

From: Thomas O'Brien
To: sns,bgu
Date: 12/18/97 11:18am

Steve-

I understand you will consolidate the UTAH review (#78326).

I reviewed sections R313-15 and R313-18. I determined that these regulations are compatible and therefore have no comments.

[As a side note, however, Utah's R313-15-1201 section adopts the proposed wording provided in the August 1997 draft SSR equivalent part. Discussion w/ Pat Larkins indicated this is acceptable even though it is much more stringent. We have not yet approved this draft SSR (D. Blanton is in the process). My concern is if we do NOT approve the draft SSR wording, this may conflict with the finding of compatible for the Utah regs.]

Per discussion w/ Bangert, O'Brien & Blanton, this section is H+S + meets the essential objectives.

Brenda-

Please close out my ticket on this task.

For your RATS:

The revisions to Utah's R313-15 implement the NRC rules listed below:

- Resolution of Dual Regulation of Airborne Effluents of Radioactive Materials (all parts)
- Criteria for Release of Individuals Administered Radioactive Material (all parts)
- Frequency of Medical Examinations for Use of Respiratory Protection Equipment (all parts)
- Radiation protection requirements: Amended Definitions and Criteria (all parts)
- Low Level Waste Manifest (Part 20 only)
- Medical Administration of Radiation and Radioactive Materials (part 20 only)
- Termination or transfer of licensed activities: Housekeeping requirements (part 20 only)

cc:

tjo



State of Utah

DEPARTMENT OF ENVIRONMENTAL QUALITY DIVISION OF RADIATION CONTROL

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November 24, 1997

Charles Hackney
State Agreements Program
Nuclear Regulatory Commission
611 Ryan Plaza Drive Suite 400
Arlington, Texas 76011

*O'Brien D'Brien
(Part 19) (Part 20)*

*Bolling
(Parts 20, 35)*

Dear Mr. Hackney:

Rollaway
The Division of Radiation Control (DRC) is currently working on a number of rulemaking actions which are compatibility items. We are proposing to change R313-12, General Provisions; R313-15, Standards for Protection Against Radiation; R313-18, Notices, Instructions and Reports to Workers by Licensees or Registrants--inspections; R313-25, License Requirements for Land Disposal of Radioactive Waste--General Provision; and R313-32 Medical Use of radioactive Material.
(Part 6) (Parts 20 + 35) Bolling

Our rulemaking process involves the Utah Radiation Control Board. After presentation to the Board, the rules are filed with the Utah Division of Administrative Rules. At this time, a 30 day public comment period is expected to open on December 1, 1997. The DRC expects to finalize and adopt the rules about January 23, 1998.

Please find enclosed a hard copy and a WordPerfect (version 6.1) electronic copy of the proposed rules. New text is shown with an underline. Deleted text is interlined and placed within brackets. If you have questions, please contact Craig Jones at (801) 536-4250.

Sincerely,

William J. Sinclair
William J. Sinclair, Director
Division of Radiation Control

Enclosure: As stated



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File Name: Hackney3.ltr
File Location: cwj/wp

R313. Environmental Quality, Radiation Control.

R313-18. Notices, Instructions and Reports to Workers by Licensees or Registrants--Inspections.

R313-18-1. Purpose and Authority.

(1) The purpose of this rule is to establish requirements for notices, instructions and reports by licensees or registrants to individuals engaged in work under a license or registration and options available to such individuals in connection with inspections of licensees or registrants.

(2) The rules set forth herein are adopted pursuant to the provisions of Sections 19-3-104(3) and 19-3-104(6).

R313-18-2. General.

The rules of R313-18 shall apply to all persons who receive, possess, use, own or transfer a source of radiation licensed by or registered with the Department pursuant to the rules in R313-16, R313-19 or R313-22.

R313-18-11. Posting of Notices to Workers.

(1) Licensees or registrants shall post current copies of the following documents:

(a) the rules in R313-15 and R313-18;

(b) the license, certificate of registration, conditions or documents incorporated into the license by reference and amendments thereto;

(c) the operating procedures applicable to work under the license or registration; and

(d) a notice of violation involving radiological working conditions, proposed imposition of civil penalty, order issued pursuant to R313-14, or any response from the licensee or registrant.

(2) If posting of a document specified in R313-18-11(1)(a), (b), or (c)

1 is not practicable, the licensee or registrant may post a notice which describes
2 the document and states where it may be examined.

3 (3) DRC-04 "Notice to Employees," shall be posted by licensees or
4 registrants wherever individuals work in or frequent a portion of a restricted
5 area.

6 (4) Documents from the Executive Secretary which are posted pursuant to
7 R313-18-11(1)(d) shall be posted within five working days after receipt of the
8 documents from the Executive Secretary; the licensee's or registrant's response,
9 if there is one, shall be posted for a minimum of five working days after
10 dispatch from the licensee or registrant. The documents shall remain posted for
11 a minimum of five working days or until action correcting the violation has been
12 completed, whichever is later.

13 (5) Documents, notices or forms posted pursuant to R313-18-11 shall appear
14 in a sufficient number of places to permit individuals engaged in work under the
15 license or registration to observe them on the way to or from any particular work
16 location to which the document applies, shall be conspicuous, and shall be
17 replaced if defaced or altered.

18
19 **R313-18-12. Instructions to Workers.**

20 (1) All individuals who in the course of employment are likely to receive
21 in a year an occupational dose in excess of 1.0 mSv (100 mrem):

22 (a) shall be kept informed of the storage, transfer, or use of sources of
23 radiation in the licensee's or registrant's workplace;

24 (b) shall be instructed in the health protection considerations associated
25 with exposure to radiation or radioactive material to the individual and
26 potential offspring, in precautions or procedures to minimize exposure, and in
27 the purposes and functions of protective devices employed;

28 (c) shall be instructed in, and instructed to observe, to the extent

1 within the worker's control, the applicable provisions of these rules and
2 licenses for the protection of personnel from exposure to radiation or
3 radioactive material;

4 (d) shall be instructed as to their responsibility to report promptly to
5 the licensee or registrant a condition which may constitute, lead to, or cause
6 a violation of the Act, these rules, and licenses or unnecessary exposure to
7 radiation or radioactive material;

8 (e) shall be instructed in the appropriate response to warnings made in
9 the event of an unusual occurrence or malfunction that may involve exposure to
10 radiation or radioactive material; and

11 (f) shall be advised as to the radiation exposure reports which workers
12 shall be furnished pursuant to R313-18-13.

13 (2) In determining those individuals subject to the requirements of R313-
14 18-12(1), licensees must take into consideration assigned activities during
15 normal and abnormal situations involving exposure to radiation or radioactive
16 material which can reasonably be expected to occur during the life of a licensed
17 facility. The extent of these instructions shall be commensurate with potential
18 radiological health protection considerations for the workplace.

19
20 **R313-18-13. Notifications and Reports to Individuals.**

21 (1) Radiation exposure data for an individual and the results of
22 measurements, analyses, and calculations of radioactive material deposited and
23 retained in the body of an individual shall be reported to the individual as
24 specified in R313-18-13. The information reported shall include data and results
25 obtained pursuant to these rules, orders, or license conditions, as shown in
26 records maintained by the licensee or registrant pursuant to R313-15-1107.
27 Notifications and reports shall:

28 (a) be in writing;

1 (b) include appropriate identifying data such as the name of the licensee
2 or registrant, the name of the individual, and the individual's identification
3 number, preferably social security number;

4 (c) include the individual's exposure information; and

5 (d) contain the following statement:

6 "This report is furnished to you under the provisions of the Utah
7 Administrative Code Section R313-18-13. You should preserve this report for
8 further reference."

9 (2) Licensees or registrants shall ~~(advise)~~ furnish to each worker annually
10 a written report of the worker's dose as shown in records maintained by the
11 licensee or registrant pursuant to R313-15-1107.

12 (3) Licensees or registrants shall furnish a written report of the
13 worker's exposure to sources of radiation at the request of a worker formerly
14 engaged in activities controlled by the licensee or registrant. The report shall
15 include the dose record for each year the worker was required to be monitored
16 pursuant to R313-15-502. The report shall be furnished within 30 days from the
17 date of the request, or within 30 days after the dose of the individual has been
18 determined by the licensee or registrant, whichever is later. The report shall
19 cover the period of time that the worker's activities involved exposure to
20 sources of radiation and shall include the dates and locations of work under the
21 license or registration in which the worker participated during this period.

22 (4) When a licensee or registrant is required pursuant to R313-15-1202,
23 R313-15-1203, or R313-15-1204 to report to the Executive Secretary an exposure
24 of an individual to sources of radiation, the licensee or the registrant shall
25 also provide the individual a written report on the exposure data included
26 therein. Reports shall be transmitted at a time not later than the transmittal
27 to the Executive Secretary.

28 (5) At the request of a worker who is terminating employment with the

1 licensee or registrant in work involving exposure to radiation or radioactive
2 material, during the current year, the licensee or registrant shall provide at
3 termination to the worker, or to the worker's designee, a written report
4 regarding the radiation dose received by that worker from operations of the
5 licensee or registrant during the current year or fraction thereof. If the most
6 recent individual monitoring results are not available at that time, a written
7 estimate of the dose shall be provided together with a clear indication that this
8 is an estimate.

9
10 **R313-18-14. Presence of Representative of Licensees or Registrants and Workers**
11 **During Inspection.**

12 (1) Licensees or registrants shall afford representatives of the Board or
13 the Executive Secretary, at reasonable times, the opportunity to inspect
14 materials, machines, activities, facilities, premises, and records pursuant to
15 these rules.

16 (2) During an inspection, representatives of the Board or the Executive
17 Secretary may consult privately with workers as specified in R313-18-15. The
18 licensee or registrant may accompany representatives during other phases of an
19 inspection.

20 (3) If, at the time of inspection, an individual has been authorized by
21 the workers to represent them during Department inspections, the licensee or
22 registrant shall notify the representatives of the Board or the Executive
23 Secretary of the authorization and shall give the workers' representative an
24 opportunity to accompany the representatives during the inspection of physical
25 working conditions.

26 (4) The workers' representative shall be routinely engaged in work under
27 control of the licensee or registrant and shall have received instructions as
28 specified in R313-18-12.

1 (5) Different representatives of licensees or registrants and workers may
2 accompany the representatives of the Board or the Executive Secretary during
3 different phases of an inspection if there is no resulting interference with the
4 conduct of the inspection. However, only one workers' representative at a time
5 may accompany the representatives of the Board or the Executive Secretary.

6 (6) With the approval of the licensee or registrant and the workers'
7 representative, an individual who is not routinely engaged in work under control
8 of the licensee or registrant, for example, a consultant to the licensee or
9 registrant or to the workers' representative, shall be afforded the opportunity
10 to accompany representatives of the Board or the Executive Secretary during the
11 inspection of physical working conditions.

12 (7) Notwithstanding the other provisions of R313-18-14, representatives
13 of the Board or the Executive Secretary are authorized to refuse to permit
14 accompaniment by an individual who deliberately interferes with a fair and
15 orderly inspection. With regard to areas containing information classified by
16 an Agency of the U.S. Government in the interest of national security, an
17 individual who accompanies an inspector may have access to such information only
18 if authorized to do so. With regard to areas containing proprietary information,
19 the workers' representative for that area shall be an individual previously
20 authorized by the licensee or registrant to enter that area.

21
22 **R313-18-15. Consultation with Workers During Inspections.**

23 (1) Representatives of the Board or the Executive Secretary may consult
24 privately with workers concerning matters of occupational radiation protection
25 and other matters related to applicable provisions of these rules and licenses
26 to the extent the representatives deem necessary for the conduct of an effective
27 and thorough inspection.

28 (2) During the course of an inspection, workers may bring privately to the

1 attention of the representatives of the Board or the Executive Secretary, either
2 orally or in writing, a past or present condition which the worker has reason to
3 believe may have contributed to or caused a violation of the Act, these rules,
4 or license condition, or an unnecessary exposure of an individual to sources of
5 radiation under the licensee's or registrant's control. A notice in writing
6 shall comply with the requirements of R313-18-16(1).

7 (3) The provisions of R313-18-15(2) shall not be interpreted as
8 authorization to disregard instructions pursuant to R313-18-12.

9
10 **R313-18-16. Request by Workers for Inspections.**

11 (1) A worker or representative of workers believing that a violation of
12 the Act, these rules, or license conditions exists or has occurred in work under
13 a license or registration with regard to radiological working conditions in which
14 the worker is engaged, may request an inspection by giving notice of the alleged
15 violation to the Executive Secretary. The notice shall be in writing, shall set
16 forth the specific grounds for the notice, and shall be signed by the worker or
17 representative of the workers. A copy shall be provided to the licensee or
18 registrant by representatives of the Board or the Executive Secretary no later
19 than at the time of inspection except that, upon the request of the worker giving
20 the notice, his name and the name of individuals referred to therein shall not
21 appear in a copy or on a record published, released, or made available by the
22 Department except for good cause shown.

23 (2) If, upon receipt of the notice, representatives of the Board or the
24 Executive Secretary, determine that the complaint meets the requirements set
25 forth in R313-18-16(1), and that there are reasonable grounds to believe that the
26 alleged violation exists or has occurred, an inspection shall be made as soon as
27 practicable to determine if the alleged violation exists or has occurred.
28 Inspections pursuant to R313-18-16 need not be limited to matters referred to in

1 the complaint.

2 3019(b) (3) A licensee, registrant or contractor or subcontractor of a licensee
3 or registrant shall not discharge or discriminate against a worker because that
4 worker has filed a complaint or instituted or caused to be instituted a
5 proceeding under these rules or has testified or is about to testify in a
6 proceeding or because of the exercise by the worker on behalf of the worker or
7 others of an option afforded by R313-18.

8
9 **R313-18-17. Inspections Not Warranted -- Informal Review.**

10 (1)(a) If the representatives of the Board or the Executive Secretary
11 determine, with respect to a complaint under Section R313-18-16, that an
12 inspection is not warranted because there are no reasonable grounds to believe
13 that a violation exists or has occurred, the Executive Secretary shall notify the
14 complainant in writing of that determination. The complainant may obtain review
15 of the determination by submitting a written statement of position with the
16 Executive Secretary. The Executive Secretary will provide the licensee or
17 registrant with a copy of the statement by certified mail, excluding, at the
18 request of the complainant, the name of the complainant. The licensee or
19 registrant may submit an opposing written statement of position with the
20 Executive Secretary. The Executive Secretary will provide the complainant with
21 a copy of the statement by certified mail.

22 (b) Upon the request of the complainant, the Board may hold an informal
23 conference in which the complainant and the licensee or registrant may orally
24 present their views. An informal conference may also be held at the request of
25 the licensee or registrant, but disclosure of the identity of the complainant
26 will be made only following receipt of written authorization from the
27 complainant. After considering written and oral views presented, the Board shall
28 affirm, modify, or reverse the determination of the representatives of the Board

1 or the Executive Secretary and furnish the complainant and the licensee or
2 registrant a written notification of the decision and the reason therefor.

3 (2) If the Executive Secretary determines that an inspection is not
4 warranted because the requirements of R313-18-16(1) have not been met, the
5 complainant shall be notified in writing of the determination. The determination
6 shall be without prejudice to the filing of a new complaint meeting the
7 requirements of R313-18-16(1).

8
9 KEY: radioactive material, inspection, radiation safety, licensing

10 [~~1993~~]1998

19-3-104

11 Notice of Continuation March 26, 1997

19-3-108

R313. Environmental Quality, Radiation Control.

R313-15. Standards for Protection Against Radiation.

R313-15-1. Purpose, Authority and Scope.

(1) R313-15 establishes standards for protection against ionizing radiation resulting from activities conducted pursuant to licenses issued by the Executive Secretary. These rules are issued pursuant to Sections 19-3-104(3) and 19-3-104(6).

(2) The requirements of Rule R313-15 are designed to control the receipt, possession, use, transfer, and disposal of sources of radiation by any licensee or registrant so the total dose to an individual, including doses resulting from all sources of radiation other than background radiation, does not exceed the standards for protection against radiation prescribed in Rule R313-15. However, nothing in Rule R313-15 shall be construed as limiting actions that may be necessary to protect health and safety.

(3) Except as specifically provided in other sections of these rules, Rule R313-15 applies to persons licensed or registered by the Executive Secretary to receive, possess, use, transfer, or dispose of sources of radiation. The limits in Rule R313-15 do not apply to doses due to background radiation, to exposure of patients to radiation for the purpose of medical diagnosis or therapy, to exposure from individuals administered radioactive material and released in accordance with Section R313-32-75, or to exposure from voluntary participation in medical research programs.

R313-15-2. Definitions.

"Annual limit on intake" (ALI) means the derived limit for the amount of radioactive material taken into the body of an adult worker by inhalation or ingestion in a year. ALI is the smaller value of intake of a given radionuclide in a year by the reference man that would result in a committed effective dose

1 equivalent of 0.05 Sv (5 rem) or a committed dose equivalent of 0.5 Sv (50 rem)
2 to any individual organ or tissue. ALI values for intake by ingestion and by
3 inhalation of selected radionuclides are given in Table I, Columns 1 and 2, of
4 Appendix B of 10 CFR 20.1001 to 20.2402, 199[3]2 ed., which is incorporated by
5 reference.

6 "Class" means a classification scheme for inhaled material according to its
7 rate of clearance from the pulmonary region of the lung. Materials are
8 classified as D, W, or Y, which applies to a range of clearance half-times: for
9 Class D, Days, of less than ten days, for Class W, Weeks, from ten to 100 days
10 , and for Class Y, Years, of greater than 100 days. For purposes of these rules,
11 "lung class" and "inhalation class" are equivalent terms.

12 "Constraint (dose constraint)" means a value above which specified licensee
13 actions are required.

14 "Declared pregnant woman" means a woman who has voluntarily informed her
15 employer, in writing, of her pregnancy and the estimated date of conception.

16 "Derived air concentration" (DAC) means the concentration of a given
17 radionuclide in air which, if breathed by the reference man for a working year
18 of 2,000 hours under conditions of light work, results in an intake of one ALI.
19 For purposes of these rules, the condition of light work is an inhalation rate
20 of 1.2 cubic meters of air per hour for 2,000 hours in a year. DAC values are
21 given in Table I, Column 3, of Appendix B of 10 CFR 20.1001 to 20.2402, 199[3]2
22 ed., which is incorporated by reference.

23 "Derived air concentration-hour" (DAC-hour) means the product of the
24 concentration of radioactive material in air, expressed as a fraction or multiple
25 of the derived air concentration for each radionuclide, and the time of exposure
26 to that radionuclide, in hours. A licensee or registrant may take 2,000 DAC-
27 hours to represent one ALI, equivalent to a committed effective dose equivalent
28 of 0.05 Sv (5 rem).

1 "Dosimetry processor" means an individual or an organization that processes
2 and evaluates individual monitoring devices in order to determine the radiation
3 dose delivered to the monitoring devices.

4 "Inhalation class", refer to "Class".

5 "Labeled package" means a package labeled with a Radioactive White I,
6 Yellow II, or Yellow III label as specified in U.S. Department of Transportation
7 regulations 49 CFR 172.403 and 49 CFR 172.436 through 440, 199[9]7 ed. Labeling
8 of packages containing radioactive materials is required by the U.S. Department
9 of Transportation if the amount and type of radioactive material exceeds the
10 limits for an excepted quantity or article as defined and limited by U.S.
11 Department of Transportation regulations 49 CFR 173.403(m) and (w) and 49 CFR
12 173.421 through 424, 199[9]7 ed.

13 "Lung class", refer to "Class".

14 "Nonstochastic effect" means a health effect, the severity of which varies
15 with the dose and for which a threshold is believed to exist. Radiation-induced
16 cataract formation is an example of a nonstochastic effect. For purposes of
17 these rules, "deterministic effect" is an equivalent term.

18 "Planned special exposure" means an infrequent exposure to radiation,
19 separate from and in addition to the annual occupational dose limits.

20 "Quarter" means a period of time equal to one-fourth of the year observed
21 by the licensee, approximately 13 consecutive weeks, providing that the beginning
22 of the first quarter in a year coincides with the starting date of the year and
23 that no day is omitted or duplicated in consecutive quarters.

24 "Reference Man" means a hypothetical aggregation of human physical and
25 physiological characteristics determined by international consensus. These
26 characteristics may be used by researchers and public health ~~[workers]~~employees
27 to standardize results of experiments and to relate biological insult to a common
28 base. A description of the Reference Man is contained in the International

1 Commission on Radiological Protection report, ICRP Publication 23, "Report of the
2 Task Group on Reference Man."

3 "Respiratory protective equipment" means an apparatus, such as a
4 respirator, used to reduce an individual's intake of airborne radioactive
5 materials.

6 "Sanitary sewerage" means a system of public sewers for carrying off waste
7 water and refuse, but excluding sewage treatment facilities, septic tanks, and
8 leach fields owned or operated by the licensee or registrant.

9 "Stochastic effect" means a health effect that occurs randomly and for
10 which the probability of the effect occurring, rather than its severity, is
11 assumed to be a linear function of dose without threshold. Hereditary effects
12 and cancer incidence are examples of stochastic effects. For purposes of these
13 rules, "probabilistic effect" is an equivalent term.

14 "Very high radiation area" means an area, accessible to individuals, in
15 which radiation levels could result in an individual receiving an absorbed dose
16 in excess of five Gy (500 rad) in one hour at one meter from a source of
17 radiation or from any surface that the radiation penetrates. At very high doses
18 received at high dose rates, units of absorbed dose, gray and rad, are
19 appropriate, rather than units of dose equivalent, sievert and rem.

20 "Weighting factor" w_T for an organ or tissue (T) means the proportion of
21 the risk of stochastic effects resulting from irradiation of that organ or tissue
22 to the total risk of stochastic effects when the whole body is irradiated
23 uniformly. For calculating the effective dose equivalent, the values of w_T are:

24
25 TABLE

26
27 ORGAN DOSE WEIGHTING FACTORS

Organ or Tissue	w_T
Gonads	0.25
Breast	0.15
Red bone marrow	0.12
Lung	0.12
Thyroid	0.03
Bone surfaces	0.03
Remainder	0.30(1)
Whole Body	1.00(2)

(1) 0.30 results from 0.06 for each of five "remainder" organs, excluding the skin and the lens of the eye, that receive the highest doses.

(2) For the purpose of weighting the external whole body dose, for adding it to the internal dose, a single weighting factor, $w_T = 1.0$, has been specified. The use of other weighting factors for external exposure will be approved on a case-by-case basis until such time as specific guidance is issued.

R313-15-3. Implementation.

(1) Any existing license or registration condition that is more restrictive than Rule R313-15 remains in force until there is an amendment or renewal of the license or registration.

(2) If a license or registration condition exempts a licensee or registrant from a provision of Rule R313-15 in effect on or before January 1, 1994, it also exempts the licensee or registrant from the corresponding provision of Rule R313-15.

(3) If a license or registration condition cites provisions of Rule R313-15 in effect prior to January 1, 1994, which do not correspond to any provisions of Rule R313-15, the license or registration condition remains in force until

1 there is an amendment or renewal of the license or registration that modifies or
2 removes this condition.

3
4 **R313-15-101. Radiation Protection Programs.**

5 (1) Each licensee or registrant shall develop, document, and implement a
6 radiation protection program sufficient to ensure compliance with the provisions
7 of Rule R313-15. See Section R313-15-1102 for recordkeeping requirements
8 relating to these programs.

9 (2) The licensee or registrant shall use, to the extent
10 ~~[practicable]~~practical, procedures and engineering controls based upon sound
11 radiation protection principles to achieve occupational doses and public doses
12 that are as low as is reasonably achievable (ALARA).

13 (3) The licensee or registrant shall, at intervals not to exceed 12
14 months, review the radiation protection program content and implementation.

15 (4) To implement the ALARA requirements of Subsection R313-15-101(2), and
16 notwithstanding the requirements in Section R313-15-301, a constraint on air
17 emissions of radioactive material to the environment, excluding radon-222 and its
18 daughters, shall be established by licensees or registrants such that the
19 individual member of the public likely to receive the highest dose will not be
20 expected to receive a total effective dose equivalent in excess of 0.1 mSv (10
21 mrem) per year from these emissions. If a licensee or registrant subject to this
22 requirement exceeds this dose constraint, the licensee or registrant shall report
23 the exceedance as provided in Section R313-15-1203 and promptly take appropriate
24 corrective action to ensure against recurrence.

25
26 **R313-15-201. Occupational Dose Limits for Adults.**

27 (1) The licensee or registrant shall control the occupational dose to
28 individual adults, except for planned special exposures pursuant to Section R313-

1 15-206, to the following dose limits:

2 (a) An annual limit, which is the more limiting of:

3 (i) The total effective dose equivalent being equal to 0.05 Sv (5 rem);

4 or

5 (ii) The sum of the deep dose equivalent and the committed dose equivalent
6 to any individual organ or tissue other than the lens of the eye being equal to
7 0.50 Sv (50 rem).

8 (b) The annual limits to the lens of the eye, to the skin, and to the
9 extremities which are:

10 (i) An eye dose equivalent of 0.15 Sv (15 rem), and

11 (ii) A shallow dose equivalent of 0.50 Sv (50 rem) to the skin or to any
12 extremity.

13 (2) Doses received in excess of the annual limits, including doses
14 received during accidents, emergencies, and planned special exposures, shall be
15 subtracted from the limits for planned special exposures that the individual may
16 receive during the current year and during the individual's lifetime. See
17 Subsections R313-15-206(5)(a) and R313-15-206(5)(b).

18 (3) The assigned deep dose equivalent and shallow dose equivalent shall
19 be for the portion of the body receiving the highest exposure determined as
20 follows:

21 (a) The deep dose equivalent, eye dose equivalent and shallow dose
22 equivalent may be assessed from surveys or other radiation measurements for the
23 purpose of demonstrating compliance with the occupational dose limits, if the
24 individual monitoring device was not in the region of highest potential exposure,
25 or the results of individual monitoring are unavailable; or

26 (b) When a protective apron is worn while working with medical
27 fluoroscopic equipment and monitoring is conducted as specified in Subsection
28 R313-15-502(1)(d), the effective dose equivalent for external radiation shall be

determined as follows:

(1) When only one individual monitoring device is used and it is located at the neck outside the protective apron, and the reported dose exceeds 25 percent of the limit specified in Subsection R313-15-201(1), the reported deep dose equivalent value multiplied by 0.3 shall be the effective dose equivalent for external radiation; or

(2) When individual monitoring devices are worn, both under the protective apron at the waist and outside the protective apron at the neck, the effective dose equivalent for external radiation shall be assigned the value of the sum of the deep dose equivalent reported for the individual monitoring device located at the waist under the protective apron multiplied by 1.5 and the deep dose equivalent reported for the individual monitoring device located at the neck outside the protective apron multiplied by 0.04.

(4) Derived air concentration (DAC) and annual limit on intake (ALI) values are ~~presented~~ specified in Table I of Appendix B of 10 CFR 20.1001 to 20.2402, 199[3]7 ed., which is incorporated by reference, and may be used to determine the individual's dose and to demonstrate compliance with the occupational dose limits. See Section F --15-1107.

(5) Notwithstanding the annual dose limits, the licensee shall limit the soluble uranium intake by an individual to ten milligrams in a week in consideration of chemical toxicity. See footnote 3, of Appendix B of 10 CFR 20.1001 to 20-2402, 199[3]7 ed., which is incorporated by reference.

(6) The licensee or registrant shall reduce the dose that an individual may be allowed to receive in the current year by the amount of occupational dose received while employed by any other person. See Subsection R313-15-205(5).

R313-15-202. Compliance with Requirements for Summation of External and Internal Doses.

1 (1) If the licensee or registrant is required to monitor pursuant to both
2 Subsections R313-15-502(1) and R313-15-502(2), the licensee or registrant shall
3 demonstrate compliance with the dose limits by summing external and internal
4 doses. If the licensee or registrant is required to monitor only pursuant to
5 Subsection R313-15-502(1) or only pursuant to Subsection R313-15-502(2), then
6 summation is not required to demonstrate compliance with the dose limits. The
7 licensee or registrant may demonstrate compliance with the requirements for
8 summation of external and internal doses pursuant to Subsections R313-15-202(2),
9 R313-15-202(3) and R313-15-202(4). The dose equivalents for the lens of the eye,
10 the skin, and the extremities are not included in the summation, but are subject
11 to separate limits.

12 (2) Intake by Inhalation. If the only intake of radionuclides is by
13 inhalation, the total effective dose equivalent limit is not exceeded if the sum
14 of the deep dose equivalent divided by the total effective dose equivalent limit,
15 and one of the following, does not exceed unity:

16 (a) The sum of the fractions of the inhalation ALI for each radionuclide,
17 or

18 (b) The total number of derived air concentration-hours (DAC-hours) for
19 all radionuclides divided by 2,000, or

20 (c) The sum of the calculated committed effective dose equivalents to all
21 significantly irradiated organs or tissues (T) calculated from bioassay data
22 using appropriate biological models and expressed as a fraction of the annual
23 limit. For purposes of this requirement, an organ or tissue is deemed to be
24 significantly irradiated if, for that organ or tissue, the product of the
25 weighting factors, w_T , and the committed dose equivalent, $H_{T,50}$, per unit intake
26 is greater than ten percent of the maximum weighted value of $H_{T,50}$ that is,
27 $w_T H_{T,50}$, per unit intake for any organ or tissue.

28 (3) Intake by Oral Ingestion. If the occupationally exposed individual

1 ~~also~~ receives an intake of radionuclides by oral ingestion greater than ten
2 percent of the applicable oral ALI, the licensee or registrant shall account for
3 this intake and include it in demonstrating compliance with the limits.

4 (4) Intake through Wounds or Absorption through Skin. The licensee or
5 registrant shall evaluate and, to the extent practical, account for intakes
6 through wounds or skin absorption. The intake through intact skin has been
7 included in the calculation of DAC for hydrogen-3 and does not need to be
8 evaluated or accounted for pursuant to Subsection R313-15-202(4).

9
10 **R313-15-203. Determination of External Dose from Airborne Radioactive Material.**

11 (1) Licensees or registrants shall, when determining the dose from
12 airborne radioactive material, include the contribution to the deep dose
13 equivalent, eye dose equivalent, and shallow dose equivalent from external
14 exposure to the radioactive cloud. See footnotes 1 and 2 of Appendix B of 10 CFR
15 20.1001 to 20.2402, 199[3] ed., which is incorporated by reference.

16 (2) Airborne radioactivity measurements and DAC values shall not be used
17 as the primary means to assess the deep dose equivalent when the airborne
18 radioactive material includes radionuclides other than noble gases or if the
19 cloud of airborne radioactive material is not relatively uniform. The
20 determination of the deep dose equivalent to an individual shall be based upon
21 measurements using instruments or individual monitoring devices.

22
23 **R313-15-204. Determination of Internal Exposure.**

24 (1) For purposes of assessing dose used to determine compliance with
25 occupational dose equivalent limits, the licensee or registrant shall, when
26 required pursuant to Section R313-15-502, take suitable and timely measurements
27 of:

28 (a) Concentrations of radioactive materials in air in work areas; or

1 (b) Quantities of radionuclides in the body; or

2 (c) Quantities of radionuclides excreted from the body; or

3 (d) Combinations of these measurements.

4 (2) Unless respiratory protective equipment is used, as provided in
5 Section R313-15-703, or the assessment of intake is based on bioassays, the
6 licensee or registrant shall assume that an individual inhales radioactive
7 material at the airborne concentration in which the individual is present.

8 (3) When specific information on the physical and biochemical properties
9 of the radionuclides taken into the body or the behavior of the material in an
10 individual is known, the licensee or registrant may:

11 (a) Use that information to calculate the committed effective dose
12 equivalent, and, if used, the licensee or registrant shall document that
13 information in the individual's record; and

14 (b) Upon prior approval of the Executive Secretary, adjust the DAC or ALI
15 values to reflect the actual physical and chemical characteristics of airborne
16 radioactive material, for example, aerosol size distribution or density; and

17 (c) Separately assess the contribution of fractional intakes of Class D,
18 W, or Y compounds of a given radionuclide to the committed effective dose
19 equivalent. See Appendix B of 10 CFR 20.1001 to 20.2402, 199[3] ed., which is
20 incorporated by reference.

21 (4) If the licensee or registrant chooses to assess intakes of Class Y
22 material using the measurements given in Subsections R313-15-204(1)(b) or R313-
23 15-204(1)(c), the licensee or registrant may delay the recording and reporting
24 of the assessments for periods up to seven months, unless otherwise required by
25 Section R313-15-1202 or Section R313-15-1203. This delay permits the licensee
26 or registrant to make additional measurements basic to the assessments.

27 (5) If the identity and concentration of each radionuclide in a mixture
28 are known, the fraction of the DAC applicable to the mixture for use in

1 calculating DAC-hours shall be either:

2 (a) The sum of the ratios of the concentration to the appropriate DAC
3 value, that is, D, W, or Y, from Appendix B of 10 CFR 20.1001 to 20.2402, 199[3]Z
4 ed., which is incorporated by reference, for each radionuclide in the mixture;
5 or

6 (b) The ratio of the total concentration for all radionuclides in the
7 mixture to the most restrictive DAC value for any radionuclide in the mixture.

8 (6) If the identity of each radionuclide in a mixture is known, but the
9 concentration of one or more of the radionuclides in the mixture is not known,
10 the DAC for the mixture shall be the most restrictive DAC of any radionuclide in
11 the mixture.

12 (7) When a mixture of radionuclides in air exists, a licensee or
13 registrant may disregard certain radionuclides in the mixture if:

14 (a) The licensee or registrant uses the total activity of the mixture in
15 demonstrating compliance with the dose limits in Section R313-15-201 and in
16 complying with the monitoring requirements in Subsection R313-15-502(2), and

17 (b) The concentration of any radionuclide disregarded is less than ten
18 percent of its DAC, and

19 (c) The sum of these percentages for all of the radionuclides disregarded
20 in the mixture does not exceed 30 percent.

21 (8) When determining the committed effective dose equivalent, the
22 following information may be considered:

23 (a) In order to calculate the committed effective dose equivalent, the
24 licensee or registrant may assume that the inhalation of one ALI, or an exposure
25 of 2,000 DAC-hours, results in a committed effective dose equivalent of 0.05 Sv
26 (5 rem) for radionuclides that have their ALIs or DACs based on the committed
27 effective dose equivalent.

28 (b) For an ALI and the associated DAC determined by the nonstochastic

organ dose limit of 0.50 Sv (50 rem), the intake of radionuclides that would result in a committed effective dose equivalent of 0.05 Sv (5 rem), that is, the stochastic ALI, is listed in parentheses in Table I of Appendix B of 10 CFR 20.1001 to 20.2402, 199[3] ed., which is incorporated by reference. The licensee or registrant may, as a simplifying assumption, use the stochastic ALI to determine committed effective dose equivalent. However, if the licensee or registrant uses the stochastic ALI, the licensee or registrant shall also demonstrate that the limit in Subsection R313-15-201(1)(a)(ii) is met.

R313-15-205. Determination of Prior Occupational Dose.

(1) For each individual ~~[who may enter the licensee's or registrant's restricted or controlled area and is]~~ likely to receive, in a year, an occupational dose requiring monitoring pursuant to Section R313-15-502, the licensee or registrant shall:

(a) Determine the occupational radiation dose received during the current year; and

(b) Attempt to obtain the records of ~~[lifetime]~~ cumulative occupational radiation dose. A licensee or registrant may accept, as the record of cumulative radiation dose, an up-to-date form DRC-05 or equivalent, signed by the individual and countersigned by an appropriate official of the most recent employer for work involving radiation exposure, or the individual's current employer, if the individual is not employed by the licensee or registrant.

(2) Prior to permitting an individual to participate in a planned special exposure, the licensee or registrant shall determine:

(a) The internal and external doses from all previous planned special exposures; and

(b) All doses in excess of the limits, including doses received during accidents and emergencies, received during the lifetime of the individual ~~[and]~~.

1 ~~[(e) All lifetime cumulative occupational radiation dose.]~~

2 (3) In complying with the requirements of Subsection R313-15-205(1), a
3 licensee or registrant may:

4 (a) Accept, as a record of the occupational dose that the individual
5 received during the current year, a written signed statement from the individual,
6 or from the individual's most recent employer for work involving radiation
7 exposure, that discloses the nature and the amount of any occupational dose that
8 the individual received during the current year; and

9 ~~[(b) Accept, as the record of lifetime cumulative radiation dose, an up-to-~~
10 ~~date form DRC-05 or equivalent, signed by the individual and countersigned by an~~
11 ~~appropriate official of the most recent employer for work involving radiation~~
12 ~~exposure, or the individual's current employer, if the individual is not employed~~
13 ~~by the licensee or registrant; and]~~

14 ~~[(e)]~~ (b) Obtain reports of the individual's dose equivalents from the most
15 recent employer for work involving radiation exposure, or the individual's
16 current employer, if the individual is not employed by the licensee or
17 registrant, by telephone, telegram, facsimile, other electronic media or letter.
18 The licensee or registrant shall request a written verification of the dose data
19 if the authenticity of the transmitted report cannot be established.

20 (4) ~~[(e)]~~ The licensee or registrant shall record the exposure history, as
21 required by Subsection R313-15-205(1), on form DRC-05, or other clear and legible
22 record, of all the information required on that form.

23 (a) The form or record shall show each period in which the individual
24 received occupational exposure to radiation or radioactive material and shall be
25 signed by the individual who received the exposure. For each period for which
26 the licensee or registrant obtains reports, the licensee or registrant shall use
27 the dose shown in the report in preparing form DRC-05 or equivalent. For any
28 period in which the licensee or registrant does not obtain a report, the licensee

1 or registrant shall place a notation on form DRC-05 or equivalent indicating the
2 periods of time for which data are not available.

3 (b) For the purpose of complying with this requirement, ~~(b)~~ licensees or
4 registrants are not required to reevaluate the separate external dose equivalents
5 and internal committed dose equivalents or intakes of radionuclides assessed
6 pursuant to the rules in Rule R313-15 in effect before January 1, 1994. Further,
7 occupational exposure histories obtained and recorded on form DRC-05 or
8 equivalent before January 1, 1994, would not have included effective dose
9 equivalent, but may be used in the absence of specific information on the intake
10 of radionuclides by the individual.

11 (5) If the licensee or registrant is unable to obtain a complete record
12 of an individual's current and previously accumulated occupational dose, the
13 licensee or registrant shall assume:

14 (a) In establishing administrative controls under Subsection R313-15-
15 201(6) for the current year, that the allowable dose limit for the individual is
16 reduced by 12.5 mSv (1.25 rem) for each quarter for which records were
17 unavailable and the individual was engaged in activities that could have resulted
18 in occupational radiation exposure; and

19 (b) That the individual is not available for planned special exposures.

20 (6) The licensee or registrant shall retain the records on form DRC-05 or
21 equivalent until the Executive Secretary terminates each pertinent license or
22 registration requiring this record. The licensee or registrant shall retain
23 records used in preparing form DRC-05 or equivalent for three years after the
24 record is made.

25
26 **R313-15-206. Planned Special Exposures.**

27 A licensee or registrant may authorize an adult worker to receive doses in
28 addition to and accounted for separately from the doses received under the limits

1 specified in Section R313-15-201 provided that each of the following conditions
2 is satisfied:

3 (1) The licensee or registrant authorizes a planned special exposure only
4 in an exceptional situation when alternatives that might avoid the higher
5 exposure are unavailable or impractical.

6 (2) The licensee or registrant, and employer if the employer is not the
7 licensee or registrant, specifically authorizes the planned special exposure, in
8 writing, before the exposure occurs.

9 (3) Before a planned special exposure, the licensee or registrant ensures
10 that each individual involved is:

11 (a) Informed of the purpose of the planned operation; and

12 (b) Informed of the estimated doses and associated potential risks and
13 specific radiation levels or other conditions that might be involved in
14 performing the task; and

15 (c) Instructed in the measures to be taken to keep the dose ALARA
16 considering other risks that may be present.

17 (4) Prior to permitting an individual to participate in a planned special
18 exposure, the licensee or registrant ascertains prior doses as required by
19 Subsection R313-15-205(2) during the lifetime of the individual for each
20 individual involved.

21 (5) Subject to Subsection R313-15-201(2), the licensee or registrant shall
22 not authorize a planned special exposure that would cause an individual to
23 receive a dose from all planned special exposures and all doses in excess of the
24 limits to exceed:

25 (a) The numerical values of any of the dose limits in Subsection R313-15-
26 201(1) in any year; and

27 (b) Five times the annual dose limits in Subsection R313-15-201(1) during
28 the individual's lifetime.

1 (6) The licensee or registrant maintains records of the conduct of a
2 planned special exposure in accordance with Section R313-15-1106 and submits a
3 written report in accordance with Section R313-15-1204.

4 (7) The licensee or registrant records the best estimate of the dose
5 resulting from the planned special exposure in the individual's record and
6 informs the individual, in writing, of the dose within 20 days from the date of
7 the planned special exposure. The dose from planned special exposures shall not
8 be considered in controlling future occupational dose of the individual pursuant
9 to Subsection R313-15-201(1) but shall be included in evaluations required by
10 Subsections R313-15-206(4) and R313-15-206(5).

11
12 **R313-15-207. Occupational Dose Limits for Minors.**

13 The annual occupational dose limits for minors are ten percent of the
14 annual occupational dose limits specified for adult workers in Section R313-15-
15 201.

16
17 **R313-15-208. Dose to an Embryo/Fetus.**

18 (1) The licensee or registrant shall ensure that the dose to an
19 embryo/fetus during the entire pregnancy, due to occupational exposure of a
20 declared pregnant woman, does not exceed five mSv (0.5 rem). See Section R313-
21 15-1107 for recordkeeping requirements.

22 (2) The licensee or registrant shall make efforts to avoid substantial
23 variation above a uniform monthly exposure rate to a declared pregnant woman so
24 as to satisfy the limit in Subsection R313-15-208(1).

25 (3) The dose to an embryo/fetus shall be taken as the sum of:

26 (a) The ~~[deep dose equivalent to the]~~ dose to the embryo/fetus from
27 radionuclides in the embryo/fetus and radionuclides in the declared pregnant
28 woman; and

1 (b) The dose that is most representative of the dose to the embryo/fetus
2 from external radiation, that is, in the mother's lower torso region. [
3 radionuclides in the embryo/fetus and radionuclides in the declared pregnant
4 woman.]

5 (i) If multiple measurements have not been made, assignment of the highest
6 deep dose equivalent for the declared pregnant woman shall be the dose to the
7 embryo/fetus, in accordance with Subsection R313-15-205(3); or

8 (ii) If multiple measurements have been made, assignment of the deep dose
9 equivalent for the declared pregnant woman from the individual monitoring device
10 which is most representative of the dose to the embryo/fetus shall be the dose
11 to the embryo fetus. Assignment of the highest deep dose equivalent for the
12 declared pregnant woman to the embryo/fetus is not required unless that dose is
13 also the most representative deep dose equivalent for the region of the
14 embryo/fetus.

D.208 (c)(ii)

15 (4) If by the time the woman declares pregnancy to the licensee or
16 registrant, the dose to the embryo/fetus has exceeded 4.5 mSv (0.45 rem) the
17 licensee or registrant shall be deemed to be in compliance with Subsection R313-
18 15-208(1) if the additional dose to the embryo/fetus does not exceed 0.50 mSv
19 (0.05 rem) during the remainder of the pregnancy.

20
21 **R313-15-301. Dose Limits for Individual Members of the Public.**

22 (1) Each licensee or registrant shall conduct operations so that:

23 (a) [The] Except as provided in Subsection R313-15-301(1)(c), the total
24 effective dose equivalent to individual members of the public from the licensed
25 or registered operation does not exceed one mSv (0.1 rem) in a year, exclusive
26 of the dose contribution from background radiation, from any medical
27 administration the individual has received, from exposure to individuals
28 administered radioactive material and released in accordance with Section R313-

1 32-75, from voluntary participation in medical research programs, and from the
2 licensee's or registrant's disposal of radioactive material into sanitary
3 sewerage in accordance with Section R313-15-1003[7], and

4 (b) The dose in any unrestricted area from external sources, exclusive of
5 the dose contributions patients administered radioactive material and
6 released in accordance with Section R313-32-75, does not exceed 0.02 mSv (0.002
7 rem) in any one hour[7], and

8 (c) The total effective dose equivalent to individual members of the
9 public from infrequent exposure to radiation from radiation machines does not
10 exceed 5 mSv (0.5 rem) in a year.

11 (2) If the licensee or registrant permits members of the public to have
12 access to controlled areas, the limits for members of the public continue to
13 apply to those individuals.

14 (3) A licensee, registrant, or an applicant for a license or registration
15 may apply for prior Executive Secretary authorization to operate up to an annual
16 dose limit for an individual member of the public of five mSv (0.5 rem). This
17 application shall include the following information:

18 (a) Demonstration of the need for and the expected duration of operations
19 in excess of the limit in Subsection R313-15-301(1); and

20 (b) The licensee's or registrant's program to assess and control dose
21 within the five mSv (0.5 rem) annual limit; and

22 (c) The procedures to be followed to maintain the dose ALARA.

23 (4) The Executive Secretary may impose additional restrictions on
24 radiation levels in unrestricted areas and on the total quantity of radionuclides
25 that a licensee or registrant may release in effluents in order to restrict the
26 collective dose.

27
28 R313-15-302. Compliance with Dose Limits for Individual Members of the Public.

1 (1) The licensee or registrant shall make or cause to be made surveys of
2 radiation levels in unrestricted and controlled areas and radioactive materials
3 in effluents released to unrestricted and controlled areas to demonstrate
4 compliance with the dose limits for individual members of the public in Section
5 R313-15-301.

6 (2) A licensee or registrant shall show compliance with the annual dose
7 limit in Section R313-15-301 by:

8 (a) Demonstrating by measurement or calculation that the total effective
9 dose equivalent to the individual likely to receive the highest dose from the
10 licensed or registered operation does not exceed the annual dose limit; or

11 (b) Demonstrating that:

12 (i) The annual average concentrations of radioactive material released in
13 gaseous and liquid effluents at the boundary of the unrestricted area do not
14 exceed the values specified in Table II of Appendix B of 10 CFR 20.1001 to
15 20.2402, 199[3] ed., which is incorporated by reference; and

16 (ii) If an individual were continu[ely]ously present in an unrestricted
17 area, the dose from external sources would not exceed 0.02 mSv (0.002 rem) in an
18 hour and 0.50 mSv (0.05 rem) in a year.

19 (3) Upon approval from the Executive Secretary, the licensee or registrant
20 may adjust the effluent concentration values in Appendix B, Table II of 10 CFR
21 20.1001 to 20.2402, 199[3] ed., which is incorporated by reference, for members
22 of the public, to take into account the actual physical and chemical
23 characteristics of the effluents, such as, aerosol size distribution, solubility,
24 density, radioactive decay equilibrium, and chemical form.

25
26 **R313-15-401. Testing for Leakage or Contamination of Sealed Sources.**

27 (1) The licensee or registrant in possession of any sealed source shall
28 assure that:

1 (a) Each sealed source, except as specified in Subsection R313-15-401(2),
2 is tested for leakage or contamination and the test results are received before
3 the sealed source is put into use unless the licensee or registrant has a
4 certificate from the transferor indicating that the sealed source was tested
5 within six months before transfer to the licensee or registrant.

6 (b) Each sealed source that is not designed to emit alpha particles is
7 tested for leakage or contamination at intervals not to exceed six months or at
8 alternative intervals approved by the Executive Secretary, an Agreement State,
9 a Licensing State, or the U.S. Nuclear Regulatory Commission.

10 (c) Each sealed source that is designed to emit alpha particles is tested
11 for leakage or contamination at intervals not to exceed three months or at
12 alternative intervals approved by the Executive Secretary, an Agreement State,
13 a Licensing State, or the Nuclear Regulatory Commission.

14 (d) For each sealed source that is required to be tested for leakage or
15 contamination, at any other time there is reason to suspect that the sealed
16 source might have been damaged or might be leaking, the licensee or registrant
17 shall assure that the sealed source is tested for leakage or contamination before
18 further use.

19 (e) Tests for leakage for all sealed sources, except brachytherapy sources
20 manufactured to contain radium, shall be capable of detecting the presence of 185
21 Bq (0.005 uCi) of radioactive material on a test sample. Test samples shall be
22 taken from the sealed source or from the surfaces of the container in which the
23 sealed source is stored or mounted on which one might expect contamination to
24 accumulate. For a sealed source contained in a device, test samples are obtained
25 when the source is in the "off" position.

26 (f) The test for leakage for brachytherapy sources manufactured to contain
27 radium shall be capable of detecting an absolute leakage rate of 37 Bq (0.001
28 uCi) of radon-222 in a 24 hour period when the collection efficiency for radon-

1 222 and its daughters has been determined with respect to collection method,
2 volume and time.

3 (g) Tests for contamination from radium daughters shall be taken on the
4 interior surface of brachytherapy source storage containers and shall be capable
5 of detecting the presence of 185 Bq (0.005 uCi) of a radium daughter which has
6 a half-life greater than four days.

7 (2) A licensee or registrant need not perform tests for leakage or
8 contamination on the following sealed sources:

9 (a) Sealed sources containing only radioactive material with a half-life
10 of less than 30 days;

11 (b) Sealed sources containing only radioactive material as a gas;

12 (c) Sealed sources containing 3.7 MBq (100 uCi) or less of beta or photon-
13 emitting material or 370 kBq (ten uCi) or less of alpha-emitting material;

14 (d) Sealed sources containing only hydrogen-3;

15 (e) Seeds of iridium-192 encased in nylon ribbon; and

16 (f) Sealed sources, except teletherapy and brachytherapy sources, which
17 are stored, not being used and identified as in storage. The licensee or
18 registrant shall, however, test each such sealed source for leakage or
19 contamination and receive the test results before any use or transfer unless it
20 has been tested for leakage or contamination within six months before the date
21 of use or transfer.

22 (3) Tests for leakage or contamination from sealed sources shall be
23 performed by persons specifically authorized by the Executive Secretary, an
24 Agreement State, a Licensing State, or the U.S. Nuclear Regulatory Commission to
25 perform such services.

26 (4) Test results shall be kept in units of becquerel or microcurie and
27 maintained for inspection by representatives of the Executive Secretary. Records
28 of test results for sealed sources shall be made pursuant to Section R313-15-

1 1104.

2 (5) The following shall be considered evidence that a sealed source is
3 leaking:

4 (a) The presence of 185 Bq (0.005 uCi) or more of removable contamination
5 on any test sample.

6 (b) Leakage of 37 Bq (0.001 uCi) of radon-222 per 24 hours for
7 brachytherapy sources manufactured to contain radium.

8 (c) The presence of removable contamination resulting from the decay of
9 185 Bq (0.005 uCi) or more of radium.

10 (6) The licensee or registrant shall immediately withdraw a leaking sealed
11 source from use and shall take action to prevent the spread of contamination.
12 The leaking sealed source shall be repaired or disposed of in accordance with
13 Rule R313-15.

14 (7) Reports of test results for leaking or contaminated sealed sources
15 shall be made pursuant to Section R313-15-1208.

16
17 **R313-15-501. Surveys and Monitoring - General.**

18 (1) Each licensee or registrant shall make, or cause to be made, surveys
19 that:

20 (a) Are necessary for the licensee or registrant to comply with Rule R313-
21 15; and

22 (b) Are necessary under the circumstances to evaluate:

23 (i) Radiation levels; and

24 (ii) Concentrations or quantities of radioactive material; and

25 (iii) The potential radiological hazards that could be present.

26 (2) The licensee or registrant shall ensure that instruments and equipment
27 used for quantitative radiation measurements, for example, dose rate and effluent
28 monitoring, are calibrated at intervals not to exceed 12 months for the radiation

1 measured, except when a more frequent interval is specified in another applicable
2 part of these rules or a license condition.

3 (3) All personnel dosimeters, except for direct and indirect reading
4 pocket ionization chambers and those dosimeters used to measure the dose to any
5 extremity, that require processing to determine the radiation dose and that are
6 used by licensees and registrants to comply with Section R313-15-201, with other
7 applicable provisions of these rules, or with conditions specified in a license
8 or registration shall be processed and evaluated by a dosimetry processor:

9 (a) Holding current personnel dosimetry accreditation from the National
10 Voluntary Laboratory Accreditation Program (NVLAP) of the National Institute of
11 Standards and Technology; and

12 (b) Approved in this accreditation process for the type of radiation or
13 radiations included in the NVLAP program that most closely approximates the type
14 of radiation or radiations for which the individual wearing the dosimeter is
15 monitored.

16 (4) The licensee or registrant shall ensure that adequate precautions are
17 taken to prevent a deceptive exposure of an individual monitoring device.

18
19 **R313-15-502. Conditions Requiring Individual Monitoring of External and Internal**
20 **Occupational Dose.**

21 Each licensee or registrant shall monitor exposures from sources of
22 radiation at levels sufficient to demonstrate compliance with the occupational
23 dose limits of Rule R313-15. As a minimum:

24 (1) Each licensee or registrant shall monitor occupational exposure to
25 radiation and shall supply and require the use of individual monitoring devices
26 by:

27 (a) Adults likely to receive, in one year from sources external to the
28 body, a dose in excess of ten percent of the limits in Subsection R313-15-201(1);

1 and

2 (b) Minors and declared pregnant women likely to receive, in one year from
3 sources external to the body, a dose in excess of ten percent of any of the
4 applicable limits in Sections R313-15-207 or R313-15-208; and

5 (c) Individuals entering a high or very high radiation area; and

6 (d) Individuals working with medical fluoroscopic equipment.

7 (i) An individual monitoring device used for the dose to an embryo/fetus
8 of a declared pregnant woman, pursuant to Subsection R313-15-208(1), shall be
9 located under the protective apron at the waist.

10 (A) If an individual monitoring device worn by a declared pregnant woman
11 has a monthly reported dose equivalent value in excess of 0.5 mSv (50 mrem), the
12 value to be used for determining the dose to the embryo/fetus, pursuant to
13 Subsection R313-15-208(3)(a) for radiation from medical fluoroscopy, may be the
14 value reported by the individual monitoring device worn at the waist underneath
15 the protective apron which has been corrected for the potential overestimation
16 of dose recorded by the monitoring device because of the overlying tissue of the
17 pregnant individual. This correction shall be performed by a radiation safety
18 officer of an institutional radiation safety committee, a qualified expert
19 approved by the Board, or a representative of the Executive Secretary.

20 (ii) An individual monitoring device used for eye dose equivalent shall
21 be located at the neck, or an unshielded location closer to the eye, outside the
22 protective apron.

23 (iii) When only one individual monitoring device is used to determine the
24 effective dose equivalent for external radiation pursuant to Subsection R313-15-
25 201(3)(b), it shall be located at the neck outside the protective apron. When
26 a second individual monitoring device is used, for the same purpose, it shall be
27 located under the protective apron at the waist. Note: The second individual
28 monitoring device is required for a declared pregnant woman.

1 (2) Each licensee or registrant shall monitor, to determine compliance
2 with Section R313-15-204, the occupational intake of radioactive material by and
3 assess the committed effective dose equivalent to:

4 (a) Adults likely to receive, in one year, an intake in excess of ten
5 percent of the applicable ALI in Table I, Columns 1 and 2, of Appendix B of 10
6 CFR 20.1001 to 20.2402, 199[3]7 ed., which is incorporated by reference; and

7 (b) Minors and declared pregnant women likely to receive, in one year, a
8 committed effective dose equivalent in excess of 0.50 mSv (0.05 rem).

9
10 R313-15-503. Location of Individual Monitoring Devices.

11 Each licensee or registrant shall ensure that individuals who are required
12 to monitor occupational doses in accordance with Subsection R313-15-502(1) wear
13 individual monitoring devices as follows:

14 (1) An individual monitoring device used for monitoring the dose to the
15 whole body shall be worn at the unshielded location of the whole body likely to
16 receive the highest exposure. When a protective apron is worn, the location of
17 the individual monitoring device is typically at the neck (collar).

18 (2) An individual monitoring device used for monitoring the dose to an
19 embryo/fetus of a declared pregnant woman, pursuant to Subsection R313-15-208(1),
20 shall be located at the waist under any protective apron being worn by the woman.

21 (3) An individual monitoring device used for monitoring the eye dose
22 equivalent, to demonstrate compliance with Subsection R313-15-201(1)(b)(i), shall
23 be located at the neck (collar), outside any protective apron being worn by the
24 monitored individual, or at an unshielded location closer to the eye.

25 (4) An individual monitoring device used for monitoring the dose to the
26 extremities, to demonstrate compliance with Subsection R313-15-201(1)(b)(ii),
27 shall be worn on the extremity likely to receive the highest exposure. Each
28 individual monitoring device shall be oriented to measure the highest dose to the

1 extremity being monitored.

2
3 **R313-15-601. Control of Access to High Radiation Areas.**

4 (1) The licensee or registrant shall ensure that each entrance or access
5 point to a high radiation area has one or more of the following features:

6 (a) A control device that, upon entry into the area, causes the level of
7 radiation to be reduced below that level at which an individual might receive a
8 deep dose equivalent of one mSv (0.1 rem) in one hour at 30 centimeters from the
9 source of radiation or from any surface that the radiation penetrates; or

10 (b) A control device that energizes a conspicuous visible or audible alarm
11 signal so that the individual entering the high radiation area and the supervisor
12 of the activity are made aware of the entry; or

13 (c) Entryways that are locked, except during periods when access to the
14 areas is required, with positive control over each individual entry.

15 (2) In place of the controls required by Subsection R313-15-601(1) for a
16 high radiation area, the licensee or registrant may substitute continuous direct
17 or electronic surveillance that is capable of preventing unauthorized entry.

18 (3) The licensee or registrant may apply to the Executive Secretary for
19 approval of alternative methods for controlling access to high radiation areas.

20 (4) The licensee or registrant shall establish the controls required by
21 Subsections R313-15-601(1) and R313-15-601(3) in a way that does not prevent
22 individuals from leaving a high radiation area.

23 (5) The licensee or registrant is not required to control each entrance
24 or access point to a room or other area that is a high radiation area solely
25 because of the presence of radioactive materials prepared for transport and
26 packaged and labeled in accordance with the rules of the U.S. Department of
27 Transportation provided that:

28 (a) The packages do not remain in the area longer than three days; and

1 (b) The dose rate at one meter from the external surface of any package
2 does not exceed 0.1 mSv (0.01 rem) per hour.

3 (6) The licensee or registrant is not required to control entrance or
4 access to rooms or other areas in hospitals solely because of the presence of
5 patients containing radioactive material, provided that there are personnel in
6 attendance who are taking the necessary precautions to prevent the exposure of
7 individuals to radiation or radioactive material in excess of the established
8 limits in Rule R313-15 and to operate within the ALARA provisions of the
9 licensee's or registrant's radiation protection program.

10 (7) The registrant is not required to control entrance or access to rooms
11 or other areas containing sources of radiation capable of producing a high
12 radiation area as described in Section R313-15-601 if the registrant has met all
13 the specific requirements for access and control specified in other applicable
14 sections of these rules, such as, Rule R313-36 for industrial radiography, Rule
15 R313-28 for x rays in the healing arts, Rule R313-30 for therapeutic radiation
16 machines, and Rule R313-~~44~~35 for ~~[particle accelerators]~~ industrial use of x-ray
17 systems.

18
19 **R313-15-602. Control of Access to Very High Radiation Areas.**

20 (1) In addition to the requirements in Section R313-15-601, the licensee
21 or registrant shall institute measures to ensure that an individual is not able
22 to gain unauthorized or inadvertent access to areas in which radiation levels
23 could be encountered at five Gy (500 rad) or more in one hour at one meter from
24 a source of radiation or any surface through which the radiation penetrates.
25 This requirement does not apply to rooms or areas in which diagnostic x-ray
26 systems are the only source of radiation, or to non-self-shielded irradiators.

27 (2) The registrant is not required to control entrance or access to rooms
28 or other areas containing sources of radiation capable of producing a very high

1 radiation area as described in Subsection R313-15-602(1) if the registrant has
2 met all the specific requirements for access and control specified in other
3 applicable sections of these rules, such as, Rule R313-36 for industrial
4 radiography, Rule R313-28 for x rays in the healing arts, Rule R313-30 for
5 therapeutic radiation machines, and Rule R313-[44]35 for [particle
6 accelerators] industrial use of x-ray systems.

7
8 **R313-15-603. Control of Access to Very High Radiation Areas -- Irradiators.**

9 (1) Section R313-15-603 applies to licensees or registrants with sources
10 of radiation in non-self-shielded irradiators. Section R313-15-603 does not
11 apply to sources of radiation that are used in teletherapy, in industrial
12 radiography, or in completely self-shielded irradiators in which the source of
13 radiation is both stored and operated within the same shielding radiation barrier
14 and, in designed configuration of the irradiator, is always physically
15 inaccessible to any individual and cannot create a high levels of radiation in
16 an area that is accessible to any individual.

17 (2) Each area in which there may exist radiation levels in excess of five
18 Gy (500 rad) in one hour at one meter from a source of radiation that is used to
19 irradiate materials shall meet the following requirements:

20 (a) Each entrance or access point shall be equipped with entry control
21 devices which:

22 (i) Function automatically to prevent any individual from inadvertently
23 entering a very high radiation area; and

24 (ii) Permit deliberate entry into the area only after a control device is
25 actuated that causes the radiation level within the area, from the source of
26 radiation, to be reduced below that at which it would be possible for an
27 individual to receive a deep dose equivalent in excess of one mSv (0.1 rem) in
28 one hour; and

1 (iii) Prevent operation of the source of radiation if it would produce
2 radiation levels in the area that could result in a deep dose equivalent to an
3 individual in excess of one mSv (0.1 rem) in one hour.

4 (b) Additional control devices shall be provided so that, upon failure of
5 the entry control devices to function as required by Subsection K313-15-
6 603(2)(a):

7 (i) The radiation level within the area, from the source of radiation, is
8 reduced below that at which it would be possible for an individual to receive a
9 deep dose equivalent in excess of one mSv (0.1 rem) in one hour; and

10 (ii) Conspicuous visible and audible alarm signals are generated to make
11 an individual attempting to enter the area aware of the hazard and at least one
12 other authorized individual, who is physically present, familiar with the
13 activity, and prepared to render or summon assistance, aware of the failure of
14 the entry control devices.

15 (c) The licensee or registrant shall provide control devices so that, upon
16 failure or removal of physical radiation barriers other than the sealed source's
17 shielded storage container:

18 (i) The radiation level from the source of radiation is reduced below that
19 at which it would be possible for an individual to receive a deep dose equivalent
20 in excess of one mSv (0.1 rem) in one hour; and

21 (ii) Conspicuous visible and audible alarm signals are generated to make
22 potentially affected individuals aware of the hazard and the licensee or
23 registrant or at least one other individual, who is familiar with the activity
24 and prepared to render or summon assistance, aware of the failure or removal of
25 the physical barrier.

26 (d) When the shield for stored sealed sources is a liquid, the licensee
27 or registrant shall provide means to monitor the integrity of the shield and to
28 signal, automatically, loss of adequate shielding.

1 (e) Physical radiation barriers that comprise permanent structural
2 components, such as walls, that have no credible probability of failure or
3 removal in ordinary circumstances need not meet the requirements of Subsections
4 R313-15-603(2)(c) and R313-15-603(2)(d).

5 (f) Each area shall be equipped with devices that will automatically
6 generate conspicuous visible and audible alarm signals to alert personnel in the
7 area before the source of radiation can be put into operation and in time for any
8 individual in the area to operate a clearly identified control device, which
9 shall be installed in the area and which can prevent the source of radiation from
10 being put into operation.

11 (g) Each area shall be controlled by use of such administrative procedures
12 and such devices as are necessary to ensure that the area is cleared of personnel
13 prior to each use of the source of radiation.

14 (h) Each area shall be checked by a radiation measurement to ensure that,
15 prior to the first individual's entry into the area after any use of the source
16 of radiation, the radiation level from the source of radiation in the area is
17 below that at which it would be possible for an individual to receive a deep dose
18 equivalent in excess of one mSv (0.1 rem) in one hour.

19 (i) The entry control devices required in Subsection R313-15-603(2)(a)
20 shall be tested for proper functioning. See Section R313-15-1110 for
21 recordkeeping requirements.

22 (i) Testing shall be conducted prior to initial operation with the source
23 of radiation on any day, unless operations were continued uninterrupted from the
24 previous day; and

25 (ii) Testing shall be conducted prior to resumption of operation of the
26 source of radiation after any unintentional interruption; and

27 (iii) The licensee or registrant shall submit and adhere to a schedule for
28 periodic tests of the entry control and warning systems.

1 (j) The licensee or registrant shall not conduct operations, other than
2 those necessary to place the source of radiation in safe condition or to effect
3 repairs on controls, unless control devices are functioning properly.

4 (k) Entry and exit portals that are used in transporting materials to and
5 from the irradiation area, and that are not intended for use by individuals,
6 shall be controlled by such devices and administrative procedures as are
7 necessary to physically protect and warn against inadvertent entry by any
8 individual through these portals. Exit portals for irradiated materials shall
9 be equipped to detect and signal the presence of any loose radioactive material
10 that is carried toward such an exit and automatically to prevent loose
11 radioactive material from being carried out of the area.

12 (3) Licensees, registrants, or applicants for licenses or registrations
13 for sources of radiation within the purview of Subsection R313-15-603(2) which
14 will be used in a variety of positions or in locations, such as open fields or
15 forests, that make it ~~[impracticable]~~ impractical to comply with certain
16 requirements of Subsection R313-15-603(2), such as those for the automatic
17 control of radiation levels, may apply to the Executive Secretary for approval
18 of alternative safety measures. Alternative safety measures shall provide
19 personnel protection at least equivalent to those specified in Subsection R313-
20 15-603(2). At least one of the alternative measures shall include an entry-
21 preventing interlock control based on a measurement of the radiation that ensures
22 the absence of high radiation levels before an individual can gain access to the
23 area where such sources of radiation are used.

24 (4) The entry control devices required by Subsections R313-15-603(2) and
25 R313-15-603(3) shall be established in such a way that no individual will be
26 prevented from leaving the area.

27
28 **R313-15-701. Use of Process or Other Engineering Controls.**

1 The licensee or registrant shall use, to the extent
2 ~~[practicable]~~practical, process or other engineering controls, such as,
3 containment or ventilation, to control the concentrations of radioactive material
4 in air.

5
6 **R313-15-702. Use of Other Controls.**

7 When it is not ~~[practicable]~~practical to apply process or other engineering
8 controls to control the concentrations of radioactive material in air to values
9 below those that define an airborne radioactivity area, the licensee or
10 registrant shall, consistent with maintaining the total effective dose equivalent
11 ALARA, increase monitoring and limit intakes by one or more of the following
12 means:

- 13 (1) Control of access; or
14 (2) Limitation of exposure times; or
15 (3) Use of respiratory protection equipment; or
16 (4) Other controls.

17
18 **R313-15-703. Use of Individual Respiratory Protection Equipment.**

19 (1) If the licensee or registrant uses respiratory protection equipment
20 to limit intakes pursuant to Section R313-15-702:

21 (a) Except as provided in Subsection R313-15-703(1)(b), the licensee or
22 registrant shall use only respiratory protection equipment that is tested and
23 certified or had certification extended by the National Institute for
24 Occupational Safety and Health and the Mine Safety and Health Administration.

25 (b) ~~[if it]~~The licensee or registrant ~~[wishes to]~~may use equipment that has
26 not been tested or certified by the National Institute for Occupational Safety
27 and Health and the Mine Safety and Health Administration, has not had
28 certification extended by the National Institute for Occupational Safety and

1 Health and the Mine Safety and Health Administration, or for which there is no
2 schedule for testing or certification, provided the licensee or registrant
3 [shall submit] has submitted to the Executive Secretary and the Executive
4 Secretary has approved an application for authorized use of that equipment,
5 including a demonstration by testing, or a demonstration on the basis of
6 [reliable] test information, that the material and performance characteristics
7 of the equipment are capable of providing the proposed degree of protection under
8 anticipated conditions of use.

9 (c) The licensee or registrant shall implement and maintain a respiratory
10 protection program that includes:

11 (i) Air sampling sufficient to identify the potential hazard, permit
12 proper equipment selection, and estimate exposures; and

13 (ii) Surveys and bioassays, as appropriate, to evaluate actual intakes;
14 and

15 (iii) Testing of respirators for operability immediately prior to each
16 use; and

17 (iv) Written procedures regarding selection, fitting, issuance,
18 maintenance, and testing of respirators, including testing for operability
19 immediately prior to each use; supervision and training of personnel; monitoring,
20 including air sampling and bioassays; and recordkeeping; and

21 (v) Determination by a physician prior to initial fitting of respirators,
22 and [at least] either every 12 months thereafter or periodically at a frequency
23 determined by a physician, that the individual user is [physically] medically
24 [able] fit to use the respiratory protection equipment.

25 (d) The licensee or registrant shall issue a written policy statement on
26 respirator usage covering:

27 (i) The use of process or other engineering controls, instead of
28 respirators; and

1 (ii) The routine, nonroutine, and emergency use of respirators; and
2 (iii) The length of periods of respirator use and relief from respirator
3 use.

4 (e) The licensee or registrant shall advise each respirator user that the
5 user may leave the area at any time for relief from respirator use in the event
6 of equipment malfunction, physical or psychological distress, procedural or
7 communication failure, significant deterioration of operating conditions, or any
8 other conditions that might require such relief.

9 (f) The licensee or registrant shall use respiratory protection equipment
10 within the equipment manufacturer's expressed limitations for type and mode of
11 use and shall provide proper visual, communication, and other special
12 capabilities, such as adequate skin protection, when needed.

13 (2) When estimating exposure of individuals to airborne radioactive
14 materials, the licensee or registrant may make allowance for respiratory
15 protection equipment used to limit intakes pursuant to Section R313-15-702,
16 provided that the following conditions, in addition to those in Subsection R313-
17 15-703(1), are satisfied:

18 (a) The licensee or registrant selects respiratory protection equipment
19 that provides a protection factor, specified in Appendix A of 10 CFR 20.1001 to
20 20.2402, 199[3] ed., which is incorporated by reference, greater than the
21 multiple by which peak concentrations of airborne radioactive materials in the
22 working area are expected to exceed the values specified in Appendix B, Table I,
23 Column 3 of 10 CFR 20.1001 to 20.2402, 199[3] ed., which is incorporated by
24 reference. However, if the selection of respiratory protection equipment with
25 a protection factor greater than the [peak concentration] multiple defined in the
26 preceding sentence is inconsistent with the goal specified in Section R313-15-702
27 of keeping the total effective dose equivalent ALARA, the licensee or registrant
28 may select respiratory protection equipment with a lower protection factor

1 provided that such a selection would result in a total effective dose equivalent
2 that is ALARA. The concentration of radioactive material in the air that is
3 inhaled when respirators are worn may be initially estimated by dividing the
4 average concentration in air, during each period of uninterrupted use, by the
5 protection factor. If the exposure is later found to be greater than initially
6 estimated, the corrected value shall be used; if the exposure is later found to
7 be less than initially estimated, the corrected value may be used.

8 (b) The licensee or registrant shall obtain authorization from the
9 Executive Secretary before assigning respiratory protection factors in excess of
10 those specified in Appendix A of 10 CFR 20.1001 to 20.2402, 199[3] ed., which
11 is incorporated by reference. The Executive Secretary may authorize a licensee
12 or registrant to use higher protection factors on receipt of an application that:

13 (i) Describes the situation for which a need exists for higher protection
14 factors, and

15 (ii) Demonstrates that the respiratory protection equipment provides these
16 higher protection factors under the proposed conditions of use.

17 (c) In an emergency, the licensee or registrant shall use as emergency
18 equipment only respiratory protection equipment that has been specifically
19 certified or had certification extended for emergency use by the National
20 Institute for Occupational Safety and Health and the Mine Safety and Health
21 Administration.

22 (d) The licensee or registrant shall notify the Executive Secretary in
23 writing at least 30 days before the date that respiratory protection equipment
24 is first used pursuant to either Subsections R313-15-703(1) or R313-15-703(2).

25
26 R313-15-801. Security and Control of Licensed or Registered [Stored] Sources of
27 Radiation.

28 ~~[The licensee or registrant shall secure from unauthorized removal or~~

1 ~~access licensed or registered sources of radiation that are stored in controlled~~
2 ~~or unrestricted areas.~~

4 ~~R313-15-802. Control of Sources of Radiation not in Storage.~~

5 ~~—— (1) The licensee or registrant shall control and maintain constant~~
6 ~~surveillance of licensed or registered radioactive material that is in a~~
7 ~~controlled or unrestricted area and that is not in storage or in a patient.~~

8 ~~—— (2) The registrant shall maintain control of radiation machines that are~~
9 ~~in a controlled or unrestricted area and that are not in storage.] (1) The~~
10 licensee or registrant shall secure licensed or registered radioactive material
11 from unauthorized removal or access.

12 (2) The licensee or registrant shall maintain constant surveillance, and
13 use devices or administrative procedures to prevent unauthorized use of licensed
14 or registered radioactive material that is in an unrestricted area and that is
15 not in storage.

16 (3) The registrant shall secure registered radiation machines from
17 unauthorized removal.

18 (4) The registrant shall use devices or administrative procedures to
19 prevent unauthorized use of registered radiation machines.

21 **R313-15-901. Caution Signs.**

22 (1) Standard Radiation Symbol. Unless otherwise authorized by the
23 Executive Secretary, the symbol prescribed by 10 CFR 20.1901, 199[3]7 ed., which
24 is incorporated by reference, shall use the colors magenta, or purple, or black
25 on yellow background. The symbol prescribed is the three-bladed design as
26 follows:

27 (a) Cross-hatched area is to be magenta, or purple, or black, and

28 (b) The background is to be yellow.

1 (2) Exception to Color Requirements for Standard Radiation Symbol.
2 Notwithstanding the requirements of 10 CFR 20.1901a, 199[3]7 ed., which is
3 incorporated by reference, licensees or registrants are authorized to label
4 sources, source holders, or device components containing sources of radiation
5 that are subjected to high temperatures, with conspicuously etched or stamped
6 radiation caution symbols and without a color requirement.

7 (3) Additional Information on Signs and Labels. In addition to the
8 contents of signs and labels prescribed in Rule R313-15, the licensee or
9 registrant shall provide, on or near the required signs and labels, additional
10 information, as appropriate, to make individuals aware of potential radiation
11 exposures and to minimize the exposures.

12
13
14 **R313-15-902. Posting Requirements.**

15 (1) Posting of Radiation Areas. The licensee or registrant shall post
16 each radiation area with a conspicuous sign or signs bearing the radiation symbol
17 and the words "CAUTION, RADIATION AREA."

18 (2) Posting of High Radiation Areas. The licensee or registrant shall
19 post each high radiation area with a conspicuous sign or signs bearing the
20 radiation symbol and the words "CAUTION, HIGH RADIATION AREA" or "DANGER, HIGH
21 RADIATION AREA."

22 (3) Posting of Very High Radiation Areas. The licensee or registrant
23 shall post each very high radiation area with a conspicuous sign or signs bearing
24 the radiation symbol and words "GRAVE DANGER, VERY HIGH RADIATION AREA."

25 (4) Posting of Airborne Radioactivity Areas. The licensee or registrant
26 shall post each airborne radioactivity area with a conspicuous sign or signs
27 bearing the radiation symbol and the words "CAUTION, AIRBORNE RADIOACTIVITY AREA"
28 or "DANGER, AIRBORNE RADIOACTIVITY AREA."

(5) Posting of Areas or Rooms in which Licensed or Registered Material is Used or Stored. The licensee or registrant shall post each area or room in which there is used or stored an amount of licensed or registered material exceeding ten times the quantity of such material specified in Appendix C of 10 CFR 20.1001 to 20.2402, 199[3]7 ed., which is incorporated by reference, with a conspicuous sign or signs bearing the radiation symbol and the words "CAUTION, RADIOACTIVE MATERIAL" or "DANGER, RADIOACTIVE MATERIAL."

R313-15-903. Exceptions to Posting Requirements.

(1) A licensee or registrant is not required to post caution signs in areas or rooms containing sources of radiation for periods of less than eight hours, if each of the following conditions is met:

(a) The sources of radiation are constantly attended during these periods by an individual who takes the precautions necessary to prevent the exposure of individuals to sources of radiation in excess of the limits established in Rule R313-15; and

(b) The area or room is subject to the licensee's or registrant's control.

(2) Rooms or other areas in hospitals that are occupied by patients are not required to be posted with caution signs pursuant to Section R313-15-902 provided that the patient could be released from ~~[confinement]~~ licensee control pursuant to Section R313-32-75.

(3) A room or area is not required to be posted with a caution sign because of the presence of a sealed source provided the radiation level at 30 centimeters from the surface of the sealed source container or housing does not exceed 0.05 mSv (0.005 rem) per hour.

(4) A room or area is not required to be posted with a caution sign because of the presence of radiation machines used solely for diagnosis in the healing arts.

1 **R313-15-904. Labeling Containers and Radiation Machines.**

2 (1) The licensee or registrant shall ensure that each container of
3 licensed or registered material bears a durable, clearly visible label bearing
4 the radiation symbol and the words "CAUTION, RADIOACTIVE MATERIAL" or "DANGER,
5 RADIOACTIVE MATERIAL." The label shall also provide information, such as the
6 radionuclides present, an estimate of the quantity of radioactivity, the date for
7 which the activity is estimated, radiation levels, kinds of materials, and mass
8 enrichment, to permit individuals handling or using the containers, or working
9 in the vicinity of the containers, to take precautions to avoid or minimize
10 exposures.

11 (2) Each licensee or registrant shall, prior to removal or disposal of
12 empty uncontaminated containers to unrestricted areas, remove or deface the
13 radioactive material label or otherwise clearly indicate that the container no
14 longer contains radioactive materials.

15 (3) Each registrant shall ensure that each radiation machine is labeled
16 in a conspicuous manner which cautions individuals that radiation is produced
17 when it is energized.

18
19 **R313-15-905. Exemptions to Labeling Requirements.**

20 A licensee or registrant is not required to label:

21 (1) Containers holding licensed or registered material in quantities less
22 than the quantities listed in Appendix C of 10 CFR 20.1001 to 20.2402, 199[3]2
23 ed., which is incorporated by reference; or

24 (2) Containers holding licensed or registered material in concentrations
25 less than those specified in Table III of Appendix B of 10 CFR 20.1001 to
26 20.2402, 199[3]2 ed., which is incorporated by reference; or

27 (3) Containers attended by an individual who takes the precautions
28 necessary to prevent the exposure of individuals in excess of the limits

1 established by Rule R313-15; or

2 (4) Containers when they are in transport and packaged and labeled in
3 accordance with the rules of the U.S. Department of Transportation; or

4 (5) Containers that are accessible only to individuals authorized to
5 handle or use them, or to work in the vicinity of the containers, if the contents
6 are identified to these individuals by a readily available written record.
7 Examples of containers of this type are containers in locations such as water-
8 filled canals, storage vaults, or hot cells. The record shall be retained as long
9 as the containers are in use for the purpose indicated on the record; or

10 (6) Installed manufacturing or process equipment, such as piping and
11 tanks.

12
13 **R313-15-906. Procedures for Receiving and Opening Packages.**

14 (1) Each licensee or registrant who expects to receive a package
15 containing quantities of radioactive material in excess of a Type A quantity, as
16 defined in Section R313-19-4 and Subsection R313-19-100(19), shall make
17 arrangements to receive:

18 (a) The package when the carrier offers it for delivery; or

19 (b) The notification of the arrival of the package at the carrier's
20 terminal and to take possession of the package expeditiously.

21 (2) Each licensee or registrant shall:

22 (a) Monitor the external surfaces of a labeled package for radioactive
23 contamination unless the package contains only radioactive material in the form
24 of gas or in special form as defined in Section R313-12-3; and

25 (b) Monitor the external surfaces of a labeled package for radiation
26 levels unless the package contains quantities of radioactive material that are
27 less than or equal to the Type A quantity, as defined in Section R313-19-4 and
28 Subsection R313-19-100(19); and

1 (c) Monitor all packages known to contain radioactive material for
2 radioactive contamination and radiation levels if there is evidence of
3 degradation of package integrity, such as packages that are crushed, wet, or
4 damaged.

5 (3) The licensee or registrant shall perform the monitoring required by
6 Subsection R313-15-906(2) as soon as ~~[practicable]~~practical after receipt of the
7 package, but not later than three hours after the package is received at the
8 licensee's or registrant's facility if it is received during the licensee's or
9 registrant's normal working hours~~(, or not later than three hours from the~~
10 ~~beginning of the next working day if it is received after working hours.)~~ or if
11 there is evidence of degradation of package integrity, such as a package that is
12 crushed, wet, or damaged. If a package is received after working hours, and has
13 no evidence of degradation of package integrity, the package shall be monitored
14 no later than three hours from the beginning of the next working day. HES

15 (4) The licensee or registrant shall immediately notify the final delivery
16 carrier and, by telephone and telegram, mailgram, or facsimile, the Executive
17 Secretary when:

18 (a) Removable radioactive surface contamination exceeds the limits of
19 Subsection R313-19-100(13)(h); or

20 (b) External radiation levels exceed the limits of Subsections R313-19-
21 100(13)(i) and R313-19-100(13)(j).

22 (5) Each licensee or registrant shall:

23 (a) Establish, maintain, and retain written procedures for safely opening
24 packages in which radioactive material is received; and

25 (b) Ensure that the procedures are followed and that due consideration is
26 given to special instructions for the type of package being opened.

27 (6) Licensees or registrants transferring special form sources in vehicles

1 owned or operated by the licensee or registrant to and from a work site are
2 exempt from the contamination monitoring requirements of Subsection R313-15-
3 906(2), but are not exempt from the monitoring requirement in Subsection R313-15-
4 906(2) for measuring radiation levels that ensures that the source is still
5 properly lodged in its shield.

6
7 **R313-15-1001. Waste Disposal - General Requirements.**

8 (1) A licensee or registrant shall dispose of licensed or registered
9 material only:

10 (a) By transfer to an authorized recipient as provided in Section R313-15-
11 1006 or in Rules R313-21, R313-22, or R313-25, or to the U.S. Department of
12 Energy; or

13 (b) By decay in storage; or

14 (c) By release in effluents within the limits in Section R313-15-301; or

15 (d) As authorized pursuant to Sections R313-15-1002, R313-15-1003, R313-
16 15-1004, or R313-15-1005.

17 (2) A person shall be specifically licensed or registered to receive waste
18 containing licensed or registered material from other persons for:

19 (a) Treatment prior to disposal; or

20 (b) Treatment or disposal by incineration; or

21 (c) Decay in storage; or

22 (d) Disposal at a land disposal facility licensed pursuant to Rule R313-
23 25; or

24 (e) Storage until transferred to a storage or disposal facility authorized
25 to receive the waste.

26
27 **R313-15-1002. Method for Obtaining Approval of Proposed Disposal Procedures.**

28 A licensee or registrant or applicant for a license or registration may

1 apply to the Executive Secretary for approval of proposed procedures, not
2 otherwise authorized in these rules, to dispose of licensed or registered
3 material generated in the licensee's or registrant's operations. Each
4 application shall include:

5 (1) A description of the waste containing licensed or registered material
6 to be disposed of, including the physical and chemical properties that have an
7 impact on risk evaluation, and the proposed manner and conditions of waste
8 disposal; and

9 (2) An analysis and evaluation of pertinent information on the nature of
10 the environment; and

11 (3) The nature and location of other potentially affected facilities; and

12 (4) Analyses and procedures to ensure that doses are maintained ALARA and
13 within the dose limits in Rule R313-15.

14
15 **R313-15-1003. Disposal by Release into Sanitary Sewerage.**

16 (1) A licensee or registrant may discharge licensed or registered material
17 into sanitary sewerage if each of the following conditions is satisfied:

18 (a) The material is readily soluble, or is readily dispersible biological
19 material, in water; and

20 (b) The quantity of licensed or registered radioactive material that the
21 licensee or registrant releases into the sewer in one month divided by the
22 average monthly volume of water released into the sewer by the licensee or
23 registrant does not exceed the concentration listed in Table III of Appendix B
24 of 10 CFR 20.1001 to 20.2402, 199[3] ed., which is incorporated by reference; and

25 (c) If more than one radionuclide is released, the following conditions
26 shall also be satisfied:

27 (i) The licensee or registrant shall determine the fraction of the limit
28 in Table III of Appendix B of 10 CFR 20.1001 to 20.2402, 199[3] ed., which is

1 incorporated by reference, represented by discharges into sanitary sewerage by
2 dividing the actual monthly average concentration of each radionuclide released
3 by the licensee or registrant into the sewer by the concentration of that
4 radionuclide listed in Table III of Appendix B of 10 CFR 20.1001 to 20.2402,
5 199[3] ed., which is incorporated by reference; and

6 (ii) The sum of the fractions for each radionuclide required by Subsection
7 R313-15-1003(1)(c)(i) does not exceed unity; and

8 (d) The total quantity of licensed or registered radioactive material that
9 the licensee or registrant releases into the sanitary sewerage system in a year
10 does not exceed 185 GBq (five Ci) of hydrogen-3, 37 GBq (one Ci) of carbon-14,
11 and 37 GBq (one Ci) of all other radioactive materials combined.

12 (2) Excreta from individuals undergoing medical diagnosis or therapy with
13 radioactive material are not subject to the limitations contained in Subsection
14 R313-15-1003(1).

15
16 **R313-15-1004. Treatment or Disposal by Incineration.**

17 A licensee or registrant may treat or dispose of licensed or registered
18 material by incineration only in the ~~[amounts and forms]~~ form and concentration
19 specified in Section R313-15-1005 or as specifically approved by the Executive
20 Secretary pursuant to Section R313-15-1002.

21
22 **R313-15-1005. Disposal of Specific Wastes.**

23 (1) A licensee or registrant may dispose of the following licensed or
24 registered material as if it were not radioactive:

25 (a) 1.85 kBq (0.05 uCi), or less, of hydrogen-3 or carbon-14 per gram of
26 medium used for liquid scintillation counting; and

27 (b) 1.85 kBq (0.05 uCi) or less, of hydrogen-3 or carbon-14 per gram of
28 animal tissue, averaged over the weight of the entire animal.

1 (2) A licensee or registrant shall not dispose of tissue pursuant to
2 Subsection R313-15-1005(1)(b) in a manner that would permit its use either as
3 food for humans or as animal feed.

4 (3) The licensee or registrant shall maintain records in accordance with
5 Section R313-15-1109.

6
7 **R313-15-1006. Transfer for Disposal and Manifests.**

8 (1) ~~[The ~~r~~]~~Requirements of Section R313-15-1006 and Appendix F and G of
9 10 CFR 20.1001 to 20.2402, 199[3] ed. [~~which is incorporated by reference,~~]

10 (a) The requirements of Section R313-15-1006 and Appendix F and G of 10
11 CFR 20.1001 to 20.2402, 1997 ed., which are incorporated into these rules by
12 reference, are designed to:

13 (i) control transfers of low-level radioactive waste [~~intended for~~
14 disposal at a licensed low-level radioactive waste disposal facility, establish
15 a manifest tracking system, and supplement existing requirements concerning
16 transfers and recordkeeping for those wastes], by any waste generator, waste
17 collector, or waste processor licensee, as defined in Appendix F or G in 10 CFR
18 20.1001 to 20.2402, 1997 ed., who ships low-level waste either directly, or
19 indirectly through a waste collector or waste processor, to a licensed low-level
20 waste land disposal facility as defined in Section R313-25-2;

21 (ii) establish a manifest tracking system; and

22 (iii) supplement existing requirements concerning transfers and
23 recordkeeping for those wastes.

24 (b) Beginning March 1, 1998, all affected licensees must use Appendix G
25 of 10 CFR 20.1001 to 20.2402, 1997 ed., which is incorporated into these rules
26 by reference. Prior to March 1, 1998, a low-level waste disposal facility
27 operator or its regulatory authority may require the shipper to use Appendix F
28 or Appendix G of 10 CFR 20.1001 to 20.2402, 1997 ed. Licensees using Appendix

1 F shall comply with Subsection R313-15-1006(2)(a). Licensees using Appendix G
2 shall comply with Subsection R313-15-1006(2)(b).

3 (2) Shipment of Radioactive Waste.

4 (a) Each shipment of radioactive waste designated for disposal at a
5 licensed low-level radioactive waste disposal facility shall be accompanied by
6 a shipment manifest as specified in Section I of Appendix F of 10 CFR 20.1001 to
7 20.2402, 199[3] ed., which is incorporated by reference.

8 (b) Any licensee shipping radioactive waste intended for ultimate disposal
9 at a licensed land disposal facility must document the information required on
10 the U.S. Nuclear Regulatory Commission's Uniform Low-Level Radioactive Waste
11 Manifest and transfer this recorded information to the intended consignee in
12 accordance with Appendix G to 10 CFR 20.1001 to 20.2402, 1997 ed., which is
13 incorporated into these rules by reference.

14 (3) Each shipment manifest shall include a certification by the waste
15 generator as specified in Section II of Appendix F or G, as appropriate, of 10
16 CFR 20.1001 to 20.2402, 199[3] ed., which is incorporated by reference. See
17 Subsection R313-15-1006(1)(b) to determine the appropriate Appendix.

18 (4) Each person involved in the transfer of waste for disposal or in the
19 disposal of waste, including the waste generator, waste collector, waste
20 processor, and disposal facility operator, shall comply with the requirements
21 specified in Section III of Appendix F or G, as appropriate, of 10 CFR 20.1001
22 to 20.2402, 199[3] ed., which is incorporated by reference. See Subsection
23 R313-15-1006(1)(b) to determine the appropriate Appendix.

24
25 **R313-15-1007. Compliance with Environmental and Health Protection Rules.**

26 Nothing in Sections R313-15-1001, R313-15-1002, R313-15-1003, R313-15-1004,
27 R313-15-1005, or R313-15-1006 relieves the licensee or registrant from complying
28 with other applicable Federal, State and local rules governing any other toxic

or hazardous properties of materials that may be disposed of pursuant to Sections R313-15-1001, R313-15-1002, R313-15-1003, R313-15-1004, R313-15-1005, or R313-15-1006.

R313-15-1008. Classification and Characteristics of Low-Level Radioactive Waste.

(1) Classification of Radioactive Waste for Land Disposal

(a) Considerations. Determination of the classification of radioactive waste involves two considerations. First, consideration shall be given to the concentration of long-lived radionuclides (and their shorter-lived precursors) whose potential hazard will persist long after such precautions as institutional controls, improved waste form, and deeper disposal have ceased to be effective. These precautions delay the time when long-lived radionuclides could cause exposures. In addition, the magnitude of the potential dose is limited by the concentration and availability of the radionuclide at the time of exposure. Second, consideration shall be given to the concentration of shorter-lived radionuclides for which requirements on institutional controls, waste form, and disposal methods are effective.

(b) Classes of waste.

(i) Class A waste is waste that is usually segregated from other waste classes at the disposal site. The physical form and characteristics of Class A waste shall meet the minimum requirements set forth in Subsection R313-15-1008(2)(a). If Class A waste also meets the stability requirements set forth in Subsection R313-15-1008(2)(b), it is not necessary to segregate the waste for disposal.

(ii) Class B waste is waste that shall meet more rigorous requirements on waste form to ensure stability after disposal. The physical form and characteristics of Class B waste shall meet both the minimum and stability requirements set forth in Subsection R313-15-1008(2).

(iii) Class C waste is waste that not only shall meet more rigorous requirements on waste form to ensure stability but also requires additional measures at the disposal facility to protect against inadvertent intrusion. The physical form and characteristics of Class C waste shall meet both the minimum and stability requirements set forth in Subsection R313-15-1008(2).

(c) Classification determined by long-lived radionuclides. If the radioactive waste contains only radionuclides listed in Table I, classification shall be determined as follows:

(i) If the concentration does not exceed 0.1 times the value in Table I, the waste is Class A.

(ii) If the concentration exceeds 0.1 times the value in Table I, but does not exceed the value in Table I, the waste is Class C.

(iii) If the concentration exceeds the value in Table I, the waste is not generally acceptable for land disposal.

(iv) For wastes containing mixtures of radionuclides listed in Table I, the total concentration shall be determined by the sum of fractions rule described in Subsection R313-15-1008(1)(g).

TABLE I

Concentration

Radionuclide	curie/cubic meter(1)	nanocurie/gram(2)
C-14	8	
C-14 in activated metal	80	
Ni-59 in activated metal	220	
Nb-94 in activated metal	0.2	

1	Tc-99	3	
2	I-129	0.08	
3	Alpha emitting transuranic		
4	radionuclides with half-		
5	life greater than five		
6	years		100
7	Pu-241		3,500
8	Cm-242		20,000
9	Ra-226		100

10

11 NOTE: (1) To convert the Ci/m³ values to gigabecquerel (GBq)/cubic meter
 12 multiply the Ci/m³ value by 37.

13 (2) To convert the nCi/g values to becquerel (Bq)/gram, multiply the nCi/g value
 14 by 37.

15

16 (d) Classification determined by short-lived radionuclides. If the waste
 17 does not contain any of the radionuclides listed in Table I, classification shall
 18 be determined based on the concentrations shown in Table II. However, as
 19 specified in Subsection R313-15-1008(1)(f), if radioactive waste does not contain
 20 any nuclides listed in either Table I or II, it is Class A.

21 (i) If the concentration does not exceed the value in Column 1, the waste
 22 is Class A.

23 (ii) If the concentration exceeds the value in Column 1 but does not
 24 exceed the value in Column 2, the waste is Class B.

25 (iii) If the concentration exceeds the value in Column 2 but does not
 26 exceed the value in Column 3, the waste is Class C.

27 (iv) If the concentration exceeds the value in Column 3, the waste is not
 28 generally acceptable for near-surface disposal.

(v) For wastes containing mixtures of the radionuclides listed in Table II, the total concentration shall be determined by the sum of fractions rule described in Subsection R313-15-1008(1)(g).

TABLE II

Radionuclide	Concentration, curie/cubic meter (1)		
	Column 1	Column 2	Column 3
Total of all radionuclides with less than 5-year half-life	700	(2)	(2)
H-3	40	(2)	(2)
Co-60	700	(2)	(2)
Ni-63	3.5	70	700
Ni-63 in activated metal	35	700	7000
Sr-90	0.04	150	7000
Cs-137	1	44	4600

NOTE: (1) To convert the Ci/m³ value to gigabecquerel (GBq)/cubic meter, multiply the Ci/m³ value by 37.

(2) There are no limits established for these radionuclides in Class B or C wastes. Practical considerations such as the effects of external radiation and internal heat generation on transportation, handling, and disposal will limit the concentrations for these wastes. These wastes shall be Class B unless the concentrations of other radionuclides in Table II determine the waste to be Class

C independent of these radionuclides.

(e) Classification determined by both long- and short-lived radionuclides. If the radioactive waste contains a mixture of radionuclides, some of which are listed in Table I and some of which are listed in Table II, classification shall be determined as follows:

(i) If the concentration of a radionuclide listed in Table I is less than 0.1 times the value listed in Table I, the class shall be that determined by the concentration of radionuclides listed in Table II.

(ii) If the concentration of a radionuclide listed in Table I exceeds 0.1 times the value listed in Table I, but does not exceed the value in Table I, the waste shall be Class C, provided the concentration of radionuclides listed in Table II does not exceed the value shown in Column 3 of Table II.

(f) Classification of wastes with radionuclides other than those listed in Tables I and II. If the waste does not contain any radionuclides listed in either Table I or II, it is Class A.

(g) The sum of the fractions rule for mixtures of radionuclides. For determining classification for waste that contains a mixture of radionuclides, it is necessary to determine the sum of fractions by dividing each radionuclide's concentration by the appropriate limit and adding the resulting values. The appropriate limits shall all be taken from the same column of the same table. The sum of the fractions for the column shall be less than 1.0 if the waste class is to be determined by that column. Example: A waste contains Sr-90 in a concentration of 1.85 TBq/m³ (50 Ci/m³) and Cs-137 in a concentration of 814 GBq/m³ (22 Ci/m³). Since the concentrations both exceed the values in Column 1, Table II, they shall be compared to Column 2 values. For Sr-90 fraction, $50/150 = 0.33.$, for Cs-137 fraction, $22/44 = 0.5$; the sum of the fractions = 0.83. Since the sum is less than 1.0, the waste is Class B.

(h) Determination of concentrations in wastes. The concentration of a radionuclide may be determined by indirect methods such as use of scaling factors which relate the inferred concentration of one radionuclide to another that is measured, or radionuclide material accountability, if there is reasonable assurance that the indirect methods can be correlated with actual measurements. The concentration of a radionuclide may be averaged over the volume of the waste, or weight of the waste if the units are expressed as becquerel (nanocurie) per gram.

(2) Radioactive Waste Characteristics

(a) The following are minimum requirements for all classes of waste and are intended to facilitate handling and provide protection of health and safety of personnel at the disposal site.

(i) Wastes shall be packaged in conformance with the conditions of the license issued to the site operator to which the waste will be shipped. Where the conditions of the site license are more restrictive than the provisions of ~~[Part D]~~ Rule R313-15, the site license conditions shall govern.

(ii) Wastes shall not be packaged for disposal in cardboard or fiberboard boxes.

(iii) Liquid waste shall be packaged in sufficient absorbent material to absorb twice the volume of the liquid.

(iv) Solid waste containing liquid shall contain as little free-standing and non-corrosive liquid as is reasonably achievable, but in no case shall the liquid exceed one percent of the volume.

(v) Waste shall not be readily capable of detonation or of explosive decomposition or reaction at normal pressures and temperatures, or of explosive reaction with water.

(vi) Waste shall not contain, or be capable of generating, quantities of toxic gases, vapors, or fumes harmful to persons transporting, handling, or

1 disposing of the waste. This does not apply to radioactive gaseous waste
2 packaged in accordance with Subsection R313-15-1008(2)(a)(viii).

3 (vii) Waste shall not be pyrophoric. Pyrophoric materials contained in
4 wastes shall be treated, prepared, and packaged to be nonflammable.

5 (viii) Wastes in a gaseous form shall be packaged at an absolute pressure
6 that does not exceed 1.5 atmospheres at 20 degrees celsius. Total activity shall
7 not exceed 3.7 TBq (100 Ci) per container.

8 (ix) Wastes containing hazardous, biological, pathogenic, or infectious
9 material shall be treated to reduce to the maximum extent [~~practicable~~]practical
10 the potential hazard from the non-radiological materials.

11 (b) The following requirements are intended to provide stability of the
12 waste. Stability is intended to ensure that the waste does not degrade and
13 affect overall stability of the site through slumping, collapse, or other failure
14 of the disposal unit and thereby lead to water infiltration. Stability is also
15 a factor in limiting exposure to an inadvertent intruder, since it provides a
16 recognizable and nondispersible waste.

17 (i) Waste shall have structural stability. A structurally stable waste
18 form will generally maintain its physical dimensions and its form, under the
19 expected disposal conditions such as weight of overburden and compaction
20 equipment, the presence of moisture, and microbial activity, and internal factors
21 such as radiation effects and chemical changes. Structural stability can be
22 provided by the waste form itself, processing the waste to a stable form, or
23 placing the waste in a disposal container or structure that provides stability
24 after disposal.

25 (ii) Notwithstanding the provisions in Subsections R313-15-1008(2)(a)(iii)
26 and R313-15-1008(2)(a)(iv), liquid wastes, or wastes containing liquid, shall be
27 converted into a form that contains as little free-standing and non-corrosive
28 liquid as is reasonably achievable, but in no case shall the liquid exceed one

1 percent of the volume of the waste when the waste is in a disposal container
2 designed to ensure stability, or 0.5 percent of the volume of the waste for waste
3 processed to a stable form.

4 (iii) Void spaces within the waste and between the waste and its package
5 shall be reduced to the extent [~~practicable~~] practical.

6 (3) Labeling. Each package of waste shall be clearly labeled to identify
7 whether it is Class A, Class B, or Class C waste, in accordance with Subsection
8 R313-15-1008(1).

9
10 **R313-15-1101. Records - General Provisions.**

11 (1) Each licensee or registrant shall use the SI units becquerel, gray,
12 sievert and coulomb per kilogram, or the special units, curie, rad, rem, and
13 roentgen, including multiples and subdivisions, and shall clearly indicate the
14 units of all quantities on records required by Rule R313-15.

15 (2) Notwithstanding the requirements of Subsection R313-15-1101(1), when
16 recording information on shipment manifests, as required in Subsection R313-15-
17 1006(2), information must be recorded in SI units or in SI units and the special
18 units specified in Subsection R313-15-1101(1).

19 ~~[(2)]~~ (3) The licensee or registrant shall make a clear distinction among
20 the quantities entered on the records required by Rule R313-15, such as, total
21 effective dose equivalent, total organ dose equivalent, shallow dose equivalent,
22 eye dose equivalent, deep dose equivalent, or committed effective dose
23 equivalent.

24
25 **R313-15-1102. Records of Radiation Protection Programs.**

26 (1) Each licensee or registrant shall maintain records of the radiation
27 protection program, including:

28 (a) The provisions of the program; and

(b) Audits and other reviews of program content and implementation.

(2) The licensee or registrant shall retain the records required by Subsection R313-15-1102(1)(a) until the Executive Secretary terminates each pertinent license or registration requiring the record. The licensee or registrant shall retain the records required by Subsection R313-15-1102(1)(b) for three years after the record is made.

R313-15-1103. Records of Surveys.

(1) Each licensee or registrant shall maintain records showing the results of surveys and calibrations required by Section R313-15-501 and Subsection R313-15-906(2). The licensee or registrant shall retain these records for three years after the record is made.

(2) The licensee or registrant shall retain each of the following records until the Executive Secretary terminates each pertinent license or registration requiring the record:

(a) Records of the results of surveys to determine the dose from external sources of radiation used, in the absence of or in combination with individual monitoring data, in the assessment of individual dose equivalents; and

(b) Records of the results of measurements and calculations used to determine individual intakes of radioactive material and used in the assessment of internal dose; and

(c) Records showing the results of air sampling, surveys, and bioassays required pursuant to Subsections R313-15-703(1)(c)(i) and R313-15-703(1)(c)(ii); and

(d) Records of the results of measurements and calculations used to evaluate the release of radioactive effluents to the environment.

R313-15-1104. Records of Tests for Leakage or Contamination of Sealed Sources.

1 Records of tests for leakage or contamination of sealed sources required
2 by Section R313-15-401 shall be kept in units of becquerel or microcurie and
3 maintained for inspection by the Executive Secretary for five years after the
4 records are made.

5
6 **R313-15-1105. Records of Prior Occupational Dose.**

7 For each individual who is likely to receive in a year an occupational dose
8 requiring monitoring pursuant to Section R313-15-502, [F]the licensee or
9 registrant shall retain the records of prior occupational dose and exposure
10 history as specified in Section R313-15-205 on form DRC-05 or equivalent until
11 the Executive Secretary terminates each pertinent license requiring this record.
12 The licensee or registrant shall retain records used in preparing form DRC-05 or
13 equivalent for three years after the record is made.

14
15 **R313-15-1106. Records of Planned Special Exposures.**

16 (1) For each use of the provisions of Section R313-15-206 for planned
17 special exposures, the licensee or registrant shall maintain records that
18 describe:

19 (a) The exceptional circumstances requiring the use of a planned special
20 exposure; and

21 (b) The name of the management official who authorized the planned special
22 exposure and a copy of the signed authorization; and

23 (c) What actions were necessary; and

24 (d) Why the actions were necessary; and

25 (e) What precautions were taken to assure that doses were maintained
26 ALAC and

27 (f) What individual and collective doses were expected to result; and

28 (g) The doses actually received in the planned special exposure.

(2) The licensee or registrant shall retain the records until the Executive Secretary terminates each pertinent license or registration requiring these records.

R313-15-1107. Records of Individual Monitoring Results.

(1) Recordkeeping Requirement. Each licensee or registrant shall maintain records of doses received by all individuals for whom monitoring was required pursuant to Section R313-15-502, and records of doses received during planned special exposures, accidents, and emergency conditions. Assessments of dose equivalent and records made using units in effect before January 1, 1994, need not be changed. These records shall include, when applicable:

(a) The deep dose equivalent to the whole body, eye dose equivalent, shallow dose equivalent to the skin, and shallow dose equivalent to the extremities; and

(b) The estimated intake of radionuclides, see Section R313-15-202; and

(c) The committed effective dose equivalent assigned to the intake of radionuclides; and

(d) The specific information used to calculate the committed effective dose equivalent pursuant to Subsection R313-15-204(3); and

(e) The total effective dose equivalent when required by Section R313-15-202; and

(f) The total of the deep dose equivalent and the committed dose to the organ receiving the highest total dose.

(2) Recordkeeping Frequency. The licensee or registrant shall make entries of the records specified in Subsection R313-15-1107(1) at intervals not to exceed one year.

(3) Recordkeeping Format. The licensee or registrant shall maintain the records specified in Subsection R313-15-1107(1) on form DRC-06, in accordance

1 with the instructions for form DRC-06, or in clear and legible records containing
2 all the information required by form DRC-06.

3 (4) The licensee or registrant shall maintain the records of dose to an
4 embryo/fetus with the records of dose to the declared pregnant woman. The
5 declaration of pregnancy, including the estimated date of conception, shall also
6 be kept on file, but may be maintained separately from the dose records.

7 (5) The licensee or registrant shall retain each required form or record
8 until the Executive Secretary terminates each pertinent license or registration
9 requiring the record.

10
11 **R313-15-1108. Records of Dose to Individual Members of the Public.**

12 (1) Each licensee or registrant shall maintain records sufficient to
13 demonstrate compliance with the dose limit for individual members of the public.
14 See Section R313-15-301.

15 (2) The licensee or registrant shall retain the records required by
16 Subsection R313-15-1108(1) until the Executive Secretary terminates each
17 pertinent license or registration requiring the record. Requirements for
18 disposition of these records, prior to license termination, are located in
19 Section R313-12-51 for activities licensed under these rules.

20
21 **R313-15-1109. Records of Waste Disposal.**

22 (1) Each licensee or registrant shall maintain records of the disposal of
23 licensed or registered materials made pursuant to Sections R313-15-1002, R313-15-
24 1003, R313-15-1004, R313-15-1005, Rule R313-25, and disposal by burial in soil,
25 including burials authorized before January 28, 1981.

26 (2) The licensee or registrant shall retain the records required by
27 Subsection R313-15-1109(1) until the Executive Secretary terminates each
28 pertinent license or registration requiring the record.

1 R313-15-1110. Records of Testing Entry Control Devices for Very High Radiation
2 Areas.

3 (1) Each licensee or registrant shall maintain records of tests made
4 pursuant to Subsection R313-15-603(2)(i) on entry control devices for very high
5 radiation areas. These records shall include the date, time, and results of each
6 such test of function.

7 (2) The licensee or registrant shall retain the records required by
8 Subsection R313-15-1110(1) for three years after the record is made.

9
10 R313-15-1111. Form of Records.

11 Each record required by Rule R313-15 shall be legible throughout the
12 specified retention period. The record shall be the original or a reproduced copy
13 or a microform, provided that the copy or microform is authenticated by
14 authorized personnel and that the microform is capable of producing a clear copy
15 throughout the required retention period or the record may also be stored in
16 electronic media with the capability for producing legible, accurate, and
17 complete records during the required retention period. Records, such as letters,
18 drawings, and specifications, shall include all pertinent information, such as
19 stamps, initials, and signatures. The licensee shall maintain adequate
20 safeguards against tampering with and loss of records.

21 "C"

22 R313-15-1201. Reports of Stolen, Lost, or Missing Licensed or Registered Sources
23 of Radiation.

24 (1) Telephone Reports. Each licensee or registrant shall report to the
25 Executive Secretary by telephone each stolen, lost, or missing source of
26 radiation immediately after its absence becomes known to the licensee or
27 registrant. This requirement does not apply to sources of radiation that are not
28 required to be licensed or registered. (as follows:--)

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1 ~~[(a) Immediately after its occurrence becomes known to the licensee or~~
2 ~~registrant, stolen, lost, or missing licensed or registered radioactive material~~
3 ~~in an aggregate quantity equal to or greater than 1,000 times the quantity~~
4 ~~specified in Appendix C of 10 CFR 20.1001 to 20.2402, 1993 ed., which is~~
5 ~~incorporated by reference, under such circumstances that it appears to the~~
6 ~~licensee or registrant that an exposure could result to individuals in~~
7 ~~unrestricted areas;~~

8 ~~----- (b) Within 30 days after its occurrence becomes known to the licensee or~~
9 ~~registrant, lost, stolen, or missing licensed or registered radioactive material~~
10 ~~in an aggregate quantity greater than ten times the quantity specified in~~
11 ~~Appendix C of 10 CFR 20.1001 to 20.2402, 1993 ed., which is incorporated by~~
12 ~~reference, that is still missing.~~

13 ~~----- (c) Immediately after its occurrence becomes known to the registrant, a~~
14 ~~stolen, lost, or missing radiation machine.]~~

15 (2) Written Reports. Each licensee or registrant required to make a
16 report pursuant to Subsection R313-15-1201(1) shall, within 30 days after making
17 the telephone report, make a written report to the Executive Secretary setting
18 forth the following information:

19 (a) A description of the licensed or registered source of radiation
20 involved, including, for radioactive material, the kind, quantity, and chemical
21 and physical form; and, for radiation machines, the manufacturer, model and
22 serial number, type and maximum energy of radiation emitted;

23 (b) A description of the circumstances under which the loss or theft
24 occurred; and

25 (c) A statement of disposition, or probable disposition, of the licensed
26 or registered source of radiation involved; and

27 (d) Exposures of individuals to radiation, circumstances under which the
28 exposures occurred, and the possible total effective dose equivalent to persons

1 in unrestricted areas; and

2 (e) Actions that have been taken, or will be taken, to recover the source
3 of radiation; and

4 (f) Procedures or measures that have been, or will be, adopted to ensure
5 against a recurrence of the loss or theft of licensed or registered sources of
6 radiation.

7 (3) Subsequent to filing the written report, the licensee or registrant
8 shall also report additional substantive information on the loss or theft within
9 30 days after the licensee or registrant learns of such information.

10 (4) The licensee or registrant shall prepare any report filed with the
11 Executive Secretary pursuant to Section R313-15-1201 so that names of individuals
12 who may have received exposure to radiation are stated in a separate and
13 detachable portion of the report.

14
15 **R313-15-1202. Notification of Incidents.**

16 (1) Immediate Notification. Notwithstanding other requirements for
17 notification, each licensee or registrant shall immediately report each event
18 involving a source of radiation possessed by the licensee or registrant that may
19 have caused or threatens to cause any of the following conditions:

20 (a) An individual to receive:
21 (i) A total effective dose equivalent of 0.25 Sv (25 rem) or more; or
22 (ii) An eye dose equivalent of 0.75 Sv (75 rem) or more; or
23 (iii) A shallow dose equivalent to the skin or extremities or a total
24 organ dose equivalent of 2.5 Gy (250 rad) or more; or

25 (b) The release of radioactive material, inside or outside of a restricted
26 area, so that, had an individual been present for 24 hours, the individual could
27 have received an intake five times the occupational ALI. This provision does not
28 apply to locations where personnel are not normally stationed during routine

1 operations, such as hot-cells or process enclosures.

2 (2) Twenty-Four Hour Notification. Each licensee or registrant shall,
3 within 24 hours of discovery of the event, report to the Executive Secretary each
4 event involving loss of control of a licensed or registered source of radiation
5 possessed by the licensee or registrant that may have caused, or threatens to
6 cause, any of the following conditions:

7 (a) An individual to receive, in a period of 24 hours:

8 (i) A total effective dose equivalent exceeding 0.05 Sv (five rem); or

9 (ii) An eye dose equivalent exceeding 0.15 Sv (15 rem); or

10 (iii) A shallow dose equivalent to the skin or extremities or a total
11 organ dose equivalent exceeding 0.5 Sv (50 rem); or

12 (b) The release of radioactive material, inside or outside of a restricted
13 area, so that, had an individual been present for 24 hours, the individual could
14 have received an intake in excess of one occupational ALI. This provision does
15 not apply to locations where personnel are not normally stationed during routine
16 operations, such as hot-cells or process enclosures.

17 (3) The licensee or registrant shall prepare each report filed with the
18 Executive Secretary pursuant to Section R313-15-1202 so that names of individuals
19 who have received exposure to sources of radiation are stated in a separate and
20 detachable portion of the report.

21 (4) Licensees or registrants shall make the reports required by
22 Subsections R313-15-1202(1) and R313-15-1202(2) to the Executive Secretary by
23 telephone, telegram, mailgram, or facsimile to the Executive Secretary.

24 (5) The provisions of Section R313-15-1202 do not apply to doses that
25 result from planned special exposures, provided such doses are within the limits
26 for planned special exposures and are reported pursuant to Section R313-15-1204.

27
28 **R313-15-1203. Reports of Exposures, Radiation Levels, and Concentrations of**

1 **Radioactive Material Exceeding the Constraints or Limits.**

2 (1) Reportable Events. In addition to the notification required by
3 Section R313-15-1202, each licensee or registrant shall submit a written report
4 within 30 days after learning of any of the following occurrences:

5 (a) Incidents for which notification is required by Section R313-15-1202;
6 or

7 (b) Doses in excess of any of the following:

8 (i) The occupational dose limits for adults in Section R313-15-201; or

9 (ii) The occupational dose limits for a minor in Section R313-15-207; or

10 (iii) The limits for an embryo/fetus of a declared pregnant woman in
11 Section R313-15-208; or

12 (iv) The limits for an individual member of the public in Section R313-15-
13 301; or

14 (v) Any applicable limit in the license or registration; or

15 (vi) The ALARA constraints for air emissions established under Subsection
16 R313-15-101(4); or

17 (c) Levels of radiation or concentrations of radioactive material in:

18 (i) A restricted area in excess of applicable limits in the license or
19 registration; or

20 (ii) An unrestricted area in excess of ten times the applicable limit set
21 forth in Rule R313-15 or in the license or registration, whether or not involving
22 exposure of any individual in excess of the limits in Section R313-15-301; or

23 (d) For licensees subject to the provisions of U.S. Environmental
24 Protection Agency's generally applicable environmental radiation standards in 40
25 CFR 190, levels of radiation or releases of radioactive material in excess of
26 those standards, or of license conditions related to those standards.

27 (2) Contents of Reports.

28 (a) Each report required by Subsection R313-15-1203(1) shall describe the

1 extent of exposure of individuals to radiation and radioactive material,
2 including, as appropriate:

3 (i) Estimates of each individual's dose; and

4 (ii) The levels of radiation and concentrations of radioactive material
5 involved; and

6 (iii) The cause of the elevated exposures, dose rates, or concentrations;
7 and

8 (iv) Corrective steps taken or planned to ensure against a recurrence,
9 including the schedule for achieving conformance with applicable limits, ALARA
10 constraints, generally applicable environmental standards, and associated license
11 or registration conditions.

12 (b) Each report filed pursuant to Subsection R313-15-1203(1) shall include
13 for each occupationally overexposed individual[~~exposed~~]: the name, Social
14 Security account number, and date of birth. With respect to the limit for the
15 embryo/fetus in Section R313-15-208, the identifiers should be those of the
16 declared pregnant woman. The report shall be prepared so that this information
17 is stated in a separate and detachable portion of the report.

18 (3) All licensees or registrants who make reports pursuant to Subsection
19 R313-15-1203(1) shall submit the report in writing to the Executive Secretary.

20
21 **R313-15-1204. Reports of Planned Special Exposures.**

22 The licensee or registrant shall submit a written report to the Executive
23 Secretary within 30 days following any planned special exposure conducted in
24 accordance with Section R313-15-206, informing the Executive Secretary that a
25 planned special exposure was conducted and indicating the date the planned
26 special exposure occurred and the information required by Section R313-15-1106.

27
28 **R313-15-1205. Reports to Individuals of Exceeding Dose Limits.**

1 When a licensee or registrant is required, pursuant to the provisions of
2 Sections R313-15-1203 or R313-15-1204, to report to the Executive Secretary any
3 exposure of an identified occupationally exposed individual, or an identified
4 member of the public, to sources of radiation, the licensee or registrant shall
5 also provide a copy of the report submitted to the Executive Secretary to the
6 individual. This report shall be transmitted at a time no later than the
7 transmittal to the Executive Secretary.

8
9 **R313-15-1207. Notifications and Reports to Individuals.**

10 (1) Requirements for notification and reports to individuals of exposure
11 to radiation or radioactive material are specified in Rule R313-18.

12 (2) When a licensee or registrant is required pursuant to Section R313-15-
13 1203 to report to the Executive Secretary any exposure of an individual to
14 radiation or radioactive material, the licensee or registrant shall also notify
15 the individual. Such notice shall be transmitted at a time not later than the
16 transmittal to the Executive Secretary, and shall comply with the provisions of
17 Rule R313-18.

18
19 **R313-15-1208. Reports of Leaking or Contaminated Sealed Sources.**

20 If the test for leakage or contamination required pursuant to Section R313-
21 15-401 indicates a sealed source is leaking or contaminated, a report of the test
22 shall be filed within five days with the Executive Secretary describing the
23 equipment involved, the test results and the corrective action taken.

24
25 **R313-15-1301. Vacating Premises.**

26 Each specific licensee or registrant shall, no less than 30 days before
27 vacating or relinquishing possession or control of premises which may have been
28 contaminated with radioactive material as a result of his activities, notify the

1 Executive Secretary in writing of intent to vacate. When deemed necessary by the
2 Executive Secretary, the licensee shall decontaminate the premises in such a
3 manner as the Executive Secretary may specify.
4

5 KEY: radioactive material, contamination, waste disposal, safety

6 [~~1993~~] 1998

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