versehen. Die obere Ofenhälfte 1 ist an einer Tragkonstruktion 6 unabhängig vom Unterteil bzw. von der unteren Ofenhällte 2 befestigt und wird von dieser Tragkonstruktion abgestützt. Zum Antrieb des einstellbaren Tuners bzw. der Abstimmvorrichtung 4 dient ein Motor mit der mit dem Tuner 4 über ein Kegelradgetriebe 20 verbunden ist, damit die Höhe der Abstimmvorrichtung 4 im Mikrowellenofen eingestellt werden kann. Die obere Ofenhälfte 1 ist ferner mit einer Abgasleitung 7 versehen, durch die Schwebstolle, d.h. Staub und Rauch, die im Schmelzolen entstehen und die Wirksamkeit der Mikrowellenbestrahlung senken, aus dem Schmelzofen abgeleitet werden. In der unteren Ofenhälfte 2 ist ein Schmelztiegel 8 (siehe Fig 2) angeordnet. Die untere Ofenhälfte 2 wird von einer
⁵ Trageinrichtung 10 getragen, die die untere Ofenhälfte 2 von der oberen Ofenhälfte 1 wegbewegen und zu dieser hinbewegen kann. Die Trageinrichtung 10 besteht aus einem Drehmechanismus 10, 1 mit einem Motor nij und einem Hubmechanismus 10,2 Zum Drehmechanismus

¹⁰ 10.1 gehört ein Trogorm 11. dessen eines Ende mit der unteren Ofenhalfte 2 verbunden ist. An seinem anderen Ende ist der Trogorm 11 mit einem Zohnrad 22 versehen, das auf einer Welle 21 befestigt ist, wie Fig. 1(1) erkennen laßt, in der die Trogeinrichtung 10 im

¹⁵ Schnitt gemäß A-A in Fig. 1(11) dargestellt ist. Das Zahnrad 22 kämmt mit einem Zahnrad 23, das zum Motor mit gehört und von diesem angetrieben wird, so daß der Tragarm 11 in einer horizontalen Ebene um die Achse der Welle 21 gedreht bzw. geschwenkt wird und

20 dabei die untere Ofenhälfte 2 von der oberen Ofenhälfte 1 wegbewegen und in eine zurückgezogene Stellung bringen kann, in der die untere Ofenhälfte mit 2' bezeichnet ist. Der Drehmechanismus 10.1 wird von einem Hubtisch 24 des Hubmechanismus 10.2 getragen.

25 Der Hubtisch 24 wird von einer hydraulischen oder anderen Antriebseinrichtung aufwärts- und abwärtsbewegt, um die untere Ofenhälfte 2 in senkrechter Richtung zur oberen Ofenhälfte 1 und von dieser wegzubewegen.

30 Während des Betriebs der vorstehend beschriebenen Schmelzvorrichtung in Gestalt eines Mikrowellenofens wird die untere Ofenhälfte 2. die den Schmeiztiegel 8 trägt, mit der oberen Olenhällte verbunden, indem die Trageinrichtung 10 die untere Ofenhälfte in ihre 35 Arbeitsstellung schwenkt und hebt, bevor der Mikrowelien-Schmelzofen mit dem Material bzw. Schmelzgut M refullt wird, das aus einer Aufgabevorrichtung B durch "uführleitung 5 zugeführt wird. Wahrend des 0.0 Schmein werden Schwebstoffe, d.h. Staub und 10 Rauch, die un Schmelzofen während des Schmelzens auftreten und die Bestrahlung mit Mikrowellen behindern, durch die Abgasleitung 7 abgeleitet, während das Material M im Schmelztiegel B mit Mikrowellen bestrahlt wird, die von einem nicht dargestellten

*5 Mikrowellengenerator erzeugt werden und durch die Mikrowellenzuleitung 3 zum Schmelzofen geleitet werden.

Es versieht sich, daß die obere Ofenhälfte 1 und die untere Ofenhälfte 2 dicht miteinander verbunden sind.

50 damit weder die in den Schmelzofen geleiteten Mikrowellen noch der Staub austreten können, der im Schmelzofen während des Schmelzens entsteht.

Damit der Erwärmungs- und Schmelzvorgang gleichmaßig und wirkungsvoll durchgeführt werden kann, ist

55 es erforderlich, das eingefüllte Schmelzgut gleichmaßig mit den Mikrowellen zu bestrahlen. Es hat sich allerdings oftmals als schwierig erwiesen, eine gleichmä-Bige Mikrowellenbestrahlung durchzuführen, wenn das Schmelzgut in den Schmelzofen derart eingeführt

40 werden mußte, daß sich an bestimmten Stellen des Schmelzofens eine größere Schmelzgutmenge ansammelte oder daß das Schmelzgut eine unebene Oberflächenkontur annahm die eine unregelmäßige Wirkung der auftreffenden Mikrowellen verursachten, so daß

⁶³ unterschiedliche Bereiche des Schmelzgutes unterschiedlich stark erwärmt wurden. Dies könnte dadurch vermieden werden, daß am Mikrowellen-Schmelzofen mehrere Mikrowellen-Bestrahlungsquellen vorgesehen



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werden: allerdings hätte das den Nachteil, daß die Schmelzvorrichtung groß und kompliziert würde. Bei der vorliegenden Mikrowellen-Schmelzvorrichtung wird dieses Problem durch eine drehbare Ofenkonstruktion, d. h. durch einen drehbaren Behalter gelöst, der den Schmelztiegel aufnimmt und im Unterteil des Ofens gelogert ist.

Wie in Fig 2 erkennbar ist, ist der Schmelztiegel 8 an und in einem drehbaren Behalter 12 ausgehängt, der auf einer drehbaren Welle 13 im unteren Abschnitt der 111 unteren Olenhallte 2 angebracht ist, so daß der drehbard Behalter 12 und der Schmelztiegel in einer horizontalen Ebene gedreht werden können. Die Welle 13 ist mit einem geeigneten, nicht dargestellten Drehantrieb verbunden, beispielsweise einem Antriebsmotor, der an 15 der unteren Ofenhälfte 2 befostigt ist. Der im drehbaren Behälter 12 aufgehängte Schmeiztingel 8 wird somit während der Mikrowellenbestraniung mit geeigneter Geschwindigkeit gedreht, so daß alle Bereiche des Materials M gleichmäßig mit Mikrowellen bestrahlt 20 werden, d.h. gleichmäßig erwärmt und geschmolzen werden, unabhängig von ungleichmäßiger Verteilung des Materials M im Schmelzolen und unabhängig von gegebenenlalls unebenen Oberflächen des Schmelzgu-85.

Der drehbare Behälter 12 ist vorzugsweise lösbar an der unteren Ofenhälfte 2 angebracht, um die Wartung des Schmelzoiens in dem Fall zu erleichtern, daß geschmolzenes Material aufgrund einer Undichtheit des Schmelztiegels 8 in den drehbaren Behälter 12 fließt.

Während der Schmelzbehandlung des Schmelzgutes dehnt sich der Schmelztiegel 8 in Längsrichtung thermisch aus. Durch diese thermische Ausdehnung des Schmelztiegels treten bei der Erfindung keine Schwierigkeiten auf, da der Schmelztiegel im bzw. am 35 drehbaren Behälter 12 aufgehängt ist.

Während des Erwärmens und Schmelzens des Schmelzgutes ist as bisweilen notwendig. Reaktionen zwischen dem Schmelzgut und der Atmosphäre im Mikrowellen-Schmelzofen zu verhindern, um nach der 40 Schmelzbehandlung erstarrtes Material mit bestimmten chemischen und physikalischen Eigenschaften zu erhalten. In diesem Fall kann der Schmelzofen mit Mitteln zum Einleiten eines Inertgases versehen sein, um auf diese Weise im Schmelzofen eine Inertgasatmosphäre 43 u erzeugen. Das Einleiten eins Inertases hat die usätzlichen Wirkungen, daß der durch Oxidation nervorgerufene Verschleiß des Schmelztiegels geringer bleibt und daß ein Kühleffekt erzielt wird, der Schäden am Schmelztiegel durch Überhitzen verhindert. 50

Bei der Ausführungsform gemäß Fig. 2 ist am Boden der unteren Ofenhälfte 2 ein Inertgasanschluß 14 vorgesehen, durch den ein Inertgas in einen Spalt G zwischen der Außenseite bzw. wand des drehbaren Behälters 12 und der Innenseite der Wand der unteren Ofenhälfte geleitet wird. Der Druck der Inertgasatmosphare im Spalt G wird auf einen Wert eingestellt, der etwas höher als der Druck der Atmosphäre im Schmelzofen ist, so daß das Inertgas im Spalt G in den Schmelzofen strömt und darin eine Inertgasatmosphäre erzeugt, wobei es gleichzeitig das Austreten von Rauch oder anderen Abgasen durch einen Spalt C verhindert.

Rauchgase, die in die Mikrowellenzuleitung 3 unbehandeltes Material in die gelangen, werden mit Mikrowellen bestrahlt und werden, bis die Füllung des Sc verringern die nutzbaren Mikrowellenenergie erheblich, 65 bestimmtes Maßangewachsen ist.

da sie zu Enilodurgeh oder onderen Phonomenen führen. Um dies zu verhindelin, ist inder Mikrowellenzuleitung 3 eine Zwischenwand Si. zw. Sy vorgesehen. wobei Luli oder ein Inerigas in de Raum auf der dem Mikrowellen Scimelzofen zugews idien Seite der Zwischenwand gele iet wird, domy: ",asstrome entstehen, die die Rauchga e und den Sin o standig in Richtung zum Schmelzofe, spülen, isbesondere bei einer Schmelzvorrichtung ... E. andlung radioaktiven Materials sind vorzugsweise zwei Zwischenwande Si und Si aus »Tefion« oder Quarzglas am inneren tzw. außeren Ende der Mikrowellenzuleitung 3 angeordnet, wie dies in Fig. 3 gezeigt ist. Für den Fall, daß die Lufidichtheit durch Ermudung der inneren Zwischenwand Si nachläßt, ist der Innenraum 5 zwischen den zwei Zwischenwänden Si und Si vorzugsweise mit einem Inerigas gefüllt, das einen Druck hat, der etwas höher als der Druck der Ofenaumosphäre ist, wodurch verhindert wird, daß Gas aus dem Schmelzofen in die Mikrowellenzuleitung 3 strömt.

Der bei der Mikrowellenolen-Schmelzvorrichtung verwendete und als Abstimmvorrichtung dienende Tuner 4 hat vorzugsweise die in Fig. 4(i) dargestellte Ausbildung er weist einen Hohlkörper aus Metall mit einer Längsbohrung 15 auf. Der Tuner ist an seinem unteren Ende mit einem Gitter bzw. Netz 16 aus leitfhäigem Material, das das Austreten von Mikrowellen unterbindet sowie an seinem oberen Ende mit einem Sichtlenster 17 aus einem plattenförmigen, lichtdurchlässigen Material wie Quarzglas versehen, das eine Beobachtung des Inneren des Mikrowellen-Schmelzofens ermöglicht und das Austreten von Gasen und Staub, die im Schmelzofen entstehen, verhindert. Beispiele für die Ausbildung des Netzes 16 sind in Fig. 4(1) dargestellt.

Das Eindringen von Staub in die Längsbohrung 15 des Tuners bzw. der Abstimmvorrichtung 4 kann durch Einleiten eines Inertgases durch einen Inertgasanschluß 18 in die Längsbohrung verhindert werden. Das eingeleitete Inertgas hat einen Druck, der etwas höher als der Innendruck des Mikrowellen-Schmelzofens ist. Gegen Strahlungswärme kann der Tuner geschützt werden, indem durch einen Wandungshohlraum 19 auf seiner Außenseite Kühl- asser umgewälzt wird.

Der Schmelztiegel kunn aus einem metallischen Material, beispielsweise röstfreiem Stahl, oder einem kohlenstolfhaltigen Material, beispielsweise 7 phil, bestehen. Vorzugsweise wird allerdings ellist gel aus Metall benutzt. Wenn das Schmelzgut einen hohen Schmelzpullt hat, kann ein Schmelztiegel benutzt werden, dessen Innenseite mit einer Lage aus wärmeisolierendem Material mit hohem Schmelzpunkt wie beispielsweise Aluminiumoxidzement beschichtet ist.

 Während des Schmelzens mittels der beschriebenen Mikrowellen-Schmelzvorrichtung kann das zu behandeinde Material kontinuierlich dem Schmelztiegel zugeführt werden, wobei dann die Erwärmungs- und Schmelzbehandlung durch Mikrowellenbestrahlung
 kontinuierlich erfolgt. Alternativ kann, nachdem eine Materialcharge geschmolzen worden ist und dadurch ihr Volumen verringert worden ist, immer wieder unbehandeltes Material in die Schmelze zugegeben werden, bis die Füllung des Schmelztiegels auf ein bestimmtes Maß angewächsen ist.



ZEICHNUNGEN BLATT 1

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Fig. 1 (II)

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Fig. 4 (11)

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4



(12) UK Patent (19) GB (11) 2067 823 B

(54) Title of Invention

System for treating radioactive waste

(S1) INT CL3 G21F 9/00

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- (73) Proprietor
 Kobe Steel Limited,
 3—18 1-chome, Wakinohamacho, Fukiai-ku, Kobe-City,
 Japan
- (72) Inventors Atsushi Tagusagawa, Yorihisa Sakaki, Yoshihisa Saweda, Fumiaki Komatsu, Masaru Hayashi
- Agent
 Elkington and Fife,
 High Holborn House, 52/54
 High Holborn, London
 WC1V 6SH
- GB 2 067 823 B

1 111



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. 4

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Fig.3



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a 3



Fig.4

Χ.,



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Fig.8





9





SPECIFICATION System for treating radioactive weste

Background of the Invention This invention relates to comprehensive system

5 for treating the radioactive waste which is discharged as a slurry in large quantities from an aromatic plant and is subsequently thickened, dried and prepared for melting.

There have already been proposed a number of 10 methods for treating radioactive waste in slurry

- form, which contains radioactive components such as the primary cooling water of an atomic pile. These proposals include that of Japanese Laid-Open Patent Specification No. 17572/78, in
- 15 which it has been ensidered advantageous from the viewpoint of the economic disposal of radioactive waste to store the latter after sedimental collection, drying, melting and solidification of the radioactive component.
- 20 However, up to the present, apparatus for putting such proposals into practical operation has not yet been developed.

Summary of the Invention

Under these circumstances, the present

- 25 invention has as its object the provision of a comprehensive system which can treat a slurry with radioactive components efficiently in one place by successively thickening and drying the radioactive waste into a form ready for charging to 30 the hopper of a melter.
- According to the present invention, there is provided an apparatus for the treatment of radioactive waste, comprising:
- a storage tank holding a slurry of the 35 radioactive waste:
- a sedimentation tank for thickening the slurry of 100 mechanism; radioactive waste received from said storage tank; a drier for drying the thickened radioactive waste; a hopper for receiving the dried radioactive 40 waste; and
 - a rotary mechanism for transferring the radioactive waste to and from the sedimentation tank, drier and hopper, said mechanism having a number of containers for holding the radioactive
- 45 waste, rotary arms for supporting said containers: a rotational drive mechanism for said rotary arms, and a lift mechanism for venically lifting said containers up and down.

In one embodiment of the apparatus of the

- 50 invention the sedimentation tank and drier are located successively above the rotational path of travel of said containers, and said hopper is located beneath said rotational path of travel in a position spaced from said drier by a
- 55 predetermined angle about the axis of rotation of said rotary mechanism. The apparatus may also comprise means for inverting and upturning said containers, provided respectively before and after said hopper in the rotational path of travel of said 50 containers.
 - In a particular embodiment of the invention, said sedimentation tank is provided with a rotary

vertical bore forming inlet and outlet openings on

- 65 the upper and lower sides thereof in alignment with an opening at the bottom of said sedimentation tark and a tapered bore formed perpendicularly to, and across, said vertical bore, a rotary shaft having a tapered body portion fitted in
- 70 said tapered bore and provided with a cavity in said tapered body portion in alignment with said inlet opening for collecting radioactive sediment, a spring urging one end of said rotary shaft in the tapered direction thereof, and a rotational drive
- 75 mechanism connected to the other end of said rotary shaft for rotating the latter through a predetermined angle, the radioactive sediment collected in said cavity being dropped through said outlet opening upon rotation of said rotary shaft.
- 80 In one embodiment of the apparatus of the invention the lift mechanism is so designed as to lift said rotary arms up and down.

In another embodiment, said lift mechanism is installed beneath said drier or said rotary collector

85 whereby said container is lifted up and down when said container comes beneath said drier or said rotary collector.

Brief Description of the Drawings Reference is now made to the accompanying

90 drawings in which:

Fig. 1 is a schematic illustration of one embodiment according to the present invention of apparatus for treating radioactive weste:

- Fig. 2 is a schematic vertical section of a rotary 95 collector:
 - Fig. 3 is a sectional view across a rotary collector shaft:
 - Fig. 4 is a plan view of the rotary collector: Fig. 5 is a schematic view of a rotary transfer
 - Fig. 6 is a plan view of the rotary transfer mechanism;
 - Fig. 7 is a schematic section of pot supporting structures:
- 105 Fig. B is a plan view of the pot supporting structures;

Fig. 9 is a view explanatory of the inversion of a pot: and

Fig. 10 is a view showing an inverted pot to be 110 upturned into initial upright position.

Description of Preferred Embodiments Referring to the accompanying drawings and first to Fig. 1, indicated at 1 is a tank which is provided in the lower half of a fixed frame 2 to

- 115 hold a slurry of radicactive waste therein, and at 3 a vessel which is supported on top of the frame 2 for thickening the slurry and which is, in the particular example shown, a sedimentation tank. Provided beside the frame 2 is a side frame 5 of a
- 120 smaller height supporting thereon a rotary mechanism which turns a number of pots 6 along a predetermined rotational path of travel. Designated at 7 is a dryer such as a microwave dryer for drying the condensate which is 125 accommodated in the pot 6, and at 108 is a
- summer which receives the dried material and

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supplies it to a melter 110 of the next stage, if necessary, by means of a feeder 109. In the upper portion of the slurry storage tank 1.

there is provided a stirrer 112 with upper and 5 lower stirring blades 114 and 114b which are

- mounted on a drive shah 113. The slurry in the tank 1 is sucked by a slurry feed pump 116 through a suction pipe 115 and quantitatively fed to the sedimentation tank 3 through a pipe 118
- 10 with an electromagnetic valve 117. The sedimentation tank 3 is provided with a level switch 120 which produces a signal when the slurry fed from the slurry feed pump 116 reaches a predetermined level, thereby stopping
- 15 the operation of the pump 116 and closing the electromagnetic valve 117, to suspend the supply of the sluny to the tank 3. The sedimentation tank 3 is supplied with a high molecular weight flocking liquid through a nozzle 121, which is mixed with
- 20 the slurry by a stirrer 122, which is mounted on top of the tank 3. The slurry contains in addition iron rust such as hematite, and magnetite, radioactive corrosion products of cobalt, manganese and the like. The suspended
- 25 radioactive components are flocked by the high molecular weight flocking liquid and gradually fall to the bottom 3a of inverted conical shape and finally in to a rotary collector 123 which is located at the pointed end of the bottom portion 3a.
- 30 Indicated at 125 is a hopper which holds a fusible additive which is to be used in the succeeding melting and solidifying stage, the fusible additive in the hopper 125 being supplied to the sedimentation tank 3 in a predetermined
- 35 quantity through a feeder 126. The fusible additive is dispersed into the sourcy by the stirrer 122 and constead in the rotary collector 123 in the form of a mixture with the radioactive substance or substances.
- 40 The above-mentioned rotary collector 123 consists of a cylindrical rotary shaft 9 which is provided with a cavity 95 in alignment with the opening 35 at the bottom end of the sedimental tank 3 to receive the condensate in the cavity 95.
- 45 The condensate received in the cavity 9b is dropped into a pot 6 as the cavity 9b is turned 180° about the axis of the shaft 9. Fig. 2 shows a more particular example of the rotary collector 123, in which the collector 123
- 50 includes a metal housing of stainless steel having an annular portion 8a fixedly fitted on the outer periphery of the sedimentation tank 3 and, contiguously to the annular portion 8a, a collecting portion of inverted conical shape
- 55 forming the bottom of the sedimentation tank 3. The housing 8 is provided with an opening 8d at the pointed bottom end of the collecting portion 8b in communication with a conical opening 8d in the bottom wall of the housing 8, through a
- 60 vertical hore 8e formed in alignment with the vertical center line of the sedimentation tank 3. A rotary shaft 9 is journalled in a horizontal bore 87 which is formed in the housing 8 of stainless steel or other metal across the vertical

intermediate portion thereof, a tapered body 9a which is fitted liquid-tight in the horizontal bore 87 to prevent leakage of the slurry. The tapered body portion 9a is centrally provided with a cup-shaped 70 cavity 9b at a position substantially in vertical

- alignment with the opening 8c in the bottom wall of the housing 8, to receive and collect flocks which gravitate through the opening 8c. One end 9c of the rotary shaft 9 is extended
- 75 exially through a cup ring 9d and keyed to a rotational sleeve 14 which is rotatably journalled in bearings 13 within a housing 12 of a rotational drive mechanism 11 fixed on a support frame 10. The rotational sleeve 14 has a worm wheel 15
- 80 fixedly fitted thereon and rotatably driven by a worm shaft 16, which is connected to a motor 10, thereby rotatably driving the collector shaft 9. The other end of the collector shaft 9 is

extended through a cylindrical spring cover 19, 85 which is rotatably and axially slidably fitted in the cylindrical hool 18 fixed on one side wall of the housing 8. An externally threaded end 9e of the shaft is engaged in an internally threaded screw

- member 20 which is supported on an end wall 90 19a of the cover 19. A compression coil spring 23
- having a large spring constant is interposed between the side wall of the housing 8 and a spring seat 22, which is supported on the end wall 19a of the cover 19. A compression coil spring 23
- 95 having a large spring constant is interposed between the side wall of the housing 8 and a spring seat 22, which is supported on the end wall 19a through a bearing 21, coaxially and rotatably relative to the shaft 9, constantly urging the shaft
- 100 9 in the direction indicated by arrow X to maintain the intimate fitting contact between the tapered body 9a of the shaft 9 and the bore 8/. In this instance, it is desirable to have a fitting surface pressure of 1 kg/cm² or greater from the 105 standpoint of secure sealing.

The cavity 9b in the tapered body 9a of the rotary shaft 9 is shaped in an oval form in section, as shown in Fig. 4, with the longer axis of the oval being disposed in the axial direction of the rotary 110 shaft, 9 to give a large allowance to its axial

- alignment with the opening 8c at the bottom of the housing 8. In an initially assembled state, the center Q of the cavity 9b is preferred to be located slightly closer to the divergent end of the bore 8/ 115 in consideration of the friction which would result
 - from the rotation of the rotary shaft 9. In Figs. 2 to 4, the reference numeral 24 denotes a passage which opens into the bottom
- portion 8b of inverted conical shape of the 120 housing 8 and is connected to a discharge pipe 25, as shown in Fig. 3, to discharge a supernatant liquid from the tank 3. Designated at 26 in Figs. 2 and 4 is a handle for manual operation of the rotary shaft 9.
- 125 In operation, the sedimentation tank 3 receives a slurry to be treated, for example, a slurry containing radipactive suspended matter (hereinafter referred to as "clud slurry"), which is filtered out from primary cooling water of a pile or





are iron oxides which are suspended in a concentration of 500—50,000 ppm. When feeding the clud slurry to the sedimentation tank 3, the rotary collector shaft 9

- 5 is retained in the collecting position shown in Figs. 3 and 3. After feeding a predetermined quantity of the Clud slurry to the tank 3, the stirrer is started to mix the clud slurry. For this purpose, the stirrer is preferred to be driven at a speed of 100-200.
- 10 r.p.m. Next, a predetermined amount of flocking agent of high molecular weight is added to the slurry in a concentration of about 2----10 ppm, to flock the suspended substances.
- After continuing the stirring at that speed for a 15 predetermined time period, the stirring speed is dropped to allow small flocks to grow into larger ones, to facilitate the sedimentation of the flocks. Upon lapse of a predetermined time, the stirrer is stopped and the suspension is left to stand for a
- 20 while for sedimentation of the flocks. The precipitated flocks go down the tank, guided by the bottom wall 85 of inverted conical shape, and finally settle in the cavity 95 in the tapered body pontion 9a of the rotary collector shaft 9.
- 25 After sedimentation of a predetermined time period, the supernatant liquid is drawn out and collected by opening a valve (not shown) of the discharge pipe 25. Thereafter, the motor of the rotational drive mechanism 11 is actuated to
- 30 rotate the rotary collecting shaft 9 by 180° through the worm shaft 16 and worm wheel 15, turning the cavity 9b upside down, to drop the sediment into a pot of the rotary transfer mechanism 4, which is located beneath the 35 collector shaft 9.
- As shown in Fig. 5, the rotary transfer mechanism 4 basically includes a center shaft 29 which is rotatably supported on the side frame 5 by a bearing 28 for rotation about a vertical axis, a
- 40 hydraulic cylinder 30 which is coaxially fixed to the upper portion of the center shaft 29, and a rotary head 31, which is attached to the upper end of a plunger 30a of the hydraulic cylinder 30. The rotary head 31 supports thereon four rotary arms
- 45 32 which have their respective base ends supported in bearings for rotation about a horizontal axis, each rotary arm 32 supporting reversibly, at its outer or front end, a pot 6 for receiving the collected sediment.
- 50 A driven gear 33, which is fixedly mounted at an intermediate position on the center shaft 29, is meshed with a drive gear 36 which is driven from a motor 34 through a reducer 35, so that the center shaft 29 is rotated upon actuating the
- 55 motor 34. If necessary, the hydraulic cylinder 30 is 120 securely received in the pot 5, operated by a hydraulic control device 37 to lift or to lower the rotary head 31 through the plunger 30a. Date the position 90° shifted from the locus of rotation of position 90° shifted from the locus of solution solution for the locus of solution solution
- Instead of a mechanism such as power cylinder, 60 e.g. the hydraulic cylinder 30, the lifting and 1
- lowering means may be installed beneath the drier 7 or the rotary collector 123, respectively. The pot 6 is lifted and lowered when, through the rotation, the pot 6 reaches positions where the es pot 6 receives the collected sediment and where inco

the collected sediment in the pot 6 is subjected to drying treatment in the drier 7.

As shown in fig. 7, each pot 6 is in a tapered form with an inside diameter increasing towards

- 70 the upper open end and a round bottom, so that its content is easily released when the pot is turned upside down. Contiguously beneath an antiular groove 6d, the pot 6 is provided with a flanged bottom wall 6d to be fitted with a recess.
- 75 38a of a seat plate 38, which is mounted on the rotary arm 32. The seat plate 38 is provided with a flanged ponion 38b around its outer periphery, on which a locking lever 40 is hinged by a hinge pin 39, rockably in a horizontal plane. As shown in Fig.
- 80 8, a pawl portion 40a at the front end of the locking lever 40 is fitted into the annular groove 6d on the outer periphery of the pot 6. In this manner, the pot 6 and the seat plate 38 are integrally connected with each other by hinge
- 85 pins, which are located at three positions on the outer periphery of the pot 6. As shown in Figs. 7 and 8, a spring 41 tensioned between a spring stop pin 42 fixed at the rear end of the locking lever 40 and a spring stop pin 43 fixed on the
- 90 circumference of the seat plate 38, urges the locking lever 40 in the locking direction. Fixedly secured to the center portion of the seatr plate 38 of the pot 6 is the upper end of a reversing rod 45 which is extended vertically
- 95 through a bore 32a in the front end-portion of the rotary arm 32, and provided with an externally threaded portion 45a at its lower end, in engagement with a weight 46 and a stop nut 47. The pot 6 is thus stably supported on the rotary
- 100 arm 32 by the weight 46, and, when the pot is turned upside down, the contents of the pot are securely released by the falling impact of the weight 46.
- In Figs. 1, 5 and 6, designated at 48 is a circular guide rail which guides the turning movements of the rotary arms 32 on the underside thereof, and which is liftable up and down and normally urged upward by guide rods 49 and r, side cylinders 50. As shown in Fig. 1, the segmentation tank 3
- 110 and drier 7 are located above the locus of rotation of the pots 6 on the rotary arms 32, and in positions spaced from each other by 90° about the axis of rotation, as particularly shown in Fig. 6. In this instance, the lower opening of the rotary 115 collector shaft 9 at the bottom of the
 - sedimentation tank 3 is positioned so that it is brought into alignment with the center axis of a pot 5 which latter pot is turned into a receiving position, ensuring that the dropped sediment is securely received in the pot 6.

On the other hand, the hopper 108 is located buneath the locus of rotation of the pots 6 at a position 90° shifted from the drier 7, so that the dried material in the pot 6 is dropped into the 125 hopper 108 when the pot 6 is turned upside down, as will be described hereinafter.

As shown in Fig. 6, in order to invert the pots 6, a pot-inverting mechanism is provided above the hopper 108, including a knock pin 52 which is



the pots 6 which are successively turned towards the hopper 8. The inverted pot 6 is upturned again by an upturning knock pin 54, similar to the potupsetting knock pin 52, which is supponed on an 5 arm 55 at a position rotationally forward of the

hopper 108.

Referring to Fig. 9 the pot-upsetting pin 52 hits the outer well surface of the pot 6 when the latter is rotated by the rotary arm 32 to a point above

- 10 the hopper 108, and, upon further rotation, the pot 6 is turned upside down together with the rotary arm 32, about the axis thereof, as indicated by the arrow R in Fig. 9. Upon inversion of the pot 6, the weight 46 is allowed to drop freely by a play of the
- 15 rod 45 relative to the rotary arm 32 and hits against the arm, the impact transmitted to the pot 6 encouraging dumping of its contents. The inverted pot 6 then abuts against a pot

upturning pin 54 which, as shown in fig. 10 is located in a position beneath, and spaced by a certain angle from, the pot-inverting pin 52,

- causing the vacant pot 6 and arm 32 to turn clockwise to assume again their original upright position.
- 25 In the operation of the above-described treatment system, with stirring by the stirrer 112, a slurry in the slurry tank 1 is fed to the sedimentation tank 3 through conduits 115 and 118 by operation of the slurry feed pump 116. As
- 30 soon as the slurry in the sedimentation tank 3 reaches a predetermined level, the level switch 120 is actuated to stop the slurry feed pump 116 so as to suspend the feed of the slurry. A predetermined amount of a high molecular weight
- 35 flocking liquid is added to the slurry in the sedimentation tank 3 through the nozzle 121 and mixed therewith by actuating the stirrer 122. Simultaneously, a fusible additive stored in the hopper 125, which is quantitatively fed by the
- 40 feeder 126, is also mixed into the slurry in the tank 3.

In this stirring and mixing stage, the radioactive substances in the slurry are flocked by the high molecular weight flocking liquid and, upon

- 45 stopping the stirrer 122, allowed to fall by gravitation onto the bottom portion 3a of the tank, finally settling in the cavity 3b of the rotary collector 123. In this flocking and settling stage, the fusible additive is also collected together with
- 50 the condensate in the form of a mixture with the radioactive substances.

At the time point when the sedimentation has proceeded to a sufficient degree, the supernatant liquid is drawn out of the tark 3 by opening the

55 electromagnetic valve in the discharge pipe 60 which communicates with the bottom portion 3a of the tank 3, sending the liquid to a waste water treatment process.

After the extraction of the supernatant liquid, 60 the rotary shaft 9 of the rotary collector 123 is

rotated by 180° to drop the condensate in the cavity 95 into a pot 6, which has been lifted to a position close to the cutlet opening 8e of the rotary collector 123 by the operation of the lift During the above-described flocking and collecting operation, a condensate collected by the preceding flocking and collecting operation is dried in the drier 7 until the succeeding mass of

- 70 condensate is received in a vacant pot 6, whereupon the cylinder 30 is lowered once and the motor 34 is actuated to rotate the rotary arm 32 through 90° to bring the received condensate to a position beneath the drier 7. The water vapour 75 which is generated by heating is passed through
- an exhaust gas treatment device like a condenser. As a result of the rotation of the arm 32, the pot 6 which holds the dried condensate is turned
- upside down above the hopper 6 by the pin 52 of 80 the pot-inverting mechanism of Fig. 6, to release the dried condensate into the hopper 8 in the manner explained hereinbefore in connection with Fig. 9. At this time, the dried material which is deposited on the inner wall surface of the pot 6 is
- 85 caused to fall off more forcibly by the impact of the dropping weight 46 than when resort is made to natural or spontaneous fall of the material. After the 90° rotation, the rotary head 31 is
- lifted by the operation of the lift cylinder 30 to 90 position a vacant pot 6 immediately beneath the rotary collector 123 of the sedimentation tank 3, while passing a pot 6 filled with a condensate to the dryer 7 from beneath for heating and drying the condensate.
- 95 During the above-described revolution of the rotary head 31, a pot 6 which passes over the hopper 108 in an inverted state is upturned into an upright position and is stopped in a stand-by position.
- 100 The dried condensate released into the hopper 8, which is a mixture of the radioactive substances and fusible additive, is fed into the melting furnace 110 by the feeder 9. The mixture which is melted in the furnace 110 is solidified and put in a storage

105 container, which is then capped, sealed and cleaned of contaminants for storage over a long time period.

The condensate in the pots, which is heated and dried by a single dryer in the foregoing 110 embodiment, may be adapted to be preheated and then fully heated by driers which are located in two separate positions in the rotational path of travel.

Further, although the fusible additive is 115 admixed to the slurry in the sedimentation tank, it may be fed to, and mixed with, the condensate in the hopper 108 prior to charging to the melting furnace 110.

As appears from the foregoing description, the 120 present invention employs a rotary mechanism which includes in its rotational path of travel a number of stages required for treating a slurry of radioactive substances, i.e., a stage of condensation of the slurry, a stage of heating and

125 drying the condensate and a stage of transfer to a melting furnace, prior to the final melting and solidifying stages, thereby allowing treatment of radio-active waste continuously in one place. According to the invention, the waste is

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respective stages of treatment by the operation of a single rotary mechanism, so that it becomes possible to enhance the capacity of treatment and to reduce the floor space required for the

- 5 treatment operation. In addition, the rotary mechanism with liftable pots simplifies the treatment operation and permits of remote control or complete automation of the treatment operation.
- 10 CLAIMS
 - Apparatus for the treatment of radioactive waste, comprising:
 - a storage tank holding a slurry of the radioactive waste;
- 15 a sedimentation tank for thickening the slurry of radioactive waste received from said storage tank; a drier for drying the thickened radioactive waste:
- a hopper for receiving the dried radioactive 20 waste; and
- a rotary mechanism for transferring the radioactive waste to and from the sedimentation tank, drier and hopper, said mechanism having a number of containers for holding the radioactive
- 25 waste, rotary arms for supporting said containers: a rotational drive mechanism for said rotary arms, and a lift mechanism for vertically lifting said containers up and down.
- 2. Apparatus as set forth in claim 1, wherein 30 said sedimentation tank and drier are located
- successively above the rotational path of travel of said containers, and said hopper is located beneath said rotational path of travel in a position spaced from said crier by a predetermined angle
- 35 about the axis of rotation of said rotation

mechanism.

 Apparatus as set forth in claim 1 or 2, further comprising means for inverting and upturning said containers, provided respectively before and after
said hopper in the rotational path of travel of said

containers. 4. Apparatus as set form in claim 1, 2 or 3, wherein said sedimentation tank is provided with a rotary collector including a housing block having

- 45 a vertical bore forming inlet and outlet openings on the upper and lower side thereof in alignment with an opening at the bottom of said sedimentation tank and a tapered bore formed perpendicularly to, and across, said vertical bore, a
- 50 rotary shaft having a tapered body portion fitted in said tapered bore and provided with a cavity in said tapered body portion in alignment with said inlet opening for collecting radioactive sediment, a spring urging one end of said rotary shaft in the
- 55 tapered direction thereof, and a rotational drive mechanism connected to the other end of said rotary shaft for rotating the latter through a predetermined angle, the radioactive sediment collected in said cavity being dropped through said
- 60 outlet opening upon rotation of said rotary shaft. 5. Apparatus as set forth in claim 1, 2, 3 or 4, wherein said lift mechanism is so designed as to lift said rotary arms up and down.

6. Apparatus as set forth in claim 1, 2, 3 or 4.

- 65 wherein said lift mechanism is installed beneath said drier or said rotary collector whereby said container is lifted up and down when said container comes beneath said drier or said rotary collector.
- 70 7. Apparatus as set forth in claim 1 substantially as hereinbefore described with reference to the accompanying drawings.

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