

INTERAGENCY AGREEMENT

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1. INTERAGENCY AGREEMENT NO. NRC-06-05-301		2. EFFECTIVE DATE 06/03/2005		3. PROJECT/REQUISITION NO. STP05301	
4. ISSUED BY U.S. Nuclear Regulatory Commission Div. of Contracts Attn: Sharlene McCubbin Mail Stop T-7-I-2 Washington, DC 20555 Sharlene McCubbin 301-415-6565			5. NAME, TITLE & PHONE NUMBER OF EACH GOVERNMENT PROJECT OFFICER FOR BOTH AGENCIES Kathleen Schneider (301) 415-2320 (Project Officer) Penny Boyce (301)594-3650 (Other Agency's Project Monitor)		
6. AGENCY PERFORMING SERVICE FOOD & DRUG ADMINISTRATION 5600 FISHERS LANE HF-1 ROCKVILLE MD 208570001					
7. PROJECT TITLE Partial Suppor for the FDA Cooperative Agreement with the recipient of the Grant for Assuring Radiation Protection					
8. PROJECT OBJECTIVE See Pages 2 through 5 for remaining pages of IA.					
9. PERFORMANCE PERIOD OF AGREEMENT 06-02-2005 through 04-30-2006					
10. ACCOUNTING AND APPROPRIATION DATA 31X0200 B&R: 57Q15344244 BOC: 252A JCN: L2337 OBLIGATE: \$110,000.00				11. DOLLAR VALUE OF AGREEMENT 110000.00	
12. FUNDING (The Issuing Agency agrees to advance/reimburse funds up to the dollar amount of this agreement, upon receipt of a properly executed Standard Form 1080 or 1081. The appropriate form must be executed in original and _____ copies, be identified with agreement number and accounting data, and transmitted to the following office for funds action.) U.S. Nuclear Regulatory Commission Payment Team, Mail Stop T-9-H-4 Washington, DC 20555 Attn: (insert contract or order nu					
(Any funds not utilized for the performance of the work described in this agreement must be returned to the Issuing Agency.)					
13. PURSUANT TO THE AUTHORITY OF The Economy Act of 1932; Sect. 301 of the PHS Act 42 USC 241					
(The Issuing Agency may enter into this agreement. The person executing this agreement has a written delegation of authority to do so on behalf of the agency.)					
14. SIGNATURE OF PERSON AUTHORIZED TO SIGN 		DATE 8/10/05	16. SIGNATURE OF PERSON AUTHORIZED TO SIGN 		DATE 6-6-05
15. TYPE NAME & TITLE OF PERSON AUTHORIZED TO SIGN			17. TYPE NAME & TITLE OF PERSON AUTHORIZED TO SIGN Mary H. Mace Contracting Officer		

TEMPLATE - ADM001

SISP REVIEW COMPLETE

ADM002

Pursuant to the Economy Act of 1932, as amended, the U.S. Nuclear Regulatory Commission (NRC) and the Food and Drug Administration (FDA), Bureau of Radiological Health (BRH) desire to enter into an agreement for the support of the Conference of Radiation Control Program Directors (CRCPD). FDA Authority: Section 301 of the Public Health Service Act (42 USC 241).

The CRCPD consists of radiation control program directors from each of the 50 states and the District of Columbia. It was created to unify the States in reducing unnecessary exposure of the public to ionizing and nonionizing radiation. The services of these radiation control experts in the conference is used to assist the Federal Government in identification of, resolution and implementation of solutions for radiation control problems.

The CRCPD coordinates State radiation control activities with the programs of the sponsoring Federal agencies (NRC, FDA-BRH, EPA), assists in maintaining a national data bank of radiation control activities and participates in formulation of regulations, standards, and suggested legislation.

Accordingly upon your agreement as provided below, the terms and conditions of this interagency agreement are as follows:

Article I - Scope of Work

Under the proposed agreement, FDAA-BRH shall perform in accordance with the Scope of Work entitled "Partial Support for the FDA Cooperative Agreement with the Recipient of the Grant for Assuring Radiation Protection," which is attached hereto and made a part of this interagency agreement.

Article II - Deliverables

Deliverables including, but not limited to, reports and newsletters shall be submitted in accordance with Article 1, of the attached Statement of Work.

Article III - Period of Performance

The period of performance of this agreement shall be from June 06, 2005 - April 30, 2006.

Article IV - Payment

Payment shall be made on a reimbursable basis. Invoices shall be submitted in accordance with Attachment 1, Billing Instructions for Interagency Agreements, which is attached hereto and made a part hereof.

Each invoice shall cite the following data:

U.S. NRC
APPN No.: 31X0200
B&R: 57Q15344244
BOC: 252A
JCN: L2337

Article V - Obligation of Funds

The amount presently obligated by the NRC for this agreement for performance of work is \$110,000.00.

Article IV - Points of Contact

The NRC contacts are:

Technical:	Kathleen N. Schneider, Project Officer (301) 415-2320, Mail Stop: O-3C10
Contractual:	Sharlene McCubbin, Contract Specialist (301) 415-6565, Mail T-712

The FDA contacts are:

Grants Officer:	Rosemary T. Springer (301) 827-7182
Administrative :	Penny Boyce (301) 594-3650

If this agreement is acceptable to FDA-BRH, please so indicate by signing in the space provided below and on Block 14 and 15 of Page 1 of the IA form. Return two fully executed copies to me at the address below. You may retain the third copy for your records.

U.S. Nuclear Regulatory Commission
ATTN: Sharlene McCubbin, Contract Specialist
Division of Contracts, MS T-712 c
Office of Administration
Washington, DC 20555

Should you have any questions regarding this Interagency Agreement, please contact Sharlene McCubbin, Contract Specialist on 301-415-6565.

Sincerely,



Mary H. Mace, Contracting Officer
Contract Management Branch 1
Division of Contracts
Office of Administration

Attachment: As stated

ACCEPTED: FOOD AND DRUG ADMINISTRATION

BY: _____

TITLE: _____

DATE: _____

SCOPE OF WORK

224.76.6006

Interagency Agreement Between the Nuclear Regulatory Commission and the Food and Drug Administration

TITLE: PARTIAL SUPPORT FOR THE FDA COOPERATIVE AGREEMENT WITH THE RECIPIENT OF THE GRANT FOR ASSURING RADIATION PROTECTION

I. BACKGROUND -

- (A) NRC will contribute approximately \$110,000 for FY05 towards a U.S. Food and Drug Administration (FDA) Cooperative Agreement with the recipient of the grant for Assuring Radiation Protection. Other participating agencies include DOE, FEMA, EPA and FDA (HHS). The FDA Project Officer will be responsible for completion of all work, and will handle all accounting and financial aspects of the agreement. NRC will receive technical reports of the committees either through distribution by the recipient or through planning meetings of the central management board or committee of the recipient.
- (B) The NRC Federal Liaison, located in the Office of State and Tribal Programs, will regularly attend the planning meetings of the central management board or committee, and will communicate with other NRC members of the task forces and related committees. The NRC Federal Liaison also regularly attends the annual meeting, which is usually held in May, to address national radiation protection issues. The annual meeting will last approximately one week with approximately 350 individuals attending. The NRC, in coordination with the FDA Project Officer, and recipient will also communicate on major policy and regulatory issues, such as suggested state regulations covering radioactive materials, low-level waste, radioactive contamination and emergency response planning.

II. DUTIES

Under the direction of the FDA Project Officer, the grant recipient will accomplish the following on behalf of the NRC:

- A. Perform all administrative management functions required for the conduct of an annual technical meeting of Federal and State officials and sponsors including site visits and selections, program planning, and facility arrangement. Conduct the annual meeting consisting of a general session, special sessions and training sessions for the following purposes:
 - A.1 General Sessions: To present reports on the status of studies conducted during previous years, technical reports on new developments and task force reports of problems defined and assigned as recommended by the committee reports, grant recipient or sponsors.

- A.2 **Special Sessions:** To study and define identified problems and/or areas of mutual concern in radiation control and recommend action as needed to resolve these problems. The special session reports are submitted to the recipient for cost benefit evaluation and further action. Problems that require extensive study are assigned to a task force or committee comprised of experts in that field for solving.
- A.3 **Training Sessions:** To demonstrate mutually beneficial techniques, procedures and systems which have been developed by the sponsoring agencies or the recipient.
- B. Appoint and arrange for standing committees and task forces to study, evaluate, develop recommended actions and/or solutions to current identified problems and provide guidance and assistance to sponsoring agencies. Although, the recipient may, at any given time, have a number of groups working on specific projects, the recipient shall set priorities annually for specific committees and products. Reports generated by these task forces will be evaluated by the central management board or committee of the recipient and reported to the participating agencies. The central management board or committee of the recipient will also set priorities for the standing committees and task forces in coordination with the FDA and its participating agencies. All administrative management functions required to support these task forces will be performed by the recipient. Work by the recipient can be broken down into the following broad categories that are related to NRC responsibilities:
 - B.1 **Suggested State Regulations:** To assist State regulatory agencies in developing radiation control regulations for radioactive materials regulatory programs which will promote uniformity between the States;
 - B.2 **Environmental Nuclear:** Radioactive waste disposal, radioactive contamination, contaminated sites, emergency response planning, bonding and surety, and decontamination and decommissioning;
 - B.3 **General Radiation Protection:** Ionizing radiation safety concerns, international radiation protection, industrial, medical and other uses of radioactive material.
 - B.4 **Training:** Conduct activities on training to evaluate the RCP training and program implementation training needs of the States. Continue to review these needs and assess the training requirements that must be approached from the national level to solve problems encountered. Work with the sponsoring or other Federal agencies, universities, manufacturers, or other resources to develop the training needed in radiation control.

- C. Conduct an orphan sources program which will include the following:
 - C.1 Clarify jurisdictions and regulatory responsibilities for addressing orphan source problems, including providing assistance for identification, handling, and disposal of orphaned sources.
 - C.2 Complete development, establishment, implementation, and management of a national orphan source management program, including:
 - (1) establishing a definition for discrete orphan sources, which would exclude diffuse radioactive material;
 - (2) establishing agreements with interested Agreement and non-Agreement States, covering identification and disposal of discrete orphan sources;
 - (3) communication and coordination with Federal agencies, State Agencies, and other stakeholders on matters related to orphan source issues;
 - (4) establishing cost guidelines for disposal of discrete orphan sources; and
 - (5) reimbursement to States for recovery, recycle, arrangements for re-use, and disposal costs of discrete orphan sources, which are subject to NRC or Agreement State jurisdiction under the Atomic Energy Act, but excluding reimbursement for disposal of sources which fall under the category of "greater than Class C" waste.
 - C.3 Provide a annual report, which should include, but not be limited to the number of agreements entered into with Agreement and non-Agreement States, the number and type of sources dispositioned, and the means and cost to recover and disposition the sources.
- D. Predecisional information provided to the recipient by the NRC will be limited to use by Agreement and non-Agreement State Radiation Control Program Directors, their staff, recipient staff, and Federal resource representatives. This information will not be disclosed to the public and non-State members of committees or task forces unless NRC through the FDA Project Officer, gives permission.
- E. Provide recipient's attendance at stakeholder meetings and Radiation Advisory Committee meetings of supporting agencies to provide State technical experience and views into the subject under advisement.
- F. The major products of the recipient are technical publications prepared by the recipient's working groups and formal positions taken by the membership and/or the central management board or committee of the recipient including:
 - 1. Proceedings of Annual Meetings
 - 2. Publication of the Suggested State Regulations for Control of Radiation Control (SSRCR) and
 - 3. Annual publication of Directory of Personnel Responsible for Radiological Health Programs.

- G. When the Directory of Personnel Responsible for Radiological Health Programs is published, without cost submit 40 copies of the Directory and a non-copyrighted electronic copy of the Directory on disk to the NRC Federal Liaison. Upon availability, delivery of documents specified in C.3, F. and G. will be coordinated through the FDA Project Officer.