

UNITED STATES OF AMERICA NUCLEAR REGULATORY COMMISSION

Title: BRIEFING BY IIT ON LOSS OF IRIIDIUM-192
SOURCE AND THERAPY MISADMINISTRATION AT
INDIANA REGIONAL CANCER CENTER, INDIANA,
PA., NOVEMBER 16, 1992

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NUCLEAR REGULATORY COMMISSION

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BRIEFING BY IIT ON LOSS OF IRIDIUM-192 SOURCE
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REGIONAL CANCER CENTER, INDIANA, PA.,
NOVEMBER 16, 1992

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PUBLIC MEETING

Nuclear Regulatory Commission
One White Flint North
Rockville, Maryland

Monday, February 8, 1993

The Commission met in open session,
pursuant to notice, at 10:00 a.m., Ivan Selin,
Chairman, presiding.

COMMISSIONERS PRESENT:

IVAN SELIN, Chairman of the Commission
KENNETH C. ROGERS, Commissioner
FORREST J. REMICK, Commissioner
JAMES R. CURTISS, Commissioner
E. GAIL de PLANQUE, Commissioner

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STAFF AND PRESENTERS SEATED AT THE COMMISSION TABLE:

WILLIAM C. PARLER, General Counsel

JAMES TAYLOR, Executive Director for Operations

ED JORDAN, Director, AEOD

DR. CARL PAPERIELLO, RIII, Team Leader

CYNTHIA JONES, NMSS, Radiological Dose Evaluation

DR. MOHAMED SHANBAKY, RI, Deputy Team Leader

DR. DANIEL FLYNN, Medical Consultant

THOMAS RICH, NMSS, Engineering Evaluation

WILLIAM DAMASKA, Acting Director, Division of
Standards and Enforcement, Office of Compliance and
Surveillance, Center for Devices and Radiological
Health, FDA

DR. THOMAS STREAMS, Indiana County, Pennsylvania
Coroner

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P-R-O-C-E-E-D-I-N-G-S

10:00 a.m.

CHAIRMAN SELIN: Good morning, ladies and gentlemen.

The Commission is meeting this morning to receive a briefing from the NRC staff on the investigation into the medical therapy misadministration and the subsequent patient death that occurred in Indiana, Pennsylvania in November 1992.

Also here today are representatives of the U.S. Food and Drug Administration and Indiana County Coroner Thomas L. Streams, both of whom will participate briefly in the meeting.

The official autopsy report of the Indiana County Coroner's Office lists the cause of death unequivocally as acute radiation exposure and consequences thereof. This fatality, as well as the other troubling aspects of this extremely unfortunate incident, are of great concern to the Commission. The staff's investigation team has borne the important responsibility of conducting a comprehensive and penetrating evaluation of the circumstances leading to these regrettable events. Copies of their report are available here in the conference room. It is this

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1 report and its findings which will be briefed to us
2 today.

3 Our goal is to make sure that where there
4 are improvements needed in the regulatory program they
5 are quickly identified and made. We must not allow
6 the sobering lessons from this experience to go
7 unlearned. The number of medical administrations is
8 certainly a small fraction of the total number of
9 nuclear medicine treatments. In the therapeutic area
10 it seems to be on the order of one in 3,000. That's
11 a figure in which the nuclear medical industry can be
12 quite proud.

13 Nevertheless, it doesn't follow that we
14 should just not work to reduce the number and the
15 consequences of avoidable mistakes. We don't want to
16 dissuade patients from receiving needed care, but the
17 report we will discuss today clearly demonstrates that
18 there are serious questions which need to be answered
19 about our own regulation in this area. We will not
20 shy away from our duty to the public to provide a
21 regulatory program which is logical, consistent and
22 effective.

23 NRC has suspended the license of the
24 company involved in this administration. It is not
25 the purpose of this briefing to discuss the merits of

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1 this suspension. Rather, the purpose is to obtain
2 information for the Commission to evaluate the need
3 for further action, both with regard for NRC
4 regulation in this field and this event in particular.
5 The suspension action will not and should not be
6 discussed at the meeting this morning.

7 Do my fellow Commissioners wish to say
8 anything to me?

9 Mr. Taylor, the floor is yours. You might
10 introduce the members of your team as a start.

11 MR. TAYLOR: Yes, sir. I chartered the
12 team, the incident investigation team, on December 4th
13 based upon the complexity of the event and involving
14 potential over exposure of members of the public as
15 well as clinic and nursing home personnel, and the
16 complexity involving the supplier of the device being
17 from an agreement state and because of FDA
18 involvement. I made that decision and directed Mr.
19 Jordan to initiate this team in order to obtain as
20 many answers as we could about this event.

21 On that date, the team was chartered to be
22 led by -- and Mr. Jordan is here at the table, of
23 course -- Doctor Carl Paperiello from Region III as
24 the Team Leader. Also Doctor Mohamed Shanbaky, Region
25 I as the Assistant Team Leader, Cynthia Jones and

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1 Thomas Rich from NMSS here at the table also, and
2 other NRC staff that Doctor Paperiello will introduce.

3 I would also introduce -- assisting the
4 IIT is NRC's medical consultant, Doctor Daniel Flynn,
5 who is here at the table. Also at the table, Mr.
6 William Damaska, Acting Director of the Division of
7 Standards and Enforcement, FDA, far right, and Doctor
8 Thomas Streams, Indiana County Coroner who came down
9 to be with us today.

10 With those opening remarks, I'll ask
11 Doctor Paperiello to continue.

12 DOCTOR PAPERIELLO: Good morning. I'm
13 Carl Paperiello and I led this IIT that investigated
14 the loss of the iridium-192 source and the therapy
15 misadministration at the Indiana Regional Cancer
16 Center.

17 (Slide) Before I begin, if I could have
18 slide number 2, I would like to introduce the rest my
19 team, recognize Penny Nessen from Region I who was
20 involved in some of the -- assisted Cynthia Jones with
21 dose calculations, reviewed the service records for
22 the PrimAlert monitor and also did most of the
23 facility surveys that we briefly discuss in the
24 report. Both Ron Lloyd and Allen Madison from AEOD,
25 who I sort of expropriated for the team when I

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1 arrived, they were there and I needed more people and
2 I sort of took them on.

3 (Slide) Could I have the next slide?

4 I have to thank the administrative staff.
5 There was a lot of work in getting this report out in
6 a short period of time and I would like to see them
7 recognized.

8 (Slide) If I could have slide 4.

9 In this investigation we centered on three
10 major issues. One was the failure of the Omnitron
11 2000 source wire. Why did it break? Secondly, given
12 the breakage, why did the Indiana Regional Cancer
13 Center fail to detect this break? Thirdly, the
14 consequences of the failure. The agenda shows the
15 sequence I wish to follow in this presentation.

16 (Slide) Could I have the fifth slide,
17 please?

18 I propose to give you a brief chronology
19 following the movement of the source from its use and
20 loss on November 16th until its retrieval on December
21 1. I'm going to discuss very briefly a subsequent
22 event involving the failure of another Omnitron 2000
23 source wire in Pittsburgh for which the prompt
24 identification and intervention prevented significant
25 consequences.

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1 (Slide) May I have slide number 6?

2 On November 16th, 1992, an elderly patient
3 was treated for an anal carcinoma at the Indiana
4 Regional Cancer Center in Indiana, Pennsylvania using
5 high dose rate brachytherapy. This center is one of
6 a number of treatment centers with HDR units operated
7 by the Oncology Services Corporation, the license
8 holder. The patient died on the evening
9 of November 21, 1992, five days after the treatment.
10 Prior to the treatment, five catheters, plastic tubes,
11 were placed in the tumor. During the treatment an
12 approximately 4.3 curie iridium-192 source was to be
13 placed at various positions in each catheter to
14 irradiate the tumor using an Omnitron 2000 remote
15 afterloader. To give you an idea, this wire is a
16 measurement wire, but it gives you the diameter of the
17 wire used in the treatment. The source is actually in
18 a cavity in the tip of the wire and the remote
19 afterloader is a computer driven device that allows
20 the source to be positioned precisely in a plastic
21 tube in the patient.

22 The treatment was conducted in the same
23 shielded facility in which the linear accelerator
24 treatments are conducted. The treatment was to be the
25 first in a series of three 600 rad treatments planned

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1 by the physician and the five catheters were to remain
2 in the patient. After a trial run through the five
3 catheters with a dummy wire, the iridium source wire
4 was placed in four catheters without difficulty.
5 After several unsuccessful attempts to insert the
6 source wire and the dummy wire into a fifth catheter,
7 the treatment was terminated.

8 An area radiation monitor, which happened
9 to have the model, the PrimAlert-10, we're going to
10 talk about PrimAlert-10, but it's an area radiation
11 monitor similar to those used in nuclear power plants,
12 except this device is a lot smaller. In the treatment
13 area, it was observed in the alarm condition at some
14 point at times when the source should have been
15 retracted. In other words, they had trouble getting
16 the source into the tube.

17 They went in the room, the source should
18 have been back into the afterloader. They went to
19 check the patient trying to find connections that were
20 blocking the ability of the wire to get in. The
21 technologists aren't sure when the light came in, but
22 somewhere in that sequence the technologist observed
23 that the PrimAlert was lit.

24 Eventually all three technologists and the
25 physician attending the patient were aware of the

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1 alarm condition, but no one conducted a survey of
2 radiation levels with the available portable Geiger
3 counter. The only action taken was to check the
4 control console of the HDR afterloader. Because the
5 console indicator showed safe and there's a panel of
6 lights there and markings on it, safe meant that the
7 source was in the safe. They called the lead
8 container in the afterloader a safe. They believed
9 the source to be fully retracted into the lead shield
10 and assumed the area radiation monitor was
11 malfunctioning.

12 (Slide) Could I have the next slide?

13 This slide shows the isodose curves
14 associated with the patient once the patient was
15 removed from the room. The patient spent roughly 50
16 minutes in the treatment room. After the attempts to
17 treat the fifth catheter, the physician decided to
18 terminate the treatment. The patient was taken from
19 the treatment room to another room outside of the
20 treatment room. One catheter was observed to be loose
21 and was removed at this point.

22 Just to give you a point of reference, at
23 one meter or about a yard from the source, the dose
24 rate is approximately two R per hour. Typically,
25 sources, HDR sources are approximately one-tenth the

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1 activity of industrial radiography sources. So, it
2 gives you an idea. If you know what industrial
3 radiography sources can do, this source is one-tenth
4 the size.

5 COMMISSIONER REMICK: Carl, a question.
6 The isodose curves there are circular. I would expect
7 if you took account of shielding and build-up and
8 scattering --

9 DOCTOR PAPERIELLO: We didn't take into
10 account -- you're right, we did not take account of
11 shielding in our dose calculations. Why? If you do
12 the calculations for shielding at a patient's body and
13 you take a look at shielding, the attenuation and
14 build-up, for the amount of tissue involved they
15 almost cancel. There's not that much attenuation in
16 the body. Most of the walls are wallboard and there's
17 very little -- when you start taking a look at your
18 calculations and when you start asking people where
19 they were three weeks ago and how long, the
20 uncertainties in location and time are much greater
21 than trying to deal with all the shielding effects
22 until you got in the concrete are ten, 20 percent
23 effects. It just wasn't worthwhile trying to pursue
24 that. We did it, we found out that they were small
25 effects and we just ignored them after that.

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1 COMMISSIONER REMICK: How about scattering
2 off the floors, which I assume were concrete?

3 DOCTOR PAPERIELLO: No, we didn't take
4 that into account. On the other hand, as we get into
5 it, the cytogenetic results and the calculated results
6 are in pretty good agreement, particularly when you
7 consider the uncertainties.

8 (Slide) Could I have slide number 8?

9 The patient was returned to the nursing
10 home where she resided with four catheters, one
11 containing the source in her body and the source
12 remained in the patient's body for almost four days.

13 (Slide) Can I have slide 9?

14 This slide shows the radiation fields in
15 the nursing home associated with the source in the
16 patient's body. The other bed in the patient's room
17 was unoccupied, but there were two residents in
18 bedroom number 6. The catheter with the source came
19 loose on the fourth day and eventually the catheter
20 fell out. This occurred early in the morning of
21 November 20th. It was placed in a medical biohazards
22 bag, which is called a red bag by the medical
23 community, in a storage room two rooms down the hall
24 from room 7.

25 (Slide) Can I have slide 10?

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1 Later on the same day, a couple hours
2 later, the catheter containing the source was moved to
3 another storage location at the nursing home and
4 placed in a box with other red bags. This slide shows
5 the radiation field in the recreational room
6 associated with the source in the catheter in this
7 outside room. The patient died on November 21st,
8 1992.

9 From November 16th through November 25th,
10 numerous residents, employees and visitors to the
11 nursing home were unknowingly irradiated. The
12 ambulance staff who returned the patient to the
13 nursing home were irradiated, along with employees at
14 patients at the Indiana Regional Cancer Center who
15 were present for the approximately ten minutes the
16 patient was outside the treatment room after
17 treatment.

18 (Slide) Can I have slide 11?

19 On November 25th, 1992, a driver for the
20 Browning-Ferris Industries picked up the nursing home
21 red bag waste as part of the driver's normal rounds.
22 The driver had an operable portable radiation survey
23 meter, but contrary to company procedures did not
24 survey the nursing home waste. The nursing home
25 waste, along with other medical waste, was taken to a

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1 BFI facility in Carnegie, Pennsylvania where it was
2 loaded on a trailer. This trailer remained in
3 Carnegie throughout the following day, which was
4 Thanksgiving, and early on November 27th was driven to
5 a BFI medical waste incinerator in Warren, Ohio. At
6 the Warren facility, radiation monitors identified
7 radiation emanating from the trailer and facility
8 personnel directed its return to Carnegie the same
9 day.

10 It was left over the weekend in Carnegie
11 and on Monday, November 30th, the BFI staff searched
12 the truck for the radiation source. They identified
13 the box with the radiation source and looked at
14 individual red bags to identify the origin of the
15 waste. On December 1, BFI successfully identified a
16 name found with the red bag waste in the box and
17 traced it to the nursing home.

18 (Slide) Slide number 12, please.

19 After being notified by BFI, the nursing
20 home called the Indiana Regional Cancer Center on
21 December 1st. The Cancer Center had not used the HDR
22 afterloader for patient treatment after the single
23 treatment on November 16th. On November 17th, the
24 medical physicist did use the afterloader, but
25 primarily to recalculate the dose to the patient

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1 because the patient was not treated in the fifth
2 catheter and to turn around and determine what
3 treatment should be -- you know, what should be the
4 successive treatments in order to achieve the final
5 dose of 1800 rad, but did not -- even though the
6 machine was exercised, he never noticed that the
7 PrimAlert didn't light. Even though the source was
8 missing from the tip of the wire, the machine does not
9 tell you that you lost the source because it measures
10 the length of the wire going out, the length coming
11 back in and if you take in what you laid out, you get
12 no notification.

13 COMMISSIONER REMICK: Carl, the PrimAlert
14 alarms only visually, not audibly?

15 DOCTOR PAPERIELLO: It's only visually.
16 Physically the device is about the size of a good
17 sized thermostat and the light on it is about the size
18 of the average bimetal thermometer, about the size of
19 my thumb maybe.

20 The cancer center hadn't used the unit.
21 Upon being informed of the source discovery, the
22 medical physicist asked a technologist to extend the
23 source wire and see if the PrimAlert alarmed. It did
24 not. The medical physicist came to the Indiana
25 Regional Cancer Center and after failing to detect

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1 radiation by both a survey of the afterloader and
2 attempting to audio radiograph the source wire with x-
3 ray film, informed Region I that the source which at
4 that time -- you'll see the source changes. It's
5 decaying away. It's a 74 day half life. There was
6 about 3.7 curies was missing. The physician and the
7 medical physicist that afternoon drove to Carnegie to
8 retrieve the source.

9 (Slide) May I have the next slide, 13?

10 I'll just jump forward several days.

11 COMMISSIONER CURTISS: Carl, before you go
12 to that, I have three questions about your chronology
13 on the event. As I read the chronology, you describe
14 seven separate occasions where the RTTA attempted to
15 reinsert the dummy wire into catheter five.

16 DOCTOR PAPERIELLO: Right.

17 COMMISSIONER CURTISS: And from this
18 chronology it looks like the same reading came up on
19 the console. Were those seven identical steps each
20 time or was the technologist essentially doing the
21 same thing seven different times?

22 DOCTOR PAPERIELLO: To my knowledge they
23 were. They went in, they fiddled with the connections
24 and looked for kinks and didn't see it, and then went
25 back and tried to drive the dummy wire in

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1 unsuccessfully.

2 COMMISSIONER CURTISS: Okay. Do
3 Omnitron's procedures address that?

4 DOCTOR PAPERIELLO: I frankly don't know.

5 COMMISSIONER CURTISS: Okay. Second --

6 DOCTOR PAPERIELLO: In any case, the
7 people weren't very well -- whether they were
8 addressed or not, they weren't that well trained in
9 the procedures to begin with. I don't think they --
10 I think they're probably silent to that. I don't
11 recall --

12 COMMISSIONER CURTISS: Okay. In your
13 chronology, again you note that at 10:08, the first
14 point in time when the PrimAlert-10 was noticed to be
15 flashing. If the PrimAlert-10 was operating as it
16 should have and given the description of events here,
17 would it be correct to infer that it would have gone
18 off at 9:59 when the first error occurred?

19 DOCTOR PAPERIELLO: We think so. We think
20 it broke. It gets kind of complicated. It's not
21 necessary for the purpose of this investigation to
22 know exactly what catheter, but an issue did arise on
23 which catheter the source broke in because the
24 physician believes he removed the fifth catheter.
25 That was the one that was loose, which would imply

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1 that it broke off earlier. For a variety of reasons,
2 some of which are discussed in the report, our
3 engineers believe that the break occurred in the fifth
4 catheter. In fact, that's consistent with one of the
5 technicians who believes that the first time she went
6 into the treatment room to attend the patient's need,
7 the PrimAlert was not lit and it was the second time
8 that she went in. But it's not clear.

9 We're talking to people three weeks later
10 about an event that occurred and the problem they had
11 during the treatment is the wire didn't go in, not the
12 break. So, there's a lot of conflicting statements or
13 "I'm not sure" statements. So, people don't have --
14 you have to be careful about relying on the peoples'
15 memory, but I don't believe it's essential. The
16 essential issue is the alarm was lit. They checked
17 the computer. The computer says you're all right. The
18 alarm says you have a problem. They had a GM counter
19 that was operable. They should have checked and they
20 didn't.

21 COMMISSIONER CURTISS: Okay. Finally, on
22 page 2-5 you note that both the physician A and RTTA
23 examined the catheter connection at the patient and
24 observed no wire. What's the significance of not
25 observing a wire? Does that suggest that it's in the

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1 patient or in the device?

2 DOCTOR PAPERIELLO: Well, the tip broke
3 off and you wouldn't see it. They just said, "We
4 didn't see the wire."

5 COMMISSIONER CURTISS: Okay. You couldn't
6 infer anything from that?

7 DOCTOR PAPERIELLO: I could not infer
8 anything from it.

9 COMMISSIONER CURTISS: Okay. That's all
10 I have.

11 DOCTOR PAPERIELLO: The second wire broke
12 on December 7th. It occurred at the Pittsburgh Cancer
13 Treatment Center which happened to also be a facility
14 operated by Oncology Services. We looked into this
15 one. The wire broke in the same approximate location
16 as the first wire, at the base of the cavity. The
17 medical physicist in this case, who was personally
18 conducting the treatment, was aware of the first
19 event, immediately recognized the problem, promptly
20 and appropriately intervened and there was no
21 significant dose consequences to either the patient or
22 the cancer center staff.

23 In this case it appears that the patient
24 stressed the wire by an arm movement when the source
25 was being retracted. It was a different type of

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1 treatment. It was a lung treatment and the catheter
2 was taped to a cushion and the patient, as the source
3 was being retracted, was seen to raise their arm and
4 cause a bend where the catheter entered a connector on
5 the device and that's exactly where the source was
6 found when it broke. It looks like we know where the
7 stress came from.

8 (Slide) Can I have slide 14?

9 Going back to the Pittsburgh, the --

10 COMMISSIONER de PLANQUE: Before you do
11 that, is there any significance to the fact that the
12 error message that came up on the console in the
13 second event was different from the error message that
14 kept coming up?

15 DOCTOR PAPERIELLO: Yes. And the reason--
16 what we believe, everybody in Indiana denies that they
17 got any kind of alarm similar to what happened in
18 Pittsburgh. What we found in investigating how this
19 device works, if you go from the normal retract mode,
20 we call it AC retract, to a DC emergency retraction
21 mode, you pull all the sensors out of the path that
22 measure wire length. So, therefore, the alarm that
23 you would get because the wire is stuck out falls
24 back on one switch. There is a sensor that when the
25 wire comes back you close a path and it tells you

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1 you've pulled the wire in. The other way of doing it,
2 you measure the amount of wire going out, you measure
3 the wire coming back. Well, you lose that measurement
4 capability when you go into the DC retract mode. Even
5 Omnitron was unaware of that until we did our
6 investigation.

7 Many of the things that happen leave a
8 permanent record. The computer records it and will
9 tell you that it happened. It turns out when you go
10 into the DC retract mode, you get a brief message on
11 the screen of your computer console, but you get no
12 written record on the computer log. So, it's
13 impossible -- if you weren't looking at the screen at
14 the time, you wouldn't know it's happening. It only
15 takes about ten seconds to go into emergency retract
16 and the way the console is oriented, if the
17 technologist is observing the patient on the screen,
18 which they generally do, you wouldn't be looking at
19 the computer console and you wouldn't see it.

20 That's our hypothesis on what's happened.
21 We looked at the machine. We could find nothing wrong
22 with the Omnitron device. So, that's our theory on
23 what actually happened and why they got no error
24 message. We're going on oral statements. I have no
25 reason to believe that they weren't telling us the

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1 truth, but I'm just saying that's where we stand right
2 now.

3 (Slide) Slide 14.

4 I've completed the events. I'm going to
5 talk briefly about the radiological consequences.

6 COMMISSIONER de PLANQUE: Carl, can I ask
7 another question?

8 DOCTOR PAPERIELLO: Sure.

9 COMMISSIONER de PLANQUE: Following up on
10 Commissioner Remick's question about not taking into
11 account the shielding. It might be useful for you to
12 tell us whether as a consequence of that the dose
13 rates that you calculate are conservative or perhaps
14 higher than they would be if you took shielding into
15 account or lower.

16 DOCTOR PAPERIELLO: The dose rates are
17 probably conservative. However, when we calculated
18 the doses, we ignored a lot of small effects. For
19 example, if a nurse was working with the patient, that
20 nurse received a substantial dose by being in a room
21 with the patient. We didn't account for the dose that
22 the nurse would have received in other parts of the
23 building or while she was walking to the patient's
24 room. So, therefore, in some cases you're
25 conservative, in others you're less conservative.

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1 On the other hand, since most of the doses
2 was received when an individual was a meter or less
3 away from the patient, in the case of the certified
4 nursing assistants, these were the people who had --
5 this was an incontinent patient. She was cleaned
6 every two hours. They were in very close contact with
7 the patient. Other than tissue shielding in a body,
8 shielding would be negligible. We calculated this
9 using microshield. Couldn't believe the results. So,
10 a couple of us did it by hand. I ran microshield as
11 a code against the NBS curves in the NBS -- not NBS,
12 the NCRP 49 and got good agreement. Granted, it was
13 with iron. They don't have tissue in there. And I
14 also did the calculation using tissue attenuation --
15 energy attenuation rather than narrow beam and build-
16 up. If you use that, you get about a 20 percent
17 effect.

18 When you look at the large uncertainties,
19 how much distance it gives you, the shielding is
20 relatively small. I'm jumping the gun. If you look
21 at the cytogenetic results and compare them to the
22 calculated results, they're remarkably in agreement.
23 We did the calculations before we did cytogenetics.
24 So, it's not like we had any interaction between them.

25 I mentioned earlier what the dose rate is

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1 at two rem an hour roughly at a yard. We're going to
2 talk about the effects to the patients, to the workers
3 and to the public. Doctor Flynn, who is here,
4 concluded that an analysis of the medical records and
5 the physical dosimetry would indicate that massive
6 radiation dose was a probable contributing cause of
7 death in the patient. The consultant added that even
8 if the exact cause of death could be attributed to
9 other causes such as cancer or heart disease, the
10 extent of the radiation received would have soon
11 caused death.

12 The licensee reported that the prescribed
13 dose at one centimeter, which is the normal way of
14 prescribing brachytherapy, was to be 1800 rad. The
15 delivered dose was approximately 1.6 million rad to
16 the same point, or an overdose of almost 1,000 times.
17 The Indiana coroner is here. I've read his report and
18 the forensic pathologist concluded that acute
19 radiation exposure caused the death. But I am a
20 health physicist. I can do the calculations myself,
21 which I did. If you compare the doses to the internal
22 organs to any of the textbooks which talk about the
23 consequences, the books on radiation oncology, once
24 you go above 6 to 7,000 rad to an internal organ,
25 you're looking at either death or total destruction of

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1 the organ. They have tables of severe morbidity and
2 mortality. For a number of organs, this dose was
3 exceeded. So, whether or not from my viewpoint when
4 I was in the middle of this thing, the patient
5 actually died from this or would have died, I think
6 from our regulatory needs, I already knew that there
7 was a severe problem.

8 In addition to the patient, we evaluated
9 radiation doses to 94 people associated with the
10 event. Some individuals had film badges and we used
11 those results. Except for the physicians, all other
12 exposures were derived by calculations based on time
13 and distance. The physician did not wear his badge at
14 the times that he was exposed to the source. It was
15 on his lab jacket hanging in his office. We did time
16 and distance calculations for the physician and we
17 also added up the exposures on badges to people that
18 associated with him in those activities. The badges
19 come in somewhat higher, but in either case he was not
20 over exposed. His dose runs in the order of between
21 400 and 800 millirad.

22 COMMISSIONER REMICK: Carl, for those
23 people who had film badges, did you independently
24 estimate doses and did you pretty well agree with the
25 result from film badges?

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1 MS. JONES: Yes, we did.

2 DOCTOR PAPERIELLO: And nobody was over --
3 none of these individuals were over exposed. That
4 means they were within 1250 millirad.

5 Data were based -- so we did calculations
6 using time and distance studies. The data were based
7 on interviews and records. Individuals visiting the
8 patient were identified through interviews with
9 nursing home staff and known visitors. We became
10 aware of -- we knew there was an unknown visitor. We
11 didn't know the person's name, but the name appeared
12 in a newspaper article. We interviewed that person
13 and we subsequently placed an advertisement in the
14 local Indiana, Pennsylvania newspaper. The
15 advertisement requested any individual who was
16 concerned about having come in contact with the
17 radiation from the source and who had not been
18 interviewed to contact the NRC. A collect phone
19 number was provided. This led to two additional
20 contacts.

21 (Slide) Can I have slide 15?

22 This summarizes the doses that we
23 calculate or measured for 94 individuals that were
24 specifically evaluated. We believe that this bounds
25 the radiological consequences. For anybody to have a

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1 dose that was higher than reported here, they would
2 have to occupy places as close to or closer to the
3 source than individuals we evaluated, or have spent a
4 longer time period in the vicinity of the source.
5 This would have been observed by individuals
6 interviewed and reported. So, I don't think we have
7 anything above these.

8 We calculated extremity doses to those
9 individuals with the highest potential for extremity
10 exposure significantly above whole body dose. These
11 were the caregivers at the nursing home who attended
12 to the body needs of the patient and a couple BFI
13 employees who searched the most for the source of
14 radiation on the trailer. The highest extremity
15 exposure was estimated to range from 74 to 160 rems to
16 the hands of a certified nursing assistant. This
17 individual also received the highest whole body dose,
18 estimated to be between 16 and 22 rem.

19 (Slide) Can I have slide 16?

20 This slide presents a summary of
21 collective doses for all locations. The nursing home
22 clearly is the location where most of the public
23 exposure occurred.

24 (Slide) Can I have slide 17?

25 Further studies were conducted by blood

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1 counts and cytogenetic studies on selected
2 individuals. Blood counts are simple to perform, but
3 have a sensitivity limited to somewhere in the order
4 of 75 to 150 rem. Cytogenetic studies are more
5 sensitive, about 20 rem for a given single isolated
6 individual, but are much more difficult to perform.
7 Cytogenetic studies were performed for the NRC by the
8 Oak Ridge group at REACTS, the Radiation Emergency
9 Assistance Center and Training Site. Individuals
10 selected for cytogenetic studies were almost always
11 those with the greatest calculated dose. Since the
12 highest calculated doses were about the limit of
13 detection of 20 rem, we expected that the cytogenetic
14 data would show if there were significant non-
15 conservative errors. In other words, there were
16 measured doses that were higher than our calculated
17 values.

18 Again, go back. We are relying on
19 memories that were several weeks old and we had to
20 make simplifications to do the calculations and we did
21 not add trivial doses to the more significant doses.
22 Cytogenetic studies were to show whether or not we had
23 made some drastic errors. The cytogenetic results are
24 consistent with the calculated doses within the limit
25 of accuracy of both techniques. No occupational

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1 worker received a dose above the NRC limit of one and
2 a quarter rem per quarter. While members of the
3 public received radiation doses above applicable
4 regulatory limits, no one received the dose at which
5 acute radiation injury or clinical signs are expected
6 to occur.

7 There were no signs that contamination
8 resulted from the event. The bare source is iridium
9 metal. You would not expect it to leak. But we did
10 survey the nursing home, the ambulance, the clinic and
11 the insides of the machine itself and we found traces,
12 two nanocuries on the guide tubes which is less than
13 the leakage limit of five nanocuries and I'm not even
14 sure that wasn't picked up when the machine was
15 assembled.

16 (Slide) Can I have the next slide, 18?

17 Let's talk a little bit about the wire and
18 why it failed. The source is basically in a cavity at
19 the end of the wire. The wire is nickel-titanium
20 alloy. It's a shaped memory alloy. In other words,
21 you can -- and the reason it's used, it has very good
22 flexibility. You can bend it and it pops right back.
23 That's the reason why the wire was chosen. It
24 maximizes the usefulness of high dose rate
25 brachytherapy.

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1 The breaks in both wires in use occurred
2 at the base of the cavity, about 13 millimeters from
3 the tip of the wire.

4 (Slide) Slide 19.

5 Now, why did the wire break? We don't
6 know for sure. We are using a consultant, Southwest
7 Research. Omnitron, the manufacturer of the device,
8 and its consultant are investigating the failures.
9 I'm going to talk to what I knew up to a couple days
10 ago.

11 In May of 1992, the wires in use were
12 changed from stainless steel to a metal called
13 Nitinol, that is your nickel-titanium alloy, for new
14 wires. We found when we looked at the breaks under a
15 microscope, looked at x-rays of the cavities, that
16 some of the cavities were eccentric. They weren't
17 centered, they were tilted. We're not sure this
18 contributed to the break, but we did see that.

19 In September of 1992, Omnitron, this is a
20 couple months before these breaks occurred, Omnitron
21 identified degradation of the teflon liner in source
22 shipping containers and started to replace teflon with
23 stainless steel. At the end of December, Omnitron
24 looked at an old used wire that had been stored in a
25 teflon-lined shipping container and found it broken at

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1 the cavity. The current hypothesis being evaluated is
2 that storage of a source wire in teflon in the
3 presence of moisture causes decomposition of the
4 teflon and formation of either fluorine or
5 hydrofluoric acid. This, either directly or
6 indirectly, results in embrittlement of the wire.

7 We and our consultants are evaluating this
8 hypothesis and conducting various studies. There's a
9 whole bunch of studies. Some are listed in the report
10 that were done that support this hypothesis. It's
11 either one that you break the oxide coating on the
12 wire and let hydrogen get at the metal and that causes
13 embrittlement or even the fluorine getting into the
14 wire directly causes the embrittlement, but we don't
15 know for sure.

16 (Slide) Can I have slide 20?

17 Given that the wire broke, why wasn't it
18 detected? I think I've alluded to this earlier.

19 At the Pittsburgh Cancer Center, the
20 medical physicist received an audible alarm on the
21 retraction of the source wire as a result of the
22 break. At Indiana Regional Cancer Center, the staff
23 stated that no such alarm occurred. We discovered
24 that if the device enters this DC retract mode, the
25 system that measures the length of wire retracted to

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1 compare it to the wire played out is disabled.
2 There's only a brief error message on the screen and
3 nothing is reported in the computer log. The staff
4 was aware of the radiation monitor alarm, but the
5 computer console said the source wasn't retracted and
6 they didn't use the available survey meter. Why?

7 (Slide) Next slide.

8 Breakage of the wire was not considered a
9 credible accident by either the vendor or the
10 licensee. Omnitron's emergency procedures were
11 directed toward an emergency retraction of a wire with
12 a source that was stuck out and required manual
13 retraction. There's a crank on the side of the
14 machine.

15 There was a prior history of spurious
16 alarms on the area radiation monitor, the PrimAlert,
17 and the staff had developed a habit of ignoring it.
18 These originated even before the HDR afterloader was
19 procured. One technician developed the habit of
20 unplugging and replugging the monitor to reset it.
21 Now, the monitor does not work this way. While we
22 were there, we checked the monitor with a sealed
23 source. It will light when the source is near the
24 monitor and when you pull the source away the monitor
25 goes off. So, it's not one of these things where you

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1 have to reset it. When irradiation goes away, the
2 monitor should go off.

3 Oncology Services as a corporation appears
4 to have provided no formal radiation safety training
5 to the staff at the Indiana Regional Cancer Center.
6 Reliance was placed on the staff's previous academic
7 training, training by Omnitron which included
8 Omnitron's emergency procedures but not radiation
9 safety, and an expectation that the medical director
10 or medical physicist at each site would provide
11 radiation safety training. This expectation was
12 neither met nor were steps taken to confirm it.

13 Given the conflict between the data
14 provided by the area radiation monitor and the
15 Omnitron control panel and the lack of radiation
16 safety training that would have conditioned the staff
17 to respond promptly to the radiation alarms, the
18 licensee staff failed to respond to the alarm.
19 Instead, a technologist unplugged and reset the alarm.
20 He doesn't recall when he did this. He didn't know
21 whether or not the alarm was still on when he
22 unplugged it. The technologist should have responded
23 by using the portable survey meter available at the
24 control console. The meter would have identified the
25 fact that the source had not returned to the

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1 afterloader shield.

2 (Slide) Can I have the next slide?

3 We've bunched our observations under six
4 major headings.

5 (Slide) Can I have the next slide?

6 I've already discussed the radiological
7 aspects and discussed some of Oncology Service
8 Corporation radiation safety shortcomings.

9 (Slide) Slide number 24.

10 A number of weaknesses at the Indiana
11 Regional Cancer Center shown on this slide were also
12 identified at other sites operated by Oncology
13 Services Corporation and inspected by Region I
14 subsequent to the initiation of this IIT and on this
15 team's recommendation.

16 (Slide) Slide 25.

17 A number of weaknesses were found in the
18 design and testing of the Omnitron 2000. Prior to the
19 incident, it appears that no engineering calculations
20 for tensile, shear or fatigue strengths were performed
21 on the Nitrinol source wire, particularly for the
22 source cavity. No tests were performed to ensure the
23 environmental effects of radiation, moisture, teflon
24 and combinations thereof would not affect the
25 integrity of the source wire.

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1 I've already discussed weaknesses in the
2 design of the emergency retraction system.

3 We and the FDA also found weaknesses in
4 Omnitron's QA/QC program for the production of the HDR
5 afterloader and source wire, but at this time these do
6 not appear to have contributed to the event.

7 (Slide) Next slide.

8 NRC regulations do not directly address
9 high dose rate brachytherapy to the extent that low
10 dose rate brachytherapy and teletherapy are addressed.
11 Licensing guidance for HDR has remained unchanged
12 since 1986 in spite of significant changes in Part 35
13 and associated medical licensing guidance. The
14 inspection guidance does not specifically address HDR.

15 (Slide) Next slide.

16 Although inspected by Region I within a
17 year of initial licensing, this slide will show you
18 how the license changed. The licensing and inspection
19 program had no provision to cause an early
20 reinspection subsequent to Amendment 2. When that
21 facility was inspected in September of 1991, they had
22 one high dose rate machine used in Harrisburg and
23 under the direct control and observation of the
24 corporate radiation safety officer. That inspection
25 found the staff there to be very knowledgeable. There

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1 was no problem with training at that particular site.

2 After that inspection, the program
3 literally exploded. They were authorized to have
4 dozens of machines at up to six sites under that
5 license and they went into other sites under agreement
6 state licenses and at least one other NRC license and
7 there was nothing in our inspection or licensing
8 program that would flag this. At this point, the
9 informal procedures that they used to control their
10 program became ineffective. So, you had a mixed bag
11 when you got to -- at the Greater Pittsburgh site, we
12 had a very conscientious medical physicist who was
13 there full-time. At other sites, many of the medical
14 physicists were only part-time employees and that's
15 where we found weaknesses in training.

16 When I talk about weaknesses in training,
17 it's not a question of not just attending a session.
18 They didn't know, the technologists didn't know how to
19 operate the Geiger counter. The technologist who
20 performed the treatment, when asked what the most
21 sensitive range of the counter was, says, "I assume
22 it's the times 1,000 range." So, it's not a tool that
23 these people use. They were unaware of what the field
24 was in the vicinity of the source. They had no idea
25 how long -- let's suppose they had to retract the

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1 source in an emergency using Omnitron's procedures.
2 They didn't know how long they could remain in that
3 shielded room. To me, that's a fundamental breakdown
4 in the training that is required by Part 19.

5 COMMISSIONER REMICK: Carl, are you saying
6 that for the amendments where new devices were
7 approved at different sites, there was no pre-
8 amendment inspection by the NRC following the initial
9 inspection?

10 DOCTOR PAPERIELLO: No.

11 COMMISSIONER REMICK: You're saying --

12 DOCTOR PAPERIELLO: There was no other
13 inspection other than the one on September '91, yes,
14 and it was two months later that the amendment request
15 came in and expanded the program.

16 (Slide) Slide 28.

17 We originally called this safety culture,
18 but we were informed that wasn't politically correct,
19 so we called it critical safety awareness. But
20 really, when you look at what was done, it was
21 complete contrast on what was done in Pittsburgh. In
22 Pittsburgh, they did survey the patient after
23 treatment with a hand held meter and the medical
24 physicist said he always did.

25 Now, it may be after the fact, but he was

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1 so well prepared when we showed up the following day
2 to do the inspection that I believe him. He seemed to
3 be a very conscientious individual. He would never
4 allow a technologist to do the treatment. The medical
5 physicist did the treatment personally and the
6 physician, at least in that treatment, was with him at
7 the console.

8 He was a full-time medical physicist and
9 not part-time. He told us he did believe the wire
10 could break. Now, granted he had the after -- he knew
11 about this afterwards, but he claimed that in August
12 when he was -- he went down to Omnitron for training,
13 that he was a gadgeteer and he stayed around in the
14 evening and had the engineers show him the inside of
15 the machine and he pressed them about the wire not
16 breaking and they admitted, "Well, it's not
17 unbreakable. It could happen."

18 There is no indication of any prior wire
19 breakage in use. They did test two prototypes or
20 early production to failure, but there's no indication
21 there was any prior knowledge by Omnitron of any prior
22 breaks. We could find no evidence to that effect.

23 But anyway, the engineers told him that,
24 "Yes, we guess they could break under certain
25 circumstances." So, there are differences between

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1 the two places.

2 (Slide) Can I have the next slide?

3 The discovery of the lost source by BFI
4 may have prevented additional significant radiation
5 exposure. However, actions by BFI employees
6 subsequent to the discovery of the radiation source
7 resulted in their receiving unnecessary exposures.
8 Assistance from radiation protection experts should
9 have been sought.

10 Now, in the scrap steel industry there are
11 a lot of guidance provided to scrap dealers who have
12 monitors on what to do when you find something. We
13 help them in this area. It appears to me that we need
14 to do something with other people in the scrap either
15 waste brokers, collectors or recyclers to ensure
16 people know how to protect themselves. So, imagine,
17 they put a four curie iridium source back on the road
18 again. We had significant exposures to people who
19 went searching for the -- they had survey instruments
20 that went off scale and they had no idea what that
21 meant. In fact, they even discovered which truck this
22 source was on in Warren. They had to use a concrete
23 building as a shield because they had micro R meters
24 that -- this source could be detected at hundreds of
25 feet with a micro R meter.

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1 Lastly, this thing is complicated. I call
2 it licensing jurisdiction for want of a better word.
3 I believe the FDA is going to talk a little bit about
4 what they do. But this is a medical device and it
5 falls under FDA regulations because it is a device.
6 The source wire and the device was licensed by the
7 State of Louisiana, that's an agreement state, and of
8 course we license the users. All I can say is it's a
9 very complicated area to get into when you try to
10 understand is there something that some regulatory
11 agency should have done to identify potential
12 weaknesses in the design and testing of the source
13 wire? That's something that the team could not
14 resolve.

15 Thank you.

16 MR. TAYLOR: Mr. Chairman, Commissioners,
17 that concludes the team's presentation. The FDA was
18 kept closely advised of the work of the team and we're
19 pleased to have William Damaska here from FDA, whom I
20 introduced earlier, and he has a few remarks to add
21 before we conclude our presentations.

22 Mr. Damaska?

23 MR. DAMASKA: Thank you very much.

24 I would just like to make a brief
25 statement describing our responsibility and authority

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1 and the status of our investigation.

2 The principal authority under which FDA
3 regulates devices such as the Omnitron 2000 high dose
4 rate remote afterloader is the Federal Food, Drug and
5 Cosmetic Act. Section 201(h) of the Act defines
6 devices as an instrument, apparatus, implement,
7 machine or implant or similar or related article
8 recognized in an official compendia, intended for use
9 in the diagnosis of disease or other conditions or in
10 the cure, mitigation, treatment or prevention of
11 disease, or intending to affect the structure or any
12 function of the body.

13 FDA has established numerous programs to
14 ensure the safety and effectiveness of devices. These
15 can be described as, number one, review of clinical
16 studies such as premarket notifications and premarket
17 approval submissions. Number two, review of voluntary
18 and mandatory problem reports. And three, enforcement
19 activities such as routine and directed inspections
20 and other enforcement activities such as product
21 removals, recalls, warning letters, seizures,
22 injunctions, prosecutions and civil penalties.

23 The extent of regulatory control over a
24 device depends on its class. Class 1 devices -- there
25 are three different regulatory classes. Class 1

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1 devices are those which can be adequately regulated by
2 what we refer to as the general control provisions of
3 the Act, such as registration, listing, premarket
4 notification and routine inspections.

5 Class 2 devices are those which must
6 conform to the general controls but also need special
7 controls in the form of standards to perform
8 effectively.

9 And class 3 devices are those which must
10 meet all the general controls and must also be
11 approved by way of a premarket approval application
12 before they can be marketed. The route to bringing a
13 device to market may include an investigational device
14 exemption. That's a procedure whereby we approved
15 based on evaluation of a submitted protocol the
16 investigational study.

17 The NRC incident investigation team
18 requested and received from FDA a number of documents
19 relating to the investigation of the Omnitron 2000 HDR
20 incident. In April 1988, Omnitron International
21 submitted a premarket notification to the FDA.
22 Omnitron was asked to provide additional information
23 in relation to specific aspects of that submission,
24 device design specifications, device safety features,
25 laboratory and clinical testing and quality control

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1 procedures.

2 On the basis of the information provided
3 by Omnitron, FDA concluded that the device was
4 substantially equivalent to similar class 2 devices
5 that were currently on the market and we allowed
6 Omnitron to market the device. We have cooperated
7 with the NRC by conducting joint inspections at both
8 the incident site and the two manufacturing plants.
9 Our Dallas district office is coordinating the ongoing
10 investigation and is working with the Texas State
11 Department of Health and the NRC.

12 The New Orleans District Office, in close
13 cooperation with the Louisiana Department of Health,
14 has completed an inspection of the wire production
15 facility in Louisiana. Under a voluntary agreement
16 between FDA and Omnitron, no shipments of the Omnitron
17 2000 afterloader system or the source wire are
18 currently being made. The company on December 17th,
19 1992 issued a safety alert to all customers, informing
20 them of the two incidents where the source separated
21 from the wire during brachytherapy. This alert also
22 provided guidance to physicians recommending strict
23 adherence to the NRC requirements for post-treatment
24 patient monitoring and use of the equipment only when
25 it is deemed necessary in light of other available

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1 modes of therapy and the patient's condition until the
2 cause of the fractures are determined and corrected.

3 Determining an appropriate regulatory
4 resolution will necessitate a thorough assessment of
5 any violations of the Act and the potential impact on
6 public health. However, because our investigation has
7 not been completed, I'm somewhat limited in the extent
8 to which I can discuss the aspects of our
9 investigation. But I'll be certainly glad to try and
10 answer any questions you may have.

11 CHAIRMAN SELIN: Well, I'd like to ask you
12 one question, not so much about the investigation and
13 the device, but Doctor Paperiello pointed out a number
14 of possible areas of either weakness or gaps or what
15 have you in the regulatory process that the NRC
16 follows and there was a generic comment about possible
17 jurisdictional questions. Are you prepared to
18 speculate about some of the questions that this has
19 shown in the interface problems between your
20 responsibilities for inspecting the devices and ours
21 for talking about the methods for using the devices?

22 MR. DAMASKA: I really -- I'd rather not
23 speculate in terms of any lapses in the regulatory
24 authority, but perhaps I could expand a little bit on
25 that in terms of just what our authority involves with

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1 respect to the manufacturing process and some of the
2 things that Doctor Paperiello mentioned with respect
3 to the wire breakage and that sort of thing.

4 One of our significant regulations, of
5 course, is what we call good manufacturing practice
6 requirements. The GMPs, as they're known, require
7 that manufacturers not only develop adequate
8 procedures for production, but that they be validated
9 and that they establish adequate quality control and
10 testing procedures to assure that their devices are,
11 in fact, safe and effective in normal routine use.
12 That's part of our current investigation in terms of
13 the adequacy of those procedures.

14 CHAIRMAN SELIN: Let me be a little bit
15 more specific. A couple of the things that turned up
16 in the IIT were that the device had modes of operation
17 that were not known to the people who bought the
18 device, presumably were not in the documentation of
19 the device. These modes don't necessarily make the
20 device unsafe or unlicenseable, but they're
21 information that has to be available to the users if
22 they're used.

23 From your description of the FDA process,
24 it doesn't sound as if somebody in a class 2 device is
25 going to exhaustively take the description, the

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1 specifications that are given to the users and check
2 that against the device. In fact, you described a
3 process whereby analogy, since another device looked
4 okay, this was essentially identical, which I would
5 say would be far short of line by line check.

6 MR. DAMASKA: Well, part of the general
7 control provisions that I mentioned include the
8 provision to provide complete and adequate directions
9 for use for any device. In other words, if the
10 instructions are not adequate, then we would consider
11 the device to be misbranded under the Act, and that
12 could be a violation, could be a serious violation if,
13 in fact, adequate directions for use were not provided
14 for all the various ramifications and capabilities of
15 the product.

16 So, it is covered, but to also answer the
17 other part of your question was, as a class 2 device,
18 the review process there, it may not include a
19 thorough enough review of the labeling to assess any
20 potential shortcomings. That's always a possibility
21 for that type of a problem.

22 CHAIRMAN SELIN: So, it's a performance
23 kind of thing. After the fact, if it was found that--

24 MR. DAMASKA: That's right.

25 CHAIRMAN SELIN: -- that information

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1 hadn't been given to the operator, then by definition
2 the manufacturer didn't do something right.

3 MR. DAMASKA: That's correct. That's
4 right.

5 COMMISSIONER ROGERS: I have a question
6 along those lines. If this wire breakage occurred but
7 the source somehow or other wasn't left in the patient
8 and this event as an over exposure didn't take place,
9 but nevertheless the system failed mechanically, would
10 there be any way in your present procedures that you
11 would notify the NRC of this as a potential failure
12 mechanism that might be something that we should be
13 alert to and see that licensees in using the device
14 are alert to it?

15 MR. DAMASKA: Yes. I think under our
16 current procedures we have a system that we call the
17 device experience network which involves various types
18 of mandatory problem reporting systems. The way we
19 assess any report we receive, I think we've
20 established a pretty good working relationship and
21 yes, we would have notified the NRC.

22 COMMISSIONER CURTISS: I have just a
23 couple of questions. When the change was made in May
24 of 1992 from stainless steel to whatever that was that
25 the new metal consisted of, is that typically

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1 something that would be brought back before FDA for
2 your consideration and approval?

3 MR. DAMASKA: I would -- that's one of the
4 things that we're looking at right now. I would
5 rather not speculate in terms of the significance of
6 that change because to a great extent it has to do
7 with the extent of validation that was done by the
8 manufacturer. So, I can't give you a complete answer
9 on that.

10 COMMISSIONER CURTISS: Let me ask the
11 question in a generic way then. When a change is made
12 in a device that you have approved, is there a
13 threshold that's been established for when the vendor
14 of that device has to come back and obtain your
15 concurrence or comment?

16 MR. DAMASKA: For class 2 devices -- let
17 me explain. For class 3 devices, those that are
18 subject to premarket approval, the threshold there in
19 terms of changes, almost any change that's made in an
20 approved device would have to be resubmitted to the
21 agency. For class 2 devices, when a manufacturer
22 makes what would be considered a significant change,
23 they must report that to the agency in a resubmission.
24 That's obviously a very broad definition and depends,
25 as I said, to a great extent on the extent of

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1 evaluation, the extent of engineering work that the
2 manufacturer has done.

3 COMMISSIONER CURTISS: Okay. Two other
4 questions. The action that you referred to that has
5 been taken to suspend the shipments of the device, is
6 that the action that was taken by the Texas Department
7 of Health or is that an FDA action?

8 MR. DAMASKA: You're correct, the Texas
9 Department of Health did place an embargo on the
10 Omnitron device, but Omnitron also agreed in a
11 voluntary agreement with FDA to suspend all shipments
12 of the device and the wires.

13 COMMISSIONER CURTISS: All right.

14 MR. DAMASKA: It's really a separate --

15 COMMISSIONER CURTISS: Okay. And the
16 suspension of the device, I guess from what I read in
17 the report, will remain in place until the findings of
18 your audits have been addressed? Do I understand that
19 correctly?

20 MR. DAMASKA: Yes, that's correct.

21 COMMISSIONER CURTISS: And is that a
22 determination that the Texas Department of Health
23 makes or that you at the FDA make?

24 MR. DAMASKA: That would be a
25 determination that FDA would make.

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1 COMMISSIONER CURTISS: Okay. One other
2 question. The inspections or audits, I guess you call
3 them audits, that were conducted out of your
4 Lafayette, Louisiana and Dallas, Texas offices of the
5 respective facilities in those two states, were those
6 audits scheduled and on the books before this event
7 occurred or were they a result of this event that they
8 were scheduled?

9 MR. DAMASKA: These were the result of
10 this event.

11 COMMISSIONER CURTISS: Okay. That's all
12 I have.

13 MR. TAYLOR: Mr. Chairman, if you don't
14 mind, I'd like to conclude the presentation. So, we
15 have the Indiana County Coroner Thomas Streams with us
16 and he has some remarks.

17 CHAIRMAN SELIN: Doctor Streams?

18 DOCTOR STREAMS: Thank you, ladies and
19 gentlemen.

20 To begin with, I wish to thank all those
21 individuals and agencies involved for their
22 outstanding cooperation and investigation into the
23 death of Mrs. Colgan. Your professionalism and
24 personal courtesy have made an otherwise regrettable
25 set of circumstances much more endurable and the

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1 Indiana County Coroner's Office is very grateful for
2 both your participation and for the loan of your
3 expertise.

4 One of the most important considerations
5 in investigating a case of this nature is the need for
6 all agencies involved to work together. I would
7 certainly like to stress the value of involving
8 appropriate local authorities early in the
9 investigative process. Local and state agencies such
10 as the coroner's office and police departments can be
11 important regional resources to agencies such as the
12 NRC and can offer a great deal of support in ensuring
13 that the investigation is not unnecessarily hindered
14 by local or state mandates and regulations.

15 Most, if not all states, including
16 Pennsylvania, specifically require that the county
17 coroner conduct an investigation into the cause and
18 manner of death in cases where radiation may be
19 reasonably suspected as a direct or related cause of
20 death.

21 There are a number of aspects of this
22 investigation which could not have been made
23 effectively pursued without the involvement of state
24 and local authorities working on conjunction with the
25 NRC. The cooperation of the county coroner's office

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1 and other authorities is essential for the exhumation
2 of an interred body for autopsy. Had the coroner's
3 office and Pennsylvania State Police not become
4 involved, an autopsy would not have been performed,
5 the exact cause of death would still be in doubt, and
6 a wealth of important scientific and statistical data
7 would not have been secured for future reference.

8 There exists very few documented cases of
9 death resulting from this type of radiation exposure
10 in the statistical record. This may be due in large
11 part to the fact that autopsies are not routinely
12 performed on the suspected victims, since local
13 authorities like the coroner's office may have been
14 inadvertently overlooked as a resource for
15 documentation of radiational injury through autopsy.
16 By involving the coroner early in the investigative
17 process, autopsy findings could be readily available
18 to NRC officials, thus eliminating the need for
19 speculation that radiation injury may have been a
20 contributing cause of death.

21 While the official cause of death in the
22 Colgan case has been listed as acute radiation
23 exposure and consequences thereof, there has been no
24 determination to date as to the actual manner of
25 death. The Indiana County District Attorney's office

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1 has referred the case to the Pennsylvania Attorney
2 General for consideration. In the near future, it is
3 reasonable to expect that either a preliminary hearing
4 before a state grand jury or a coroner's inquest will
5 be forthcoming to determine the exact manner of death
6 and to make recommendations regarding the filing of
7 criminal charges in this case.

8 From the onset, the Indiana County
9 Coroner's Office has worked very diligently to ensure
10 that the forensic investigation into Mrs. Colgan's
11 death was performed in conjunction with the most
12 highly qualified experts available. The autopsy
13 itself was performed by Doctor Isidore Mihalakis,
14 forensic pathologist for Lehigh Valley Hospital in
15 Allentown, Pennsylvania, a nationally recognized and
16 respected forensic pathologist. Doctor Mihalakis
17 performed an exhaustive examination of the body before
18 ultimately determining radiation to have been the
19 primary cause of death in this case. Other
20 pathologists may challenge his findings, but we are
21 satisfied that there was sufficient documentation to
22 support his determination that radiation exposure
23 alone was responsible for Mrs. Colgan's death.

24 At the direction of the coroner's office,
25 Eric J. Lee, M.Sc., a medical radiation physicist,

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1 Chief Physicist at the Division of Radiation Oncology
2 at Allegheny General Hospital and Director of
3 Allegheny Singer Research Institute Accredited
4 Dosimetry Calibration Laboratory in Pittsburgh,
5 prepared a dosimetry table indicating the distribution
6 and radiation levels that would have been delivered
7 from the radiation source that was left in Mrs.
8 Colgan's body.

9 Doctor E. Day Werts, a radiation
10 biologist, Assistant Professor of Radiologic Services
11 at the Medical College of Pennsylvania, Director of
12 Radiation Oncology Research at Allegheny General
13 Hospital, and Adjunct Assistant Professor at the
14 School of Public Health at the University of
15 Pittsburgh, subsequently interpreted the dosimetry
16 report compiled by Mr. Lee. Doctor Werts states, in
17 part:

18 "Based on the dosimetry findings, a
19 distance of 25 centimeters from the radioactive source
20 would have included most of the small and large
21 intestine of Mrs. Colgan, which would have received at
22 least 2800 cGy, a dose sufficient to result in death
23 from gastrointestinal failure.

24 "Similar radiation doses received by
25 healthy normal individuals to the bowel or bone marrow

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1 would result in their death within a few days from GI
2 toxicity or four to five weeks from unattended bone
3 marrow failure. While her cancer could have
4 contributed to general debilitation, in my opinion the
5 radiation exposure this patient received would have
6 severely affected her survivability. The doses of
7 radiation received by the gastrointestinal tract alone
8 are sufficient to cause death and histologic findings
9 of the GI epithelium consistent with the description
10 given above would suggest that this tissue was subject
11 to radiation exposure similar to those estimated.
12 Furthermore, one cannot rule out either contributory
13 factors, including the potentially lethal doses
14 received by major portions of the bone marrow and the
15 possibility of multi-organ failure."

16 It is important to note that Doctor Werts
17 had no communication with Doctor Mihalakis throughout
18 the investigation of this case and had no knowledge of
19 the official autopsy findings at the time his report
20 was filed. His conclusions were, in fact, separate
21 and independent judgments based solely on the
22 radiation levels depicted in Mr. Lee's dosimetry
23 table.

24 In conclusion, this is neither the first
25 nor the last such case of this nature to occur in the

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1 United States. As unfortunate as the events that led
2 to Mrs. Colgan's death may be, it is important that
3 all agencies involved learn from this experience that
4 cooperation among the diverse agencies and
5 jurisdictions may be of utmost priority if we are to
6 prevent future mishaps of this kind. The information
7 gathered by this investigation, made possible by the
8 synergistic partnership of all agencies involved will
9 undoubtedly lend itself to the improvement of
10 safeguards within the radiation oncology industry and
11 will help to ensure that patients can again be
12 confident of quality medical therapy through the
13 controlled use of radioactive substances.

14 Finally, I believe that although an
15 investigation of this type must be conducted with a
16 certain degree of detachment and seemingly sterile
17 mechanization, it is very important that we all
18 remember that Mildred Colgan was more than just a case
19 that we jointly investigated. We will all do well to
20 bear in mind that while much controversy and many
21 questions may surround her death, it is her life that
22 matters most to those whom she loved and who held her
23 in the highest esteem. We appreciate the cooperation
24 and forthrightness of Mrs. Colgan's family throughout
25 these difficult and trying circumstances and generally

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1 hope that the recollection of Mildred Colgan will not
2 be encumbered by the necessary discharge of our joint
3 responsibilities in this case.

4 Thank you very much.

5 CHAIRMAN SELIN: Thank you very much,
6 Doctor Streams. I would like to make a comment and
7 then ask you a question.

8 The focus of this meeting today is
9 specifically the regulatory process, the coordination,
10 et cetera. Obviously, the same fact-finding report,
11 the same IIT report will also support enforcement of
12 any civil or criminal actions that might come up. But
13 those implications are not the subject of discussion
14 today. There's a separate enforcement process and any
15 other process would also be separate.

16 So, the question I'd like to ask you is a
17 generic question. It doesn't have to do with the
18 conclusions, the medical conclusions in this case.
19 You said very nice things about the cooperation with
20 the different agencies, but I still would like to ask
21 you if your experience would lead you to suggest
22 anything be done differently in the coordination
23 between the NRC, the coroner's office, say our medical
24 consultant or any other procedural conclusions you
25 would care to draw from this experience?

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1 DOCTOR STREAMS: Yes, sir. I would like
2 to suggest to the Commission that immediately upon
3 notification of an event of this nature occurring,
4 that the appropriate local authorities would be
5 notified immediately of such instance, including the
6 coroner/medical examiner's office and the local police
7 jurisdictions. In this case, my office first became
8 aware about it when we read it in the newspaper that
9 the NRC had an investigative team in Indiana
10 investigating this death. It delayed the process of
11 obtaining a court order to exhume the body for
12 autopsy, et cetera.

13 CHAIRMAN SELIN: All right. Thank you.
14 Anything else that you'd care to add to that?

15 DOCTOR STREAMS: No, sir.

16 CHAIRMAN SELIN: Thank you very much for
17 your appearance.

18 Commissioner de Planque?

19 COMMISSIONER de PLANQUE: Yes, I just have
20 a couple of general questions.

21 Doctor Paperiello, could you elaborate a
22 little bit on the problem of whether or not a portable
23 survey meter is required for use in these
24 circumstances? What are the discrepancies --

25 DOCTOR PAPERIELLO: Yes. There's an issue

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1 with high dose rate brachytherapy on whether or not
2 the area radiation monitors which the licensing
3 guidance requires is a substitute for a survey by a
4 portable radiation survey instrument. It is the
5 opinion of OGC that it is -- and both NMSS that you
6 have to do a separate survey. However, in the course
7 of investigating this event, it's clear that our
8 regional staff, not only in Region I but in Region
9 III, my own region, believed that the area monitor met
10 the requirement. I would point out that even if the
11 area monitor met the requirement, in this case the
12 area monitor did what it was supposed to do, it
13 alarmed. Nobody believed it. On the other hand, if
14 there was a separate survey done, it certainly would
15 have caught this thing.

16 Part of the problem is that the licensing
17 guidance was issued in February of 1986 and the
18 current Part 35 with the survey requirements was
19 issued in say, fall, either August or October of '86.
20 The licensing guidance was not revised to reflect what
21 was in the new Part 35, so creating an impression in
22 the minds of the staff that was confused.

23 I'll be honest. When I started this
24 thing, I had my staff tell me that this device wasn't
25 even licensed under Part 35. I'm talking about my

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1 regional staff. I put them to rights as a result of
2 this investigation, but there certainly was confusion
3 in the minds of the staff.

4 COMMISSIONER de PLANQUE: Would this
5 extend to agreement states?

6 DOCTOR PAPERIELLO: It could. I don't
7 know.

8 COMMISSIONER de PLANQUE: Okay. Another
9 question. In the chronology there is reference made
10 to the therapy technician and others as being
11 registered. What does that registration mean?

12 DOCTOR PAPERIELLO: They're registered by
13 the American -- it's not by the state, it by a
14 national -- I assume it's a voluntary accredited --
15 no, let me check. I have a backup. I did check that
16 up because I read each of their certificates. I could
17 get that to you.

18 COMMISSIONER de PLANQUE: That's okay.

19 DOCTOR PAPERIELLO: I thought I had it
20 here and I did at one time look it up.

21 MR. TAYLOR: We'll provide that.

22 COMMISSIONER de PLANQUE: That's fine, but
23 that brings me to a more general question. Do you
24 have any observation about any credential programs
25 with respect to the training aspects that you

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1 discussed earlier?

2 DOCTOR PAPERIELLO: The licensee depended
3 strongly on credentialing. They told us that in the
4 interviews we had with the RSO. Our licensing manual,
5 Regulatory Guide 10.8, does not allow licensees to
6 take credit for it. I'll have to tell you, in the
7 transcripts of these technologist's training or the
8 examinations they took to become registered
9 technologists, every one of them had training in
10 radiation protection. Obviously whatever they had was
11 inadequate to condition them to do the right thing.

12 But this is not unlike nuclear power
13 plants. It is not enough to have a nuclear power
14 plant operator have good academic training. You have
15 to have good emergency procedures and you need
16 drilling and you need conditioning. There was
17 probably only a period of 30 seconds for these people
18 to make the right decision and they made the wrong
19 one.

20 COMMISSIONER de PLANQUE: Okay. Thank
21 you.

22 CHAIRMAN SELIN: Commissioner Remick?

23 COMMISSIONER REMICK: I'd like to follow
24 up on that question of Commissioner de Planque having
25 to do with our own staff confusion over whether a

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1 survey instrument is needed or not. I looked up our
2 regulations and to me it seems very clear. 35.404(a)
3 says, "Immediately after removing the last temporary
4 implant source from a patient, the licensee shall make
5 a radiation survey of the patient with a radiation
6 detection survey instrument," which to me is not an
7 area monitor, "to confirm that all sources have been
8 removed."

9 Then going on to (b), it says, "The
10 licensee shall retain a record of patient surveys for
11 three years. Each record must include the date of the
12 survey, the name of the patient, the dose rate from
13 the patient expressed as millirem per hour and
14 measured at one meter from the patient," which you
15 couldn't get from an area radiation monitor, "the
16 survey instrument used and the initials of the
17 individual who made the survey."

18 So, I was surprised to find in the report
19 that our own inspectors are confused by that wording.
20 It seems to me that it's emphasized the need for
21 training of our own people on the difference of an
22 area monitor and a radiation survey instrument.

23 DOCTOR PAPERIELLO: I know what you're
24 saying. I know that. I'm just -- but I'll tell you
25 the reason for the -- if you read Subpart G, first you

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1 will not find under the list of devices there, you
2 won't find a high dose rate afterloader.

3 Most of the instructions and requirements
4 in there clearly don't apply to afterloaders because
5 you talk about instructing nurses and visitors to the
6 room. You have none of that. There are critical
7 aspects of this device which aren't addressed at all.
8 Unlike conventional brachytherapy, in high dose rate
9 brachytherapy the licensee user has to measure the
10 source output, much like you do in teletherapy. So
11 really, HDR requirements and what you should do for
12 safety fall midway between teletherapy and low dose
13 rate brachytherapy. Those regulations clearly map one
14 on one everything you need to do for low dose rate
15 brachytherapy and most of them are inapplicable and
16 that's how I believe the staff had the opinion.

17 Besides, the licensing guidance is very
18 explicit about having to monitor and most licensing
19 guides will repeat the regulations that are applicable
20 and they're silent to what's in there. And of course
21 the reason is that these things were written after the
22 licensing guidance was written and, you know, the
23 inspector in the field is the lowest person on the
24 totem pole. Whoever put out the licensing guidance,
25 in my view, should have revised it when Part 35 was

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1 revised to reflect the new requirement, the current
2 requirements.

3 COMMISSIONER REMICK: I don't differ with
4 you on that. I have some other questions related to
5 this.

6 When our inspectors go out, do they audit
7 the records to see if the survey type of records are
8 being kept? Do we know if our inspectors --

9 DOCTOR PAPERIELLO: The answer is they do,
10 but in this particular case the inspector had written
11 on top of the field notes where it talks about
12 brachytherapy, it said, "HDR only," and therefore that
13 inspector did not believe that the requirements in
14 there were applicable and instead looked at the
15 license conditions which the inspector -- we're
16 talking about an experienced inspector. It wasn't
17 somebody that was relatively new, so I'm just telling
18 you I know what the staff was thinking in this
19 particular case.

20 COMMISSIONER REMICK: But in general you
21 feel that inspectors are looking to see if surveys are
22 being conducted?

23 DOCTOR PAPERIELLO: If it's low dose rate
24 brachytherapy, they're being looked at. I don't know.
25 I mean, I wasn't able to -- I didn't have time to look

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1 at other -- you can't -- unless you read the license
2 directly, there's no way I'm aware of that you can go
3 through our files quickly and identify those people
4 who are doing HDR from those who are doing just
5 conventional brachytherapy.

6 COMMISSIONER REMICK: I see.

7 Will the IIT report be provided to
8 agreement states, since there have been some
9 allegations that we don't share information with one
10 another?

11 MR. TAYLOR: It is being publicly released
12 today. It should go. We will ensure it does to all
13 agreement states. I would add at this point, of
14 course, one of the purposes of an IIT is to identify
15 our own internal operational problems and regulatory
16 issues and of course I'll be following up. At this
17 stage the information is new. We'll be providing
18 further reports to the Commission on that subject
19 before we -- as the weeks proceed ahead.

20 COMMISSIONER REMICK: Carl, going back to
21 the question of the isodose curves and so forth, in no
22 way do I differ with what you have done. I realize
23 the biggest uncertainty is how long did people spend
24 near the source and at what distance and those are
25 overwhelming uncertainties. However, the point I

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1 would make for the future, because it's not clear to
2 me whether your figures would be conservative or not,
3 the fact that the film badges and the cytogenetic
4 examinations confirmed it satisfies me, but for the
5 future I would think that Floorshine and Skyshine in
6 institutions like this where I assume they're concrete
7 it seems to me could make isodose curves possibly non-
8 conservative.

9 DOCTOR PAPERIELLO: I agree with you. I
10 really believe that's another direction. We used
11 microshield. We did have a relatively basic computer
12 code to do the calculations. We really need more
13 effort in the Agency for incident and emergency
14 response in terms of computer programs and people
15 trained on them to use them. There are codes that can
16 do it. It's finding people. It's getting the codes
17 and training people on them.

18 COMMISSIONER REMICK: I'm sure. Okay.

19 DOCTOR PAPERIELLO: Understood.

20 COMMISSIONER REMICK: If I could refer you
21 to page 5-10 of the report, I thought that all of the
22 locations of Oncology Services Corporation were under
23 one license, but just last week we got a copy of a
24 confirmatory action letter on the Marlton, New Jersey,
25 which makes me feel that's a separate license, so I'm

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1 confused.

2 DOCTOR PAPERIELLO: It is.

3 COMMISSIONER REMICK: Which one of these
4 are covered by the license in question and which ones
5 therefore are affected by the Commission order?

6 DOCTOR PAPERIELLO: It is my understanding
7 that facilities in Pennsylvania that have HDR are
8 covered under -- well, you can see that we have the
9 license in the back of the report as an attachment and
10 you can -- it lists the places that are covered. Not
11 all the places in this organizational chart have HDR,
12 I believe, and of course the facility in Florida is an
13 agreement state. I'm aware of a facility in New York.
14 It's been difficult. We don't understand the
15 corporate structure of Oncology Services and who owns
16 what and the like.

17 COMMISSIONER REMICK: Let me make sure I
18 understand what you're saying. The locations in
19 Pennsylvania that have the high dose rate units are
20 affected by -- are under the license and affected by
21 the order?

22 DOCTOR PAPERIELLO: That's right.

23 COMMISSIONER REMICK: But the Florida and
24 New Jersey locations would not be, then?

25 DOCTOR PAPERIELLO: They were a separate

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1 license.

2 COMMISSIONER REMICK: Separate license,
3 okay.

4 DOCTOR PAPERIELLO: But Region I did the
5 inspection last week of the facility in New Jersey
6 with similar findings.

7 COMMISSIONER REMICK: Okay.

8 Something that wasn't clear to me. In the
9 report it indicates that there were two individuals
10 pregnant at the time, identified as CNA-I and
11 Dietitian.

12 DOCTOR PAPERIELLO: Right.

13 COMMISSIONER REMICK: And it's indicated
14 that both of them had an average dose of about 500
15 millirem?

16 DOCTOR PAPERIELLO: It ranged from below
17 500 to slightly above, I think between 400 to 600,
18 somewhere in there.

19 COMMISSIONER REMICK: Right. But then
20 later on it indicates that the dietitian was one of
21 the nine people to receive the cytogenetic
22 evaluations, indicating that the nine were identified
23 as having the highest potential for exposure, and yet
24 the dietitian was a very low exposure.

25 DOCTOR PAPERIELLO: Well, because when we

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1 started this event we knew the dietitian was pregnant
2 and we deliberately had her done for the cytogenetics.
3 I think somewhere I said "generally." I tried --

4 COMMISSIONER REMICK: In your presentation
5 you said "almost always."

6 DOCTOR PAPERIELLO: Almost always. That
7 was the one case. The other individual we were not
8 aware of until very late. She was not around at the
9 time that we did the investigation and in both of
10 these cases the physicians -- they have had medical
11 consultation on the thing and we try to avoid getting
12 into people's private medical lives in terms of the
13 report.

14 COMMISSIONER REMICK: The thing that stuck
15 out is one had the cytogenetic evaluation and the
16 other did not.

17 DOCTOR PAPERIELLO: No. The other one we
18 didn't get early on. We found out very, very late.

19 COMMISSIONER REMICK: I see.

20 DOCTOR PAPERIELLO: Had we known early on,
21 we probably would have done that individual too.

22 COMMISSIONER REMICK: I see. Okay.

23 MS. JONES: If I could just add one thing,
24 there was -- the individual who did not have
25 cytogenetic studies done was not one of the nursing

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1 assistants who was on duty at all during that week.
2 She was just concerned and she contacted us later.
3 And because she was only assisting patients
4 periodically throughout that week and was not on
5 schedule at all with that patient, her probability for
6 exposure was very low.

7 COMMISSIONER REMICK: Right, although her
8 exposures on the average were about the same as the
9 dietitian who did receive something.

10 MS. JONES: That was coincidental.

11 COMMISSIONER REMICK: It just seemed to be
12 a curious inconsistency and that explains it.

13 Another thing, the report indicates that
14 your evaluations, your dose assessments, are about a
15 factor of three higher than the licensee. Are there
16 any obvious reasons that you're aware of?

17 DOCTOR PAPERIELLO: Yes.

18 MS. JONES: In some cases.

19 DOCTOR PAPERIELLO: Yes. I thought
20 somewhere I discussed it. They range and I did do a
21 side by side comparison, but we did a number of people
22 they didn't do. The licensee's consultant took credit
23 for attenuation, but they used narrow beam
24 attenuation. You can't use narrow beam attenuation in
25 this case, so therefore we feel they're in error.

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1 Besides, their results are not as in agreement with
2 the cytogenetic as ours.

3 COMMISSIONER REMICK: Okay. Are there any
4 facts -- I think the report is up to the 27th of
5 January. Are there any facts that you're aware of
6 since then, since the finalizing of the report, that
7 is of interest?

8 DOCTOR PAPERIELLO: If we would, Tom, it
9 would be in your area, in Southwest Research work on
10 the wire that's ongoing.

11 MR. RICH: They're just continuing to test
12 the wire at this time. They have another sample
13 received from an Omnitron that was in a shipping
14 container that had the same type of fracture surface,
15 same type of fractographical results. They're going
16 to continue to do testing with Omnitron consultants
17 and NRC consultants to determine if hydrogen is the
18 cause and, if so, how does it get absorbed to the
19 Nitinol wire and then how can they prevent it and
20 isolate it.

21 COMMISSIONER REMICK: I see. Thank you.

22 I want to commend the staff for an
23 excellent IIT team report. Very good.

24 DOCTOR PAPERIELLO: Thank you.

25 CHAIRMAN SELIN: Before we go on with the

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1 questioning, Mr. Taylor, you talked about follow-up in
2 sort of a generic sense. Have you thought out yet
3 whether there will be specific follow-up on the IIT
4 report more specifically than general follow-up on
5 medical regulation?

6 MR. TAYLOR: I think we'll probably issue
7 a supplement to this report -- we haven't scheduled
8 that -- as additional information develops.

9 CHAIRMAN SELIN: Would there be follow-up
10 of the type "lessons learned" and --

11 MR. TAYLOR: Yes. We will provide that.
12 That's in our procedures whenever we conduct this type
13 of review.

14 CHAIRMAN SELIN: Commissioner Curtiss?

15 COMMISSIONER CURTISS: I don't have any
16 questions. Let me just pick up on the point that
17 Commissioner Remick raised. I had not focused on this
18 before and I don't want to get too deeply in terms of
19 the order, but the two facilities that are not covered
20 under the order, if they are in fact as presented here
21 in the organizational chart reporting through the
22 corporate structure which in turn under this report
23 has come under some sharp criticism for various
24 reasons, perhaps it would be worth looking at what the
25 status of those two facilities is and whether they are

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1 similarly situated in terms of the problems that led
2 the staff to take the actions they did with respect to
3 the eight facilities that are covered by the order
4 itself.

5 I'd add my voice to what Commissioner
6 Remick has said. I thought the IIT was a first-rate
7 report. I'd like to commend everybody who was
8 involved in it. I know a lot of long hours and hard
9 work went into this document. I've read the thing
10 from front to back and found it extremely readable and
11 I thought quite balanced in terms of its perspective,
12 as is the case with an IIT.

13 In fact, the purpose of an IIT is to
14 permit us to step back from the line responsibilities
15 for regulating these activities. I thought it
16 presented a pretty comprehensive road map to the kinds
17 of things that not just others but that we at the NRC
18 need to think about doing in several areas and it's
19 identified some shortcomings that ought to be followed
20 up on in the near future.

21 I guess I just have one question in terms
22 of the procedure, perhaps for Mr. Taylor.

23 What is your current thinking in terms of
24 how you will coordinate the response that you propose
25 for the IIT findings with the ongoing review of the

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1 medical program? Are those two going to be brought
2 together in some fashion or have you gotten to that
3 point yet?

4 MR. TAYLOR: We really haven't gotten to
5 that point, although since Doctor Paperiello is going
6 to -- there's nothing like experience to help him out,
7 since he will be leading that review, so maybe that's
8 my answer at this point.

9 COMMISSIONER CURTISS: Okay.

10 MR. TAYLOR: But certainly this has been
11 a very important experience for us.

12 COMMISSIONER CURTISS: And, as I say, I
13 think it identifies a number of things that we need to
14 focus on. We need to coordinate that with the ongoing
15 effort so that we don't get too many things going in
16 separate directions at the same time.

17 I'd be especially interested in further
18 follow-up on our relationship with the FDA and the
19 circumstances that are the subject of their current
20 audits. I think that's an area where frankly we might
21 be able to take a look at a mechanism for closer
22 coordination on both sides.

23 So, again, I'd like to commend you for
24 what I thought was a first-rate report and thank you.

25 MR. TAYLOR: Commissioner, I'd like to

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1 note -- it hasn't been publicly noted, although the
2 Commission is aware, this is our first IIT of a
3 medical therapy type treatment event.

4 I believe that's correct, Mr. Jordan?

5 MR. JORDAN: Yes.

6 CHAIRMAN SELIN: Well, you've set a very
7 difficult standard for those, I hope, few incidents
8 that follow as far as the investigation.

9 Commissioner Rogers?

10 COMMISSIONER ROGERS: Yes, well, I'll just
11 second the superb report that was done. I think it
12 was really an outstanding job.

13 Just one question with respect to the
14 administration of the procedure. Your report
15 discovered that the source was inserted into the
16 catheters in the opposite order to the plan?

17 DOCTOR PAPERIELLO: We suspect that
18 occurred. We can't -- the people give you an argument
19 on that. We're trying to get around the issue on
20 which catheter was removed by the physician. That
21 would account -- it does seem logical that the
22 catheter that you were tugging at when you were going
23 through this might be the one the physician removed.
24 When we started asking people, "Well, how sure are you
25 which catheter was removed and which one fell out of

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1 the patients body," we got diverse answers. So that,
2 I think, is partly an open issue.

3 But I would point out you could miswire,
4 for want of a better term, "miswire" this thing.
5 There is no color-coding. There is numbering.
6 There's a treatment plan. The technologist has, you
7 might say, a wiring diagram in hand and has to connect
8 these tubes from the device to the patient, but you
9 could in fact misconnect these devices when dealing
10 with multiple catheters.

11 We discuss what we think may have
12 happened. We're not absolutely sure. It's not quite
13 clear it has bearing on the principal findings of the
14 team, but, again, you're dealing with people's
15 memories several weeks later and, if they didn't think
16 it was significant at the time, they may not recall
17 that.

18 COMMISSIONER ROGERS: Well, I wonder
19 whether FDA has any thoughts about any requirements
20 that would help to ensure that that kind of
21 misidentification, if it did occur, to prevent it in
22 the future.

23 MR. DAMASKA: That's one of the issues
24 that we're looking at and I think that would clearly
25 come under the adequate directions for use aspect,

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1 yes.

2 COMMISSIONER ROGERS: Right.

3 That's all for me, thank you.

4 CHAIRMAN SELIN: Doctor Paperiello,
5 obviously the Commission is very impressed and very
6 pleased with the thoroughness and the timeliness of
7 the report, the fact that the job seems to be so
8 thorough and was done on a very demanding schedule.

9 I would like to emphasize that this report
10 is a beginning, not an end, that you've pointed out a
11 fair number of regulatory issues that need to be
12 followed up on. We've had some incidental discussion
13 about how that might be followed up.

14 I'd also like to say something to the
15 licensee, that it's clear, I hope it's clear, that the
16 object of this meeting is not to concentrate on the
17 specific actions that deal with the licensee, that
18 there are separate forums for that, but rather to
19 follow up on the regulatory impact of the report.
20 Nevertheless, the report does make findings and
21 assertions that are of importance to the licensee.

22 If the licensee chooses to respond to this
23 report, wishes to respond to this report outside of
24 the enforcement forum, the Commission of course would
25 entertain a request from the licensee to appear before

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1 us at a later date with whatever comments they care to
2 make, but those would be in addition to the remarks
3 that the licensee would make in the enforcement forum.

4 We thank you, ladies and gentlemen, for
5 this fine effort and look forward to an early proposal
6 from the staff or set of proposals on how to implement
7 the findings of the investigation.

8 Thank you very much.

9 (Whereupon, at 11:41 a.m., the above-
10 entitled matter was concluded.)
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BRIEFING BY IIT ON LOSS OF IRIDIUM-192
TITLE OF MEETING: SOURCE AND THERAPY MISADMINISTRATION AT
INDIANA REGIONAL CANCER CENTER, INDIANA,
PA., NOVEMBER 16, 1992
PLACE OF MEETING: ROCKVILLE, MARYLAND
DATE OF MEETING: FEBRUARY 8, 1993

were transcribed by me. I further certify that said transcription
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Carol Lynch

Reporter's name: PETER LYNCH

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COMMISSION BRIEFING

INCIDENT INVESTIGATION

OF

LOSS OF AN IRIIDIUM-192 SOURCE
AND THERAPY MISADMINISTRATION AT
INDIANA REGIONAL CANCER CENTER
INDIANA, PENNSYLVANIA,
ON NOVEMBER 16, 1992

DR. CARL J. PAPERIELLO
Incident Investigation Team Leader

FEBRUARY 8, 1993

ONCOLOGY SERVICES INCORPORATED INCIDENT INVESTIGATION TEAM

Carl J. Paperiello, RIII	-	Team Leader
Mohamed M. Shanbaky, RI	-	Deputy Team Leader
Cynthia G. Jones, NMSS	-	Radiological Dose Evaluation
Penny A. Nessen, RI	-	Radiation Protection
Thomas W. Rich, NMSS	-	Engineering Evaluation
Ronald L. Lloyd, AEOD	-	General Engineering
Alan L. Madison, AEOD	-	Sequence of Events
Daniel F. Flynn, M.D.	-	NRC Medical Consultant

M. Marcia Karabelnikoff, AEOD		On-Site Admin. Coordinator
Cherie Siegel, AEOD	-	Administrative Coordinator
June L. Garland, NMSS	-	Team Secretary
Juanita F. Beeson, ADM	-	Technical Editor
Lionel J. Watkins, IRM	-	Visual Information Specialist
John Orban, IRM	-	Visual Information Specialist

AGENDA

- **Sequence of Events**
- **Radiological Consequences**
- **Failure to Detect Broken Source Wire**
- **Potential Cause of Source Wire Failure**
- **Findings and Conclusion**

SEQUENCE OF EVENTS

- **Indiana Regional Cancer Center**
- **Scenery Hill Manor Nursing Home**
- **Browning-Ferris Industries**
- **Source Retrieval**
- **Greater Pittsburgh Cancer Center**

SEQUENCE OF EVENTS

November 16, 1992

- **Indiana Regional Cancer Center**

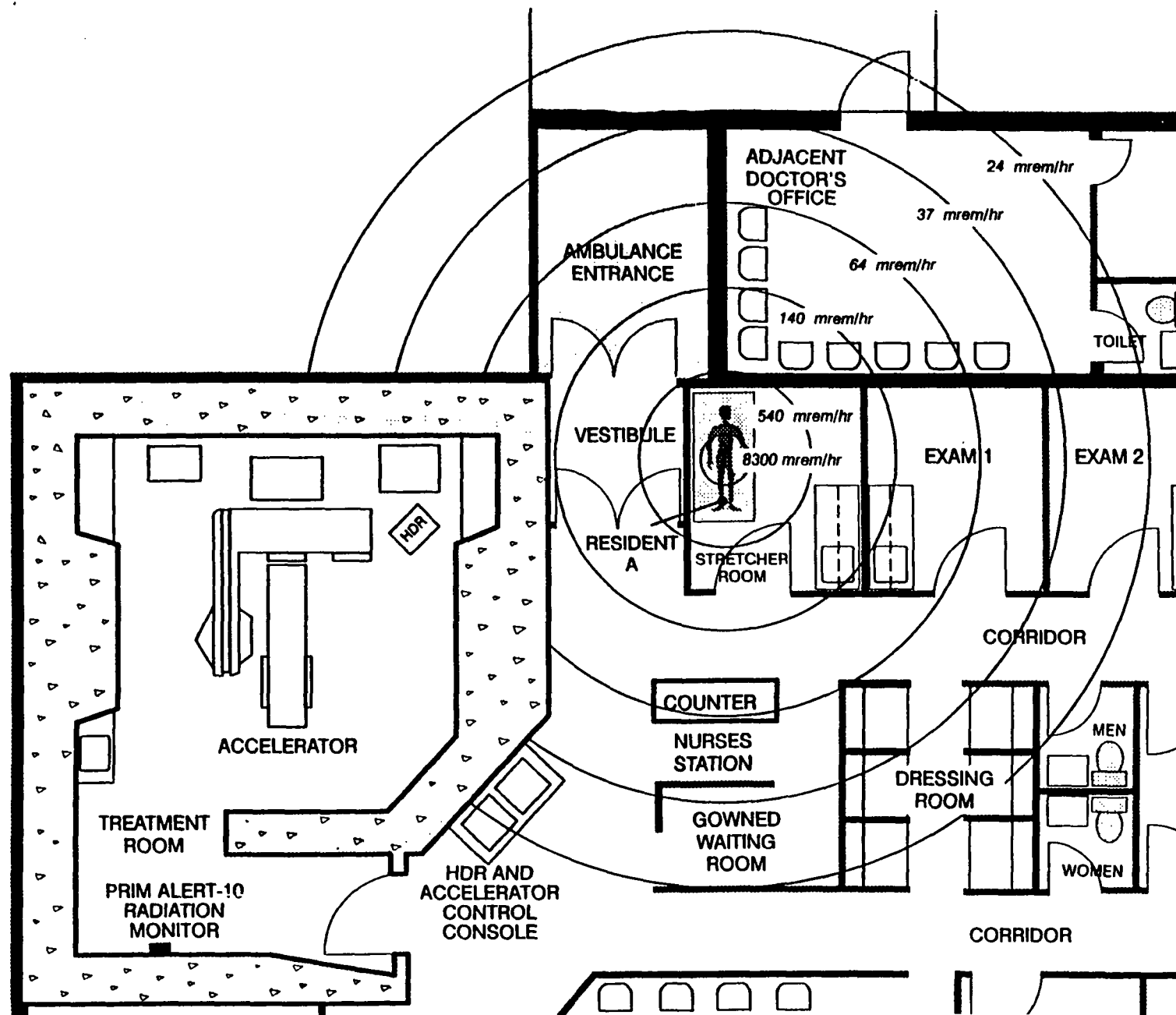


Diagram of the Indiana Regional Cancer Center Showing 2 Meter Isodose Curves from the Iridium-192 Source on November 16, 1992.

SEQUENCE OF EVENTS

(Continued)

November 16-25, 1992

- **Scenery Hill Manor Nursing Home**

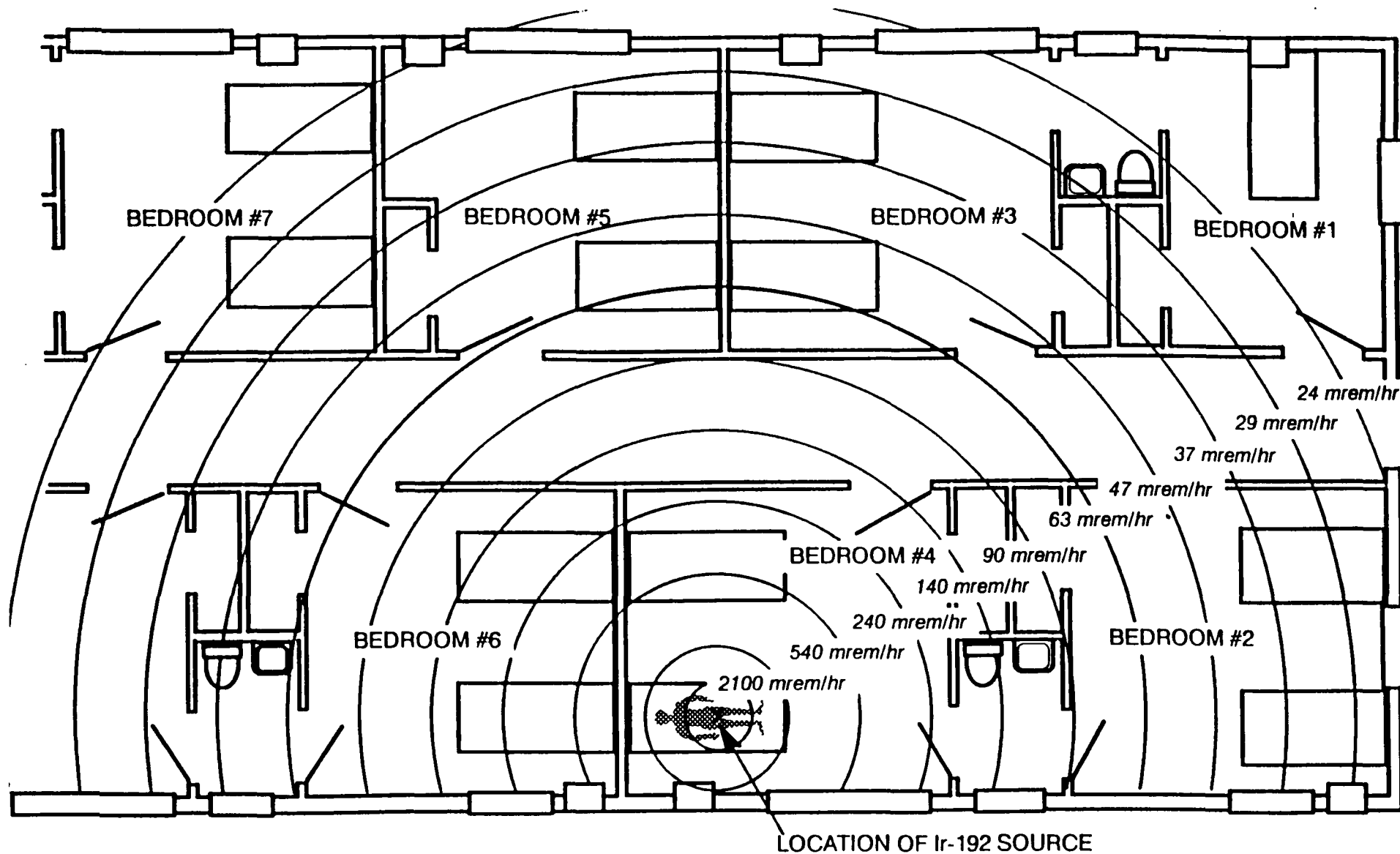
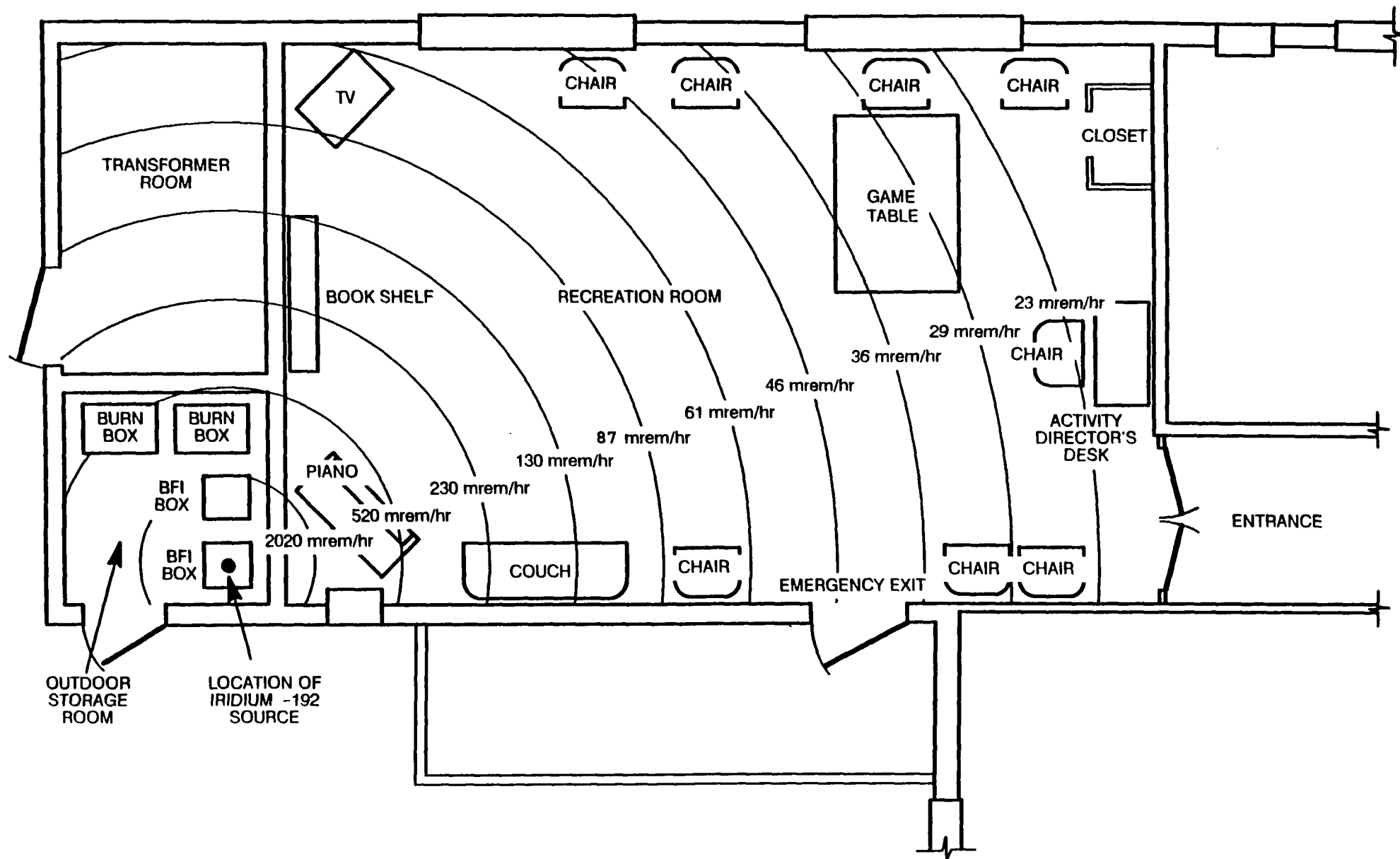


Diagram of Rooms Adjacent to Patient Room 4B Showing 1-Meter Isodose Curves from the Iridium-192 Source on November 16, 1992



Detailed Diagram of the Scenery Hill Manor Nursing Home Recreation Room Showing 1-meter Isodose Curves from the Iridium-192 Source Located in the Outside Storage Room from November 20 to 25, 1992

SEQUENCE OF EVENTS

(Continued)

November 25 - December 1, 1992

- **Browning-Ferris Industries**

SEQUENCE OF EVENTS

(Continued)

December 1, 1992

- **Source Retrieval at Browning-Ferris Industries at Carnegie, Pennsylvania**

SEQUENCE OF EVENTS

(Continued)

December 7, 1992

- **Omnitron 2000 Source Wire Break at the Greater Pittsburgh Cancer Center**

RADIOLOGICAL CONSEQUENCES

- **Patient**
- **Occupational Workers**
- **Nonoccupational Workers and the Public**

**Table A Summary of Doses Received as a Result of an
Iridium-192 Source Lost
from November 16 to December 1, 1992**

Dose Range (rem)	Estimated Number of Individuals Exposed
0.0 - 0.5	42
0.5 - 1	11
1 - 5	20
5 - 10	13
10 - 15	7
15 - 20	1
TOTALS	94

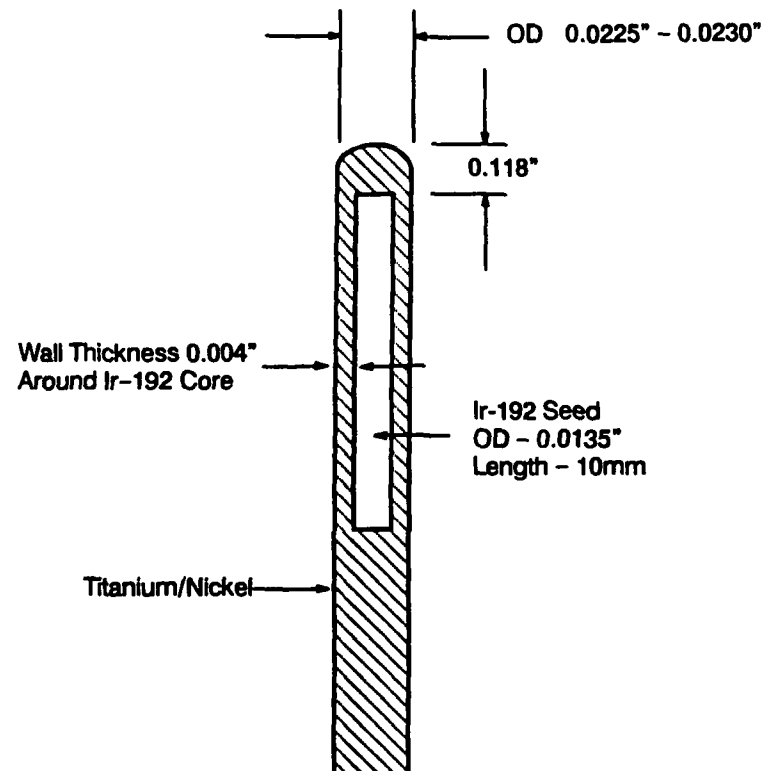
Table B Summary of Collective Doses for All Locations

Location	Estimated Number of Individuals Exposed	Dose Range (rem)	Collective Dose (Person-rem)
IRCC Clinic	6	0.1 - 0.86	2.66
IRCC-Other	17	0.01 - 1.4	0.94 - 2.6
Ambulance	2	0.5 - 2.6	1.8 - 3.4
SHM-Staff	21	0.4 - 22.3	110 - 166
SHM-Visitors	7	2.2 - 16.6	21 - 46
SHM-Residents	22	0.2 - 20.5	35 - 73
BFI-Carnegie	19	0.0004 - 9.0	8.3 - 22
TOTALS	94		180 - 316

RADIOLOGICAL CONSEQUENCES (Continued)

- **Blood Counts and Cytogenetics**
- **No Contamination**

**IR-192 Seed in Titanium/Nickel
Model #SL -777**



Ir-192 Seed in Titanium/Nickel

An Ir-192 Seed is encapsulated firmly inside a solid titanium/nickel wire.
The overall length of the Ir-192 Seed is 10mm

The above composite has passed all ANSI requirements for sealed sources.

**Iridium-192 Seed in Titanium/ Nickel
(Reproduced from Registration
Certificate LA-0760-S-102-S)**

POTENTIAL CAUSE OF SOURCE WIRE BREAK

- **Design Change Made to Source Wire Material - May 1992**
- **Source Wire Failures Occurred Near the Bottom of Cavity**
- **Cavity Manufacturing Irregularities**
- **Degradation of Teflon Liner in Source Shipping Container**
- **Source Wire Failure Hypothesis**

FAILURE TO DETECT SOURCE WIRE BREAK

- **Brief Computer Screen Error Message for dc Motor Retract**
- **Staff was Aware of PrimAlert-10 Alarm on November 16, 1992**
- **Omnitron 2000 Computer Console Indicated "Safe"**
- **Survey Meter Available but Not Used**

HUMAN FACTORS CONSIDERATIONS

- **Believed Source Wire Would Not Break**
- **Previous Spurious PrimAlert-10 Alarms**
- **Safety Awareness**

FINDINGS AND CONCLUSIONS

- **Radiological Consequences Were Serious to the Patient and Significant to Many Members of the Public**
- **Weaknesses in Corporation Radiation Protection Program**
- **Weaknesses in the Design and Testing of the Omnitron 2000 Afterloader System and Its Source Wire**

FINDINGS AND CONCLUSIONS

(Continued)

- **OSC and IRCC Lacked Critical Safety Awareness with Respect to High Dose Rate Brachytherapy**
- **Overall Regulatory Oversight Was Weak**
- **No Regulatory Guidance Exists for Nonradioactive Waste Collectors; BFI failed to Follow their Existing Radiation Control Policies**

WEAKNESSES IN CORPORATE RADIATION SAFETY PROGRAM

- **Program for HDR Was Weak and Ineffective**
- **No Formal Implemented Radiation Safety Training**
- **Reliance on Previous Academic Training**
- **No One from IRCC Acknowledged Responsibility for Training**
- **Initially No One Knew Who Was the Radiation Safety Officer**

OMNITRON WEAKNESSES

- **Testing and Validation of Source-Wire Design**
- **Emergency Retraction Information**
- **Omnitron QA/QC**

REGULATORY OVERSIGHT WAS WEAK

- **HDR Brachytherapy Was Not Clearly Addressed by Regulations**
 - **Not Explicitly Recognized by Part 35**
 - **Obsolescent Licensing Guidance**
 - **No Explicit Inspection Guidance**
 - **Not Addressed by Current Regulatory Guide (10.8, Rev. 2)**

REGULATORY OVERSIGHT WAS WEAK

(Continued)

- **HDR Inspection Program Not Responsive To Major Amendments**
 - **August, 1990 - Original License**
 - **November, 1990 - Amendment No. 1**
 - **September, 1991 - Initial Inspection**
 - **January, 1992 - Amendment No. 2 (Expansion)**
 - **August, 1992 - Amendment No. 3**
 - **September, 1995 - Routine Inspection**

CRITICAL SAFETY AWARENESS

- **No Patient Survey**
- **Treatment by Technologist**
- **No M.D. at Treatment Console**
- **Part-Time Medical Physicist**
- **Did Not Believe Wire Could Break**
- **Improper Response to PrimAlert-10**
- **No Daily Checks of Omnitron Door Interlocks**

NO REGULATORY GUIDANCE FOR NON RADIOACTIVE WASTE COLLECTORS

- **No Regulatory Guidance Exists for Waste Collectors, Brokers, and Recyclers Similar to the Steel Industry**
- **BFI Has an Aggressive Radiation Monitoring Program**
- **BFI Not Prepared to Handle Highly Radioactive Material**
- **BFI Driver Did Not Perform Required Survey of Nursing Home Waste**

LICENSING JURISDICTION FOR THE OMNITRON 2000 HDR AFTERLOADER

- **FDA Has the Pre-Market Approval and Good Manufacturing Practices Inspections for the Source Wire and HDR Afterloader**
- **State of Louisiana Approved the Source Wire and HDR Afterloader for Distribution and Use, and Issues the Sealed Source and Device Certificates**
- **NRC Responsible for Licensing of Oncology Services Corporation to Use the Omnitron 2000 HDR Afterloader**

Loss of an Iridium-192 Source and Therapy Misadministration at Indiana Regional Cancer Center Indiana, Pennsylvania, on November 16, 1992

U.S. Nuclear Regulatory Commission



Loss of an Iridium-192 Source and Therapy Misadministration at Indiana Regional Cancer Center Indiana, Pennsylvania, on November 16, 1992

Manuscript Completed: February 1993
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Washington, DC 20555



ABSTRACT

On December 1, 1992, the Indiana Regional Cancer Center reported to the U.S. Nuclear Regulatory Commission's (NRC) Region I that they believed a $1.37 \text{ E}+11$ becquerel (3.7-curie) iridium-192 source from their Omnitron 2000 high dose rate remote brachytherapy afterloader had been found at a biohazard waste transfer station in Carnegie, Pennsylvania. After notifying the NRC, this cancer center, one of several operated by the licensee, Oncology Services Corporation, retrieved the source, and Region I dispatched an inspector and a supervisor to investigate the event. The source was first detected when it triggered radiation alarms at a waste incinerator facility in Warren, Ohio. The licensee informed the NRC that the source wire had apparently broken during treatment of a patient on November 16, 1992, leaving the source in the patient. On the basis of the seriousness of the incident, the NRC elevated its response to an Incident Investigation. The Incident Investigation Team initiated its investigation on December 3, 1992. The investigation team concluded that the patient received a serious misadministration and died on November 21, 1992, and that over 90 individuals were exposed to radiation from November 16 to December 1, 1992. In a press release dated January 26, 1993, the Indiana County Coroner stated that the cause of death listed in the official autopsy report was "Acute Radiational Exposure and Consequences Thereof." An almost identical source wire failure occurred with an afterloader in Pittsburgh, Pennsylvania, on December 7, 1992, but with minimal radiological consequences. This incident was included in the investigation. This report discusses the Omnitron 2000 high dose rate afterloader source-wire failure, the reasons why the failure was not detected by Indiana Regional Cancer Center, the potential consequences to the patient, the estimated radiological doses to workers and the public, and regulatory aspects associated with this incident.

CONTENTS

Section	Page
Abstract	iii
Executive Summary	xiii
Team Members	xxi
Acknowledgements	xxii
Abbreviations	xxiii
1 INTRODUCTION	1-1
2 SEQUENCE OF EVENTS	2-1
2.1 Patient Treatment Plan	2-1
2.2 Indiana Regional Cancer Center Incident	2-2
2.3 Greater Pittsburgh Cancer Center Incident	2-12
3 OMNITRON 2000 HIGH DOSE RATE REMOTE AFTERLOADER SYSTEM	3-1
3.1 Description of the Afterloader System	3-1
3.2 Description of the Source Wire	3-6
3.3 Description of the Omnitron 2000 Afterloader System Software	3-8
3.4 Omnitron's Quality Assurance and Quality Control Program	3-10
3.5 Training, Operating, and Emergency Procedures for the Omnitron 2000 High Dose Rate Afterloader	3-11
4 EQUIPMENT PERFORMANCE	4-1
4.1 Failure Analysis Pertaining to the Source Wire	4-1
4.2 Immediate Cause of Wire Break	4-9
4.3 Potential Root Cause of Wire Break	4-10
4.4 Failure of Device To Detect Loss of Part of the Wire	4-11
4.5 Performance of PrimAlert-10	4-11

CONTENTS (continued)

Section	Page
5	HUMAN FACTORS ANALYSIS 5-1
5.1	Organizational Factors 5-1
5.2	Emergency Operating Procedures 5-7
5.3	Training 5-7
5.4	Quality Management Program for High Dose Rate Brachytherapy at the Indiana Regional Cancer Center 5-8
6	RADIOLOGICAL DOSE EVALUATIONS 6-1
6.1	Overview of Team's Methodology 6-1
6.2	NRC's Medical Evaluation of the Dosimetry and Effect of Radiation on Patient A 6-3
6.3	Evaluation of Radiation Doses for Occupational Workers at the Indiana Regional Cancer Center 6-10
6.4	The Public 6-14
6.5	Blood Studies 6-38
7	LICENSEE AND VENDOR RESPONSE TO INCIDENT 7-1
7.1	Misadministration 7-1
7.2	Licensee Actions 7-1
7.3	Vendor Actions 7-8
8	REGULATORY OVERSIGHT 8-1
8.1	High Dose Rate Afterloader Device Review by the Food and Drug Administration 8-1
8.2	Source and Device Registration by the State of Louisiana 8-1
8.3	NRC's Regulations and Guidance for Quality Assurance and Quality Control Programs for Sealed Source and Device Vendors . . . 8-2
8.4	NRC Licensing of Oncology Services Corporation 8-3
8.5	NRC Inspection Program and Process 8-4
8.6	NRC Regulatory Activities 8-5
9	PRECURSORS 9-1
9.1	High Dose Rate Remote Afterloader Incidents 9-1
9.2	Omnitron 2000 Performance History 9-2

CONTENTS (continued)

Section	Page
10 FINDINGS AND CONCLUSIONS	10-1
10.1 Radiological Consequences Were Serious to the Patient and Significant to Many Members of the Public.	10-1
10.2 Weaknesses in Oncology Services Corporation's Radiation Protection Program Were a Contributing Cause of the Seriousness of the Event and Radiation Exposure Consequences	10-2
10.3 Weaknesses Existed in the Design and Testing of the Omnitron 2000 Remote Afterloader System and Its Source Wire	10-3
10.4 Oncology Services Corporation and Indiana Regional Cancer Center Lacked Critical Safety Awareness with Respect to High Dose Rate Brachytherapy	10-3
10.5 Overall Regulatory Oversight Was Weak	10-4
10.6 No Regulatory Guidance Exists for Nonradioactive Waste Collectors. In Addition, Browning-Ferris Industries Failed To Follow Their Existing Radiation Control Policies	10-5
Glossary	G-1

Appendices

Section	Page
A Incident Investigation Team Charter	A-1
B Properties of Iridium-192	B-1
C Low Dose and High Dose Rate Brachytherapy	C-1
D Oncology Services Corporation Materials License	D-1
E Interviews and Meetings the Incident Investigation Team Conducted	E-1

CONTENTS (continued)

Tables

Number	Page
A	Summary of Doses Received as a Result of an Iridium-192 Source Lost from November 16 to December 1, 1992 xix
B	Summary of Collective Doses for All Locations xx
4.1	Summary of Test Samples 4-6
4.2	Summary of Tests Performed on Wire Samples 4-8
6.1(a)	Normal Tissue Distances from Possible Source Location in Patient A 6-7
6.1(b)	Normal Tissue Dose (cGy) Versus Possible Source Location in Patient A 6-8
6.2	Tolerance of Different Tissues to Radiation Exposure 6-9
6.3	Summary of Radiation Doses for Indiana Regional Cancer Center Occupational Workers 6-13
6.4	Summary of Radiation Doses for Indiana Regional Cancer Center Nonoccupational Workers and Patients 6-16
6.5	Summary of Radiation Doses for Adjacent Physician's Personnel and Patients . . 6-18
6.6	Summary of Radiation Doses for Ambulance Service Employees 6-20
6.7	Schedule for Nursing Staff Assigned to Patient A Week of November 16-20, 1992 6-22
6.8	Summary of Whole Body Dose Estimates for SHM Nursing Home Staff 6-26
6.9	Summary of Extremity Dose Estimates for Selected SHM Nursing Home Staff . 6-27
6.10	Summary of Radiation Doses for Relatives and Friends During November 16-20, 1992 6-29
6.11	Summary of Radiation Doses for SHM Residents During November 16-20, 1992 6-30

CONTENTS (continued)

Tables

Number	Page
6.12	Summary of Radiation Doses for SHM Nursing Home Residents While in Recreation Room During November 20-25, 1992 6-32
6.13	Summary of Whole Body Dose Estimates for BFI Employees 6-36
6.14	Summary of Extremity Dose Estimates for Selected BFI Employees 6-37
6.15	Comparison of Calculated Whole Body Doses with Cytogenetic Evaluation . . . 6-42
B.1	Calculated Dose Rate (mSv/hr) at Selected Distances for 3.3085 E+11 Bq (8.942-Ci) Iridium-192 Source Certified August 28, 1992 . . . B-4
B.2	Iridium-192 Dose Rates vs. Distance With and Without Berger Buildup Factors B-6

Figures

Number	Page
1.1	Indiana Regional Cancer Center, Indiana, Pennsylvania 1-3
2.1	Isodose Graph for Initial High Dose Rate Patient Treatment Plan 2-16
2.2	Summary of Initial High Dose Rate Treatment Plan 2-17
2.3	Back of High Dose Rate Afterloader 2-18
2.4	Front of High Dose Rate Afterloader 2-18
2.5	Diagram of Indiana Regional Cancer Center 2-19
2.6	Front of Scenery Hill Manor Nursing Home 2-20
2.7	Soiled Utility Room with Biohazard Waste Container 2-21
2.8	Waste Storage Room at Scenery Hill Manor Nursing Home 2-22

CONTENTS (continued)

Figures

Number	Page
2.9	Browning-Ferris Industries Straight Truck Used To Collect Waste from Scenery Hill Manor Nursing Home 2-23
2.10	Aerial Photograph of Browning-Ferris Industries Transfer Station in Carnegie, Pennsylvania 2-24
2.11	BFI Tractor-Trailer [14.63-m (48-ft)] Long Showing Location of Source Container in the Trailer 2-25
2.12	Route of Browning-Ferris Industries Semitruck 808 on November 25, 1992 . . 2-26
2.13	Diagram of Tractor-Trailer Route Carrying Iridium-192 Source at BFI-Warren on November 25, 1992 2-27
2.14	Boxes on Conveyor Feedline Adjacent to Radiation Monitor 2-28
2.15	Console for Monitoring Radiation at Incinerator Conveyor Feedline 2-28
2.16	Survey Meter Used at BFI-Warren 2-29
2.17	Location of Where Manager and Supervisor Detected Radiation (~400 Feet) from BFI Trailer 808 2-30
2.18	BFI Truck Parking Lot, Carnegie, Pennsylvania 2-31
2.19	Location of Isolated Biohazard Box Containing Source 2-32
2.20	BFI Locking Recycle Container 2-33
2.21	Interior of the BFI Waste Container Where Radioactive Source Was Stored . . . 2-33
2.22	Location of Radioactive Source Recycle Container at BFI-Carnegie 2-34
3.1	Omnitron 2000 High Dose Rate Remote Afterloader 3-12
3.2	Left Side of High Dose Rate Remote Afterloader, Indiana, Pennsylvania 3-13
3.3	Turret Drive Assembly, Indiana, Pennsylvania 3-14

CONTENTS (continued)

Figures

Number	Page
3.4	Control Panel for High Dose Rate Remote Afterloader, Indiana, Pennsylvania 3-15
3.5	Arrangement of the Computer Console Status Panel, Door Panel, System Printer, Linear Accelerator Controls, Computer Monitor, and Monitor for Patient Surveillance Camera 3-16
3.6	Computer Console Status Panel of High Dose Rate Remote Afterloader System . 3-17
3.7	Door Status Panel of High Dose Rate Remote Afterloader System 3-17
3.8	Connecting Catheter and Flexineedle Unassembled 3-18
3.9	Iridium-192 Seed in Titanium/Nickel 3-19
4.1	Source Wire—Possible Failure Areas 4-14
4.2	Track Where Source Tip May Have Contacted Parts of the Afterloader 4-15
4.3	Connecting Catheter and Flexineedle—Assembled 4-16
4.4	Shipping Container for Iridium-192 Source 4-17
4.5	Testing of a Sample Wire 4-18
5.1	Oncology Services Corporation Organizational Chart 5-10
6.1	Diagram of the Indiana Regional Cancer Center Showing Isodose Curves from the Iridium-192 Source on November 16, 1992 6-43
6.2	Location of Ambulance Staff and Patient A in Citizens' Ambulance 6-44
6.3	Diagram of Room 4B at the Scenery Hill Manor Nursing Home Showing Isodose Curves from the Iridium-192 Source on November 16, 1992 6-45
6.4	Diagram of the SHM Nursing Home Soiled Utility Room Showing Isodose Curves from the Iridium-192 Source on November 20, 1992 6-46

CONTENTS (continued)

Figures

Number	Page
6.5	Diagram of Rooms Adjacent to Patient Room 4B Showing 1-Meter Isodose Curves from the Iridium-192 Source on November 16, 1992 6-47
6.6	Detailed Diagram of the Scenery Hill Manor Nursing Home Recreation Room Showing Isodose Curves from the Iridium-192 Source Located in the Outside Storage Room from November 20 to 25, 1992 6-48
6.7	Portable Radiation Survey Meter from BFI-Carnegie Straight Truck 6-49
6.8	Aerial View of the Positions of Supervisor A, Safety Technician A, and Safety Technician B Around the Box Containing the Iridium-192 Source on December 1, 1992, Showing Isodose Curves in mSv per Hour 6-50
7.1	Treatment Plan Catheter Location 7-10
7.2	Actual Catheter Location 7-10
B.1	Simplified Decay Scheme for Iridium-192 B-2
B.2	Microshield 3.13 Program Showing Point Source and Shields B-3

EXECUTIVE SUMMARY

On November 16, 1992, an elderly patient was treated for anal carcinoma at the Indiana Regional Cancer Center (IRCC) in Indiana, Pennsylvania, using high dose rate (HDR) brachytherapy. The IRCC is one of ten cancer treatment centers using HDR units operated by the Oncology Services Corporation (OSC), the license holder. The patient died on the evening of November 21, 1992, five days after the treatment. Before the treatment, five catheters were placed in the tumor. During the treatment, an approximate $1.6 \text{ E}+11$ becquerel (4.3-curie) iridium-192 source was placed at various positions in each catheter to irradiate the tumor by use of a remotely controlled Omnitron 2000 afterloader. The treatment was conducted in the same shielded facility in which linear accelerator treatments were conducted. This treatment was the first of a series of three 600-centigray (rad) treatments planned by the physician, and the five catheters were to remain in the patient for subsequent treatments.

After a trial run through the five catheters with a dummy wire, the iridium source wire was placed in four catheters without difficulty. After several unsuccessful attempts to insert the source wire and the dummy wire into a fifth catheter, the treatment was terminated. An area radiation monitor in the treatment area was observed in an alarm condition at various times when the source should have been retracted during the unsuccessful attempts to insert the source wire through the catheter. Although three technologists and the physician attending the patient were aware of the alarm condition, no one conducted a survey for radiation levels with the available portable radiation survey instrument. The only action taken was to check the control console of the HDR remote afterloader. Because the console indicator showed "safe," they believed the source to be fully retracted into the lead shield and assumed the area radiation monitor was malfunctioning. They were unaware the source wire had broken, leaving the source in one of the catheters in the patient. The staff at the IRCC stated they had experienced difficulties with the area radiation monitor that had alarmed after patient linear accelerator treatments and that a survey after a previous false alarm had shown no radiation was present.

Before the incident, breakage of the wire encapsulating the iridium source was not considered credible by the vendor or the licensee. Omnitron International, Inc.'s (Omnitron's), emergency procedures were directed toward emergency retraction of a wire with a source that was stuck out and required manual retraction. The only individual interviewed who stated he believed that wire breakage was credible was the medical physicist at the Greater Pittsburgh Cancer Center (GPCC), another facility operated by OSC.

OSC appears to have provided no systematic radiation safety training to the staff at the IRCC. Dependence was placed on the staff's previous formal training; training by Omnitron on the HDR remote afterloader, which included Omnitron emergency procedures but did not include radiation safety; and an expectation that either the medical director or the medical physicist at each site would provide radiation safety training. This expectation was neither met nor were steps taken to confirm it.

Given the conflict between the data provided by the area radiation monitor and the Omnitron 2000 control panel, and the lack of radiation safety training that should have conditioned the staff to respond properly to radiation alarms, the licensee staff failed to respond to the alarm. Instead, a technologist unplugged and reset the radiation monitor. The technologist should have responded by using the portable survey meter available at the control console for the afterloader. This meter would have identified the fact that the source had not returned to the afterloader shield.

The patient spent 50 minutes in the treatment room. After the patient was removed from the treatment room, one catheter was observed to be loose and it was removed. The patient was returned to the nursing home where the patient resided with four catheters, one containing the iridium-192 source, in the patient's body. The source remained in the patient's body for almost four days. The catheter with the source came loose on the fourth day and, eventually, the catheter fell out (early on the morning of November 20, 1992). It was placed in a medical biohazards bag (red-bag) in a storage room by nursing home personnel who did not know it contained the radioactive source.

Later, on the same day, the catheter containing the source was moved to another storage location at the nursing home and placed in a box with other red bags. From November 16 through November 25, 1992, numerous residents, employees, and visitors to the nursing home were unknowingly irradiated. The ambulance staff who returned the patient to the nursing home were irradiated along with employees and patients at the IRCC who were present for the approximately 10 minutes while the patient was outside the treatment room after the treatment.

On November 25, 1992, a driver for Browning-Ferris Industries (BFI) picked up the nursing home red-bag waste as part of the driver's normal rounds. The driver had an operable portable radiation survey meter but, contrary to company procedures, did not survey the waste from the nursing home. The nursing home waste along with other medical waste was taken to a BFI facility in Carnegie, Pennsylvania, where it was loaded onto a trailer. This trailer remained in Carnegie throughout the following day, which was Thanksgiving, and early on November 27, 1992, was driven to a BFI medical waste incinerator in Warren, Ohio. At the Warren facility, fixed radiation monitors identified radiation emanating from the trailer, and, on facility personnel direction, the trailer was returned to Carnegie the same day. It was left over the weekend and on Monday, November 30, 1992, the BFI staff searched the truck for the radiation source. They identified the box with the radiation source and looked at individual red bags to identify the origin of the waste. On December 1, 1992, BFI successfully identified a name found with the red-bag waste in the box, and traced it to the nursing home.

After being notified by BFI, the nursing home called the IRCC on December 1, 1992. The cancer center had not used the HDR afterloader after the single treatment on November 16, 1992. Upon being informed of the source discovery, the medical physicist determined that

no source was present in the HDR afterloader and informed the NRC Region I office of this fact. The physician and the medical physicist drove to Carnegie to retrieve the source.

A second Omnitron 2000 source wire broke at the GPCC on December 7, 1992. This failure was included in the scope of this investigation. This wire broke in the same approximate location as the first wire. The GPCC medical physicist who was conducting the treatment was aware of the first incident and immediately recognized the problem and promptly and appropriately intervened, thereby preventing significant dose consequences to the patient or the cancer center staff. Apparently, the patient stressed the wire by an arm movement when the source was being retracted.

The NRC issued a Bulletin on December 8, 1992, to all licensees authorized to use the Omnitron 2000 afterloader requesting that they ensure patients are surveyed immediately after completing each therapy treatment; provide for prompt surgical intervention, if needed, in the event a source does not retract into the shielded container after treatment; and ensure that training on routine and emergency procedures is provided to licensee staffs initially and semiannually. In addition, an NRC Information Notice was issued December 17, 1992, informing all NRC medical licensees of this incident, reminding them of NRC requirements for positive assurance that all implanted sources have been removed before patients are released after treatment and informing them of the information provided in the Bulletin.

The cause of the source wire failures was investigated by the vendor, Omnitron, and their consultant and by the NRC team and its consultant laboratory, Southwest Research Institute. The Food and Drug Administration (FDA), which has responsibilities for the HDR afterloader as a medical device, conducted an independent inspection. The team shared its data with the FDA. The FDA issued its Form 483 with 37 findings to Omnitron on January 11, 1993. These findings were primarily in the quality assurance and quality control (QA/QC) area. The NRC team also interacted with the State of Louisiana because certain of Omnitron's activities are also overseen by this Agreement State.

Although the team identified a number of weaknesses in Omnitron's QA/QC program in addition to concerns with the design and prototype testing of the source wire, the cause of the wire failure is not known with certainty at this time. The wire broke at the bottom section of the source-wire cavity. The vendor had not calculated the maximum allowable stress in the design. The wire is made from Nitinol, a shape-memory alloy of nickel and titanium in approximately equal atomic percentages. The vendor believes it has evidence to show that storage of the source wire in teflon, if moisture is present, causes degradation of the teflon with release of fluorine or hydrogen fluoride that causes degradation of the Nitinol wire. The NRC and its consultant are still evaluating this hypothesis and conducting further studies.

An NRC medical consultant, who is a radiation oncologist, assisted the team in evaluating the effect of the source on the patient. The medical consultant concluded that an analysis of the medical records and physical dosimetry would indicate that the massive radiation dose

was a probable contributing cause of death in this patient. The consultant added that even if the exact cause of death could be attributed to other causes such as cancer or heart disease, the extent of the radiation received would have soon caused death. The consultant stated the radiation accident would probably have eventually resulted in a fatality in a younger, healthier patient within days or weeks of an equivalent accident.

The licensee reported the prescribed dose at one centimeter was $1.8 \text{ E}+03$ centigray (rad) to be delivered in three treatments and that the delivered dose was $1.6 \text{ E}+06$ centigray (rad) to the same point, an overdose of about three orders of magnitude. The licensee stated the effect on the patient would be significant local tissue damage and possible significant tissue damage to organs outside the treatment area, depending upon the progression of radiation damage with time before the patient expired. The licensee stated the dose was of sufficient magnitude that it believed it was highly probable that the radiation exposure was at least a contributing factor to the patient's subsequent death. In a press release dated January 26, 1993, the Indiana County Coroner stated that the cause of death listed in the official autopsy report was "Acute Radiation Exposure and Consequences Thereof."

In addition to the patient, the team evaluated the radiation doses to 94 persons associated with the IRCC event. Exposures of individuals wearing film badges were taken from the badge reading. Except for the physician, all other exposures were derived by calculations based on time-motion studies. These data were based on interviews and records. Individuals visiting the patient were identified through interviews with nursing home staff and known visitors. Becoming aware of a previously unidentified visitor from a newspaper article, the team interviewed that person and placed an advertisement in the local Indiana, Pennsylvania, newspaper. The advertisement requested any individual who was concerned about having come into contact with radiation from the source and who had not been interviewed to contact the NRC. A collect telephone number was provided. This led to two additional contacts. Table A presents a summary of the whole body doses received by the 94 individuals specifically evaluated. Table B presents a summary of collective doses for all locations. Extremity (i.e., hands and forearm) doses were calculated for those individuals who were judged to have the highest exposures because of the proximity of their hands to the source when caring for the patient or searching for the source. The highest extremity dose was calculated to be between 0.73 to 1.6 sieverts (73 to 160 rem) to the hands of one of the Certified Nursing Assistants.

Further studies were conducted by blood counts and cytogenetic studies on selected individuals. Blood counts are simple to perform but have sensitivity limited to 0.75 to 1.5 sieverts (75 to 150 rem). Cytogenetic studies are more sensitive [0.2 sieverts (20 rem)] but are much more difficult to perform. Cytogenetic studies were performed for the NRC by the Radiation Emergency Assistance Center/Training Site at Oak Ridge. Individuals selected for cytogenetic studies were almost always those with the greatest calculated dose. Because the highest calculated doses were at about the limit of detection [0.2 sieverts (20 rem)], the team expected that the cytogenetic data would show if there were significant nonconservative errors in the calculated doses (i.e., measured doses were higher than calculated doses).

Cytogenetic results were consistent with calculated doses within the limits of accuracy of both techniques.

No occupational worker received a radiation dose above the NRC occupational limit of 0.0125 sieverts (1.25 rem). While members of the public received radiation doses above applicable limits, no one received a dose at which acute radiation injury or clinical signs are expected to occur.

In addition to the radiological consequences and the cause of the wire failure the team made a number of findings:

1. OSC had weaknesses in their radiation safety program that were a major contributing cause of the seriousness of the event and radiation exposure consequences. Some of these were a result of a rapid expansion in their HDR brachytherapy program from one facility to ten facilities in less than a year. The Radiation Safety Officer (RSO) failed to ensure that the staffs at all facilities received adequate radiation safety training and that all management instructions relating to HDR were being followed. Informal and unwritten procedures that may have been adequate when the licensee possessed one HDR unit under the direct control of the RSO were ineffective for the expanded program.
2. A number of weaknesses were found in the design and testing of the Omnitron 2000. Weaknesses were identified in the testing and validation of source-wire design, and in the design of certain safety features of the HDR afterloader. These could allow the undetected retraction and further use of a broken wire with no warning to the user. Although not contributing to this event, weaknesses were found in Omnitron's QA/QC program.
3. The safety culture at IRCC contributed significantly to the event. Technologists routinely ignored the PrimAlert-10 alarm. Its problems were worked around and not fixed. Technologists did not survey patients, the afterloader, or the treatment room following HDR treatments. No one was sure who was responsible for radiation safety training or the radiation safety program. The authorized user failed to wear a film badge on both occasions when the source was encountered.
4. Overall regulatory oversight was weak. NRC regulations do not directly address HDR brachytherapy to the extent that teletherapy and low dose rate brachytherapy are addressed. Licensing guidance for HDR has been unchanged since 1986 in spite of significant changes in medical regulations and other medical licensing guidance. Inspection guidance for medical licensees does not specifically address HDR brachytherapy. Although inspected by the NRC Region I office within a year of initial licensing, the inspection program does not require early reinspection in cases where licensees significantly expand the scope of their program through license amendments. The regulatory interaction between the NRC, the FDA, and the

involved Agreement States in the regulation and authorization of the Omnitron 2000 HDR afterloader is poorly defined.

5. NRC guidance for scrap dealers needs to be given to waste brokers and collectors. Discovery of the lost iridium-192 source by BFI may have prevented additional significant radiation exposures; however, subsequent actions taken by BFI employees led to their receiving unnecessary exposures when they moved and searched for the source. Assistance from radiation protection experts should have been sought.

**Table A Summary of Doses Received as a Result of an
Iridium-192 Source Lost
from November 16 to December 1, 1992**

Dose Range (Sv)	Dose Range (rem)	Estimated Number of Individuals Exposed
0.0 - 0.005	0.0 - 0.5	42
0.005 - 0.01	0.5 - 1	11
0.01 - 0.05	1 - 5	20
0.05 - 0.1	5 - 10	13
0.1 - 0.15	10 - 15	7
0.15 - 0.20	15 - 20	1
TOTALS		94

Table B Summary of Collective Doses for All Locations

Location	Estimated Number of Individuals Exposed	Dose Range (mSv)	Collective Dose (Person-Sv)	Dose Range (rem)	Collective Dose (Person-rem)
IRCC Clinic	6	1.1 - 8.6	0.027 ^a	0.1 - 0.86	2.66 ^a
IRCC-Other ^b	17	0.08 - 13.9	0.009 - 0.026	0.01 - 1.4	0.94 - 2.6
Ambulance	2	4.8 - 25.7	0.018 - 0.034	0.5 - 2.6	1.8 - 3.4
SHM-Staff	21	3.6 - 223	1.1 - 1.7	0.4 - 22.3	110 - 166
SHM-Visitors	7	21.9 - 166	0.21 - 0.46	2.2 - 16.6	21 - 46
SHM-Residents	22	11.2 - 205	0.35 - 0.73	0.2 - 20.5	35 - 73
BFI-Carnegie	19	0.004- 90	0.08 - 0.22	0.0004 - 9.0	8.3 - 22
TOTALS	94		1.8 - 3.2		180 - 316

^a Established from film badge results

^b IRCC Nonoccupational workers and individuals from the adjacent physician's office

THE TEAM MEMBERS

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ABBREVIATIONS

ADO	adjacent doctor's office
AP	anterior-posterior
BFI	Browning-Ferris Industries
CAL	confirmatory action letter
CDRH	Center for Devices and Radiological Health, FDA
CFR	Code of Federal Regulations
CNA	certified nurse assistant
CTPS	Computerized Treatment Planning System
FDA	Food and Drug Administration, U.S.
GAO	Government Accounting Office, U.S.
GMP	Good Manufacturing Practices
GPCC	Greater Pittsburgh Cancer Center
GPN	graduate practical nurse
HDR	high dose rate
IIT	incident investigation team
IP	inspection procedure
IRCC	Indiana Regional Cancer Center
LED	light-emitting diode
LPN	licensed practical nurse
LS	lumbar-sacral
MC	manual chapter
NMSS	Nuclear Materials Safety and Safeguards (Office of)
NRC	Nuclear Regulatory Commission, U.S.
ORISE	Oak Ridge Institute for Sciences and Education
OSC	Oncology Services Corporation
QA	quality assurance
QC	quality control
QM	quality management

REAC/TS	Radiation Emergency Assistance Center/Training Site
RN	registered nurse
RSO	radiation safety officer
RTT	registered therapy technician
RTR	registered technologist radiographer
SHM	Scenery Hill Manor
S/N	serial number

1 INTRODUCTION

On November 16, 1992, an 82-year-old female cancer patient was undergoing radiation therapy at the Indiana Regional Cancer Center (IRCC) in Indiana, Pennsylvania (Figure 1), one of several operated by Oncology Services Corporation (OSC), an NRC licensee. The radiation therapy was to be administered by a high dose rate (HDR) afterloader with five connecting catheters. For that day's treatment, a dose of 6 Gy (600 rad) was to be administered through five catheters implanted as a single-plane perineal (rectal) implant encompassing the tumor. After a trial run through the five catheters with a dummy source, the iridium-192 source was placed in four catheters without difficulty. After several unsuccessful attempts to insert the source into the fifth catheter, the physician directed termination of the treatment. An area radiation monitor in the treatment room was observed in an alarm condition--flashing red light--at some point during the unsuccessful attempts to insert the source in the fifth catheter. Although three technologists and the physician were aware of the alarm, no one conducted a survey with the available portable survey meter to detect whether radioactivity was present. Believing that the area radiation monitor was malfunctioning, they reset the area radiation monitor and returned the patient to a local nursing home without performing any radiological surveys. The IRCC staff was unaware that the iridium-192 source had remained in the patient.

On December 1, 1992, the IRCC Medical Physicist notified NRC Region I that a $1.37 \text{ E}+11$ Bq (3.7-Ci) iridium-192 sealed source was missing from the licensee's HDR afterloader. The Medical Physicist believed that a radioactive source that was discovered by Browning-Ferris Industries (BFI) at their nonradioactive medical waste incinerator facility in Warren, Ohio, and later returned to another BFI facility in Carnegie, Pennsylvania (BFI-Carnegie), could be the same source that was missing from the HDR afterloader at IRCC. Later on December 1, 1992, the IRCC retrieved the source from BFI-Carnegie and placed it in a shielded container at their cancer center. NRC Region I inspectors went to the IRCC to ascertain the facts surrounding the loss of the iridium-192 source.

Region I inspectors determined that BFI had collected the source in the medical waste from the local nursing home where the patient had resided and later died on November 21, 1992. The location of the source between the patient's treatment and death and its subsequent transportation exposed a number of individuals to radiation. In addition to the patient, those exposed were certain members of the licensee's staff, the ambulance staff, the nursing home staff and residents, BFI personnel, and other members of the general public. Consequently, the NRC chartered an Incident Investigation Team (the team) to comprehensively review this incident.

On December 7, 1992, a similar source-wire break on an Omnitron 2000 afterloader occurred at the Greater Pittsburgh Cancer Center (GPCC). The team also examined the events associated with this second incident.

The team's report of these two incidents is organized into the following sections and appendices:

Section 2 presents a chronology of events for the two source-wire-break incidents involving an HDR afterloader.

Section 3 describes Omnitron International, Inc.'s, 2000 HDR Afterloader System, which was used to treat patients in both incidents.

Section 4 discusses the performance of equipment involved in the IRCC incident, including the HDR afterloader system and a fixed radiation monitor, PrimAlert-10.

Section 5 analyzes the human factors that could have contributed to these incidents, including the organization and performance of the licensee.

Section 6 presents the team's methodology for calculating estimated and actual doses of radiation exposures to the staffs of the cancer centers, the nursing home, visitors to any of these organizations, BFI, and the public. It also presents the assessment of Patient A's dose as determined by NRC's Medical Consultant. In addition, cytogenetic results for those believed to have had the greatest exposure to the iridium-192 source are presented.

Section 7 presents the team's findings about each involved organization's response to the incident.

Section 8 discusses which Federal or State regulatory agencies are responsible for the materials, devices, or procedures involved in these two incidents and evaluates NRC regulations and guidance.

Section 9 briefly describes any precursors to these incidents.

Finally, Section 10 presents the team's findings and conclusions for these two incidents.

Appendix A is a copy of the team's Charter for investigating these incidents; Appendix B presents the properties of the iridium-192 source; Appendix C describes both conventional and HDR brachytherapy; Appendix D is a copy of the licensee's, OSC's, NRC license; and Appendix E lists the interviews and meetings the team conducted.

The facts and data in this report are current as of January 27, 1993.

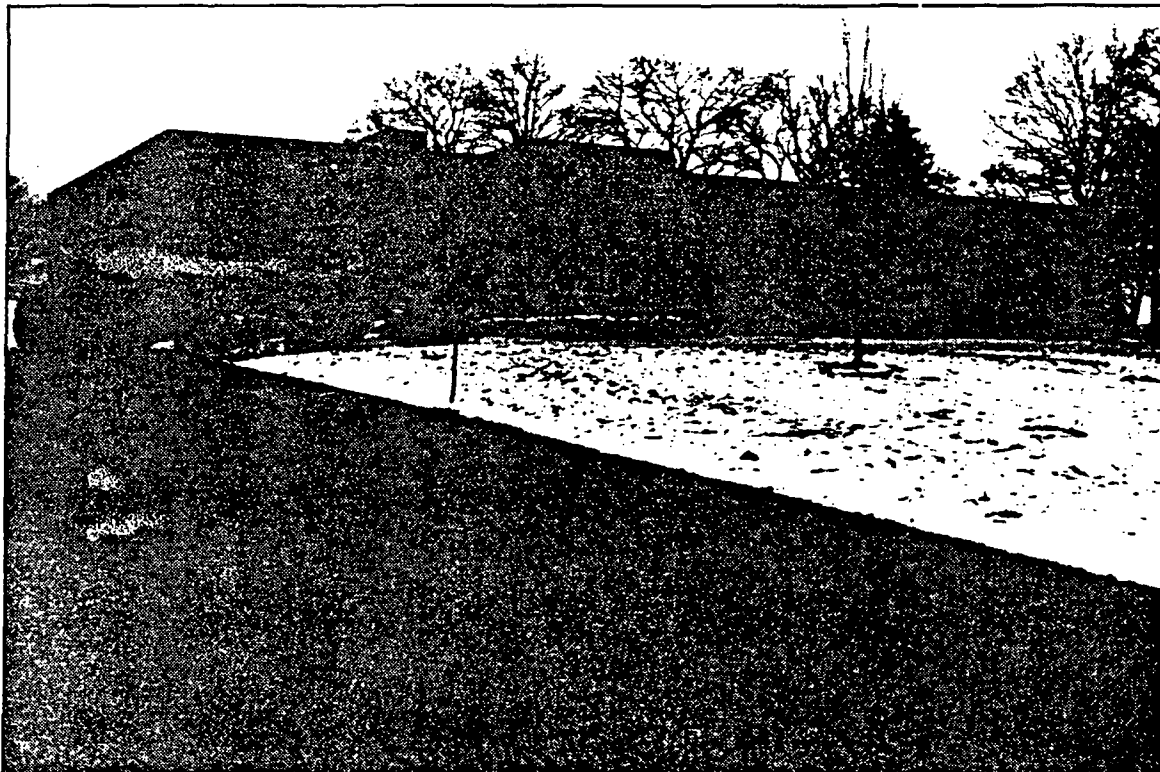


Figure 1.1 Indiana Regional Cancer Center, Indiana, Pennsylvania

2 SEQUENCE OF EVENTS

This section describes the patient treatment plan, the sequence of events associated with the brachytherapy misadministration that occurred at the Indiana Regional Cancer Center (IRCC) on November 16, 1992, the transportation of the patient from IRCC to Scenery Hill Manor (SHM) nursing home, and the subsequent chronology of events at the SHM nursing home. It also describes the sequence of events involving the loss of the iridium-192 source from IRCC and events associated with the transportation, discovery, and retrieval of the source from the Browning-Ferris Industries (BFI) facility in Carnegie, Pennsylvania (BFI-Carnegie). This section also describes events associated with the second Omnitron International, Inc., (Omnitron) 2000 high dose rate (HDR) afterloader source-wire break at another Oncology Services Corporation (OSC) cancer center in Pittsburgh, Pennsylvania.

2.1 Patient Treatment Plan

On November 16, 1992, an 82-year-old female patient was treated for anal carcinoma at the IRCC in Indiana, Pennsylvania, using HDR brachytherapy. The total dose was to be administered in three fractions of 6 Gy (600 rad) each. The HDR treatment plan included the interstitial insertion of the HDR iridium-192 source into five catheters that were strategically implanted in the treatment site. The catheters were implanted on November 13, 1992, and a localization study (simulation) was performed to ascertain the location of the five catheters within the patient. Catheter placement is further discussed in Sections 6.2 and 7.2.

On November 16, 1992, before the patient was treated, Medical Physicist A generated an Omnitron HDR dosimetry drawing with 40 dosimetric points (Figure 2.1). The calculated source strength upon initiating the treatment was $1.56 \text{ E}+11 \text{ Bq}$ (4.219 Ci). The dwell time calculations for the five implanted catheters were generated by computer (Figure 2.2). On November 16, 1992, at 9:35 a.m., the implanted catheters were connected to the HDR afterloader connecting catheters* and patient treatment was initiated. Difficulties were encountered during the source wire insertion into implanted Catheter 5, and Physician A directed termination of treatment without treating the patient through implanted Catheter 5. The patient was disconnected from the HDR afterloader (Figures 2.3 and 2.4). After the patient was removed from the treatment room, one loose catheter was removed, and the other four catheters were left in the patient for subsequent treatments.

On the basis of the licensee's assumption that the fifth catheter was not treated, Medical Physicist A generated, on November 17, 1992, a second Omnitron dosimetry plan, including different dosimetric drawings and dwell time for each location within each catheter. The licensee's dosimetric calculation, considering that no dose had been administered through the

* The connecting catheters are flexible tubes each with two connectors, one at each end. One end is connected to the HDR afterloader and the other end is connected to the implanted catheter (see Section 3.1.5 for further details).

fifth catheter, was 4.8 Gy (480 rad). The subsequent two fractions were adjusted to 6.6 Gy (660 rad) each to result in a total administered dose of 18 Gy (1800 rad).

2.2 Indiana Regional Cancer Center Incident

This section describes the sequence of events that pertain to the brachytherapy misadministration and accidental loss and transport of an approximate $1.56 \text{ E}+11 \text{ Bq}$ (4.2- Ci) HDR iridium-192 source from the IRCC to BFI, in Warren, Ohio, (BFI-Warren) between November 16 to December 1, 1992. Events preceding the actual loss of the source are also described to more fully explain the incident. Times are approximated.

Friday, November 13, 1992

At a local hospital, five catheters were surgically inserted adjacent to the patient's rectum to prepare for an HDR treatment. The patient was taken to the IRCC where a simulation study was performed to ascertain the location of the catheters. The patient was subsequently returned to the SHM nursing home. Catheter numbers in this chronology are those taken from the HDR afterloader error log given to NRC by Omnitron.

Monday, November 16, 1992

7:12 a.m.	The patient departed the nursing home in a local Citizens Ambulance Service ambulance to be transported to the IRCC.
7:30 a.m.	The patient arrived at the IRCC (Figure 2.5).
9:15 a.m.	Medical Physicist A completed the patient treatment plan.
9:20 a.m.	Physician A signed a printout of the patient treatment plan, indicating his approval of the plan.
9:30 a.m.	The patient was moved to the treatment room.
9:35 a.m.	Registered Therapy Technician A (RTT-A), in the presence of Medical Physicist A, connected the patient's five implanted catheters to the HDR afterloader connecting catheters.
9:40 a.m.	RTT-A successfully completed the insertion of the HDR dummy wire into the patient's five implanted catheters.
9:40 a.m.	Medical Physicist A left the IRCC for the day.
9:43 a.m.	RTT-A was present at the HDR afterloader computer console. Registered Technologist Radiographer (RTR) was observing the conduct

of patient treatment at the HDR computer console. The insertion of the HDR source wire into the patient's catheters was initiated by RTT-A. The computer log indicated the active source wire was successfully inserted into and retracted from the patient's implanted Catheters 1 through 4.

The following error messages were received on the HDR computer monitor for Channel 5:

- 9:59 a.m.
- Treatment halted due to Error Class 2: Error can be reset by console operator;
 - Error Code 1A: Active wire path constriction detected at 82.4 cm;
 - Treatment halted due to Error Class 2: Error can be reset by console operator;
 - Error Code 55: Console STOP pressed

RTT-A reset HDR computer error message. The following two messages were displayed on the HDR computer monitor:

- Treatment halted due to Error Class 2: Error can be reset by console operator;
- Error Code 6F: Attempt to treat with door open.

RTT-A entered the HDR treatment room and checked the implanted and connecting catheters. RTR entered the HDR treatment room to see to patient's needs. RTT-A and RTR then left the HDR treatment room. RTT-A reset HDR computer error message.

- 10:01 a.m.
- RTT-A attempted to insert the dummy wire into Catheter 5. The following two messages were displayed on the computer monitor:
- Treatment halted due to Error Class 2: Error can be reset by console operator;
 - Error Code 2A: Dummy wire path constriction detected at 90.1 cm; Dummy wire check on Channel 5 failed.

RTT-A reset HDR computer error message.

- 10:02 a.m.
- RTT-A attempted to insert the dummy wire into Catheter 5. RTR was observing the conduct of patient treatment at the HDR computer console. The following messages were displayed on the HDR computer monitor:

- Treatment halted due to Error Class 2: Error can be reset by console operator;
- Error Code 2A: Dummy wire path constriction detected at 90.0 cm; Dummy wire check on Channel 5 failed.

RTT-A reset HDR computer error message. RTT-A entered the HDR treatment room, disconnected and reconnected Catheter 5 in an attempt to remove constriction. RTR and RTT-B entered the HDR treatment room to assist RTT-A. The three individuals then left the HDR treatment room.

10:07 a.m.

RTT-A attempted to insert the dummy wire into Catheter 5. RTR was observing the conduct of patient treatment at the HDR computer console. The HDR computer monitor indicated the following:

- Treatment halted due to Error Class 2: Error can be reset by console operator;
- Error Code 2A: Dummy path constriction detected at 90.0 cm; Dummy wire check on Channel 5 failed.

RTT-A reset HDR Computer error message.

10:08 a.m.

RTT-A attempted to insert the dummy wire into Catheter 5. RTR was observing the conduct of patient treatment at HDR computer console. The HDR computer monitor indicated the following:

- Treatment halted due to Error Class 2: Error can be reset by console operator;
- Error Code 2A: Dummy wire path constriction detected at 90.1 cm; Dummy wire check on Channel 5 failed.

RTR noticed the PrimAlert-10 red light alarm flashing, and RTR informed RTT-A and RTT-B. RTT-A was concerned that the wire was still out of the HDR. RTR was instructed to inform Physician A. RTR informed Physician A, who was in his office in the IRCC, of a problem with inserting the source in Catheter 5. RTT-A observed that the indicator light on the HDR computer monitor was green, indicating that the source was "safe" (see Section 3). RTR returned to the HDR treatment room—the door to the room was open—and entered the room but did not walk all the way to the patient. RTR also informed Physician A that the PrimAlert-10 red light alarm was flashing. Physician A, RTT-A, and RTT-B entered the room while the PrimAlert-10 was flashing. Physician A and RTT-A examined the Catheter 5 connection at the HDR afterloader but observed no source

wire. RTT-A disconnected the implanted catheter that he believed to be implanted Catheter 5 from the HDR afterloader connecting Catheter 5, and both Physician A and RTT-A examined the catheter connection at the patient and observed no wire. RTT-B left the HDR treatment room. RTT-A reconnected Catheter 5 to the patient and to the HDR afterloader. Physician A and RTT-A left the HDR treatment room. Physician A directed RTT-A to again try treating through Catheter 5. RTT-A reset the HDR computer error message. Physician A returned to his office.

10:10 a.m.

RTT-A attempted to insert the dummy wire into Catheter 5. The following messages appeared on the monitor:

- Treatment halted due to Error Class 2: Error can be reset by console operator;
- Error Code 2A: Dummy wire path constriction detected at 90.1 cm; Dummy wire check on Channel 5 failed.

10:13 a.m.

RTT-A attempted to insert dummy wire into Catheter 5. RTR was observing the conduct of patient treatment at the HDR computer console. The HDR computer monitor indicated the following:

- Treatment halted due to Error Class 2: Error can be reset by console operator;
- Error Code 2A: Dummy wire path constriction detected at 90.0 cm; Dummy wire check on Channel 5 failed.

RTT-A reset the computer error message.

10:14 a.m.

RTT-A again attempted to insert the dummy wire into Catheter 5 but received the same computer error message previously received on the monitor.

10:16 a.m.

RTT-B informed Physician A of the failure to insert the dummy wire into Catheter 5, and Physician A directed RTT-A to discontinue the treatment.

10:20 a.m.

RTT-A disconnected the patient's implanted catheters from the HDR afterloader. RTT-A removed the patient from the HDR treatment room, and the patient was taken to the stretcher room (see Figure 2.5).

RTT-A unplugged and replugged the PrimAlert-10 power supply to reset the alarm sometime during the preceding events.

10:27 a.m. RTT-A informed Nurse A and Physician A that the stitches were loose on one of the patient's catheters, after which they went to the stretcher room to examine the patient.

10:28 a.m. The local ambulance was called to transport the patient from the clinic to the nursing home.

10:29 a.m. Physician A, assisted by Nurse A, removed the loose implanted catheter, which they assumed at that time to be Catheter 5. The local ambulance arrived at the clinic.

10:35 a.m. Patient A was transferred from the IRCC by ambulance to the SHM Nursing Home (Figure 2.6). IRCC Nurse A helped two ambulance assistants with placing the patient into the ambulance.

10:48 a.m. The ambulance arrived at the nursing home and Patient A was transferred to Room 4B.

10:56 a.m. The ambulance drivers left the nursing home.

Tuesday, November 17, 1992

3:00 p.m. Patient A requested that the second radiation therapy treatment scheduled for November 18 be canceled, owing to her inability to "tolerate the radiation therapy again." Staff at the SHM nursing home contacted the IRCC and rescheduled the treatment for Monday, November 23, 1992.

Thursday, November 19, 1992

7:00 p.m. Certified Nurse Assistant E (CNA-E) was performing perineum care and "removed a piece of gray-black tissue about 1-inch long that was stuck to one of the implants."

Friday, November 20, 1992

4:30 a.m. CNA-C noticed that one of the four remaining catheters (later determined to be the one containing the radioactive source) had become dislodged from the patient and protruded approximately 2.54 cm (1 inch) from the body.

6:15 a.m. During regular patient rounds, Licensed Practical Nurse B (LPN-B) and CNA-C discovered that the catheter that had been protruding earlier on the shift had fallen out of the patient onto the bedding.

LPN-B picked up the catheter containing the source, placed it in a red bag, which the nursing home uses for medical and biohazardous waste (typically referred to as "red-bag" waste), and transferred this small, red bag to a larger container for medical waste in the soiled utility room where red-bag waste is stored daily (Figure 2.7). LPN-B was unaware the catheter contained the iridium-192 source.

8:10 a.m. Registered Nurse-D (RN-D) called the IRCC to determine what method of disposal was needed for the catheter that had fallen out of the patient during the earlier shift. IRCC informed RN-D that what they had done was appropriate and that the catheter could be disposed of in the red-bag waste.

After discussing the situation with the IRCC, RN-D instructed Maintenance Man A to remove the large red bag of waste from the soiled utility room, which he normally does at 7:30 a.m., and transfer it to the outside waste storage room (see Figure 2.8).

8:30 a.m. Maintenance Man A took the large red bag of medical waste from inside the soiled utility room to the outside waste storage room, placed it inside a BFI cardboard box and locked the room.

The waste remained in this location for an additional 5 days, awaiting pickup by BFI from BFI-Carnegie, which usually occurs the last Wednesday of each month.

Saturday, November 21, 1992

11:10 p.m. Patient A dies. The remaining three catheters were subsequently removed and disposed of in red-bag waste.

Wednesday, November 25, 1992

4:30 a.m. BFI Driver A began picking up medical waste from the first of 22 stops for that day.

9:25 a.m. BFI Driver A arrived at his 12th stop, SHM Nursing home; picked up three boxes of red-bag waste from this facility; and placed it in a straight truck (Figure 2.9). Although BFI Driver A had a portable survey meter in the truck, he stated that he did not use it at the SHM nursing home. He continued with his regularly scheduled stops throughout the day, stopping at an additional 10 facilities before returning to BFI-Carnegie (Figure 2.10).

- 2:30 p.m. BFI Driver A arrived at BFI-Carnegie.
- 3:30 p.m. BFI Driver A unloaded all the boxes by himself from the straight truck onto BFI Trailer 808, which is a 14.6-meter (48-foot) trailer and left for the day. The box containing the source was one of the last ones to be loaded. The box was positioned in the rear left-hand corner of the trailer, approximately 1.8 meters (6 feet) off the floor (Figure 2.11).

Thursday, November 26, 1992

Because Thursday, November 26, 1992, was Thanksgiving Day, BFI scheduled no transfers to the BFI-Warren Medical Waste Incinerator, and the trailer remained on the site until November 27, 1992.

Friday, November 27, 1992

- 12:00 midnight BFI semitruck Driver B began his routes for the day.
- 6:15 a.m. BFI Driver B began a review of paperwork for his second tractor-trailer shipment (808) that day to BFI-Warren. As is customary, Driver B signed and dated all shipping manifests and checked the bottom and top latches at the back of the trailer before placing a padlock on the trailer.
- 6:30 a.m. BFI Driver B left BFI Carnegie via I-79, to I-680N, to I-80W to Hwy-46N, and on to Hwy-169N (Figure 2.12). He made no stops between Carnegie and Warren.
- 8:30 a.m. Driver B arrived at BFI-Warren, drove over to the unloading area (see Position A, Figure 2.13), disconnected Trailer 808, hooked up his cab to an empty trailer, drove to the front office, completed the paperwork for this shipment, and left BFI-Warren. Two fixed radiation monitors inside the facility alarmed, reading above their normal limits of 0.2 μSv (20 μrem) per hour. Employees working that shift began trying to locate the cause of the radiation alarms.
- 8:50 a.m. These employees notified both the Plant Manager and the Supervisor on the day shift who also began to attempt to find the source. They surveyed all packages on the conveyor belt (Figures 2.14 and 2.15) and those near the loading dock.

Because they could not determine the source of the radiation, they began reviewing their shipping records to determine what shipments they had received that morning. They identified two tractor-trailer

shipments, 806 and 808, that had arrived that morning at 4 a.m. and 8:30 a.m., respectively, from BFI-Carnegie.

After identifying the possible cause of the radiation alarms, both the Plant Manager and the Supervisor began to look for the source with their portable survey meters (Figure 2.16) outside of the parked trailers (Figure 2.17). When they came within approximately 121.9 meters (400 feet) of the trailer, both portable survey meters immediately alarmed and registered at their highest levels over 5 μSv (500 μrem) per hour.

9:45 a.m. Because the Plant Manager and the Supervisor could not determine which trailer had the radioactive material in it, they decided to drive one trailer at a time behind the main building, (see Figure 2.13), using it as a large concrete shield, to see if any of the portable survey meters alarmed as a trailer approached Position B in Figure 2.13.

Before the tractor-trailer drove behind the main building, the Plant Manager stood at Position C in Figure 2.13 of the facility with his portable survey meter turned on. At this location, there was no indication of radioactivity. As the tractor-trailer came from behind the building, however, the portable survey meter registered its highest level.

10:00 a.m. As soon as BFI-Warren identified the trailer containing the radioactive material, they parked it as far away on their property as they could from the main building and immediately called BFI-Carnegie to come pick up the trailer as soon as possible. The facility was fenced and secured.

2:30 p.m. BFI Driver C arrived at BFI-Warren to pick up the tractor-trailer from the Ohio facility, which was then driven to BFI-Carnegie. No radioactive material placards were placed on the tractor-trailer.

4:45 p.m. BFI Driver C arrived at BFI-Carnegie. Because it was growing late, the driver unhooked and parked the trailer in a back lot where it remained for the rest of the weekend in a fenced, secured area (Figure 2.18).

Monday, November 30, 1992

1:00 p.m. BFI-Carnegie Supervisor A and two safety assistants (Safety Technicians A and B) put gloves on and began to survey boxes from the trailer for radioactivity. When approaching the trailer, each safety

technician noticed that the portable survey meter registered its highest level [5 μ Sv (500 μ rem) per hr].

- 2:30 p.m. After surveying approximately 40 boxes, they identified the box containing the radioactive material (Figure 2.19). Because the box containing the source had no generator identification labels on the outside, Safety Technicians A and B opened the box and began to go through individual medical red bags looking for information to enable them to identify the originator of the waste. Supervisor A left for an appointment offsite but requested that the safety technicians continue to search for the originator's identification.
- 3:00 p.m. One of the two safety technicians found a portion of a prescription in the waste that had an individual's name on it. With this information, they sealed up the box, placed it in a green recycle container (Figures 2.20 and 2.21), and locked it. They returned to their office with this information and began calling the list of facilities from which waste was picked up on November 25, 1992.
- 4:00 p.m. Safety Technician B began calling hospitals and nursing homes to see if any of these facilities could recognize the name they found.
- 5:00 p.m. After contacting approximately 15 facilities to identify the location of the individual on the prescription without success, they stopped for the day and went home.

Tuesday, December 1, 1992

- 8:30 a.m. Supervisor A reopened the trailer containing the other medical waste boxes and began to look for the other two containers that had arrived with the box containing the radioactive material.
- 9:30 a.m. Supervisor A and Safety Technicians A and B again went through the red bags in these two containers trying to find identifying information. This time, they were successful and found an individual's name associated with the SHM nursing home.
- 9:45 a.m. Immediately, Supervisor A called the SHM nursing home to inform them that radioactive material had been discovered in the waste that they had picked up from the home on November 25, 1992. The SHM nursing home staff informed BFI-Carnegie that they did not have any radioactive material at their facility, but they did identify a resident (Patient A) that had recently undergone cancer treatment therapy at IRCC.

10:00 a.m.	The staff at the SHM nursing home immediately called the IRCC and spoke with Nurse A, who contacted Physician A. Physician A suspected a possible source loss.
11:00 a.m.	Nurse A called Medical Physicist A in Johnstown, Pennsylvania, to notify him of the source loss. During the telephone conversation, Medical Physicist A asked RTT-B to evacuate the treatment vault, use remote control to extend the iridium-192 source into a connecting catheter, and observe the PrimAlert-10 radiation monitor to verify the presence of a radiation reading. RTT-B informed Medical Physicist A that the PrimAlert-10 did not detect any radiation levels.
11:40 a.m.	Medical Physicist A arrived at the IRCC and verified the absence of the iridium-192 source by performing an autoradiograph of the source wire, monitoring the PrimAlert-10, and by performing portable survey meter measurements.
11:44 a.m.	Medical Physicist A called BFI-Carnegie to inform them that they would arrive shortly to retrieve the radioactive material.
11:45 a.m.	Medical Physicist A notified OSC's Radiation Safety Officer (RSO) in Harrisburg, Pennsylvania.
11:50 a.m.	Medical Physicist A notified NRC, Region I.
3:15 p.m.	<p>Medical Physicist A and Physician A, arriving in separate vehicles at BFI-Carnegie, were met by Safety Technician A, who had previously supervised and participated in the unloading, identification, and subsequent isolation of the BFI box containing the radioactive material.</p> <p>Upon approaching the green recycle container (Figure 2.22), Safety Technician A unlocked it and stood approximately 2 meters (6.6 feet) away with a portable survey meter turned on. Medical Physicist A also had a portable survey meter and noted that the radiation reading at about 1.5 to 1.8 meters (5 to 6 feet) away from the container was above 7.8 mSv (780 mrem) per hour.</p> <p>Medical Physicist A placed the lead container in which the iridium-192 source was originally shipped on the ground next to the green recycle container. Wearing surgical gloves, Medical Physicist A lifted the box containing the source out of the recycle container, and both Medical Physicist A and Physician A opened the box and began taking plastic bags out. These plastic red bags were removed one at a time, and carried toward Medical Physicist A's portable survey meter located</p>

approximately 4.6 meters (15 feet) away. The first two bags contained no radioactive material. The third bag contained radioactive material. Using long-handled [about 30 cm (11.8 inches)] forceps, Medical Physicist A opened the red bag and saw several smaller red bags inside. One contained three catheters and one contained a single catheter. The bag with the single catheter was surveyed and indicated the presence of radioactivity. Medical Physicist A quickly walked to the lead container and placed the single catheter containing the source inside.

Medical Physicist A estimated that it had taken approximately 70 seconds from the time they located the red bag containing the source to the time it took to secure the source in its protective shield.

After securing the source, Medical Physicist A placed the container inside his truck and secured its movement by placing rubber-covered sandbags around it. Medical Physicist A surveyed the source container; it read 0.35 mSv (35 mrem) per hour. In addition, he surveyed the cab of his truck and obtained a reading of 6 μ Sv (0.6 mrem) per hour.

- 3:45 p.m. Once the source was secured in place, Medical Physicist A transported it back to the IRCC for storage. No radioactive material placards were placed on the vehicle.
- 5:10 p.m. Medical Physicist A arrived at the IRCC and placed the container with the iridium-192 source in the Treatment Room.
- 5:12 p.m. Medical Physicist A surveyed the source with his survey instrument and obtained a reading of 0.35 mSv (35 mrem) per hour.

2.3 Greater Pittsburgh Cancer Center Incident

On Monday, December 7, 1992, Medical Physicist B, from the Greater Pittsburgh Cancer Center (GPCC), reported that a 1.28 E+11 Bq (3.45-Ci) iridium-192 sealed source apparently broke off from the end of the source wire while being removed from a patient following a completed endobronchial HDR treatment. The system being used was an Omnitron 2000 HDR afterloader identical to the afterloader involved in the IRCC incident. Further, GPCC was operated by OSC, the same licensee that operated the IRCC at which the November 16, 1992, incident occurred.

The following is a chronology of events at the GPCC. Times are approximated.

Monday, December, 7, 1992

11:30 a.m. Medical Physicist B acquired keys from a locked cabinet and performed daily operational and safety checks on the HDR afterloader. These checks included positional accuracy (via x-ray films); emergency stops; functionality of the treatment room door and door interlock; PrimAlert-10 operability; afterloader and console indicator lights; and a survey of the treatment room, using a portable survey meter.

12:00 noon Operational and safety checks were completed.

2:30 p.m. Patient B arrived at the GPCC.

The GPCC staff made simulation films to determine the correct extent of the treatment volume. The catheter path was determined to be clear by inserting a measurement wire.

3:10 p.m. Physician E approved the treatment plan.

Patient B was moved to the treatment room and connected to the HDR afterloader. To reduce exposure to the source, Physicist B reduced the travel distance outside of the patient to 93.3 cm (36.7 inches), and the connecting catheter was secured by taping it to a 15.2 cm by 15.2 cm by 7.6 cm (6 inch x 6 inch x 3 inch) foam rubber pad, which was then taped to the patient's left arm.

3:45 p.m. Treatment commenced. Dummy wire verified that the catheter was clear.

3:50 p.m. Medical Physicist B initiated the HDR treatment.

Patient B stirred slightly, coughed several times, and moved left arm slightly to reach for a paper tissue.

Medical Physicist B noted that the treatment was completed and that the source wire was retracting.

3:52 p.m. The staff heard an audible alarm at the HDR afterloader and its console. Also, a visual alarm indicated on the console: "Emergency Condition, Manual Retract, Check Source Status."

Medical Physicist B entered the treatment room carrying a portable survey meter and noted that the PrimAlert-10 was alarming, which indicated that the source had not retracted into the HDR afterloader.

Medical Physicist B also noted that the portable survey meter was pegged high on the X1 scale, further indicating that the source had not fully retracted.

Medical Physicist B noted that Patient B had raised left arm, which produced an approximate 45-degree angle in the catheter at the quick disconnect nearest the HDR afterloader. Concentrating his attention on that quick disconnect, Medical Physicist B observed "a piece of wire" in the catheter.

Assuming that this was the source wire, Medical Physicist B exited the treatment room, acquired a pair of scissors, reentered the treatment room, and cut the catheter approximately 20 to 30 cm (7.9 inches to 11.8 inches) from the quick disconnect. Medical Physicist B then removed the patient from the treatment room, assisted by Physician E.

Medical Physicist B and Physician E surveyed Patient B, using the same portable survey meter to verify that the source had not remained with the patient.

4:00 p.m.

Having obtained 10-inch (25.4-cm) forceps, Medical Physicist B, accompanied by RTT-C with the portable survey meter, reentered the treatment room to secure the source.

Using the forceps to hold the source, Medical Physicist B disconnected the remaining portion catheter from the HDR afterloader and then placed the source and catheter into the lead container previously stationed next to the HDR afterloader. The PrimAlert-10 stopped flashing red, and RTT-C noted that the survey meter also responded by indicating a drop in the radiation level.

Medical Physicist B placed the lead container in a shipping container, and stored it in a small room behind the treatment room. Medical Physicist B then surveyed the treatment room to verify that no further radioactivity existed.

4:05 p.m.

Medical Physicist B and RTT-C exited the treatment room, and the physicist, assisted by Physician E, once again surveyed Patient B and the implanted catheter that the physician had removed. They found no radioactivity.

4:24 p.m.

Medical Physicist B called Omnitron to notify their Senior Vice President of the incident.

4:50 p.m. Medical Physicist B notified the OSC RSO.

5:30 p.m. Medical Physicist B notified the NRC.

Calc Date: 08:09:22 November 18, 1992

Depth of cut (cm): 0.7

Angles from AP Plane:

X: 42.3 Y: 24.6 Z: 41.6

Matrix size (CM): 16.9

Plot Scale: 1.00

Site: Rectal

Comment:

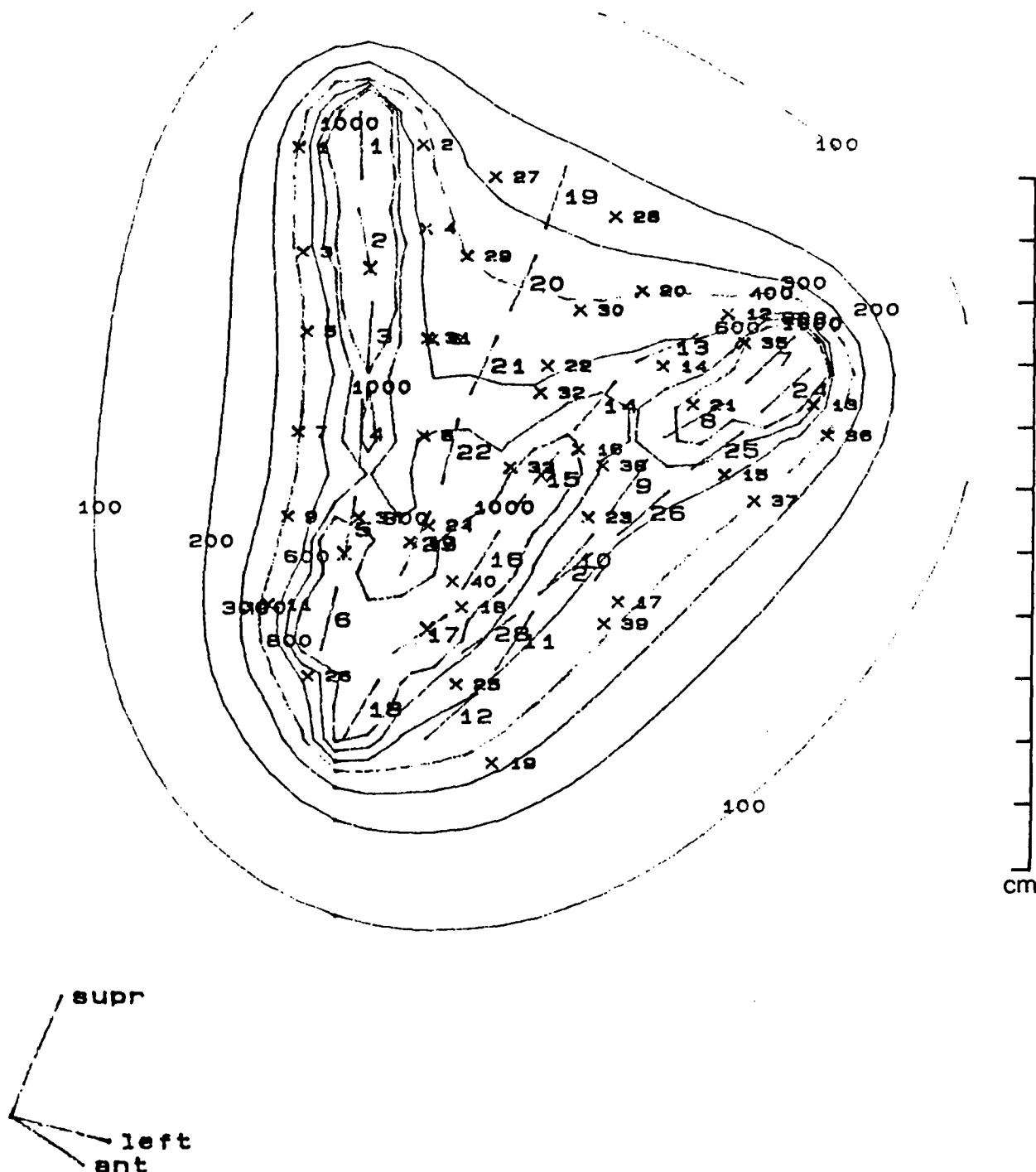


Figure 2.1 Isodose Graph for Initial High Dose Rate Patient Treatment Plan

Time of printout: 09:15:26 November 16, 1992
 Time of plan: 09:09:22 November 16, 1992

Age: 82
 Sex: F

The source 01-01-9282-001-082892-08942-08 was used in these calculations
 The original calibration was: 8.942 Curies
 The date of original calibration was: Friday August 28, 1992
 The date used for these calculations was: Monday November 16, 1992
 The half life used was: 73.83 days
 The decay factor is: 0.4719
 The present strength is: 4.219 Curies
 The source description is: Omnitron
 The sourcetype is: 1
 Calculations are for a dose fraction 600.0 cGy

Total Doses to calculation points (cGy):

1	403.5	15	846.4	29	464.3
2	443.1	16	672.8	30	500.0
3	449.0	17	650.5	31	534.6
4	523.4	18	827.0	32	656.6
5	415.9	19	446.0	33	821.8
6	543.4	20	444.2	34	794.6
7	402.6	21	591.5	35	867.1
8	655.1	22	633.9	36	388.6
9	398.1	23	719.8	37	457.2
10	853.8	24	830.8	38	818.6
11	445.3	25	663.1	39	604.6
12	471.9	26	694.9	40	799.2
13	823.1	27	417.6		
14	610.8	28	415.2		

Mean dose: 600.0 cGy
 Standard deviation: 162.6
 per cent standard deviation: 27.1

Dwell times:

Position	Cath#	Distance	seconds
1	1	90.0	49.2
2	1	88.4	42.9
3	1	86.8	26.8
4	1	85.2	26.8
5	1	83.6	16.5
6	1	82.0	45.9
7	2	90.0	24.7
8	2	88.4	25.4
9	2	86.8	16.3
10	2	85.2	26.8
11	2	83.6	29.0
12	2	82.0	45.7
13	3	90.0	28.3
14	3	88.4	26.8
15	3	86.8	22.6
16	3	85.2	26.8
17	3	83.6	26.8
18	3	82.0	29.0
19	4	90.0	45.9
20	4	88.4	26.8
21	4	86.8	27.6
22	4	85.2	22.6
23	4	83.6	8.5
24	5	90.0	29.1
25	5	88.4	28.3
26	5	86.8	26.8
27	5	85.2	29.2
28	5	83.6	15.5

Catheter
 LONDRING

3 - ○
 ○ - 1
 4 - ○
 ○ - 2
 5 - ○

Physician approval signed Physics review initialed
 The data presented by this plan must NOT be used clinically without approval of a qualified person

Figure 2.2 Summary of Initial High Dose rate Treatment Plan

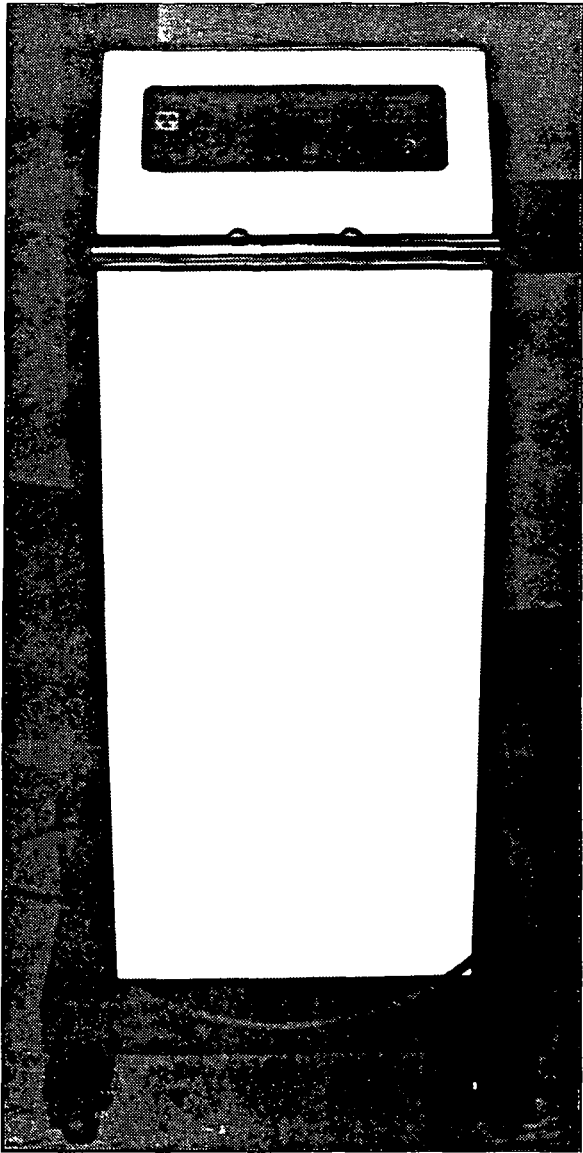


Figure 2.3 Back of High Dose Rate Afterloader

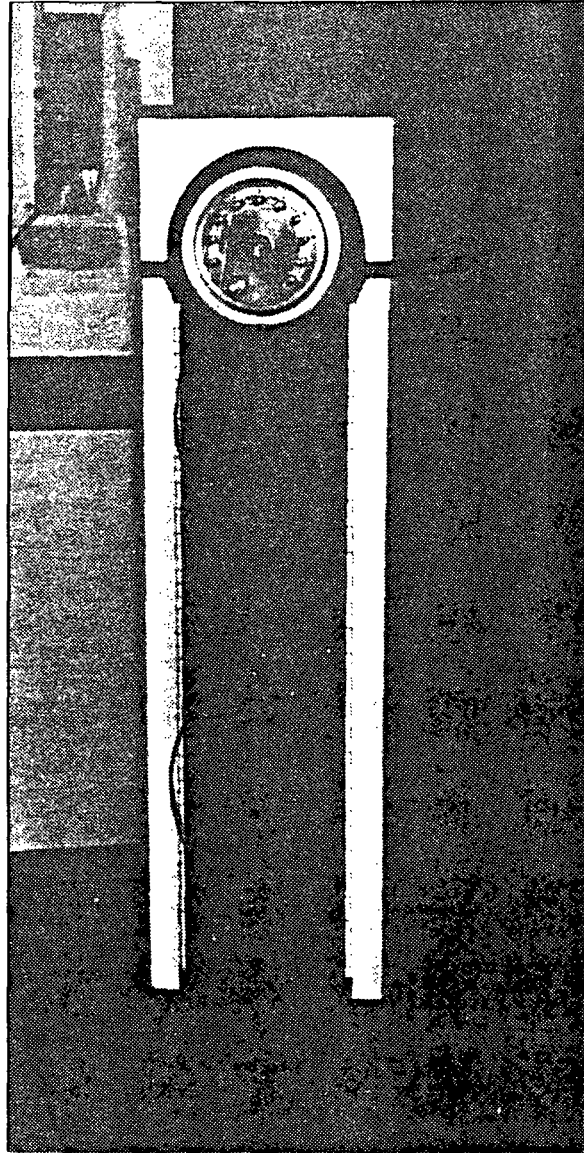


Figure 2.4 Front of High Dose Rate Afterloader

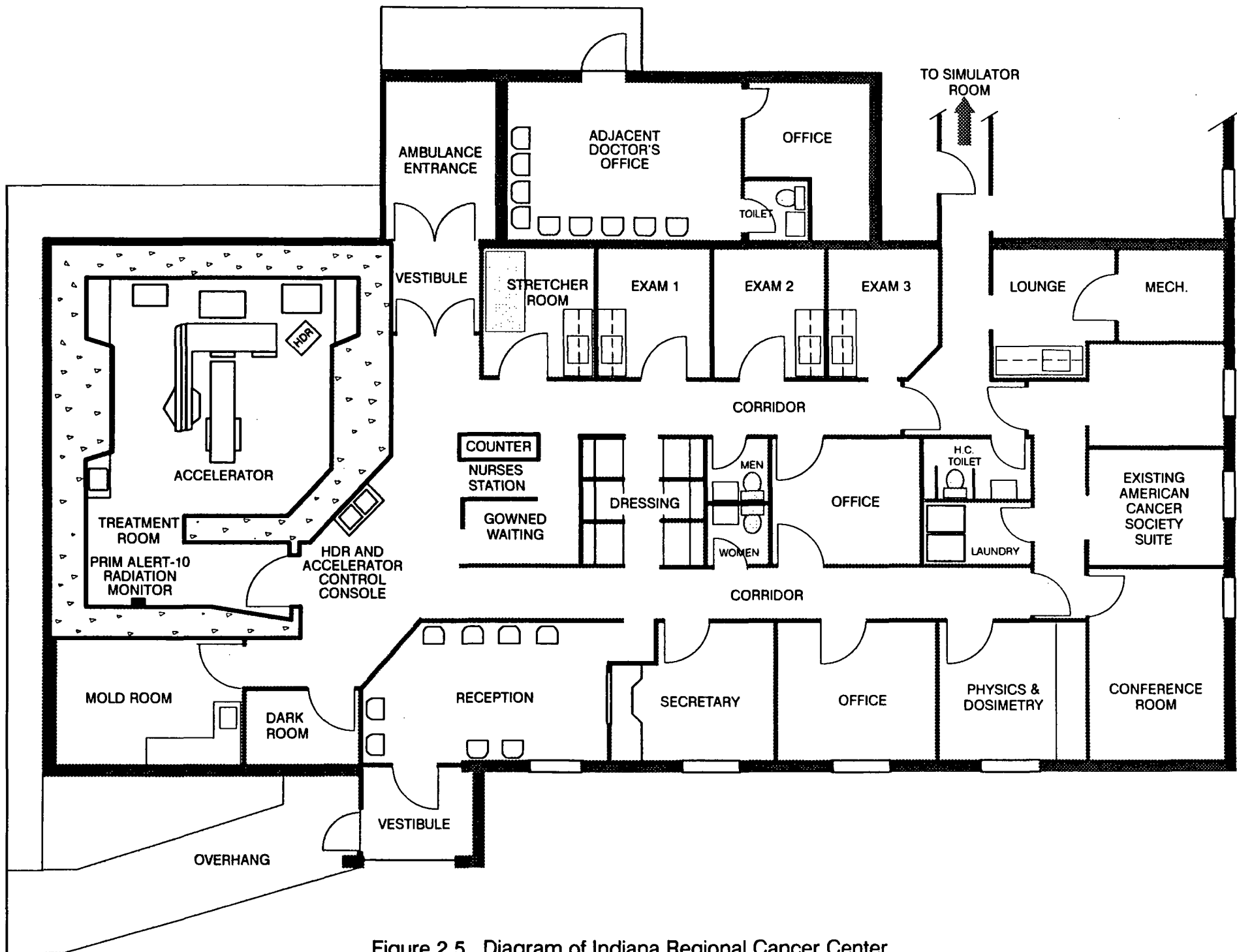


Figure 2.5 Diagram of Indiana Regional Cancer Center

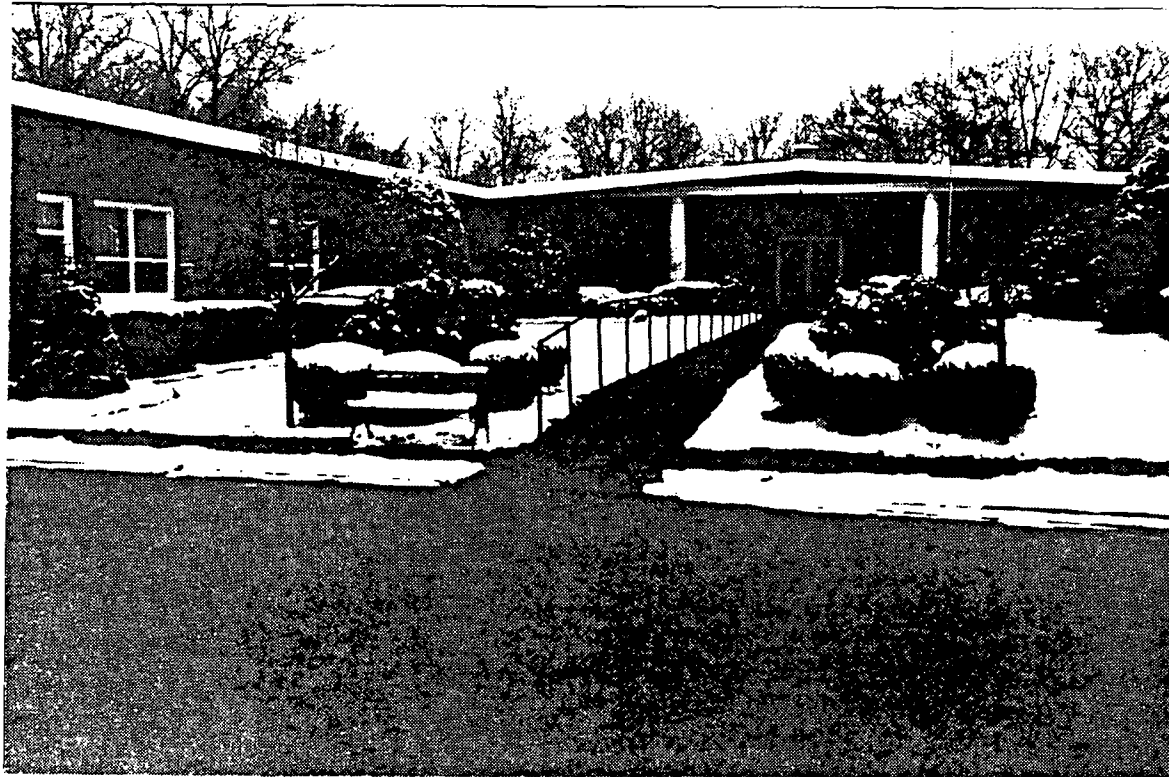


Figure 2.6 Front of Scenery Hill Manor Nursing Home

Red-Bag Waste
Containing Iridium-192
Source on 11/20/92

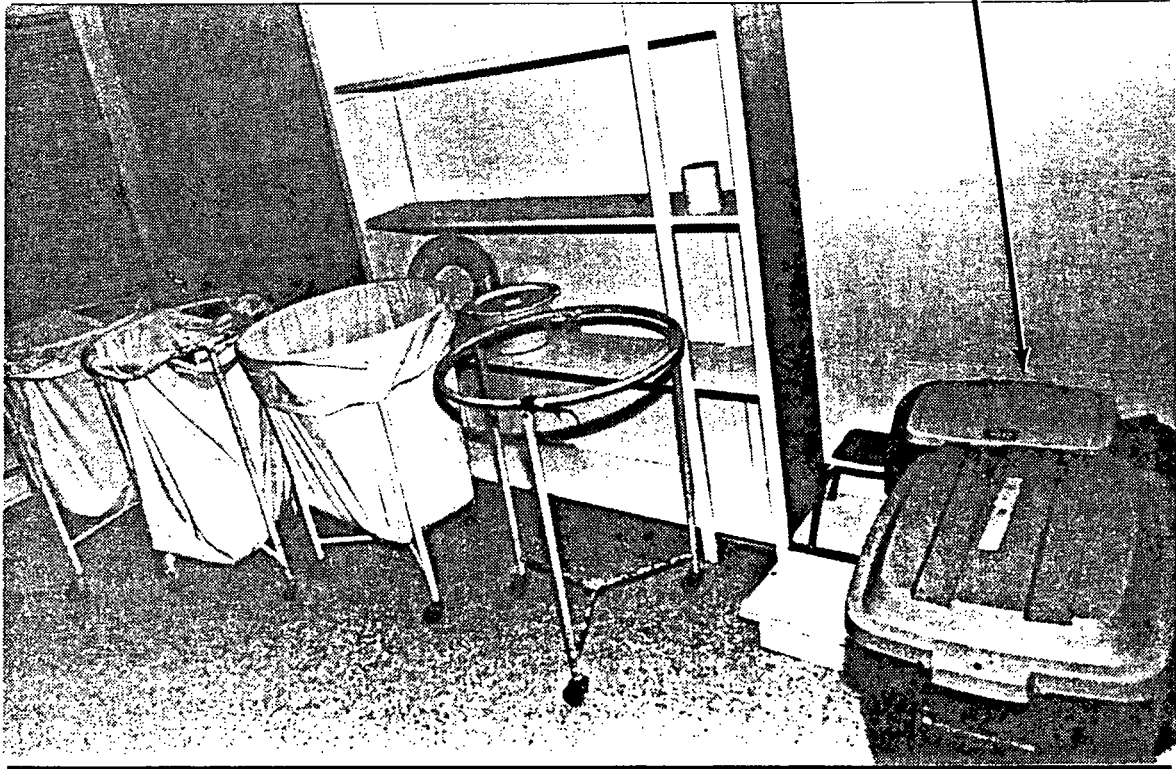


Figure 2.7 Soiled Utility Room with Biohazard Waste Container

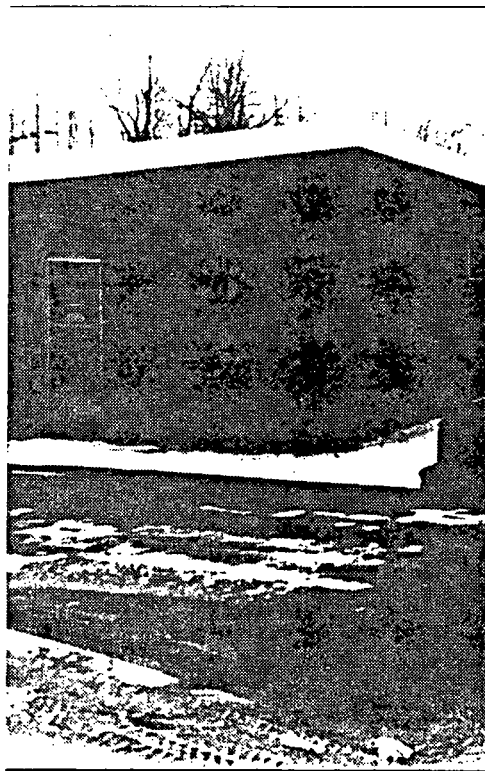


Figure 2.8 Waste Storage Room at Scenery Hill Manor Nursing Home

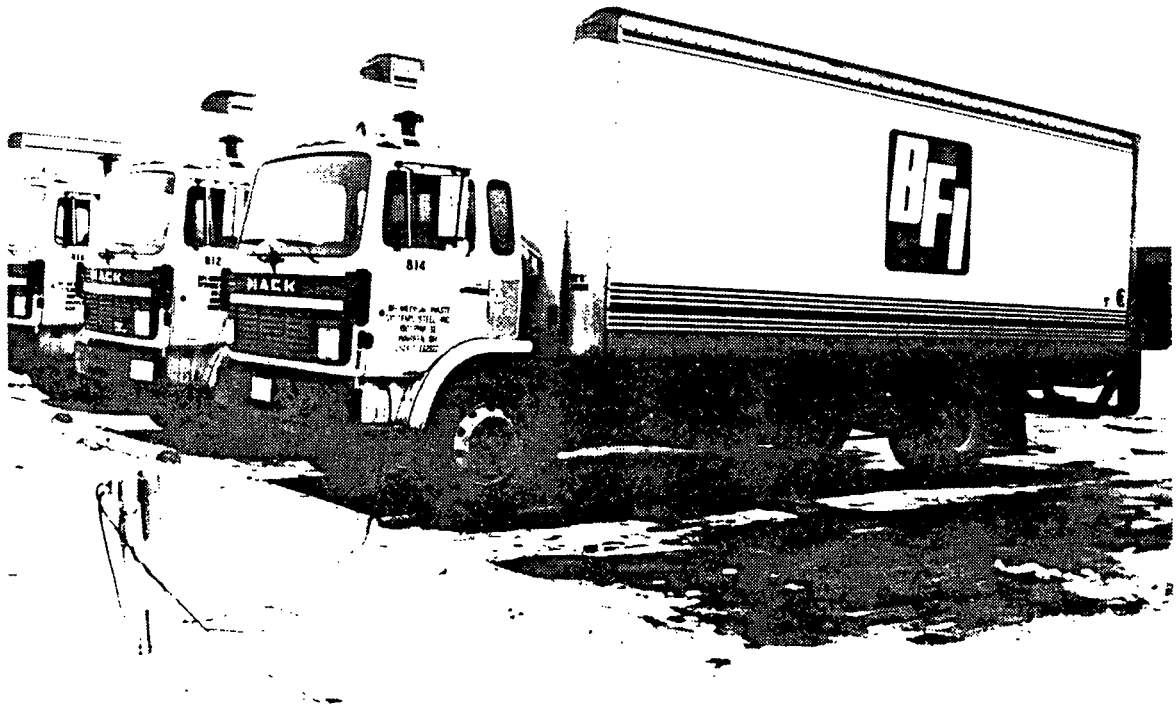


Figure 2.9 Browning - Ferris Industries Straight Truck Used to Collect Waste from Scenery Hill Manor Nursing Home

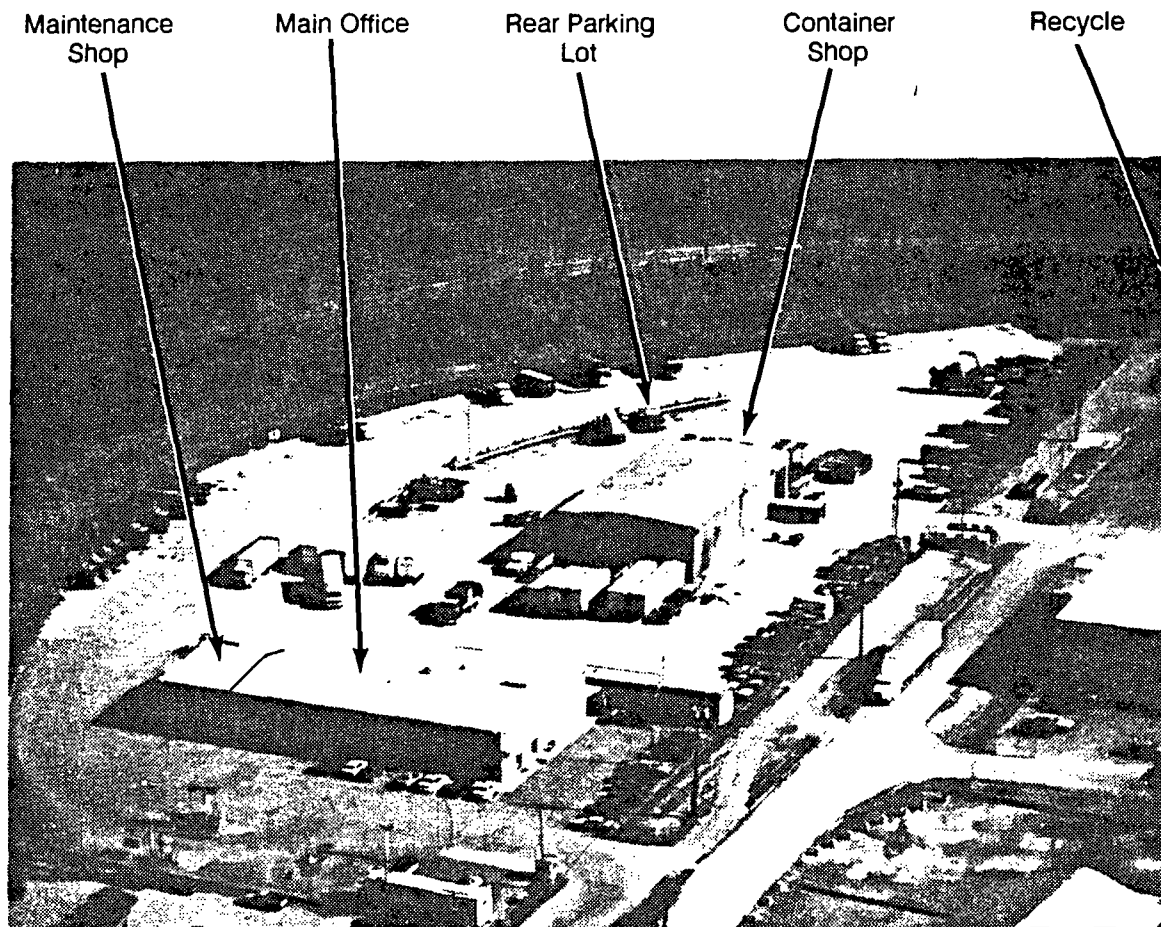


Figure 2.10 Aerial Photograph of Browning - Ferris Industries Transfer Station in Carnegie, Pennsylvania

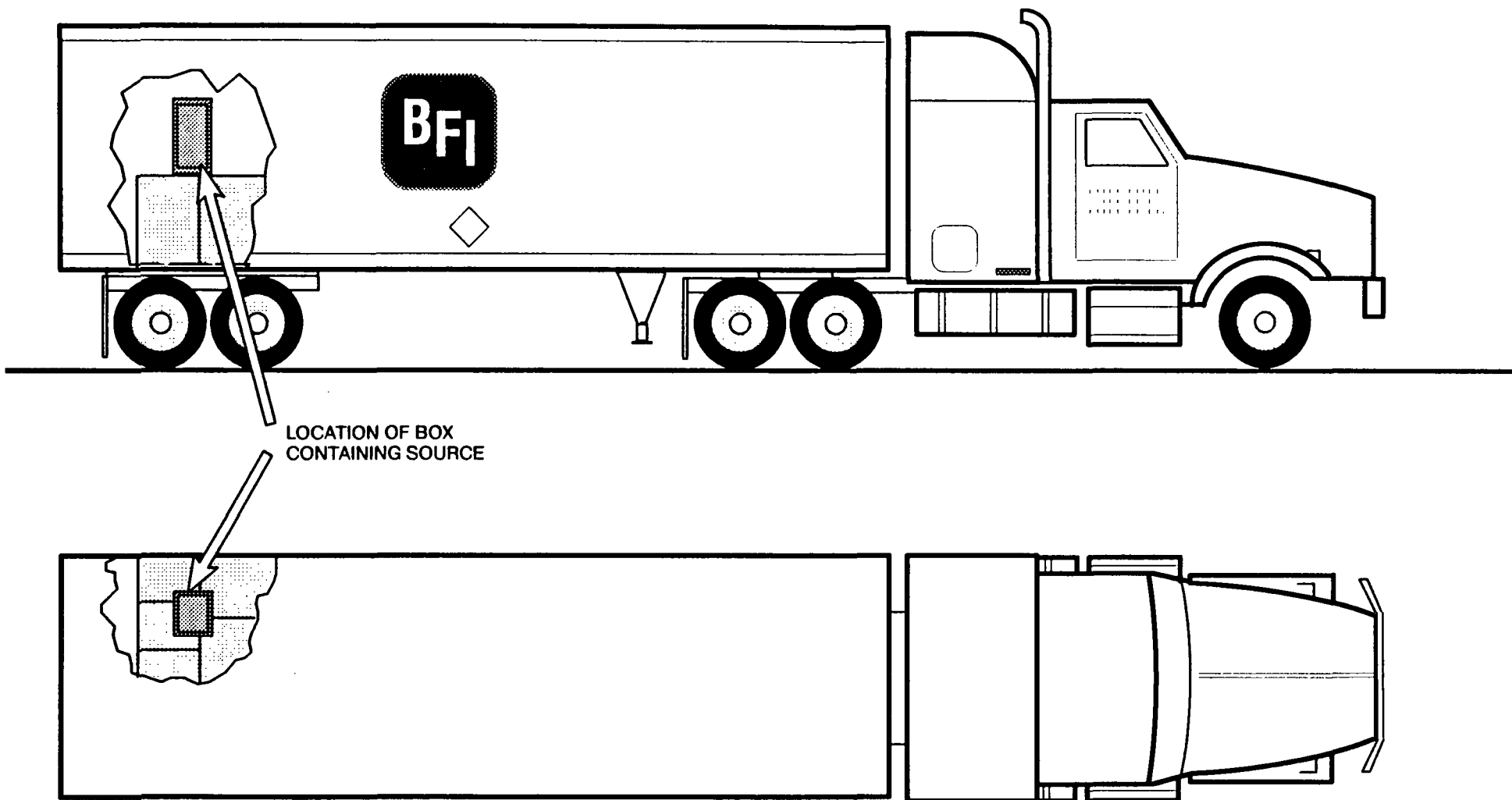


Fig. 2.11 BFI Tractor Trailer 14.63 m (48') Long Showing Location of Source Container in the Trailer.

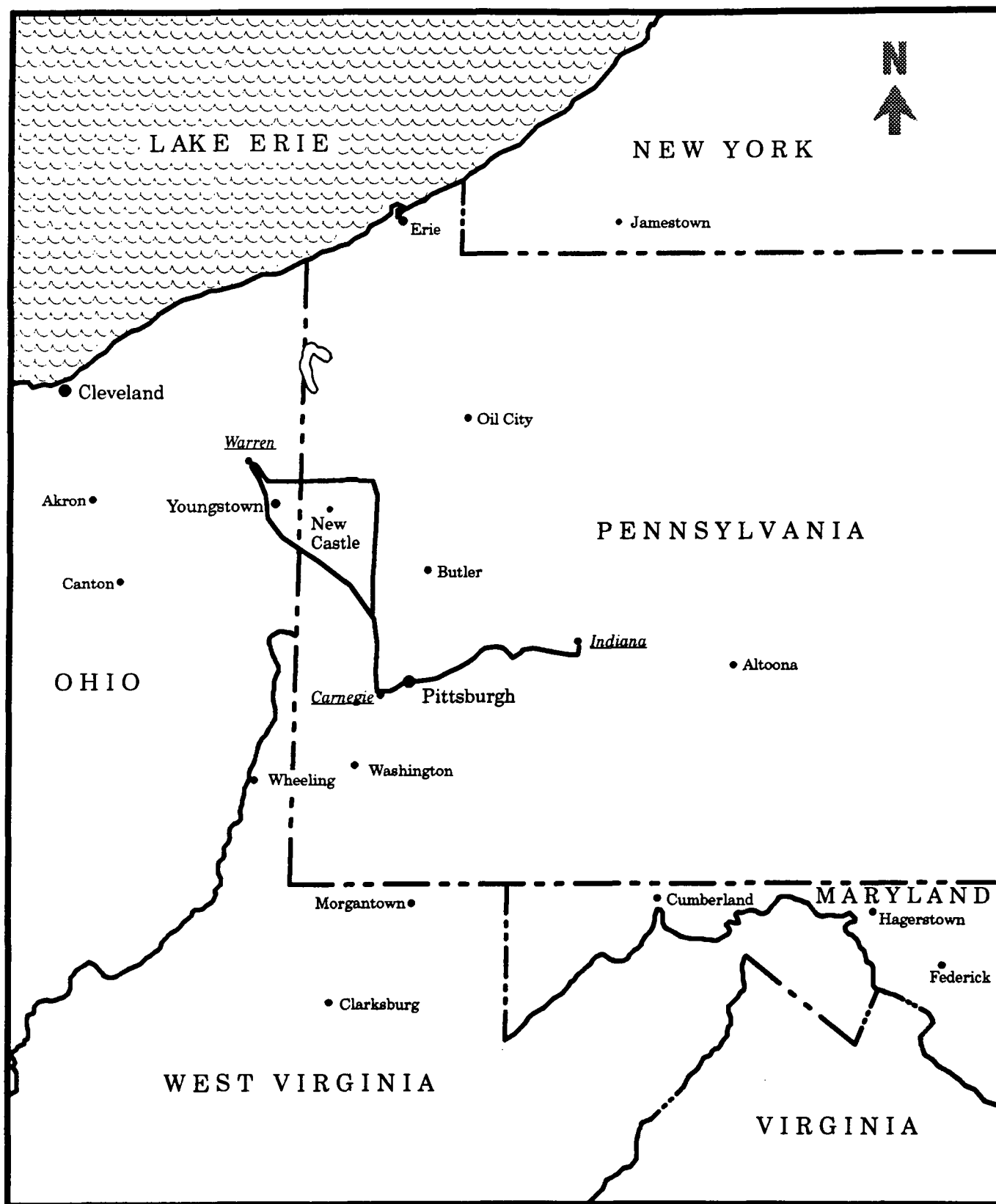


Figure 2.12 Route of Browning - Ferris Industries Semitruck 808 on November 25, 1992

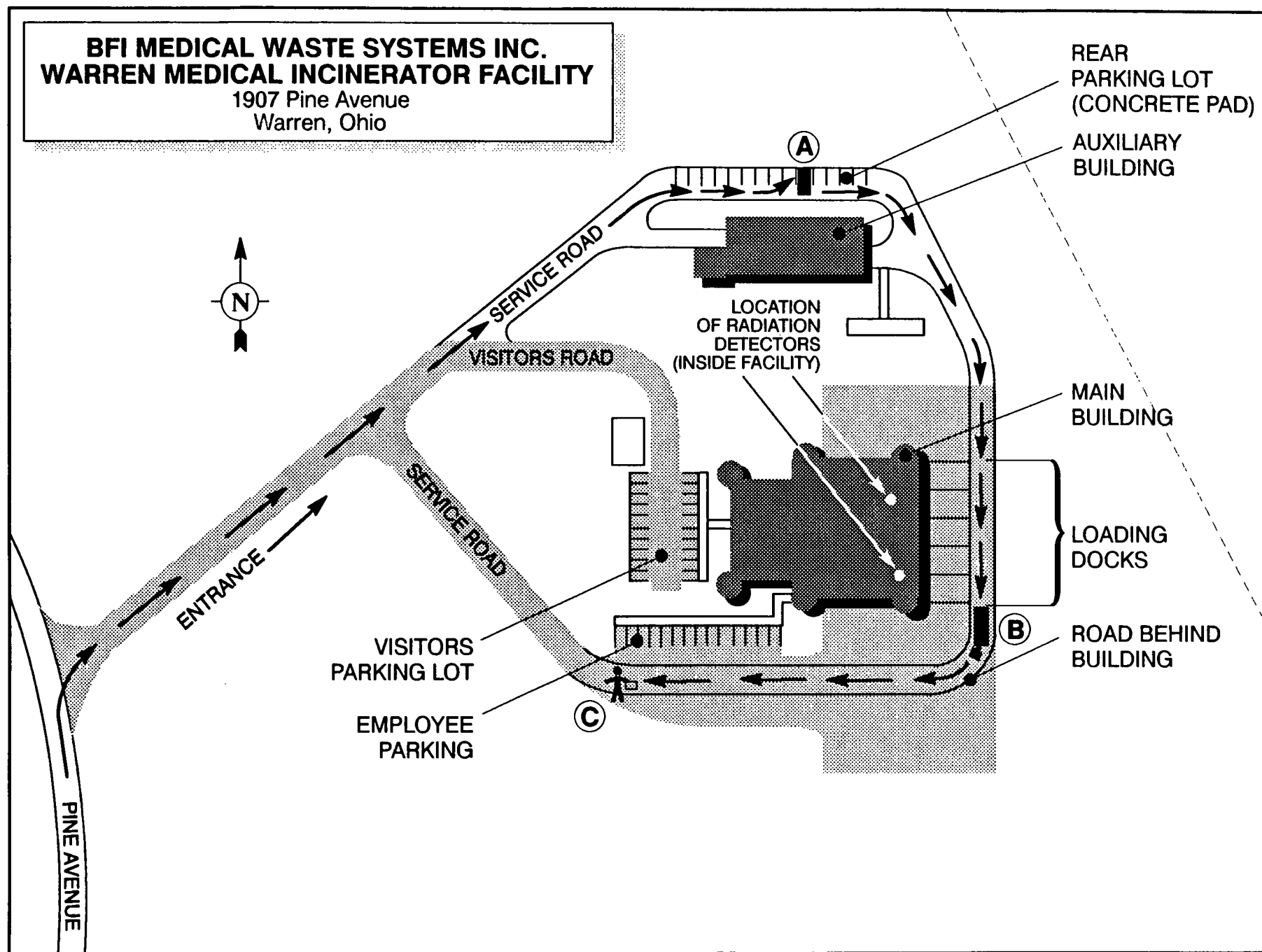


Figure 2.13 Diagram of Tractor-Trailer Route Carrying Iridium-192 Source at BFI-Warren on November 25, 1992

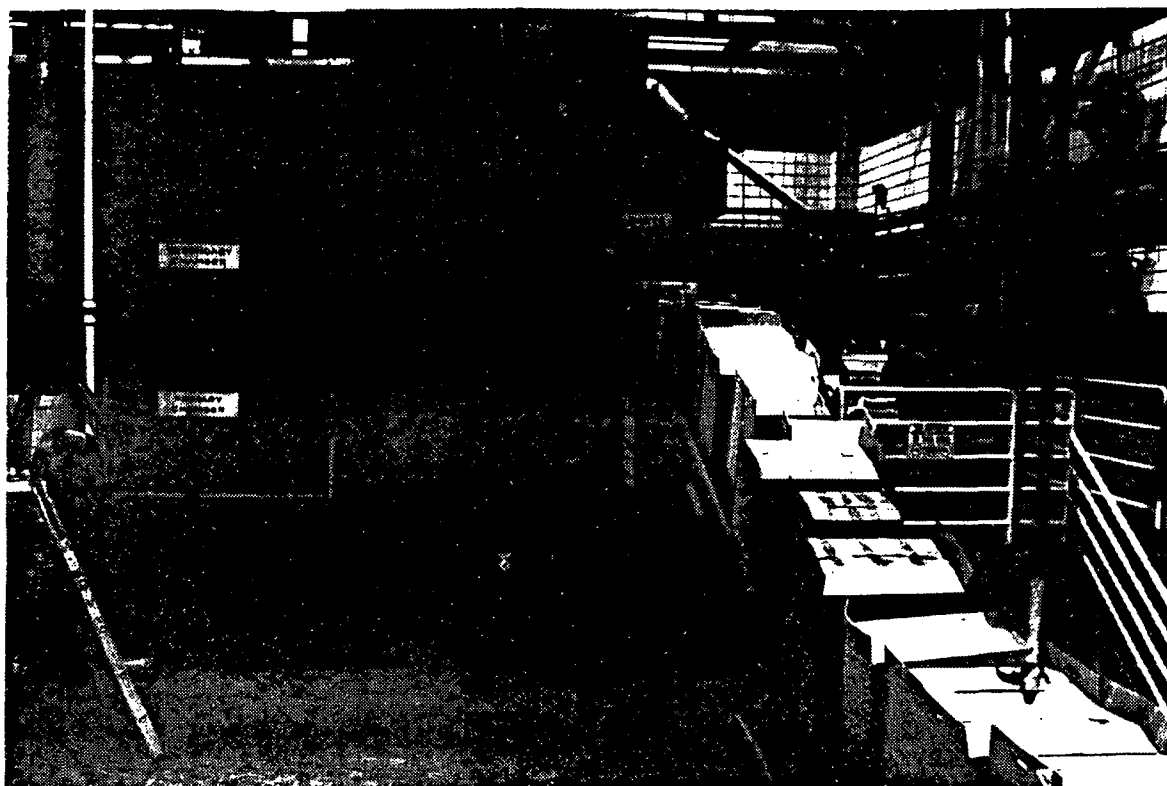


Figure 2.14 Boxes on Conveyor Feedline Adjacent to Radiation Monitor



Figure 2.15 Console for Monitoring Radiation at Incinerator Conveyor Feedline

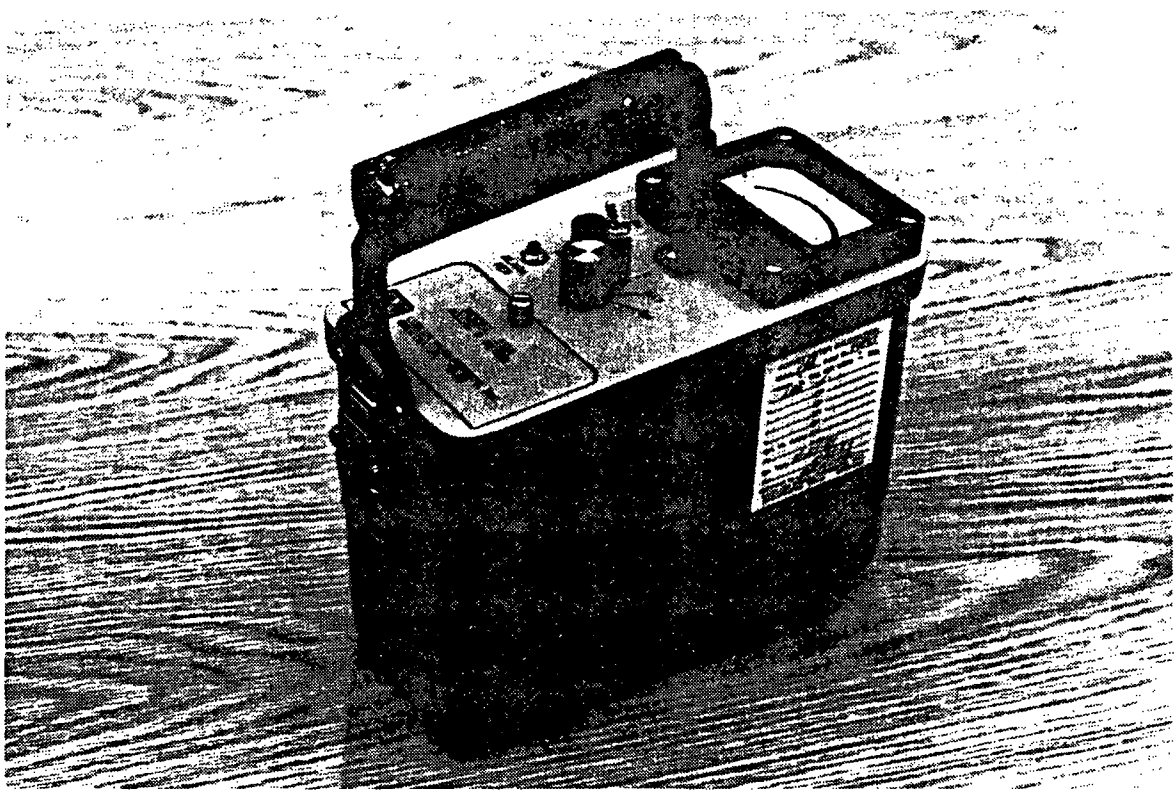


Figure 2.16 Survey Meter Used at BFI - Warren



Figure 2.17 Location of Where Manager and Supervisor Detected Radiation (~ 400')
from BFI Trailer 808

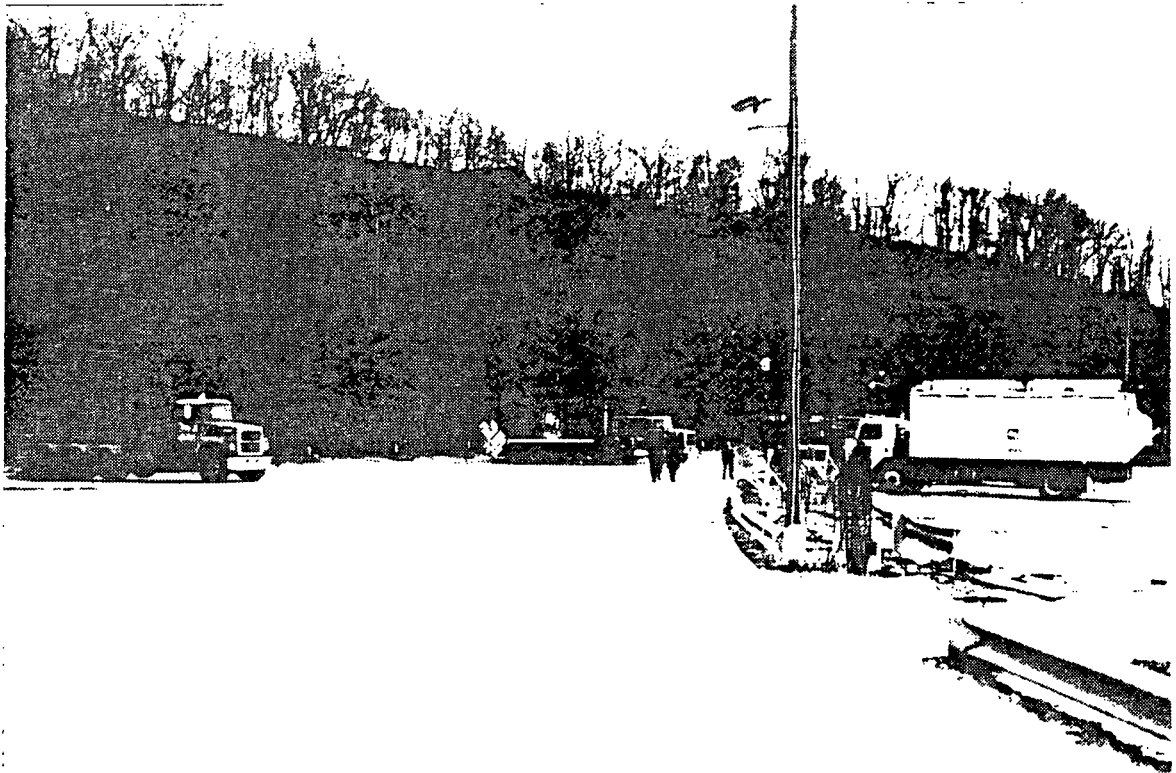


Figure 2.18 BFI Truck Parking Lot, Carnegie, Pennsylvania



Figure 2.19 Location of Isolated Biohazard Box Containing Source

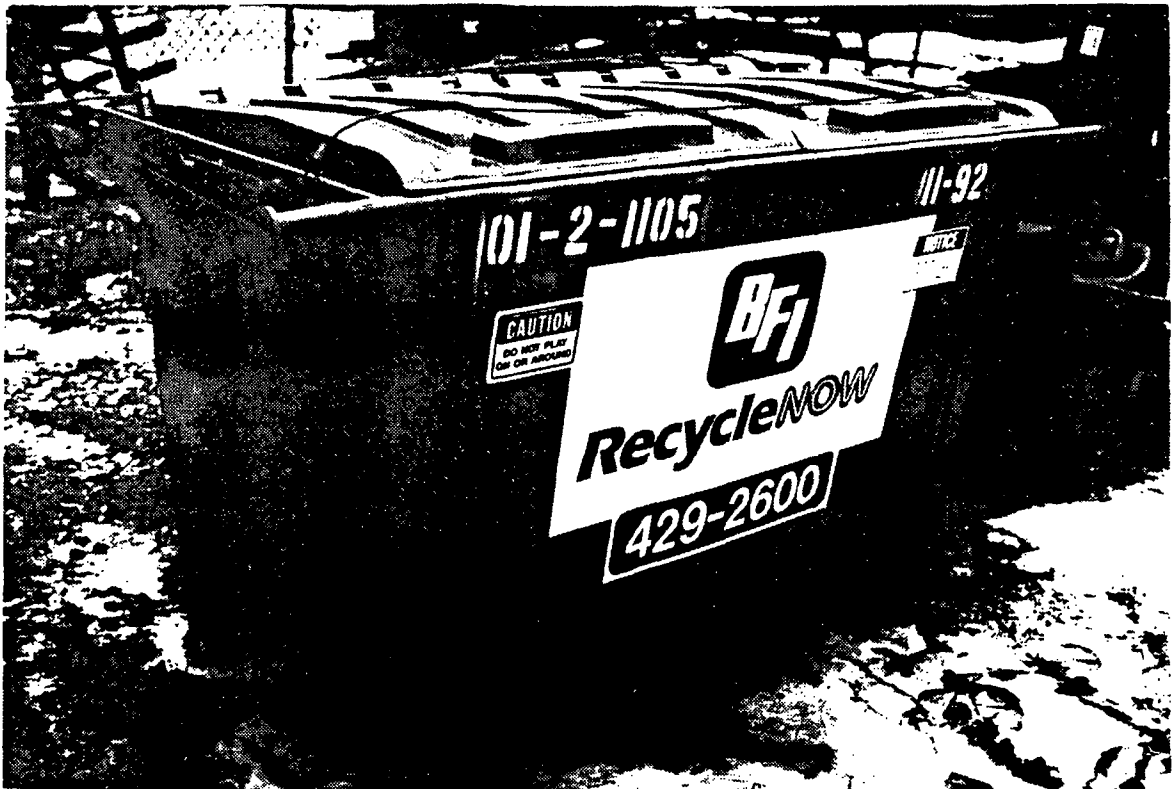


Figure 2.20 BFI Locking Recycle Container

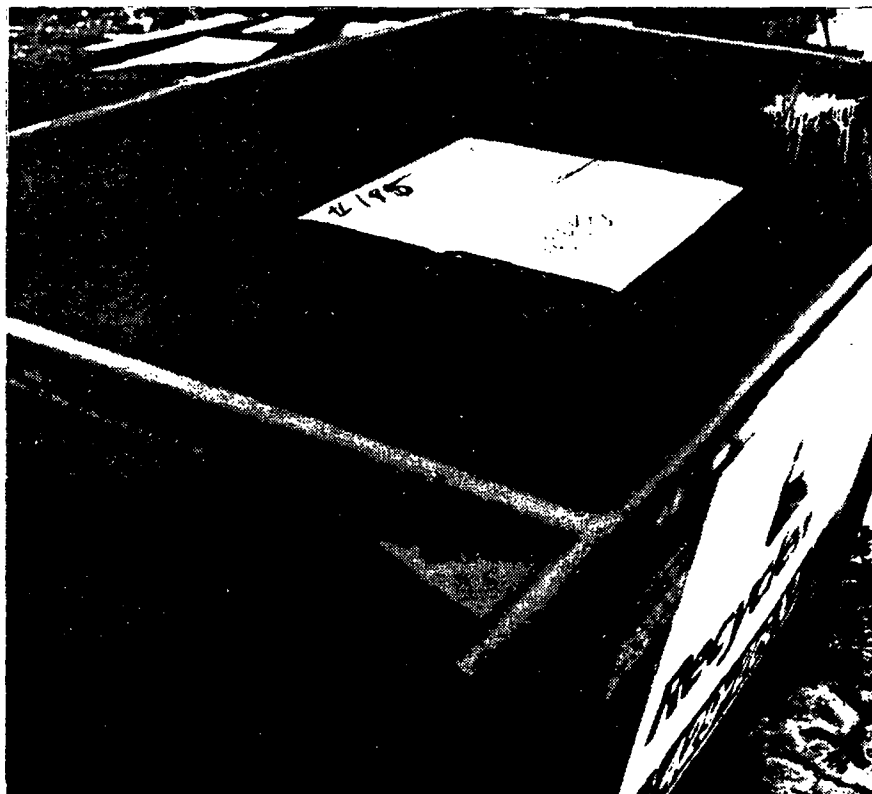


Figure 2.21 Interior of BFI Waste Container Where Radioactive Source Was Stored

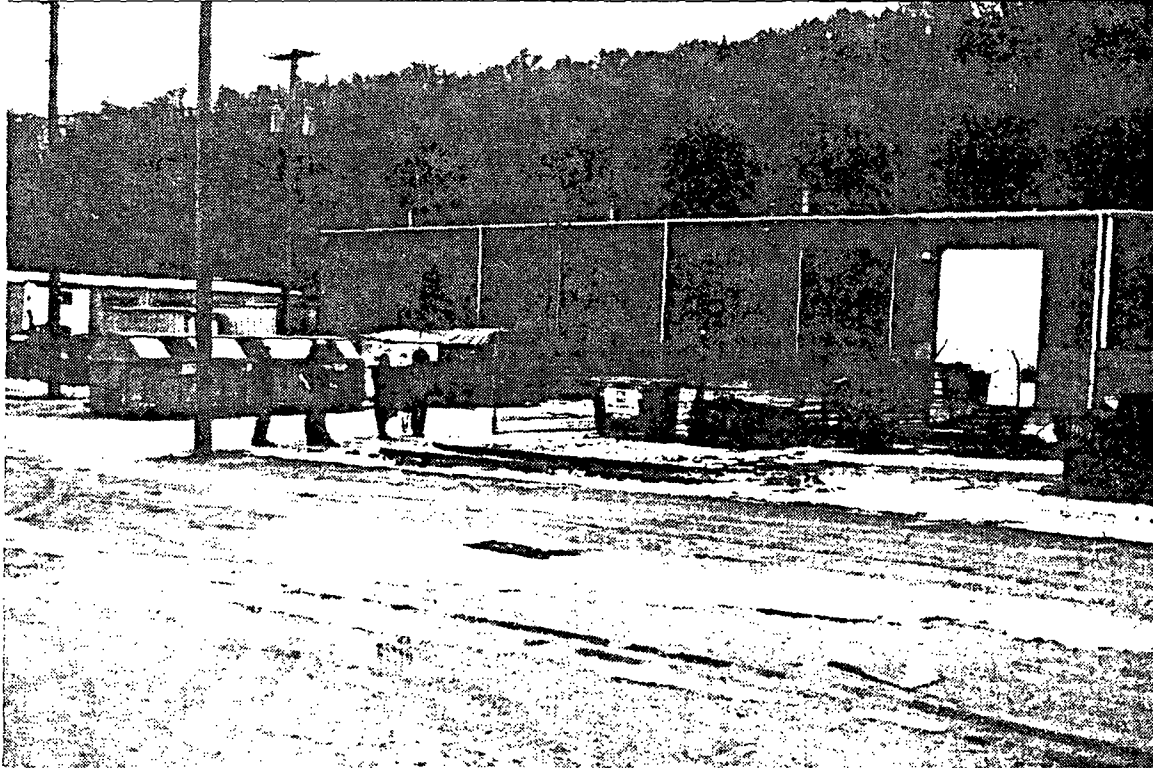


Figure 2.22 Location of Radioactive Source Recycle Container at BFI - Carnegie

3 OMNITRON 2000 HIGH DOSE RATE REMOTE AFTERLOADER SYSTEM

This section describes the Omnitron International, Inc.'s (Omnitron's) 2000 High Dose Rate (HDR) Remote Afterloader System; the nickel-titanium source wire; the HDR afterloader software; and the manufacturer's quality assurance and quality control (QA/QC) program. In addition, this section discusses the training, operating, and emergency procedures Omnitron gives to its customers.

The Incident Investigation Team (the team) obtained information from Omnitron's brochures and manuals; safety evaluations of the source and device (Registration Certificates); letters provided to the State of Louisiana to support the safety evaluations; and interviews with the persons involved. Additionally, the Food and Drug Administration (FDA) gave the NRC information they obtained from their investigation of the incident at the Indiana Regional Cancer Center (IRCC).

The scope of this investigation included only those mechanical or electrical components that could have contributed to the break in the wire and reflects the team's observations and review of documents and interviews with Omnitron personnel. The afterloader is operated by a computer; however, it is designed so that all low-level safety systems (e.g., hardware, interlocks, watchdog timers) are independent of the computer. The following description summarizes the overall operation of the afterloader, presenting specific details about the areas the team felt could have contributed to the wire break.

3.1 Description of the Afterloader System

This section discusses the main components of Omnitron's Model 2000 HDR afterloader system, which are the afterloader, the main console, the door status panel, the afterloader system safety features, and the implanted catheters and connecting catheters.

3.1.1 High Dose Rate Afterloader

The HDR afterloader (Figures 3.1 and 3.2) (see also Figures 2.3 and 2.4) contains the mechanical and electrical hardware necessary to execute a treatment. The afterloader contains a microcomputer that communicates with the main console through an RS-422 data link. The unit is approximately 107-cm high, 56-cm wide, and 61-cm deep (42-inches H x 22-inches W x 24-inches D) and weighs approximately 148 kg (325 pounds). Casters are mounted on the unit so that it is easy to move. The main components are the friction drive mechanisms, active and inactive wires, optical and mechanical switches, storage safe, treatment channel turret, stepping motors, emergency retract motors, backup power supply, and console interface card.

The turret (Figure 3.3) allows for connection of up to ten treatment channels. These channels are numbered on the turret head. Each treatment channel can have up to 20 dwell

positions with each position having dwell times of 0.1-second to 3-minute increments. For each channel, the source can be moved in 1.1-cm increments (the minimum) to a maximum treatment distance of 21 centimeters.

The implanted and connecting catheters are joined to the HDR afterloader by appropriate fittings. The turret rotates to allow the single active wire to extend and enter each implanted catheter for the desired treatment. The computer can determine the location of the turret by use of optical sensors and unique coding on the edge of the turret wheel.

The storage safe provides shielding for the tip of the source wire where the source is located. When the source wire is in the "safe" position (i.e., in the shielded position), the source is located within the center of the safe, and radiation levels at 1 meter (39.4 inches) from the surface of the device are below 0.001 mSv/hr (0.1 mR/hr).

Two independent drive systems are used in the afterloader. One drive system drives the source wire used for patient treatment and the other drives the dummy wire used to ensure that the catheter path is not constricted or obstructed. The dummy wire is also used during service and maintenance to ensure that the device is operating and functioning correctly.

Each drive system uses a pinch roller attached to a solenoid. When the solenoid is energized, the pinch roller puts pressure on the wire so that the wire comes in contact with the stepping motor's drive roller. The stepping motor moves the wire a predetermined distance (a step) for each signal sent from the computer to the motor. When the drive roller rotates, the wire travels through the guide tubes, owing to the friction between the wire, pinch roller, and drive roller.

An optical encoder mechanism is used for independent wire tracking and is located below the drive mechanism. Pinch rollers that are attached to solenoids are used for each system to keep pressure on the wire so that the wire is in contact with the optical encoder.

During a treatment, the appropriate motors and solenoids for the source-wire path or the dummy wire path are energized causing friction that allows movement of the wire and optical encoder roller.

As a stepping motor drives the source wire forward from the lead safe, a microswitch is tripped (the parked switch, which indicates the safe position, is located at the bottom of the source wire path) that resets a counter to zero (electric pulses are counted). As the wire travels, it reaches a microswitch (home position sensor) just before entering the turret and the distance is tracked by the pulses sent to the computer from the optical encoder.

The source wire travels past the first dwell position and then is pulled backwards, to remove any slack, to the first dwell position. The wire remains in this position for the prescribed dwell time and then travels to the next dwell position for that prescribed dwell time until all dwell positions and times have been completed. During each movement of the wire the

distance is recorded. After all dwell positions and times are completed, the wire is retracted. The number of pulses are counted during the retraction process until the home switch is deactivated. When the wire reaches the safe position, the solenoids are de-energized.

A computer controls the afterloader through an interface board in the computer and a signal multiplexer. Separate circuits on the interface board count the number of pulses in each direction. By using the optical encoders in combination with the stepping motors, the HDR can account for the slippage of the wire. In addition, timers are located on the interface board to time the stepping motors (used for error detection).

An emergency drive system is located in the afterloader that is separated from the main drive systems (stepping motors). The emergency drive system consists of a solenoid, pinch roller, and battery-operated dc motor. In the event of an emergency retract, the dc retract motor is energized and power is removed from the solenoids associated with the stepping motors and optical encoders, thus removing any friction forces caused by these systems. The dc motor continues to operate until the end of the source wire (opposite the source-end) contacts the parked switch.

The back of the afterloader contains a status panel, shown in Figure 3.4, which consists of the following.

- The "SAFE," green, light-emitting diode (LED) indicates that the end of the wire opposite the source end has been detected by the sensor (i.e., the parked switch is tripped).
- The "IN PROGRESS," amber, LED indicates that treatment has been initiated.
- The "DUMMY PARKED," green, LED indicates that the end of the wire has been detected by the sensors (i.e., parked switch is tripped).
- The "ERROR," red, LED indicates that an error has occurred. The error LED is lit whenever an error condition occurs. These error conditions can be caused by malfunctioning equipment, power failures, catheter restrictions, etc., and the cause of the error is displayed on the computer screen.
- The "DIAGNOSTIC MODE," amber, LED indicates that the system is undergoing diagnostics. Only Omnitron personnel are authorized and have access to this mode. Under this mode of operation, direct movement of the source and dummy wires can be controlled through the console.
- The "DOOR OPEN," red, LED indicates that the switch on the treatment room door has been tripped. Either the switch is broken or the door is open.

- The "ARMED," amber, LED indicates that the system is ready for treatment (i.e., passwords have been entered correctly, keylocks have been set, etc.).
- The "MANUAL RETRACT," red, LED indicates that both the ac stepping motors and the dc emergency retract motor failed to retract the wire, and, therefore, a manual retraction is necessary. If this condition occurs, the LED will be lit and an alarm on the console will sound.
- The "RESET," yellow push button has a hinged cover to prevent accidental activation. For resettable errors, pushing the button will erase the error message displayed on the computer console and turn off the "ERROR" LED. For nonresettable errors, a message will appear on the screen that Omnitron needs to be called to fix the error.
- The "STOP," red push button is used to stop treatment. If this button is pushed when the active wire is out, the active wire will automatically retract. Treatment information is saved, and, therefore, treatment can be resumed from the point where it was stopped when this button was pushed.
- A keyswitch is provided on the console to allow only authorized personnel to control the treatment.

A label containing a radiation symbol and warning appear on the HDR afterloader next to the status panel.

The afterloader is locked, preventing users from accessing the mechanical, electrical, and software maintenance portions of the HDR afterloader system. Entry can only be accessed by the manufacturer.

3.1.2 Main Console

The main console consists of a microprocessor, color monitor, printer, disk drive, uninterruptable power supply, and treatment unit interface card (Figure 3.5). The microprocessor is a personal computer (PC) that operates the afterloader and performs first-level safety functions (e.g., dummy wire check, door interlock). The PC controls all wire movement signals and controls. It is used to enter patient data and the patient treatment plan, to initiate treatment, and to perform service diagnostics.

The computer console contains a status panel, shown in Figure 3.6, that consists of the same LEDs, controls, and keyswitch as those described in Section 3.1.1 for the afterloader.

The system printer provides hard copies of treatment printouts, treatment logging and diagnostic reports, and error messages.

Also provided with the control console is an uninterruptable power supply, which allows treatment to continue in the event of a power failure.

3.1.3 Door Status Panel

The treatment room door status panel (Figure 3.7) provides additional indication to personnel performing the treatment as to the status of the HDR afterloader. This panel contains the error LEDs as described for the afterloader and control console, an emergency stop button to allow the user to halt treatment, and an alarm horn. The panel is installed in close proximity to the door of the treatment room.

3.1.4 Afterloader System Safety Features

The manufacturer incorporated numerous safety features into the HDR afterloader system. The team observed the performance of those safety features incorporated into the HDR afterloader relevant to the constriction or obstruction of the source wire. Team members requested that Omnitron simulate the constriction condition that occurred at the IRCC with the active source wire. Team members observed that during a constriction in which the dc motor is energized, no audible alarms were activated. However, an error message did appear on the computer console for the entire duration of the wire retraction, and an error message indicating that a constriction occurred was reported to the error log. Other safety features are described in the manufacturer's manuals and brochures.

3.1.5 Patient Applicators and Treatment Tubes

Implanted catheters, known as patient applicators or treatment applicators are inserted into the patient during a surgical operation. The active source encased in the cavity of the source wire will reside inside this catheter during treatment time. Typical implant catheters (applicators) include rigid needles, flexineedles, and custom applicators. The flexineedles must be used with appropriate accessories (e.g., needle obturator, treatment tubes).

In the HDR treatment of November 16, 1992, at the IRCC, five 10-cm flexineedles were used. Each of the 10-cm flexineedles has a 20-gauge polyester tubing needle shaft, a stainless tip, aluminum coupling, and a nylon suture button. The internal diameter of the 10-cm flexineedle is 0.813 millimeter (mm).

Connecting catheters are known as treatment tubes; these are the tubes between the HDR afterloader and the implanted catheter.

In the HDR treatment of November 16, 1992, at IRCC, five 80-cm connecting catheters were used. Each of the catheters was made of teflon tubing with aluminum coupling hardware. The nominal coupled length of the inside lumen of the 10-cm flexineedle connected to an 80-cm coupling catheter is 90.34 cm plus or minus 0.2 cm. Figure 3.8 represents a schematic of this arrangement.

3.2 Description of the Source Wire

The registration certificate for the source wire, Model SL-777 (Figure 3.9), was issued by the Louisiana Radiation Protection Division. The source wire is constructed of Nitinol (nickel-titanium alloy). The source wire was produced in the following way.

Originally the wire comes in a roll from the supplier of the Nitinol wire. Lengths of wire are cut from the roll (cut intentionally long) and sent to another company to produce a cavity in one end of the wire approximately 0.34 mm (0.014 inch) in diameter and 11-mm (0.43-inch) deep (wall thickness approximately 0.089 to 0.102 mm [0.0035 to 0.004 inch]). Previous production required a 13-mm cavity. Instructions are sent with the wire telling the company in which end of the wire to put the cavity.

The wire is sent with a traveler to another company where the cavity is x-rayed. At this time, the x-ray vendor assigns a serial number to the source wire. The wire and videotape of the x-ray are shipped back to Omnitron's Houston, Texas, office.

The wire is inspected at the Houston office for gross defects in material or workmanship and the wire's critical dimensions are checked. The wire is then sent with its traveler to Omnitron's source production facility in Edgerly, Louisiana.

At the Edgerly facility, the critical dimensions are again checked against what is stated on the traveler. If these dimensions are not the same, the discrepancy is either handled over the telephone, the wire is scrapped, or the wire is returned to Houston for rework. Once the personnel are satisfied with the wire, a sample piece of the wire is placed in the welding fixture located in the hot cell and a test weld is performed on a sample piece of the wire. The weld is inspected for an even-flow, uniform heat distribution (heat ring), and a "shiny" surface. If the weld is satisfactory, the remaining portion of the wire is cut to length. The wire is placed in the hot cell, and a weld is performed on the end opposite the cavity. All welds are performed, using a tungsten inert gas process. This weld is inspected as mentioned above, and if found to be satisfactory, the production continues.

Active iridium wires 5-mm (0.2-inch) long are received from the reactor and are placed in the hot cell. Two active iridium wires are placed in the source wire cavity and a 1-mm plug is placed on top. Previous production required a 3-mm (0.12-inch) plug. This 3-mm (0.12-inch) plug was the size used in the IRCC source wire. This end is welded to form a closure weld. The weld is inspected as previously described, and the outside diameter is measured. If the weld and outside diameter of the wire are acceptable, the wire is leak tested and placed in a transport container. The approximate dimensions of the finished wire are 2201 mm (86.7 inches) in length and 0.58 mm (0.023 inch) in diameter.

The transport container is moved to an HDR afterloader in the laboratory. The wire is loaded in the afterloader and cycled through the connecting catheters (1) to a wipe-test fixture where a wet and dry wipe are taken; (2) to a critical bend test fixture to ensure the source

wire will make the curves without failure; and (3) to a calibration station where the source is calibrated. The critical bend test consists of two curves, attached together to form a smooth "S" curve. The two radii of the test fixture were chosen to simulate actual patient treatment.

The finished wire is then packaged with the calibration and shipping papers and shipped to the customer. Omnitron staff install the source at the customer's site. Because of the source's half-life, it is replaced approximately every three months.

3.2.1 Prototype Testing Performed on Nickel-Titanium Source Wire.

Letters in support of the sealed source certification stated that the manufacturer subjected the source to the tests discussed in this section. Two prototype nickel-titanium wires were tested to the requirements of the International Organization for Standardization (ISO) 2919, "Sealed Radioactive Sources--Classification." This guidance does not provide specific tests for HDR afterloader brachytherapy sources. However, this guidance does provides specific tests, test procedures, and test equipment for other radioactive source usages (e.g., radiography, teletherapy, calibration). The classification of the source defines the testing conditions that the source has met.

Common practice has been to test brachytherapy wire sources to the same specifications as sealed sources used in medical interstitial and intracavitary appliances. The recommended tests for these sources are temperature, pressure, and impact. In addition, common practice has been to perform cycle and tensile tests on source wires where capsules are welded or swaged onto the end of the source as with radiography wires.

In a letter dated January 16, 1992, to the State of Louisiana Radiation Protection Division, Omnitron stated it subjected two prototype sources to the minimum classification recommended for the temperature, pressure, and impact tests. The prototype sources passed the tests in accordance with ISO 2919.

The manufacturer also subjected prototype sources to a bending fatigue test. Catheters connected to an afterloader were curved to represent pathways encountered during treatment. Different curved paths were used during the cycling of the source wire. The paths chosen and number of cycles tested for each path, as documented by Omnitron, are listed below:

<u>Pathway</u>	<u>Cycles</u>
4-cm (1.6-inch) radius followed by a 3.5-cm (1.4-inch) radius ("S" curve)	2500
2.5-cm (1-inch) radius (180° turn)	1
2.3-cm (0.91-inch) radius (180° turn)	1

2.2-cm (0.87-inch) radius (180° turn)	1
2.0-cm (0.79-inch) radius (180° turn)	21
1.5-cm (0.6-inch) radius (180° turn)	2

The wire was then subjected to 974 additional cycles, using the 4-cm (1.6-inch) radius followed by a 3.5-cm (1.4-inch) radius. The wire was subjected to a total of 3500 cycles.

With the use of a microscope (magnification 38 times), Omnitron reported that the source examination "showed wire tip, iridium encapsulation and body of wire to be in pristine condition. Dye penetrant test of tip and encapsulation area showed no material cracks."

An Omnitron corporate manager stated in an interview that the first production wire was tested to the breaking point and broke after 4000 cycles at the bottom of the cavity. However, the same manager's notes showed a break after 4070 cycles on one wire and 3490 cycles on another wire.

Omnitron performed no calculations to validate the design of the nickel-titanium source wire. Tensile or shear stresses were not calculated and no associated tests were performed on the wire. Through discussions with the vendor of the wire and Omnitron personnel, Omnitron decided that cycling testing simulating clinical situations would be the best way to test the source wire.

Further, Omnitron performed no metallurgical tests before the wire broke to validate the vendor's material certification. No metallurgical tests were performed after the cavity was produced to ensure that production of the cavity did not affect the overall integrity of the wire.

3.3 Description of the Omnitron 2000 Afterloader System Software

Two groups of software programs were available with the Omnitron 2000 afterloader system: (1) the Computerized Treatment Planning System (CTPS) software used to develop a treatment plan (isodose computations) for a patient and (2) the software required to run afterloader functions.

The Omnitron 2000 HDR afterloader has a dedicated software system used in isodose computations. This software system, called the CTPS, was developed by a medical physics software company independent of Omnitron. Omnitron's medical physics personnel performed specific QA checks of the CTPS software before releasing it to its customers. The CTPS resides on a stand-alone computer hardware system and has a menu-driven interface. Treatment data is transferred to the Omnitron 2000 control console, via a standard 3.5-inch floppy disk drive. The CTPS also automatically corrects for the decay of the source. The CTPS has a full three-dimensional dose optimization feature that plans the HDR

brachytherapy doses delivered by the radioactive source. Dose histograms and three-dimensional wire frame displays are produced. Separate programs exist for entering patient data and for entering or editing iridium-192 radioactive source data.

The computer software installed on the Omnitron 2000 control console was developed by Omnitron, with the assistance of a consultant. The Omnitron 2000 computer console at IRCC used Version 3.0 of the software. This software has a menu-driven operator interface and is organized in a modular manner.

The operator must (1) have the Omnitron key in TREAT mode and (2) enter the Omnitron console program system Access Password before any access to patient or afterloader information is allowed. Some of the software programs in the afterloader console are only accessible to the Omnitron personnel.

The console and afterloader software were designed to monitor all major electrical and mechanical systems, detect the proper operation of these systems, and initiate any afterloader safety functions if necessary. The software is also designed to record and print out patient treatment information, treatment interruption causes (e.g., "Active wire path constriction detected at XX cm"), whether the detected condition can be reset by the HDR afterloader operator, and any device check failures detected (e.g., "Dummy wire check on Channel X failed").

The team identified the following features of the Omnitron 2000 hardware and software configuration.

1. The dummy wire executes programmed treatment to test for proper catheter routing, programming, and to ensure that no obstructions or kinks are in the catheters. For patient treatments requiring multiple catheters or applicators (similar to the 5-channel treatment of November 16), the operator can choose between having--
 - dummy wire checks of all programmed channels (i.e., catheters) before source wire treatment of each channel, or
 - dummy wire check of Channel 1 (one catheter), followed by the source wire entering that channel; dummy wire check of Channel 2, followed by the source wire entering that channel; and so on.
2. If the source wire jams while being extended or while being retracted from the device, one message is generated "Active wire path constriction detected at XX cm." However, the two conditions are acted upon differently by the afterloader. When the source wire jams while retracting and the pull force necessary to retract the wire exceeds that of the stepper motor upper limit, the emergency dc retract motor is engaged to retract the source wire. Otherwise, if no wire jam occurs, the stepping motor retracts the wire.

3. When the emergency dc motor engages, a message stating that the dc motor engaged is displayed momentarily (for approximately 10 seconds) on the console's screen, with no written information being stored in memory or printed out.
4. The software was configured so that when the emergency dc retract motor starts retracting the source wire, all devices that monitor and record the wire length information immediately disengage and all wire length information is lost. In addition, activation of the emergency dc retract motor is not recorded on the printout,
5. Following an emergency dc retract of the source wire, the afterloader operator is allowed to reset this error and restart treatment without taking any corrective actions.

3.4 Omnitron's Quality Assurance and Quality Control Program

Omnitron International, Inc., located in Houston, Texas, was responsible for the overall QA/QC program. However, although not directly stated, Omnitron's source-wire fabrication facility, located in Edgerly, Louisiana, was responsible for the QA/QC of the source wire once it was received from the Houston office. Therefore, the team separately reviewed the QA/QC program pertaining to (1) the production of the HDR afterloader and the initial fabrication of the nickel-titanium wire and (2) the assembly of the source wire. Any part of Houston's QA/QC program that pertains to the source wire was considered during the review of the QA/QC program for the Edgerly facility. The NRC does not have specific regulations or guidance that relate to QA/QC programs for vendors of sealed sources and devices. The team used a draft NRC QA/QC guide to review Omnitron's QA/QC program. The team's findings were essentially the same as those reported in the FDA audits as noted below.

The Lafayette, Louisiana, office of the FDA performed an audit of Omnitron's Edgerly, Louisiana, facility during December 1992. Audit findings were presented to Omnitron on December 23, 1992, and documented on FDA Form 483. The inspection report findings included multiple Omnitron deficiencies relating to their (1) quality assurance program, (2) audits, (3) device history records, (4) reworking of components, (5) device master records, (6) written procedures for in-process and finished device testing, (7) cleaning and leak testing, (8) calibration of equipment, (9) validation of the work process, (10) wire receipt inspection, and (11) use of white-out on QA documents .

The Dallas, Texas, office of the FDA performed an audit of Omnitron's Houston, Texas, facility during December 1992 and January 1993. Audit findings were presented to Omnitron on January 11, 1992, and documented on a Form 483. The inspection report findings included multiple deficiencies relating to (1) their QA audits and design changes, (2) complaint file shortcomings, (3) medical device reporting, (4) software (afterloader and console) validation and testing, (4) manufacturing specifications and manufacturing process control, (5) device history records, and (6) component receipt inspection. Both FDA audits indicated a lack of procedures, component QA records, and tests. In addition, where

procedures, component QA records, and tests existed, numerous examples indicated that standard QA/QC practices were poorly implemented.

Findings documented in the recent FDA audits are similar to the overall team observations. On the basis of FDA audit findings at the Edgerly and Houston offices, the Texas Department of Health, assisted by Houston police, embargoed the Omnitron 2000 HDR afterloader on January 12, 1993. To have the embargo lifted, Omnitron must complete the corrective actions needed to resolve the issues raised during the FDA audits.

3.5 Training, Operating, and Emergency Procedures for the Omnitron 2000 High Dose Rate Afterloader

Upon request, Omnitron provides a two-day, inhouse, or onsite, training course to customers. The course in its current form became available May 1, 1992. In addition, Omnitron provides training at the customer's facility before treatment of the first patient.

Omnitron chose four trainers on the basis of their knowledge and experience. They have not been formally trained or received training to become trainers. Omnitron had (1) no written policies or procedures on what training and experience is required before an individual can become a trainer and (2) no written procedures specifying who should be trained at the customer's facility. Typically, an individual responsible for training calls or writes the customer's facility and asks the responsible physicist who they feel should be trained. Typically, the responsible physicist and the person who will most likely perform the treatment are trained. The director of training told the team that Omnitron will periodically call the customers who have not had two people trained in order to schedule future courses for them.

Omnitron provides the customer with training and reference material. The material includes system specifications, applicator and accessory specifications, treatment planning, HDR software and hardware, an instruction manual, and a one-page emergency procedure. The objectives of the two-day training course are to teach the customer how to perform treatment planning and dose calibration, execute treatment using the HDR afterloader, use emergency procedures in the event of a source retract failure, and discuss uses of accessories and applicators. The emergency procedures, which are provided to the customer, do not address breakage of the wire. Training provided before treatment of the first patient consists of a review of the operation and safety features of the HDR afterloader. Basic radiation safety training is not provided.

According to Omnitron's Sr. Vice President of Research and Development, the safety features and error messages that appear on the console monitor screen are discussed as part of the training; however, these error messages are not addressed in Omnitron's written procedures. Errors that cannot be reset are not discussed in the training, and the customers are told to call Omnitron if an error of this type occurs.

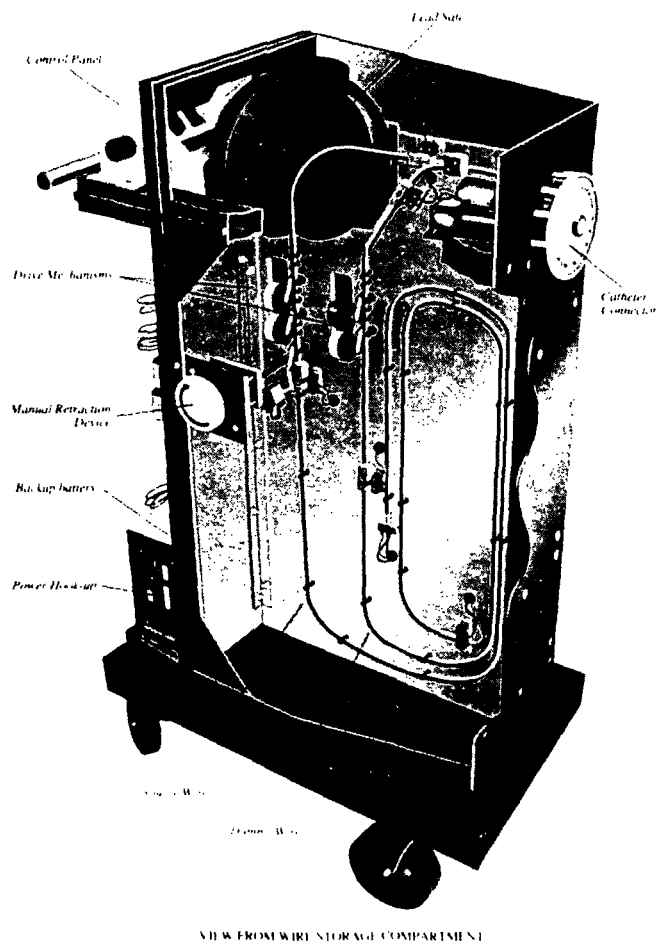


Figure 3.1 Omnitron 2000 High Dose Rate Remote Afterloader

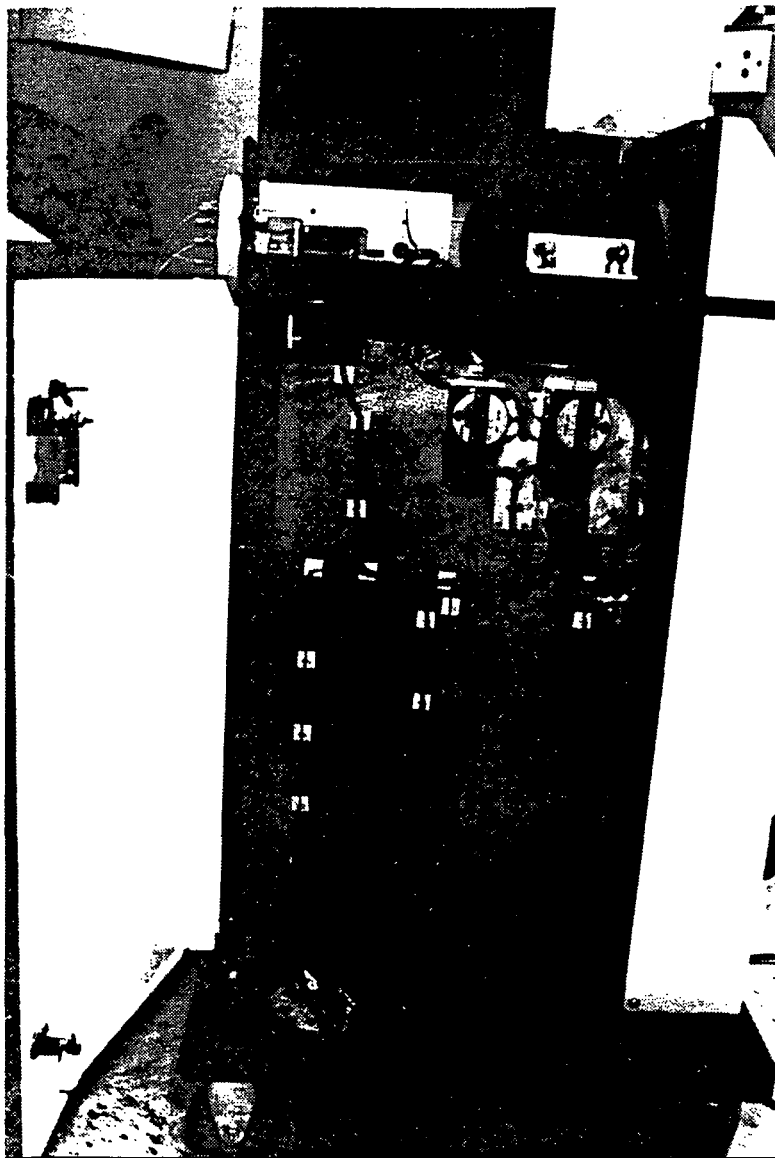


Figure 3.2 Left Side of High Dose Rate Remote Afterloader, Indiana, Pennsylvania

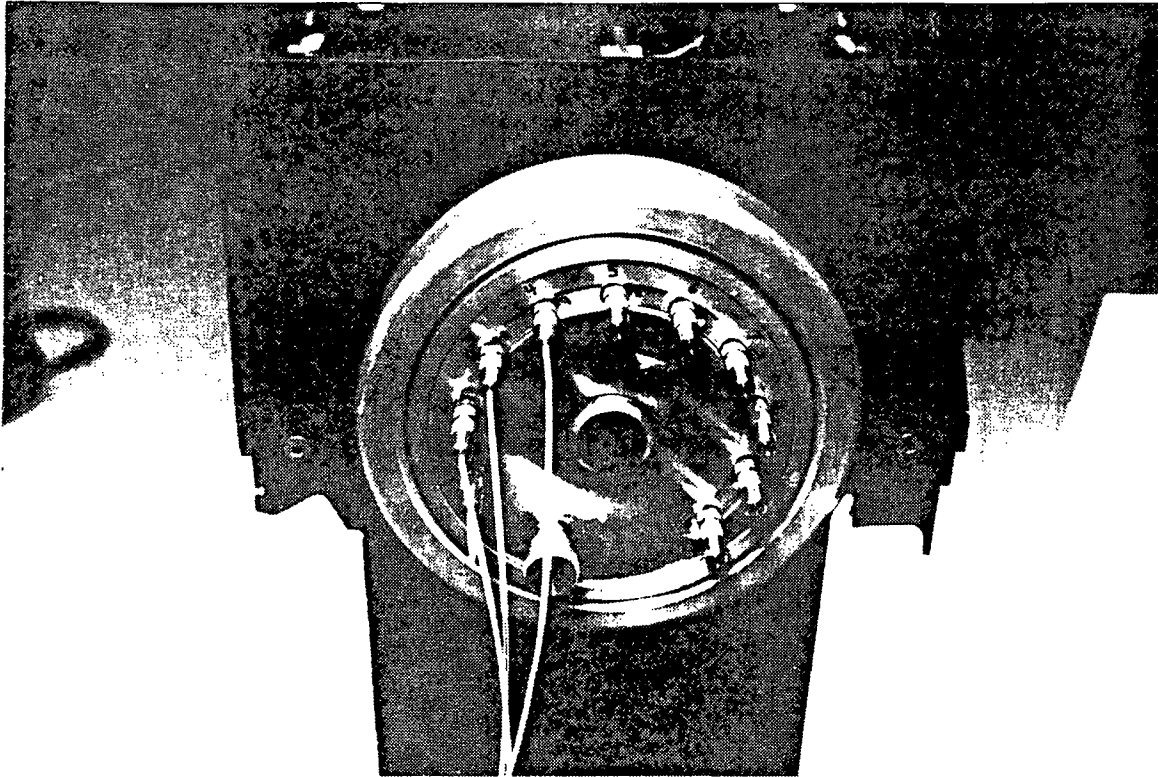


Figure 3.3 Turret Drive Assembly, Indiana, Pennsylvania

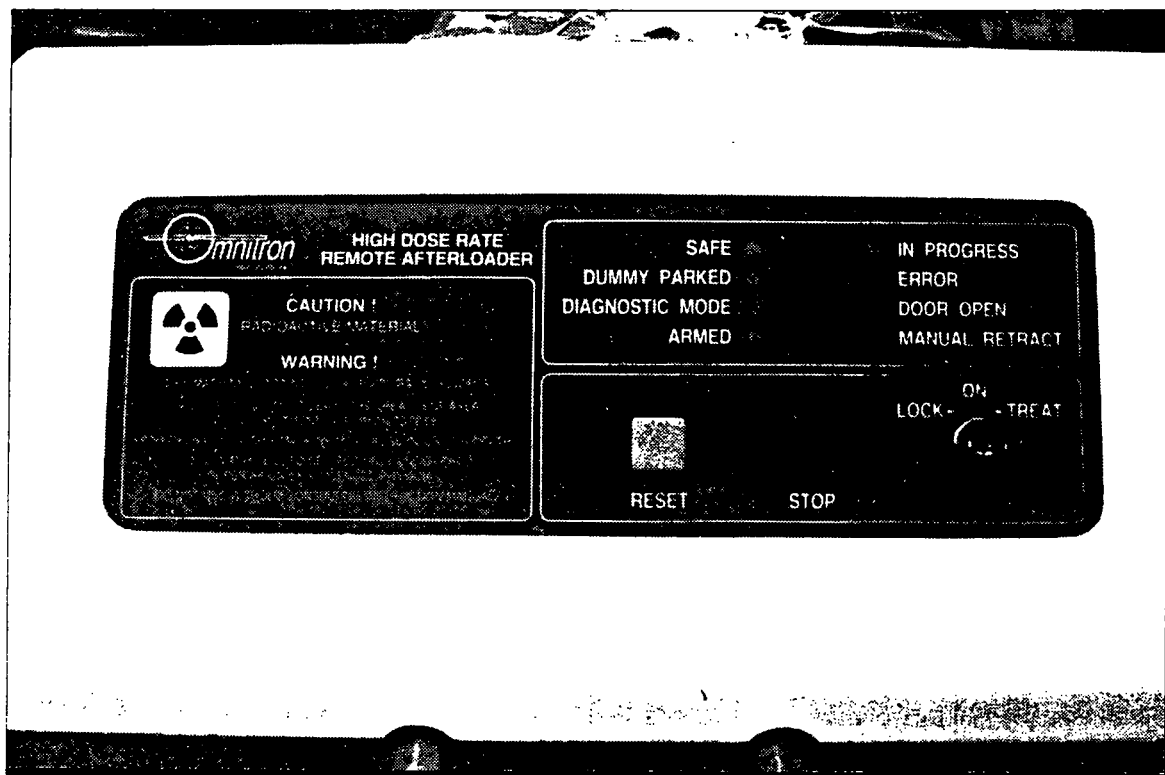


Figure 3.4 Control Panel for High Dose Rate Remote Afterloader, Indiana, Pennsylvania

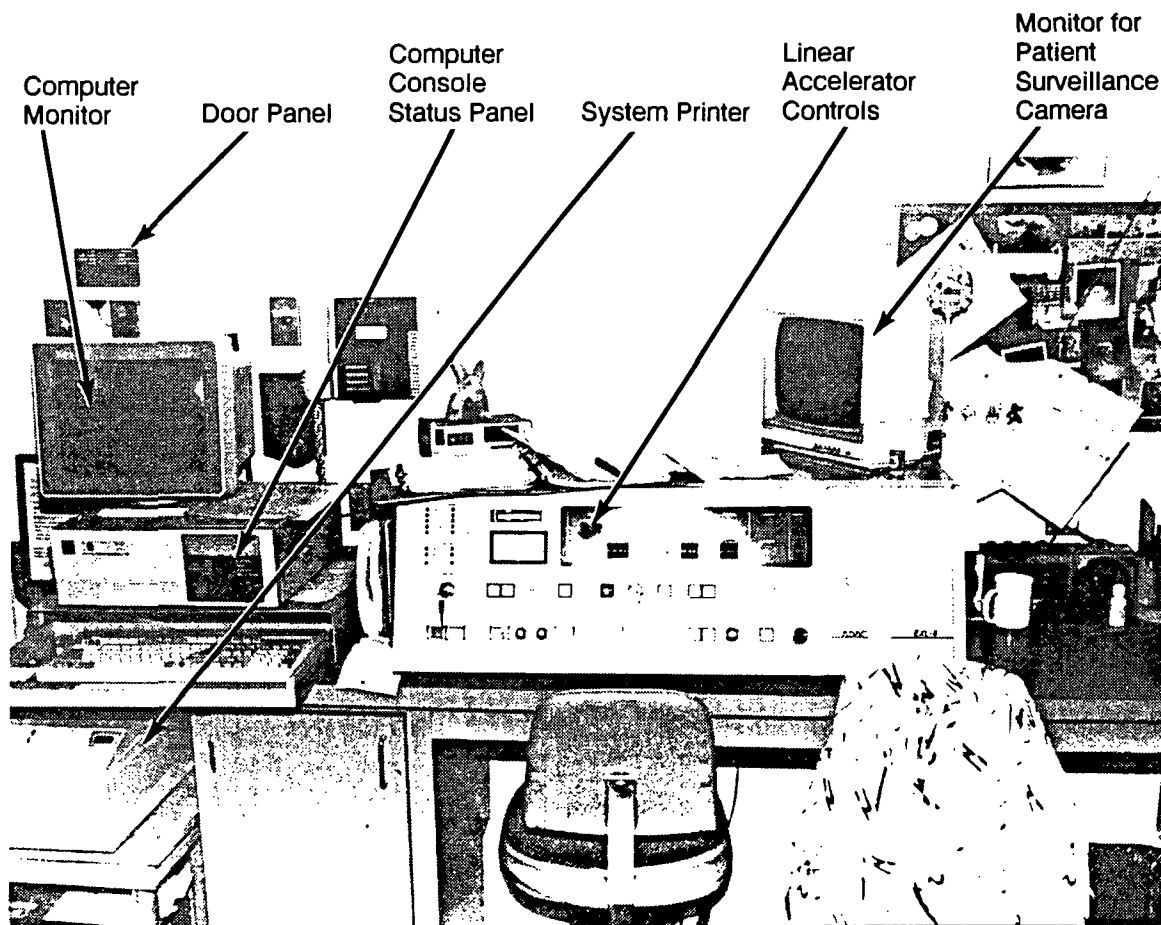


Figure 3.5 Arrangement of the Computer Console Status Panel, Door Panel, System Printer, Linear Accelerator Controls, Computer Monitor, and Monitor for Patient Surveillance Camera

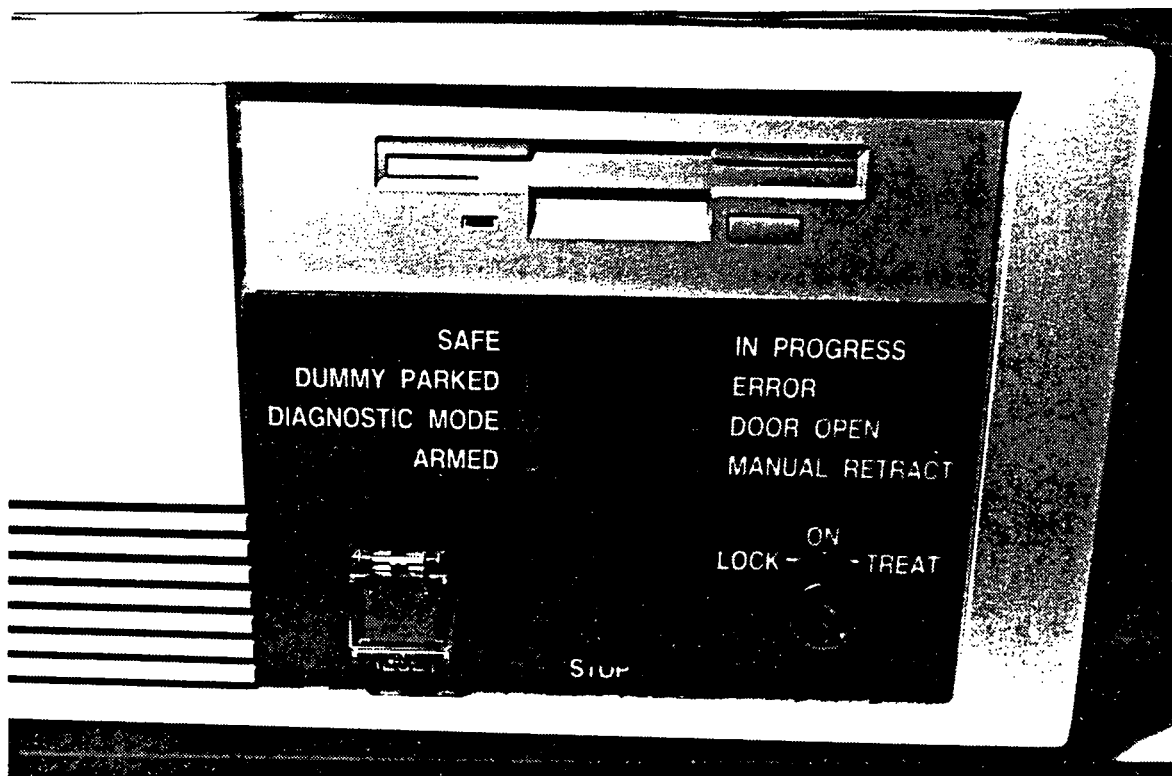


Figure 3.6 Computer Console Status Panel of High Dose Rate Remote Afterloader System

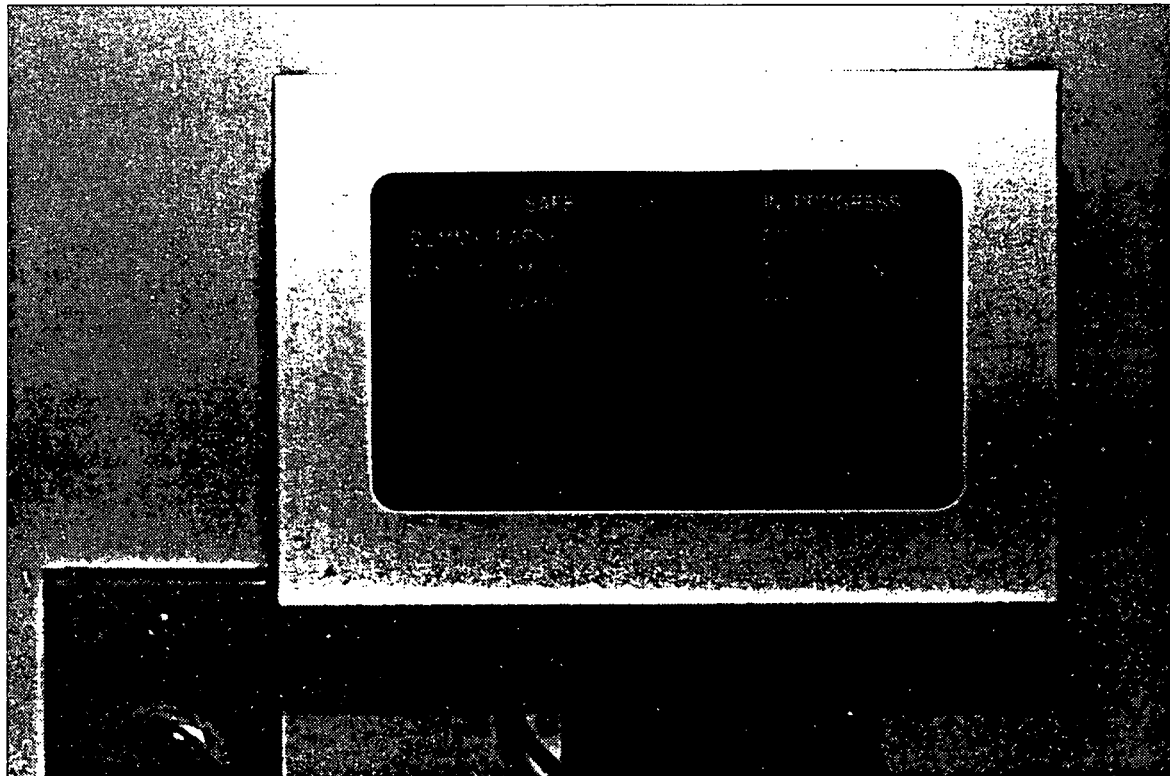
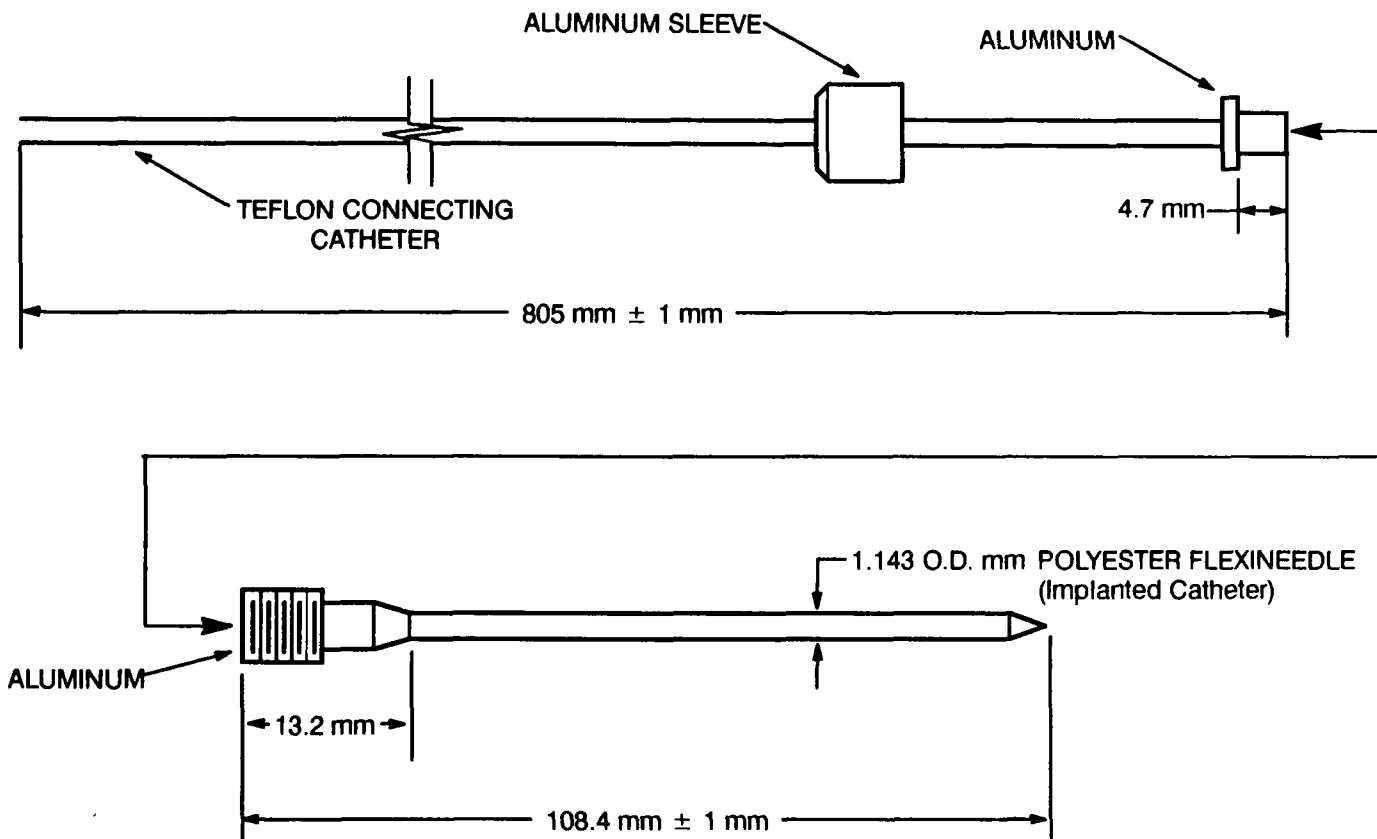


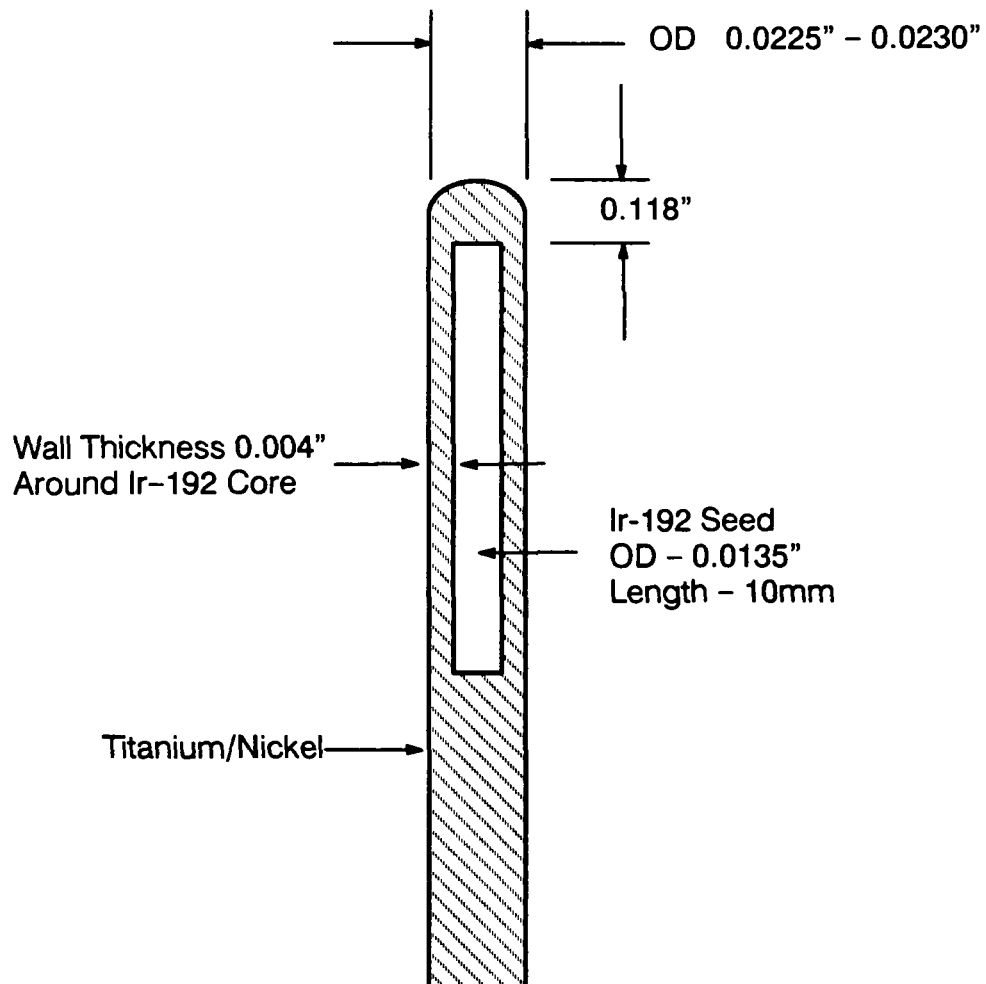
Figure 3.7 Door Status Panel of High Dose Rate Remote Afterloader System



NOTE: Drawing not to scale and all dimensions are approximate

Figure 3.8 Connecting Catheter and Flexineedle Unassembled

IR-192 Seed in Titanium/Nickel Model #SL -777



Ir-192 Seed in Titanium/Nickel

An Ir-192 Seed is encapsulated firmly inside a solid titanium/nickel wire.
The overall length of the Ir-192 Seed is 10mm

The above composite has passed all ANSI requirements for sealed sources.

**Figure 3.9 Iridium-192 Seed in Titanium/ Nickel
(Reproduced from Registration
Certificate LA-0760-S-102-S)**

4 EQUIPMENT PERFORMANCE

This section presents (1) the potential analysis of the source-wire breaks, (2) the immediate cause of the wire break, (3) the root cause of the wire break, (4) the failure of the high dose rate (HDR) afterloader system to detect the length of the broken wire, and (5) the performance of the PrimAlert-10 area radiation monitor.

The team obtained its knowledge of the source wire and HDR afterloader from Omnitron International, Inc.'s (Omnitron) brochures, visits to the HDR afterloader and source wire facilities, direct observation of the broken wire and afterloader, engineering reference material, material certification, prototype test results, and other information provided by Omnitron. Most of the testing was performed by Southwest Research Institute (SwRI), San Antonio, Texas, under an existing contract with the NRC. The team also reviewed the performance history of the PrimAlert-10 that was installed in the treatment room at the Indiana Regional Cancer Center (IRCC).

4.1 Failure Analysis Pertaining to the Source Wire

This section describes the failure analysis performed on the broken source wire at IRCC, and the Greater Pittsburgh Cancer Center, (GPCC). It also includes an analysis of other sample wires used for testing purposes and describes (1) possible areas where the source wire failure may have occurred; (2) observations of equipment used during assembly, testing, and shipping that may have caused wire degradation; (3) description of the test samples; (4) the test performed of the samples; (5) observations of the test results; (6) implications of these observations; and (7) further testing.

4.1.1 Possible Failure Areas

On December 4, 1992, the quarantine of IRCC's Omnitron 2000 HDR afterloader was lifted by the NRC, and team members observed the disassembly of the device by Omnitron. The team asked Omnitron personnel specific questions pertaining to the source wire and afterloader.

Upon examination of the inactive portion of the source wire, it appeared that the wire broke just above the bottom of the cavity. The entire source wire length as specified on drawings and the associated traveler was 2201 mm (86.65 inches). The inactive portion of the wire was measured, and found to be approximately 2188 mm (86.14 inches). Therefore, the remaining portion of the source wire containing the iridium-192 source was approximately 13 mm (0.51 inch). The cavity produced in the end of the wire was also approximately 13 mm (0.51 inch) in depth.

The team took wipe samples of the active wire and inactive dummy wire and analyzed them on site to determine leakage of radioactivity from the source. The wipe samples were below the regulatory limit of 185 Bq (0.005 μ Ci) but were sent to Region I for analysis. The wipe

samples were divided between Omnitron and NRC, both having representative samples of the dummy and source wires. The results from the gamma spectroscopy test performed in NRC's Region I office indicated a photopeak having an energy corresponding to iridium-192 for both wires. However, the levels obtained were below the regulatory limits of 185 Bq (0.005 μ Ci).

To examine the HDR afterloader system for conditions that could damage or break the source wire, the afterloader was divided into six areas where the break of the wire could have occurred (Figure 4.1):

- **Area 1—Inside the afterloader:** The team examined the entire track where the source tip came in contact with parts of the afterloader (Figure 4.2). This track consisted of the lead-safe stainless-steel guide tubes, the home sensor, the "y" block, and the turret assembly. The team examined all the guide tubes for flattening, elongation, crimps, cracks, eccentricity and other signs of visible damage. These wipe samples previously mentioned were sent to SwRI for chemical analysis of the deposit material. The chemical analysis indicated that no lead was present in the deposit material. Minor wear of the source wire and machine components could have produced the small quantities of metallic elements found in the deposit material. Sulfur and silicon were present in small quantities, however, the origin of these elements has not been established. The home sensor was taken apart and examined; the "y" block was disassembled and inspected for wear; and the drive rollers, encoder, and corresponding pinch rollers were examined. The turret head assembly was not disassembled and inspected owing to a limited selection of available tools. No significant wear was observed on any of the accessible Area 1 components.
- **Area 2—Inside the turret connector:** The five turret connectors installed in the turret head (believed to be the five used in the November 16, 1992, treatment) were examined for significant wear (see Figure 3.3). The inside holes were not obstructed, no burs were noticed, and the holes appeared to be straight. The outside surface showed no signs of damage, and the connectors did not appear to be bent. No significant wear was observed on any Area 2 components.
- **Area 3—Inside the connecting catheter (see Figure 3.8):** Four of the five connecting catheters (believed to be used during the November 16, 1992, treatment) were examined for wear. All appeared to be in good condition with no visible signs of significant damage. A catheter that was believed to be the fifth connecting catheter (not connected at the time of the inspection) was also examined. The fifth catheter was cut, and there were some signs of visible damage. Damage to the catheter could have occurred after treatment.
- **Area 4—Inside the connecting catheter connector:** Area 4 consists of an aluminum fitting pressed onto the teflon catheter. An aluminum sleeve, which is threaded

inside, fits over the implant catheter and allows the connecting catheter to be fitted to the flexineedle (Figure 4.3 and see Figure 3.8). For damage to the source wire to occur in Area 4, a misalignment of the connecting catheter and flexineedle and/or an external force causing the source to bend would have had to occur. Misalignment was not noticed by either RTT-A or RTT-B and no constriction error was reported on the error log for the dummy wire as it passed this area during treatment.

- **Area 5—Inside the flexineedle connector:** A polyester tube containing a stainless steel tip was pressed into a threaded aluminum fitting to form the flexineedle. Because the break occurred near the bottom of the source-wire cavity and a constriction error occurred at 82.4 cm, the tip of the source would have had to be placed approximately 13 mm past the end of the fitting. (See Figures 3.8 and 4.3). The fitting supplies support to the wire, and when the wire is bent and retracting, places a stress point on the wire. The team was unable to examine the fitting for wear.
- **Area 6—Inside the flexineedle:** The actual break may have occurred within this area because the error log reported a constriction at 82.4 cm. The team does not believe that the break was caused by forces on the teflon tube because most of the needle was inside the patient. The flexineedle used in Patient A was disposed of and was not examined by the team.

4.1.2 Observations of Equipment Used During Assembly, Testing, and Shipping.

On December 3, 1992, team members observed equipment at the Edgerly facility used in the manufacturing and assembly of the source wire to evaluate if the equipment could cause unnecessary stress or failure of the wire.

In reviewing the assembly procedures, the team noticed several areas that may subject the source wire to unnecessary stress. During loading and welding, part of the coiled nickel-titanium wire is unwound and inserted through a semicircular tube. The wire is clamped with rubber shoes attached to a pneumatic system. The other end is left in a coil and placed in a tray. During inspection of the weld, they rotate the wire 90 degrees by picking up the tray and rotating the tray 90 degrees, which forces the wire to rotate 90 degrees in the tube. If the pneumatic force holding the wire in place is not released, the wire would be subjected to torsional forces that could affect the reliability of the wire.

Omnitron stated that nitrogen was used to reduce moisture in their pneumatic systems. Sufficient amounts of nitrogen can cause embrittlement in titanium if the oxide layer is breached. During final inspection of each wire, Omnitron uses an afterloader to cycle the source wire to a calibration station, wipe test station, and critical bend test station (i.e., 3 full cycles). Team members examined the inside of the HDR afterloader and found signs of wear. Team members saw degradation and debris of the drive wheels and misaligned stainless steel guide tubes.

In examining the wipe station, the clamps used to hold the wire were not accurately aligned and contained some sharp edges. Cotton wads were placed on the clamps and the wire traveled through the cotton wads. When the team asked Omnitron if they ever inspected the cotton wads to see if they were worn, they stated that they never have. In addition, Omnitron stated that a wet wipe test and dry wipe test are performed on the wire. If the oxide coating on the wire were breached, hydrogen could cause embrittlement of the wire.

The failure of the wire did not directly occur because of the manufacturing equipment. If the wire break had occurred at the manufacturing facility, it probably would have been detected at the facility during the calibration or packaging of the source.

The potential for stresses or environmental conditions imposed on the source wire at the manufacturing facility contributing to the source wire failure has to be considered.

Omnitron noticed during disposal of returned source wires from their customers that some of the teflon tube lining on the inside of the wire shipping container (Figure 4.4) was degraded in the area where the active source contacted the teflon. In late September, Omnitron stated that they changed the design of the shipping container to use a stainless steel liner for the wire instead of a teflon liner. Omnitron conducted no tests to see if the degradation of the teflon affected the integrity of the source wire. Omnitron assumed that the degradation of the teflon was due to the radiation.

4.1.3 Description of Test Samples

As of January 7, 1992, nine samples of the nickel-titanium wire have been tested as indicated in Table 4.1. Some samples consist of more than one test specimen.

The following paragraphs briefly describe the samples as they were received by SwRI, tests performed by Omnitron and Omnitron observations of the failure analysis. Once the failure analysis has been completed, SwRI will provide the NRC with a more detailed report regarding the samples, tests performed, testing procedures, equipment used, and their observations and probable cause of failure.

Sample 1 consisted of the broken active source wire (inactive portion) that was used during treatment at IRCC.

Sample 2 consisted of the broken active source wire (inactive portion) that was used during treatment at GPCC.

Sample 3 consisted of a sample, provided by Omnitron, of solid wire cut from a spool of nickel-titanium wire. All wires manufactured to date were cut from this spool of wire.

Sample 4 consisted of a wire manufactured in the same manner as the active source wire except that nonradioactive iridium was placed inside the cavity. Omnitron used an afterloader and similar catheters to simulate the equipment used at IRCC.

According to Omnitron, they performed their test on Sample 4 in the following way and obtained test results before its shipment to SwRI. The afterloader was used to extend the wire to the approximate position where the source wire broke (bottom of cavity at end of flexineedle connector) at IRCC. The source was then bent over more than 90 degrees and the source was retracted. After eight attempts of the above procedure, Omnitron broke the sample by hand. The sample was observed to have permanent deformation to the source cavity, and the source wire broke at the bottom section of the cavity. The sample was sent to Omnitron's contractor in San Diego, California. Metallography was performed, and Omnitron's contractor stated that it "showed distinct necking of the material at the fracture point which is indicative of a ductile overload breakage."

Sample 5 consisted of a wire that was manufactured in the same manner as the source wire except that nonradioactive iridium was placed in the cavity.

Omnitron stated that before shipping Sample 5, they performed the following test. The iridium was loaded into the cavity with 4 pounds of thrust, instead of the typical 1.8 pounds, to intentionally create hoop stress at the bottom of the cavity. The wire was placed in a pin vise held in the tailstock of a lathe. A flat plate was held in the lathe's chuck. The tip of the clamped wire was brought in contact with the flat plate and advanced slowly until the wire bent beyond 90 degrees from the axis of the lathe (Figure 4.5). The plate was then removed and the wire was examined under a microscope. Omnitron states that no permanent deformation or yielding of the material was observed.

Sample 6 consisted of multiple specimens of nickel-titanium wire that was subjected to hydrofluoric acid. According to Omnitron, their contractor soaked the samples for 12 hours in 0.1 percent hydrofluoric solution at room temperature in different configurations (i.e., straight, 90°, and 180°). Omnitron stated in a facsimile dated January 9, 1993, that their contractor observed that oxide film on the nickel-titanium wire was attacked by the solution, and that the wires "showed brittle behavior, cracking with little deformation." In addition, "Metallographic examination showed no signs of pitting or significant corrosion or stress corrosion cracks...The experiment yielded samples with characteristics of hydrogen embrittlement." The samples were sent to SwRI for examination and analysis.

Sample 7 was a source wire that was installed on June 30, 1992, at a customer's facility. It was used for 3 months and then stored on site for approximately 3 months. It was shipped back to Omnitron. According to Omnitron, on December 26, 1992, during removal of the wire from the shipping container in order to dispose of the source, their personnel observed that the source was not attached to the end of the wire but that it was in the bottom of the shipping container. The shipping container and wire were sent to SwRI for examination.

Sample 8 was an active wire containing approximately $3.5 \text{ E}+11 \text{ Bq}$ (9.5 Ci) of iridium-192. Omnitron stated that they performed the following test on Sample 8. The sample was stored in a shipping container lined with moist teflon. The wire was removed and bend tested by bending the wire 90 degrees in a 2.6-mm radius on December 23 and on December 26, 1992. On December 30, 1992, during the bend test, the wire broke after only being bent approximately 15 degrees. The wire was shipped to SwRI for examination and analysis.

Sample 9 consisted of sample pieces of wire. Omnitron subjected one-half of each wire to a 0.1 percent hydrofluoric solution for varying amounts of time after which the samples were bend tested using the same procedure as that used for Sample 5. The samples were sent to SwRI for examination and analysis.

Table 4.1 Summary of Test Samples

Sample number	SwRI ID. number	Sample identification
1	410	Indiana service failure
2	411	Pittsburgh service failure. (Received 12/12/92)
3	412	SwRI solid wire sample
4	413	First dummy wire (Omnitron contractor)
5	414	Second dummy wire (Omnitron-Houston)
6	415	Multiple HF immersion specimens (0.1%)
7	416	Returned source after 3 months of service (treatment) and approx. 3 months storage (Received 12/30/92)
8	417	Fractured wire from wet Teflon test with active source (performed in storage pig) (Received 01/05/93)
9	418	Multiple dilute HF immersion test specimens (labeled 24, 25, and 26) (Received 01/05/93)

4.1.4 Testing performed at Southwest Research Institute

A summary of the tests performed on the different wire samples is summarized in Table 4.2.

4.1.5 Summary of Observations of Tests Performed

NRC's contractor, SwRI, made the following observations from the tests performed and their visual examinations of the samples.

- (1) No inherent material deficiencies. Composition, hardness, and transition temperature are within material specifications.
- (2) Two service failures, Samples 1 and 2, show similar features.
- (3) Failures occurred by brittle fracture in bending.
- (4) Fractures occurred consistently near bottom of cavity.
- (5) New and returned wires survived bending in Omnitron tests, presumably more severe than any clinical situation.
- (6) Material is subject to cracking and embrittlement in dilute hydrofluoric solutions.
- (7) Sample 7 completely disintegrated in storage pig (a lead container).
- (8) Sample wire 8 experienced brittle failure after wet teflon test with active source.
- (9) Sample wire failures have occurred in afterloader system service (Samples 1 and 2) and in storage facility (Samples 7 and 8).

4.1.6 Implications of Observations to Date

SwRI stated that their observations as listed in Section 4.1.5, implied the following:

- All factors to date point to failure caused by environmentally induced degradation of properties on nickel-titanium wire in the vicinity of the iridium source.
- The timing and location of initial properties degradation (in-storage versus in-service) are uncertain.
- Possible environmental factors include (1) moisture entrapment (condensation, etc.) (2) hydrogen fluoride from teflon deterioration, and (3) body fluids.

Table 4.2 Summary of Tests Performed on Wire Samples

Wire	Conditions	Metal- lography	Chem- istry	Hardness	Fracto- graphy	EDX	DSC	Comments
No. 1 Ind. wire	Service Failure	✓*	x	*✓	✓	✓	x	Two distinct fractures
No. 2 Pitts. wire	Service Failure	✓	✓	✓	✓	✓	✓	Two distinct fractures
No. 3 Bend test	Test Failure - Wire No. 1	x	x	x	✓	x	x	No comment
No. 4 1st dummy	From San Diego	✓*	x	✓*	x	x	x	Fissures on cavity
No. 5 2nd dummy	From Houston	✓	✓	✓	x	x	✓	No internal fissures No fracture
No. 6 0.1% HF	Bend test failures	x	x	x	✓	x	x	Very brittle
No. 7 Failed wire	3 mos. in service, 3 mos. in storage	x	x	x	✓	✓	x	Completely deteriorated
No. 8 Failed wire	Test failure- wet teflon & active source	x	x	x	✓	✓	x	No comment
No. 9 Dilute HF	Bend test failures	x	x	x	✓	x	x	Cracks on the surface

✓ = Test Result
 x = No test results
 * = Performed in San Diego

4.1.7 Ongoing Testing

SwRI, Omnitron's Houston and Edgerly facility, and Omnitron's contractor are continuing to test and analyze the wire. They are testing to determine which, if any, environmental factors listed above caused the source wire to embrittle.

The data suggests that radiation deteriorates the teflon lining in the shipping container producing hydrofluoric acid that attacks the wire's protective surface, allowing hydrogen to embrittle the nickel titanium. Textbooks indicate that teflon degrades from radiation when oxygen is present. However, other environmental factors such as body fluids and moisture entrapment on the wire or within the shipping container are being considered.

Further examinations of the samples received are being analyzed. Chemical analysis of the material within the GPCC wire cavity is being conducted, as well as of material found in the bottom of the shipping container received from Omnitron. Microscopic fissures found within the inside cavity wall on one of the service wire failures is being studied. SwRI will keep NRC apprised of any new observations and will keep them abreast of all testing that is being performed. In addition, as mentioned before, SwRI will provide a more detailed report of the failure analysis once the failure analysis has been completed.

4.2 Immediate Cause of Wire Break

The nickel titanium source wire used in the Omnitron Model 2000 HDR afterloader in Indiana, Pennsylvania, was shipped from the Edgerly, Louisiana, facility on August 31, 1992.

According to the traveler the source wire had a "Pit at hole entrance .010 inch long." The source wire was reworked by shortening the tip area. The wire was reinspected and passed. All other inspection steps passed with no reported problems. The team reviewed the video tape containing the x-ray of the IRCC source wire. The team determined that sections of the source cavity had a minimum wall thickness of less than 0.089 mm (0.0035 inch). In addition, the team observed from the videotape that the cavity was off center through various portions of the wire.

SwRI and contractors working with Omnitron continue to test the wire. The definitive cause of the wire break is not final.

Test results to date indicate that the source may have been weakened from environmental factors (see Section 4.1.5). The weakest section of the wire is assumed to be at the bottom of the hole because of a void space surrounded by a thin wall and stress concentration produced by the hole. Also, if the source were bent, the torque generated within the cavity section would be most significant at the bottom of the hole. The known constriction distance (82.4 cm) places the cavity section of the wire in the vicinity of the flexineedle connector. During retraction, if the source wire was bent, it would be constricted against the end of the

connector. The resulting stress (bending moment and tensile force) acting on a weakened wire (embrittlement) may cause the wire to break. This effect could be exacerbated by the observed eccentricity of the source wire cavities.

The most likely hypothesis at this time is that the wire was subjected to environmentally induced embrittlement and that during treatment the source wire was bent (probably by the patient moving), causing a constriction at the flexineedle connector. The stresses generated by the bending moment and retraction of the wire (tensile stress) were significant enough to break the weakened source.

During the team's inspection of the afterloader and treatment log, the team determined that this source was cycled 65 times.

4.3 Potential Root Cause of Wire Break

The potential root cause of the source wire breaks was weaknesses in design validation of the Nitinol source wires.

In particular, the team found that Omnitron before the incident—

- performed no engineering calculations on the source wires, especially in the area of the cavity;
- performed a bend fatigue test on two wires, but did not validate the test results by engineering calculations or proper evaluation of the results. The bend fatigue test consisted of smooth, full radii. During treatment, a patient, or equipment, could cause a sharp bend in the source, and Omnitron performed no tests to simulate this condition;
- failed to determine whether the operating environment of the equipment could affect the integrity of the source wire. Environmental embrittlement was not considered, although titanium alloys are subject to hydrogen embrittlement when their protective oxide film is breached. In the absence of this oxide film, other atoms such as carbon, oxygen, and nitrogen can embrittle the titanium;
- failed to perform tests to determine if the catheters would interfere with the integrity of the wire and,
- performed no test to ensure that the degradation of the teflon would not affect the integrity of the source wire. Although, before the incident, Omnitron was aware that there was a degradation of the teflon lining in their shipping container.

A more definitive root-cause evaluation needs to be performed once the immediate cause of the source wire failure is determined.

4.4 Failure of Device To Detect Loss of Part of the Wire

Omnitron states in its instruction manual for the Omnitron 2000 HDR afterloader that when the source wire is retracted in the safe position, the inactive tail of the source wire reaches a park switch sensor, indicating that the center of the source is located at the center of the lead safe. However, the park switch sensor for the source wire only detects the end of the source wire opposite the source end.

In addition, this manual states:

3. Applicator wire lengths are checked each time the wires are retracted into the machine to ensure the entire wire has been retrieved with no break.
7. Fail-safe retract system ensures that applicator wire has been fully retracted.

Omnitron stated that the length of the source wire is checked before the source wire is returned to the safe position (parked) to ensure that no length errors have occurred during treatment. This checking is done by comparing the wire position when the home sensor is activated during extension to the position when the home sensor is deactivated during retraction.

However, this design feature does not allow for a comparison against the physical length of the source wire (a known parameter). This design weakness was clearly demonstrated again on November 17, 1992, when Physicist A at IRCC reran the treatment sequence of the November 16 session. During the November 17 simulation, although the source had already detached from the wire, no errors were detected by the afterloader system because the source wire length leaving (as measured by the afterloader) was the same as the source wire length returned.

Omnitron stated that during a treatment, any source wire length errors are reported to the operator at the main console and will cause a manual retract alarm to sound. However, this design feature is only effective if the source wire is being retracted by the stepping motor. When the emergency dc retract motor is activated, all optical detection mechanisms disengage, and source wire length information is lost. In the November 16, 1992, incident at IRCC, the emergency dc retract motor returned the source wire back into the afterloader. The device configuration did not allow for the detection of the missing source.

4.5 Performance of PrimAlert-10

The IRCC possessed and used a PrimAlert-10 Area Alarm Monitor (Model Number 05-433, Serial Number C393) manufactured by Victoreen to detect the presence of radiation in the HDR remote afterloader and linear accelerator treatment room. The licensee indicated, as

discussed in Sections 2 and 5 of this report, that although the PrimAlert-10 alarmed, the staff did not heed the alarm because some staff members had experienced previous problems with this monitor. On the basis of this information, the NRC team investigated the performance of the PrimAlert-10 monitor.

The PrimAlert-10 uses an energy-compensated Geiger-Mueller (G.M.) detector to provide a means of monitoring the radiation levels in an area. The PrimAlert-10 monitor is activated when the radiation level exceeds a preset value. The alarm level [0.025 mGy (2.5 mrad) or 0.2 mGy (20 mrad) per hour] is selected by means of a slide switch on the front panel. When the selected alarm level is reached, a pair of bright red bulbs, [approximately 2.54 cm x 2.54 cm (1 inch x 1 inch)] on the face of the instrument flash a warning and continue to flash until the radiation level drops below the preset value or the monitor is reset by unplugging the power supply. This alarm responds—begins to flash—within 2 to 3 seconds after the radiation level exceeds the preset value.

The PrimAlert-10 continuously monitors the background radiation and provides visible proof that the instrument is functioning by a flashing green operation indicator light. The manufacturer's literature states that in high radiation fields, those over 1 Gy (100 rad) per hour, the unit will not jam and will continue to alarm.

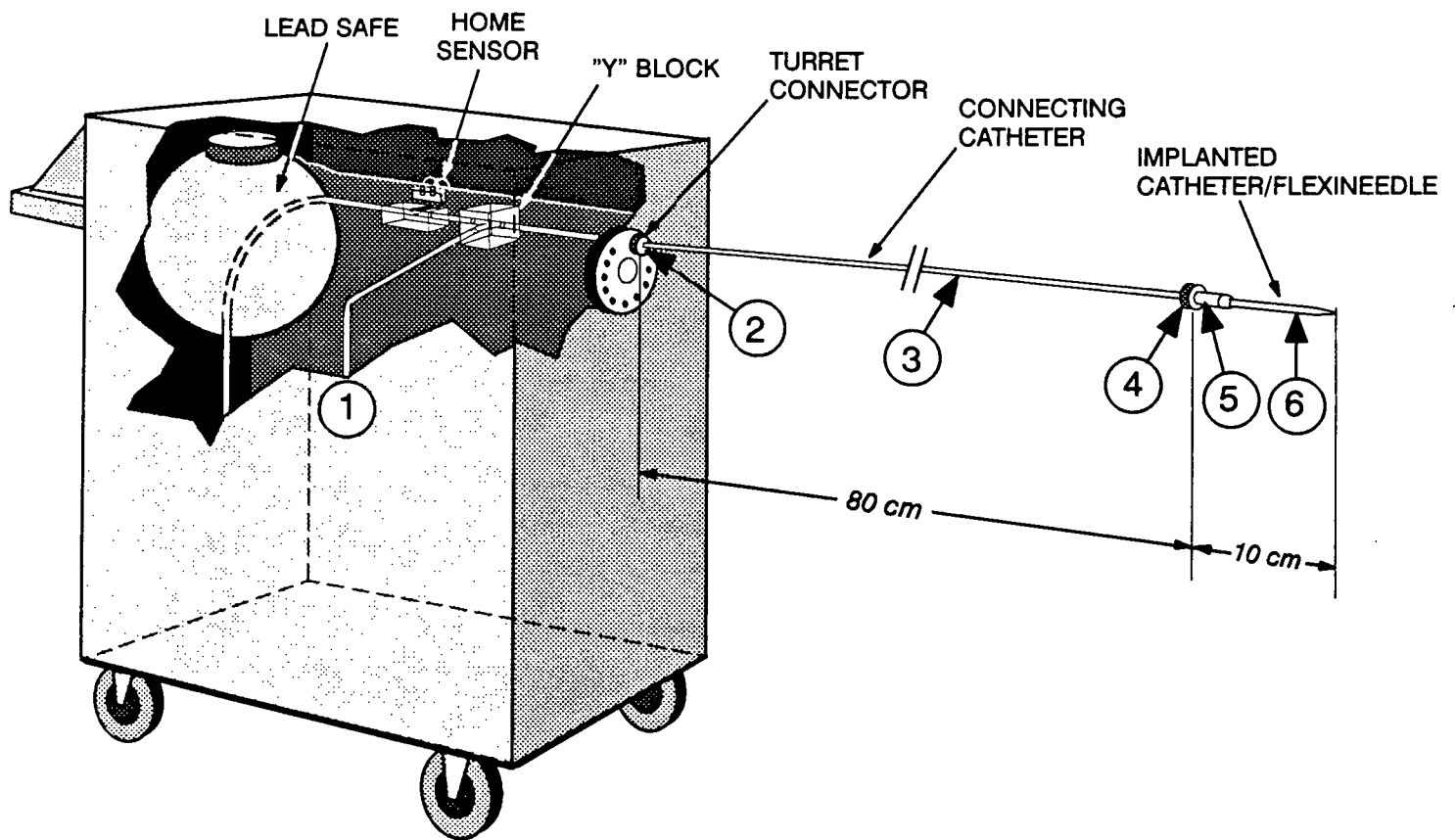
The manufacturer suggests that battery and other operational checks be performed before each use to ensure that the instrument is functioning properly. These checks include placing a check source on the top of the front panel in the position indicated to check the alarm. The manufacturer also cautions that the instrument should be used only by persons who have been trained to properly interpret its readings and to appropriately follow the safety procedures required in the presence of radiation. From discussions with the IRCC staff, the team determined that the PrimAlert-10 was not checked with the check source before each use to ensure that the instrument was functioning properly. However, the PrimAlert-10 was observed to alarm as it should when the linear accelerator was on. In addition, they learned that although the IRCC staff had been trained to know that the alarming of the monitor indicated the presence of radiation, they did not understand the alarm setpoint (the preset value) and, as discussed in Section 5, did not have confidence in the alarm indicating a radiation level higher than the alarm setpoint.

On December 2, 1992, the Deputy IIT Team Leader witnessed a test of the PrimAlert-10, using a test source. On December 5, 1992, the IIT Team Leader also witnessed a test of the PrimAlert-10, using Physician A's strontium-90 eye applicator as a radiation source. Physician A conducted the test. These tests confirm that the PrimAlert-10 at IRCC would properly alarm in the presence of radiation.

The team contacted the manufacturer to investigate the performance of the PrimAlert-10 monitor. The team reviewed service data supplied by the manufacturer. The data showed that at the end of the G.M. detector life, the PrimAlert-10 will remain in the alarm mode. The data failed to show a history of intermittent spurious alarms. Further, the team

interviewed the IRCC staff and the staffs of two cancer centers not operated by Oncology Services Corporation (OSC). Inspectors from Region I obtained information from two other OSC facilities. The staff at the IRCC, one other OSC facility, and at the other cancer centers stated that the PrimAlert-10 monitor would sometimes alarm when radiation was not present. In addition, a separate OSC facility experienced a similar problem with a different manufacturer's monitor. Each of these facilities uses a linear accelerator.

The cause of these PrimAlert-10 monitors alarming without the presence of radiation has not been determined. However, an employee of a cancer center not managed by OSC stated that when the PrimAlert-10 monitor was moved further away from the accelerator in their treatment room, the problem with the PrimAlert-10 monitor alarming without the presence of radiation was alleviated. The staff at some of the cancer centers believe that accelerator ionizing radiation levels are so high that these levels may cause some type of radiation damage to a monitor. The NRC team suggested that nonionizing radiation or electromagnetic fields associated with the linear accelerator power source may cause a spurious alarm. Some portable radiation survey meters exhibit this characteristic. The team was unable to resolve this matter.



- | | |
|---|----------------------------------|
| ① Inside HDR Afterloader | ④ Connecting catheter connector |
| ② Turret connection (catheter connection) | ⑤ Flexineedle catheter connector |
| ③ Connecting catheter (teflon tube) | ⑥ Flexineedle catheter |

Figure 4.1 Source Wire – Possible Failure Areas

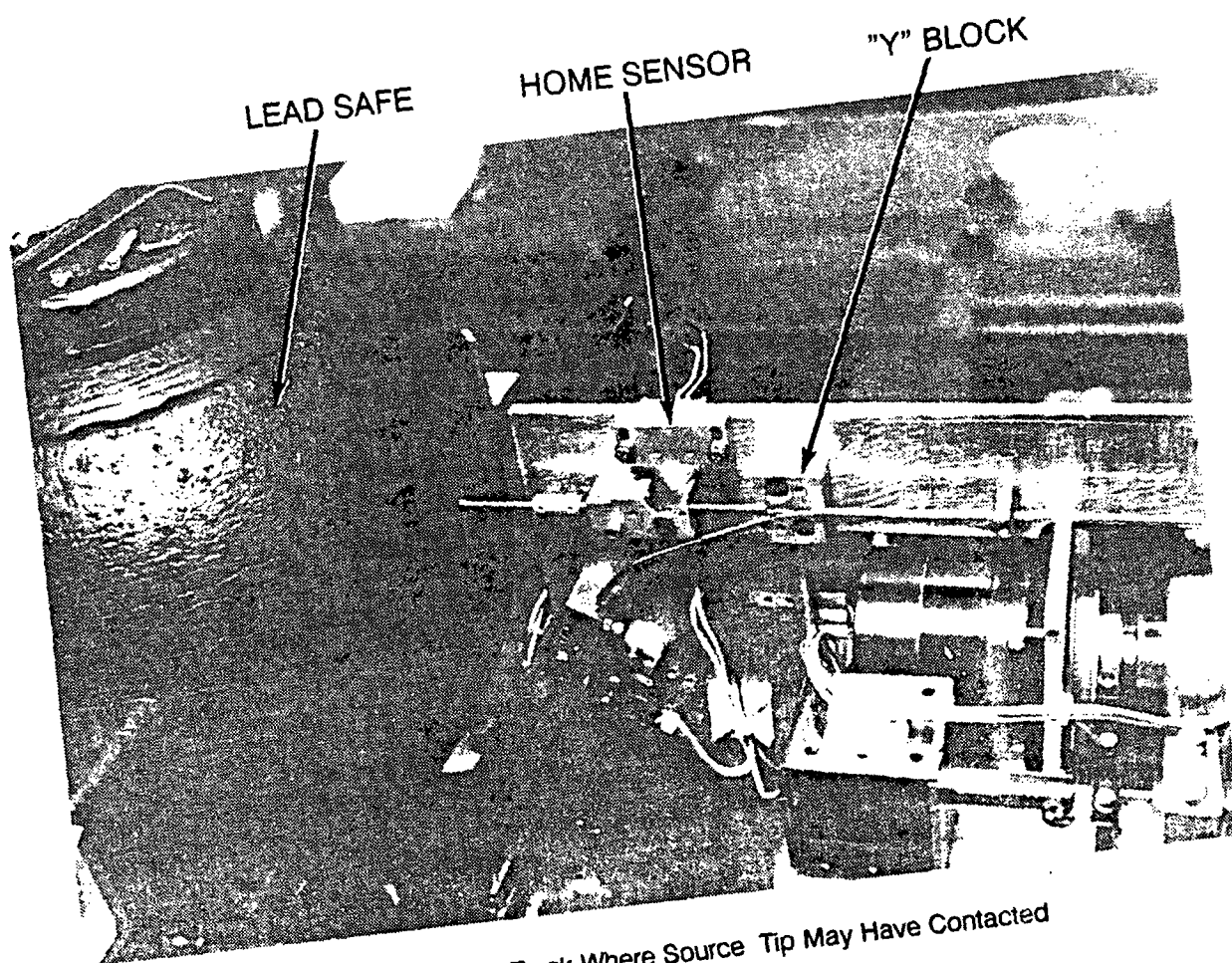
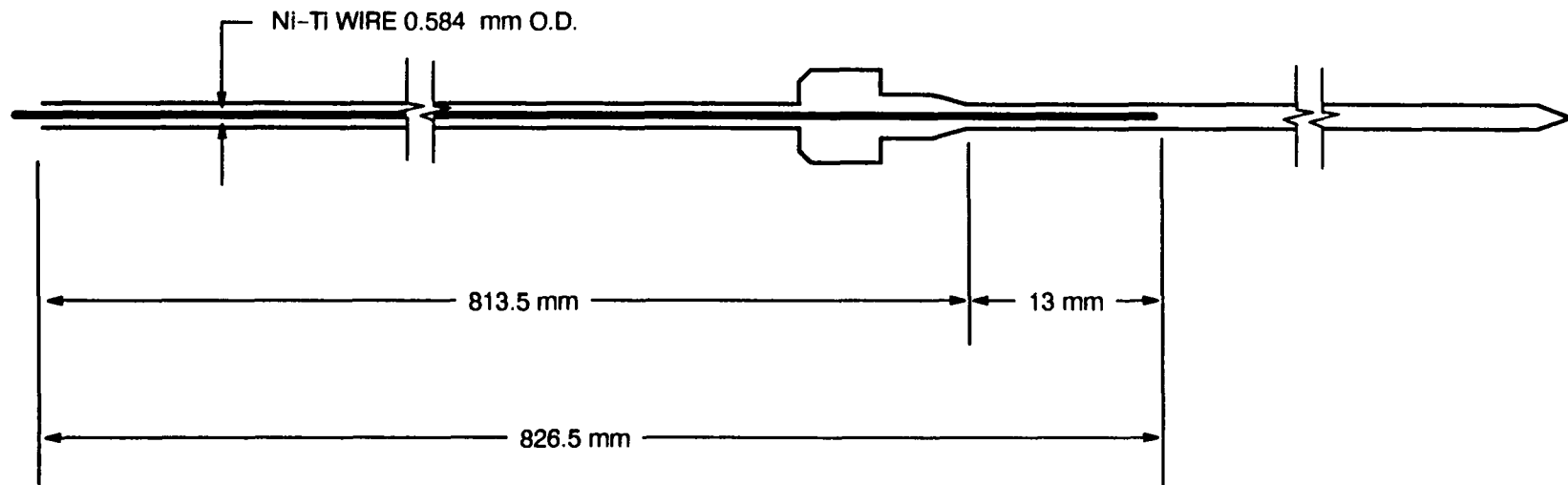


Figure 4.2 Track Where Source Tip May Have Contacted Parts of the Afterloader



NOTE: Drawing not to scale and all dimensions are approximate

Figure 4.3 Connecting Catheter and Flexineedle-Assembled



Figure 4.4 Shipping Container for Iridium-192 Source

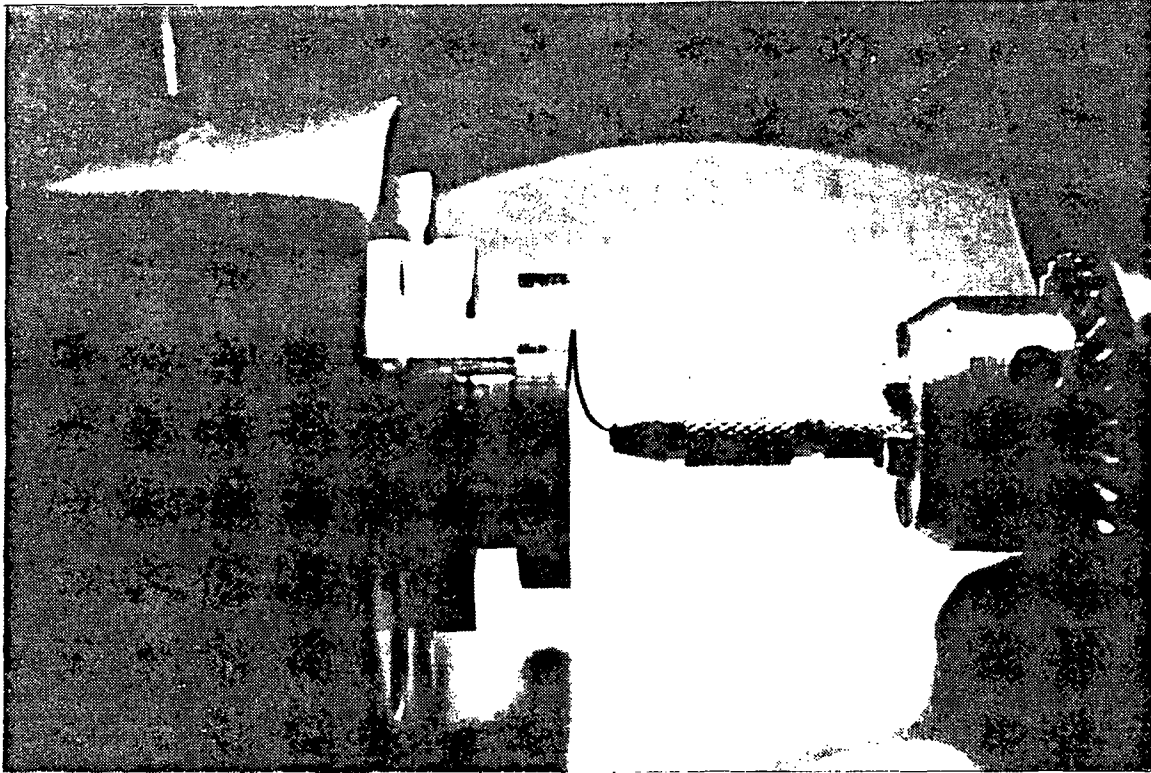


Figure 4.5 Testing of a Sample Wire

5 HUMAN FACTORS ANALYSIS

This section discusses the human factors associated with the two previously described source-wire-break incidents (see Section 2). Specifically, it discusses --

- the organizational factors for the licensee, Oncology Services Corporation (OSC); management oversight at the Indiana Regional Cancer Center (IRCC); a comparison of the safety cultures for OSC and IRCC and the Greater Pittsburgh Cancer Center (GPCC), and the physical arrangement of the treatment room and the HDR afterloader system at the two cancer centers; and
- the IRCC's emergency operating procedures, training, and the quality management program for HDR brachytherapy at IRCC.

5.1 Organizational Factors

5.1.1 Organization of Oncology Services Corporation

The corporate headquarters of OSC is located in State College, Pennsylvania. OSC managed 10 cancer treatment centers that use HDR afterloader systems in five states, including six located in Pennsylvania that are licensed by NRC Region I. Figure 5.1 presents an organization chart for OSC. OSC also managed a number of other cancer centers that do not use HDR afterloaders. All OSC cancer centers used linear accelerators.

OSC corporate headquarters provided support to the cancer centers in clinical and quality assurance areas. The corporate Radiation Safety Officer (RSO) was located at the Harrisburg Cancer Center and was responsible for providing radiological safety oversight to the cancer centers, including radiological safety program development and implementation, radiological safety audits, and radiation safety training.

Physician C was the chief executive officer of the OSC. Physician B was their medical director and primarily dealt with the physicians at the various centers. Physician D was director of all brachytherapy programs. Managers in the corporate office developed policies and expected the various centers to develop the specific procedures needed to follow these policies.

Physician A was the Medical Director of the IRCC and had full authority in all clinical areas and medical care of patients. Physician A was an authorized medical user on OSC's NRC license.

Medical Physicist A was a contract employee who worked two evenings a week at the IRCC to perform treatment planning and review therapy quality control records. Medical Physicist A reported to Physician A and the RSO. During the day he worked at another hospital that had no affiliation with OSC.

Registered Therapy Technician RTT-B was the lead technologist at the IRCC and supervised the other technologists, RTT-A, and the registered technologist radiographer (RTR). As lead technologist, RTT-B assigned and scheduled work for other technologists. Nurse A reported to Physician A.

5.1.2 Management Oversight

Indiana Regional Cancer Center. As indicated in Section 5.1.1, the IRCC staff was headed by Physician A who contracted with OSC. Neither Physician A nor the full-time staff at the IRCC were aware of who the RSO was. Physician A was not certain whether he or the corporation had the responsibility of providing radiation safety training. Physician A stated that he was not aware of any formal radiation safety training that the corporation provided to the technologists and other center staff. During interviews, and after being informed by the team who the RSO was, the staff said that the RSO came to the IRCC about once a year.

The RSO stated that Medical Physicist A was responsible for conducting training at the IRCC. Medical Physicist A said he was not responsible for radiological safety training, and, indeed, the physicist's contract did not explicitly list radiological safety training as one of his duties. However, Medical Physicist A's contract listed technical supervision of technologists as a duty. This responsibility was limited to briefings on individual treatment plans, dosimetry, and/or technical aspects of patient treatment. The medical physicist's contract did not specifically list responsibility for HDR brachytherapy support although Medical Physicist A had verbally agreed with OSC to provide this support. The contract had not been updated to reflect this verbal agreement.

5.1.3 Safety Culture

The team identified the following overall differences in the safety culture of OSC, IRCC, and GPCC:

OSC staff, including corporate managers, believed that the source wire would not break.

IRCC staff believed that the source wire would not break. Therefore, the IRCC staff were neither conditioned nor prepared to appropriately respond to a source-wire-break incident. This was demonstrated by IRCC's inadequate response to the November 16 IRCC's incident (see Section 2).

Medical Physicist B at the GPCC told the team that after his August 1992 training he believed that a source wire break was a credible, "worst case scenario," accident. He responded appropriately during the source-break incident at the GPCC.

The **IRCC** technologists were not familiar with the operation of the portable survey meter.

Medical Physicist B at the **GPCC**, upon receiving constriction error messages on the computer screen and hearing the audible alarm, entered the treatment room with a portable survey meter. He performed appropriate radiological measurements and assessment and ascertained the location of the source inside the connecting catheter and responded accordingly.

The **IRCC** technologists and Physician A either noticed or were informed that the PrimAlert-10 was flashing red. The technologists stated that the PrimAlert-10 had alarmed multiple times without the presence of radiation in the treatment room, and, they, therefore, assumed the PrimAlert-10 was malfunctioning during the November 16, 1992, incident.

Medical Physicist B at **GPCC** upon receiving an audible alarm of an error and a visual alarm on the computer screen indicating "Emergency Condition, Manual Retract, Check Source Status," immediately entered the treatment room with a portable survey meter and observed that the PrimAlert-10 was flashing red. Medical Physicist B's assessment of the alarming PrimAlert-10 was that the source was not in its shielded configuration and the physicist confirmed this assessment with portable radiation survey meter measurements and responded accordingly.

The **IRCC** technologists had limited knowledge and experience working with radioactive materials. On the basis of their day-to-day experience working with a linear accelerator, they were conditioned to believe that by turning the linear accelerator off, no radiation would be present. This conditioning and the fact that the HDR afterloader indicated that the source was "safe" caused them to believe that the source was inside the machine and that the PrimAlert-10 alarm was spurious. After completing the dummy wire insertions, Medical Physicist A had left the IRCC and Physician A was not continuously present at the HDR console during patient treatment. IRCC staff had not received a copy of draft procedures: "Oncology Services Corporation, Department of Physics, HDR Treatment Manual."

Medical Physicist B at **GPCC** had prior extensive experience working with radioactive materials, including cobalt-60 teletherapy units and an HDR afterloader. Medical Physicist B performed planning and administration of HDR treatments and was continuously present at the HDR console during patient treatment. In addition, Physician E, who is an authorized user, was watching the patient surveillance camera during the treatment. OSC gave draft procedures, "Oncology Services Corporation, Department of Physics, HDR Treatment Manual," to GPCC before the IRCC event.

Specific aspects of OSC's, IRCC's and GPCC's safety culture were revealed during the team's personal observations and interviews of personnel.

Oncology Services Corporation:

OSC staff believed that the source wire could not break. OSC's Physician D said he assumed the source wire was safe because the Federal Government had licensed it.

Indiana Regional Cancer Center:

Source Wire. IRCC staff believed the source wire could not break and that if the machine malfunctioned the wire would be stuck outside the machine and would need to be manually retracted. The IRCC technologists knew that if the wire was stuck outside the HDR afterloader that it could be manually retracted by using the wheel on the unit or by just pulling the HDR afterloader away from the patient, effectively pulling the source wire out of the patient.

Survey Meter. A portable survey meter was positioned close to the HDR afterloader computer console. All of the technologists knew that the meter was there and that it detected radiation. None of the technologists routinely used the survey meter. From direct observation of technologists handling the survey meter, the team found that the technologists were not very familiar with its use. Medical Physicist A said he had shown the technologists how to use the survey meter but this was not documented.

Alarm Response. RTR noticed the PrimAlert-10 alarm upon the third entry into the treatment room. RTR notified the other two technologists (RTT-A and RTT-B), who assumed the PrimAlert-10 was not working properly rather than that the device was detecting radiation in the room. The PrimAlert-10 had previously alarmed when no radiation was present. Specifically, it would flash for no apparent reason, presumably indicating radiation in the treatment room. On one occasion, over a year before this November 16, 1992, incident, it had flashed all day. On this occasion, RTT-A brought a portable survey meter into the room, as suggested by Medical Physicist A, to ensure that no radiation was in the room. The survey meter detected no radiation, and the technologists concluded that the PrimAlert-10 was not working properly. The IRCC staff stated they believed that the malfunctioning PrimAlert-10 was sent for repair. The PrimAlert-10 vendor stated that the unit currently used at the IRCC had never been returned for service. Neither the IRCC staff nor the team could determine whether the PrimAlert-10 in the IRCC treatment room at the time of this incident was the instrument that caused confusion a year ago or whether it was a different instrument.

Since the PrimAlert-10 had malfunctioned in the past, the technologists assumed that, if the PrimAlert-10 flashes when the linear accelerator is not turned on or the HDR afterloader is not being used, the monitor was malfunctioning rather than that radiation was present in the treatment room. They, therefore, did not use a portable survey meter upon entering the room after the PrimAlert-10 alarmed on November 16, 1992. RTT-A resolved the problem with the flashing monitor by unplugging the PrimAlert-10 thereby allowing it to reset. RTT-A stated that this had occurred multiple times in the past.

The RTR had not had this mental conditioning because the RTR had not worked at the IRCC as long as the other technologists and because the staff at the other facility where the RTR worked believed their radiation monitors were reliable.

HDR Afterloader Usage. Physician B during an interview discussed the fact that most of the OSC facilities had been linear accelerator facilities until the HDR afterloader systems were brought on site. The IRCC received their HDR afterloader in late 1991 and did not begin using it until February 1992. The IRCC did not have a cobalt-60 teletherapy unit. When linear accelerators are turned off no radiation is present, therefore, the IRCC technologists were not used to dealing with a device that contained a radioactive isotope. This lack of experience was exacerbated by the fact that the IRCC staff had only performed about 30 treatments on 10 patients since February 1992.

RTT-A who treated the patient on November 16, 1992, did not know the activity of the source and did not appear to understand its potential radioactive hazard. RTT-A was unaware of the magnitude of exposure received on entering the treatment room if the HDR afterloader source was unshielded.

The IRCC staff believed that if the HDR afterloader failed, they would hear an alarm and see an indication on the HDR computer monitor of the failure. Medical Physicist A believed that if the HDR afterloader failed there would be all kinds of "bells and whistles." The HDR afterloader software is designed to produce visible error message indications on the computer console monitor during dc emergency retraction of the source wire but is not designed to produce any other record. This visible error message indication should have caused a portion of the upper half of the screen to flash red. However, the afterloader system gives no audible alarm for an emergency retraction. The IRCC technologists did not see the error message indications on the computer console monitor. The IRCC staff revealed that they were not very familiar with the error messages associated with the HDR afterloader because Omnitron had not supplied all the meanings of all the error messages to them.

IRCC technologists stated that they did not verify the operation of the treatment room door interlock associated with the HDR afterloader before operating the unit. Additionally, IRCC technologists stated they did not use a check source to verify the operability of the PrimAlert-10 area radiation monitor before HDR afterloader operation.

Greater Pittsburgh Cancer Center:

Source Wire. When Medical Physicist B received his training at the Omnitron facility in Houston, Texas, this physicist asked Omnitron to demonstrate how the HDR afterloader operated. This demonstration lasted about an hour, during which Medical Physicist B discussed the possibility of the wire breaking with the Omnitron representative. Physicist B, upon hearing of the HDR source-wire break at the IRCC, placed a shielded container near the HDR afterloader to be used to contain the source should the source wire break.

Survey Meter. Medical Physicist B entered the treatment room with a portable survey meter to check for radiation after hearing the audible alarm. When the portable survey meter indicated radiation, Medical Physicist B acted to protect the patient and to secure the source.

Medical Physicist B said that GPCC has an unwritten policy requiring surveys of the patient after every brachytherapy procedure. Further, OSC managers stated that corporate policy requires the staff to survey each patient with a portable survey meter upon completing treatment and before discharging the patient. Medical Physicist B surveyed the patient with a portable survey meter before the patient's release.

Alarm Response. During the GPCC incident, the HDR afterloader gave an audible alarm when the source wire was retracted. Medical Physicist B responded appropriately to this alarm and the alarming PrimAlert-10 and indicated that the PrimAlert-10 alarm stopped when the source was placed in the shielded container.

HDR Afterloader Usage. The GPCC had performed 36 HDR afterloader treatments in the 4 months they had possessed the afterloader. Medical Physicist B was the only person authorized to use the HDR afterloader at GPCC. He performed all the planning for the treatments and administered all except one of the treatment procedures. In addition, Physician E was present at the HDR computer console when the December 7, 1992, treatment was being conducted.

Medical Physicist B performed operational and safety checks on a daily basis before operating the HDR afterloader. These included testing the treatment room door interlock and the PrimAlert-10 via a check source.

5.1.4 Physical Arrangement of the Treatment Room and HDR Afterloader System at the Indiana Regional Cancer Center

At the time of the incident at the IRCC, RTT-A was operating the HDR treatment system from the computer console, and the RTR was also present in the area of the computer console.

Owing to the placement of the computer console and the patient surveillance camera monitor (Figure 3.5), it was difficult for one operator to watch both simultaneously. Additionally, the PrimAlert-10 was located inside the treatment room and was visible through a window in the treatment room door. Therefore, the operator would have to look away from the console and monitor to observe any alarm on the PrimAlert-10. Physician A was not present at the computer console during the treatment, and Medical Physicist A was off the site. In contrast, at the GPCC, Medical Physicist B was watching the computer console and Physician E was watching the patient surveillance camera monitor.

The control console panel may be positioned so that it is in as prominent a position as the computer monitor. The panel and the monitor need to be positioned so that when the console panel alarms, the operator would automatically direct attention to the error messages appearing on the computer monitor.

If the door alarm display panel is used for alarm information, the operator's attention is not automatically directed to error messages appearing on the computer monitor.

While the emergency dc motor is retracting the active source, no alarm is audible, and the error message only stays on the screen until the end of the active wire contacts the park switch. It normally takes approximately 10 seconds for the emergency motor to retract the source. Therefore, the saliency of the alarm indications may be less than optimal.

5.2 Emergency Operating Procedures

Omnitron gave framed operating procedures to be used in an emergency involving the HDR afterloader to IRCC and GPCC. These procedures were not comprehensive and did not address the incidents that occurred at IRCC or GPCC or suggest an appropriate response to either incident. Each center had posted these procedures by the HDR afterloader console, but they were not used during the incident. The procedures did not provide instructions to workers as to "how to" perform any radiological safety functions, including radiological surveys and precautions.

5.3 Training

5.3.1 Omnitron Provided Training at the Indiana Regional Cancer Center

Omnitron provided the team a letter describing the training they had given the staff at the IRCC from which the following information is summarized:

Medical Physicist A had notified Omnitron that Medical Physicist A, RTT-B, and Physician A would be operating the HDR afterloader. On December 9 and 10, 1991, Trainer A from Omnitron trained Physician A, Medical Physicist A, and RTT-B. The training included --

- instruction in the use of the treatment planning computer to plan patient treatments,
- instruction in operation of the HDR afterloader system, a demonstration of the safety features and emergency procedures to be followed, a review of the equipment warranty, and support services that Omnitron provides, and
- a demonstration of the use of the various catheters provided with the system.

On February 27, 1992, Trainer A from Omnitron conducted a review of the operation and safety features of the system. The treatment of the first patient was to be conducted on

February 28, 1992. From the interviews, the team learned that Physician A, Medical Physicist A, and RTT-B had attended this review and that RTT-A and Nurse A were intermittently present when the review was being conducted.

5.3.2 Radiation Safety Training at the Indiana Regional Cancer Center

RTT-A, RTT-B, and RTR stated that during their tenure at IRCC they had not received any formal radiation safety training. All informal training they received was conducted by Medical Physicist A. However, the team could not verify the content of this training. Two technologists stated that the radiation safety training at IRCC was not very good, but the team could not verify whether they had communicated this opinion to their managers. The RSO was under the impression that Medical Physicist A was responsible for training in radiation safety at the IRCC. Medical Physicist A said that the contract with OSC did not explicitly list radiation training or other types of training as one of his duties. The RSO did not consider it appropriate for the RSO to provide radiation safety training to someone with Physician A's qualifications and, further, stated that if Physician A needed this type of training, Physician A should not be approved by the NRC as an authorized user. NRC Region I Inspectors identified similar training weaknesses during a recent inspection at OSC cancer centers in Exton and Mahoning Valley Cancer Centers in Pennsylvania.

5.4 Quality Management Program for High Dose Rate Brachytherapy at the Indiana Regional Cancer Center

OSC's RSO submitted the quality management (QM) program for HDR brachytherapy to the NRC on January 22, 1992. The licensee's QM program was two pages long and addressed the following 10 points:

1. The authorized user was to date and sign each prescription before treatment.
2. The patient's identity was to be verified.
3. The treatment setup was to be verified against the treatment plan before treatment.
4. The staff was required to ask questions concerning the treatment procedure before treatment if they were unsure what to do.
5. Radiographs were required to be taken with dummy sources to verify source positions before treatment.
6. Dose calculation checks were required before patient treatment. Two individuals were to verify input data for the HDR afterloader.
7. The physician was required to sign and date a written record after treatment, indicating the dose administered.

8. The staff was allowed to treat patients without checking the dose calculation before treatment if the delay in treatment would jeopardize the patient's health.
9. A 'radiological physicist' was to perform acceptance tests on each treatment planning or dose calculating computer program for use with the HDR afterloader system.
10. Periodic reviews were required of the QM program for HDR afterloader treatments.

The team noted that the QM program did not provide, and NRC QM requirements may not have required, guidance or any procedural requirements that could have, upon implementation, prevented the occurrence of this incident. The following problems do not appear to be addressed by the QM program:

- Patient left the treatment room with the source inside. The QM program did not address surveying the patient after treatment. If the program had required a patient survey and the staff had followed the program, the radioactive source left in the patient would have been detected and could have been removed before the patient left the cancer center.
- Identification of the implanted catheters was not accomplished. The QM program did not specifically address the need to identify these catheters and, apparently, the IRCC did not label them. Therefore, the implanted catheters could be incorrectly connected to the HDR afterloader.
- The correct emergency response to most error messages was unclear. The QM program did not specifically address how to respond to error messages from the HDR afterloader system.

Lines of Authority

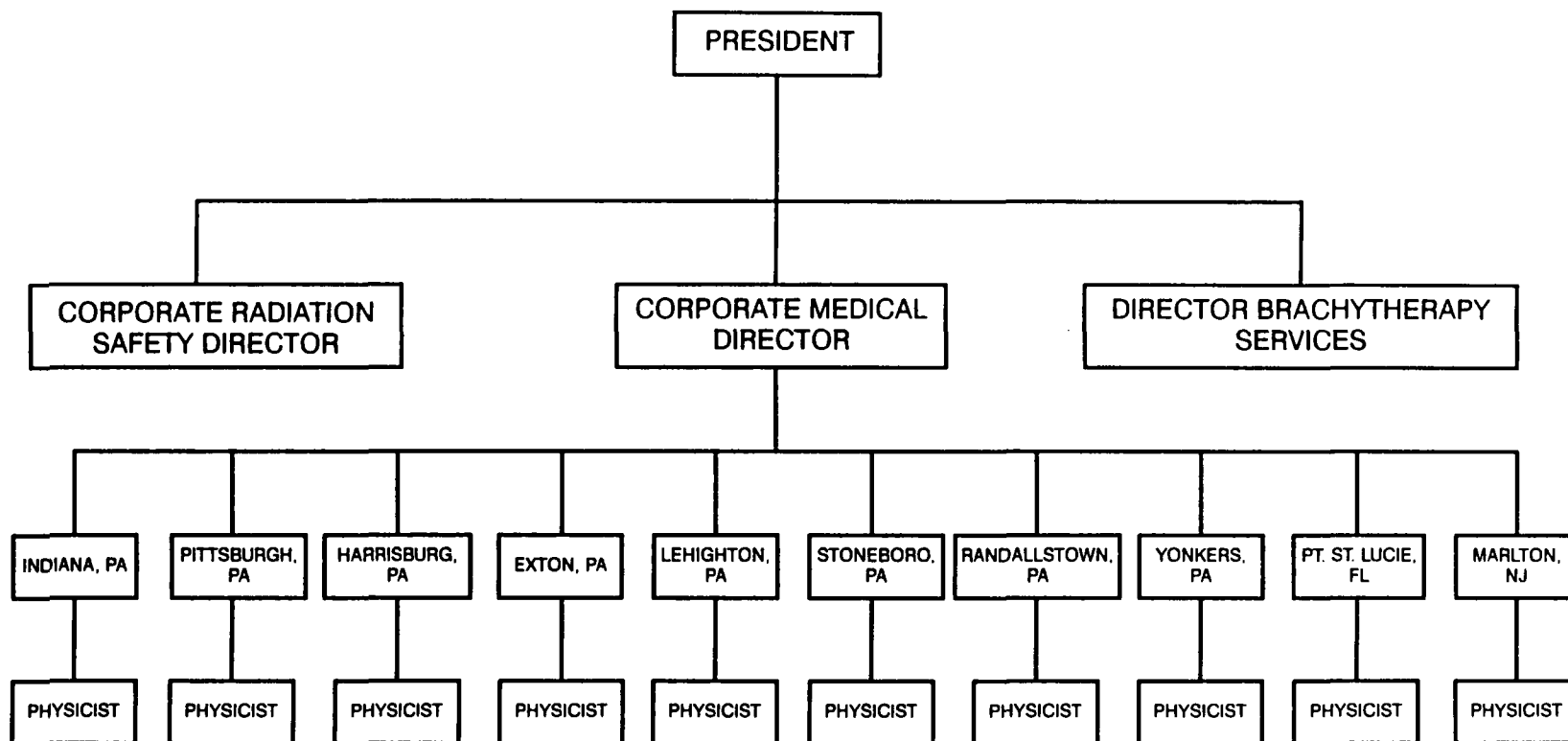


Figure 5.1 Oncology Services Corporation Organizational Chart

6 RADIOLOGICAL DOSE EVALUATIONS

The Incident Investigation Team (the team) estimated whole-body and extremity (i.e., hand, forearm) radiation doses for all persons who were known or postulated to have been exposed to radiation from the time the source wire broke at the Indiana Regional Cancer Center (IRCC) on November 16, 1992, until the time the source was eventually recovered at the Browning-Ferris Industries (BFI) in Carnegie, Pennsylvania (BFI-Carnegie) at 4 p.m. on December 1, 1992. The team performed contamination surveys of the IRCC, the Scenery Hill Manor (SHM) nursing home where the patient resided, and the ambulance used to transport the patient after the high dose rate (HDR) treatment. In addition, the Pennsylvania Department of Environmental Resources performed radiation surveys at the SHM nursing home. None of the surveys showed any indication of contamination. Because the source was not in its original shielded container after the source wire broke on November 16, 1992, and because Patient A's body provided little or no attenuation, all radiation dose estimates are based on the assumption that the source remained unshielded until its recovery on December 1, 1992 (see Appendix B).

6.1 Overview of Team's Methodology

Radiation doses to individuals (occupational and nonoccupational workers and the general public) were determined from data gathered from interviews with the following individuals:

- the IRCC staff;
- the Scenery Hill Manor (SHM) nursing home staff, residents and visitors;
- employees of Citizens Ambulance Service;
- employees at BFI-Carnegie facility, Pennsylvania;
- employees at BFI-Warren facility, Ohio; and
- other individuals who may have been near the source at any time from November 16 through December 1, 1992.

The team used data from personnel dosimeters worn by occupational workers and gathered information from individuals who were not wearing dosimeters to develop time and motion histories for those with possible exposures. The team used these histories to determine the time spent and the distances involved when in the vicinity of the source. For the purpose of this report, the words "exposure" and "dose" are exchangeable. To confirm the range of estimated radiation exposures to the nine individuals identified as having the highest potential for exposure, the team arranged for cytogenetic studies.

Radiation exposures that the team calculated for individuals who may have received radiation exposure from the iridium-192 source are described in detail in the following sections. Included in each section are the time, distance, shielding factors, and other assumptions that were used in supporting each calculated dose estimate. Each estimate of radiation exposure is presented as a dose range received by that individual based on interviews, re-enactments,

and measurements of probable or known source locations for each of the various facilities where the source was located between November 16 and December 1, 1992.

For a perspective on the magnitude of the range of doses determined from personnel dosimeter measurements, calculations, and cytogenetic studies, the values obtained can be compared to NRC regulatory limits (10 CFR Part 20), recommendations of international and national radiation protection advisory bodies, and normal exposures received by the public from natural background and man-made sources of radiation. The exposure estimates may also be compared with the maximum whole-body dose of 12.5 mSv (1.25 rem) per calendar quarter allowed by the NRC for occupationally exposed individuals. Current NRC regulations have an implied limit of 5 mSv (500 mrem) per year for members of the general public. This limit will become explicit and drop to 1 mSv (100 mrem) per year when the new 10 CFR Part 20 becomes effective on January 1, 1994.¹ This value is in accordance with the recent recommendations of the National Council on Radiation Protection and Measurements.

The estimated total effective dose equivalent rate for an average person's exposure from natural background radiation is 3 mSv (300 mrem) per year.² None of the above limits apply to or include natural background or radiation exposures prescribed by a physician. A dose that slightly exceeds the annual dose limit has little biological significance for the individual because any lifetime risk will be only slightly perturbed and can be readily offset by no exposure or reduced exposure to radiation in future years.³ The biological significance of radiation exposure is discussed in References 3,4,5, and 6.

As described on the manufacturer's certification of calibration, the unshielded source was determined to be a sealed source containing iridium-192, Serial Number (S/N) 01-01-9282-001-082892-08942-08. This source was manufactured by Omnitron International, Inc. (Omnitron), on August 28, 1992. According to the manufacturer's calibration and leak-test certificate, the source activity was $3.3085 \text{ E}+11 \text{ Bq}$ (8.942 Ci) on August 28, 1992. On November 16, 1992, when the wire containing the source broke, the source activity was $1.579 \text{ E}+11 \text{ Bq}$ (4.267 Ci).

Iridium-192 decays with a half-life of 74.02 days by electron capture to osmium-192 (4.7 percent) and by the emission of 0.67 MeV and 0.54 MeV beta particles to platinum-192 (95.3 percent). Approximately 88 percent of the beta particles are absorbed by the source capsule and the remainder are absorbed by the plastic catheter. Therefore, the beta contribution presented little hazard to anyone during the handling and storage of the source in the transportation chain. Eighteen gamma photons are also emitted during the decay of iridium-192, with energies from 0.879 MeV to 0.133 MeV. Appendix B provides several tables that give the activity and dose rate for several distances from the unshielded source.

Because the majority of the gamma rays from iridium-192 are of relatively low energy (from 0.3 to 0.6 MeV), buildup in air results in approximately 5 percent of the exposure rate in air for distances greater than 1 meter. Therefore, the buildup factor cannot be ignored in these

calculations and is included in calculating the radiation dose estimates for each person exposed to the radiation source. Table B-2 in Appendix B calculates the total dose rate from 0.02 to 15 meters, taking into account the buildup of scattered radiation in air and compares the total dose rate to the level with no buildup. Buildup factors used in these evaluations are based on the Berger method (see Appendix B).

Radiation dose estimates for all individuals evaluated during the investigation were based on exposure rates that were calculated using MICROSHIELD⁷, Version 3.13, S/N 327.004, which is a microcomputer adaptation of the mainframe code ISOSHLD⁸. This code was used to calculate the dose contribution for specific distances to occupied areas from an unshielded iridium-192 source. Correction factors for radioactive decay, attenuation in materials surrounding the source, and buildup, were included in these calculations. Because of the relatively large magnitude of certain errors associated with the analysis of radiation exposures to individuals, the dose equivalent, sievert (Sv), for gamma radiation is assumed to be essentially equal to the calculated absorbed dose, gray (Gy). Therefore, doses are reported in either units of Sv, cSv (one-hundredth of a sievert), and mSv (one-thousandth of a sievert), or Gy, cGy, and mGy. The team performed dose calculations by hand as a quality check on the MICROSHIELD results. For additional information about MICROSHIELD, see Appendix B.

6.2 NRC's Medical Evaluation of the Dosimetry and Effect of Radiation Dose on Patient A

Radiation absorbed doses to normal healthy tissues and vital organs were calculated by NRC's Medical Consultant assuming a 92.75-hour exposure from a $1.56 \text{ E}+11 \text{ Bq}$ (4.22-Ci) iridium-192 stationary source located at the needle-tip of a 10-cm catheter. Values obtained from the Omnitron 2000 HDR afterloader printout for the half-life of iridium-192 (73.83 days) and decay time (80 days) were used for Patient A's dosimetry calculations. The source location was determined by using the anterior-posterior (AP) and lateral planning films, which had dummy source markers. The information in Section 6.2 was obtained from NRC's Medical Consultant's final report dated January 13, 1993. This section was reviewed and incorporates comments from the NRC Medical Consultant as of January 27, 1993.

6.2.1 Radiation Dosimetry on Patient A

Doses to normal tissues were calculated separately, assuming that each of the five catheters contained the stationary source. The source is 1.0 cm long with a 0.3-cm dead space at the end. Therefore, the source location of the midpoint of the source in each catheter was taken as 0.8 cm from the end of the catheter as marked by the end of the dummy sources on the planning films. These points were translated to the premortem and postmortem patient computerized tomography (CAT) scans from the IRCC planning films. For the dose calculations presented in Tables 6.1(a) and 6.1(b), catheters were arbitrarily labeled A, B, C, D, and E from left to right on the IRCC planning films. Tables 6.1(a) and 6.1(b) provide the normal tissue distances and doses in cGy (rad) for major organs in the body. For

purposes of this report, 1 cGy (rad) is equal to 1 cSv (rem). Each of the five possible source location points was assigned an X,Y,Z coordinate and distances were calculated to a series of normal tissue and X,Y,Z coordinates. All possible source locations were in the posterior pelvis in the perirectal area just superior to the anus.

Physical dosimetry was performed using the Nucletron Planning System (NPS) in the HDR program mode. The following equation was used to adjust each of the calculation points for the actual activity, time, and radioactive decay:

$$D = \frac{T}{t} \frac{A}{a} dX$$

where D	=	Dose in tissue [cGy (rad)]
T	=	Total irradiation time in seconds (92.75 hours = 333,900 sec)
t	=	Dwell time of the NPS system to deliver 6000 cGy (rad) at 1.0 cm (979.9 sec)
A	=	Activity of the source in units of Ci [4.22 Ci (1.56 E+11 Bq)]
a	=	Activity in units of Ci for the NRC Medical Consultant's NPS system iridium-192 source on 01/05/93 [4.844 Ci (1.79 E+11 Bq)]
d	=	Decay factor (0.982, the decay of Ir-192, with a half-life of 74.02 days for 0.5 of the irradiation time)
X	=	Dose calculated at points of interest by the NPS system in units of cGy (rad)

The NPS uses the Van Kleffen, Star algorithm (1979) to correct for tissue absorption and scatter (buildup) in tissue. The majority of dose algorithms differ from each other as distances increase beyond 10 cm.

6.2.2 Radiation Effects on Patient A

Table 6.2 shows the tolerance of different tissues for total or partial organ irradiation. It should be emphasized that Table 6.2 is for fractionated radiation at the rate of 1000 cGy (rad) per week as might be given for external beam treatment. The tolerance doses would be much lower if the dose were given at a higher rate or all at one time. A similar comparison can be made between low dose rate (LDR) brachytherapy and HDR brachytherapy. The HDR doses are at rates that are high enough for the exposure to be shorter than the radiobiological normal tissue repair half-time of sublethal damage. This results in more normal tissue damage if the same dose is delivered by HDR as would have been delivered by

LDR. For this reason, in cancer treatment HDR doses are substantially less than LDR treatment doses for equivalent biological effect.

The radiation effects on normal tissue are dependent on a number of factors, including (1) total absorbed dose, (2) time over which a dose is administered, (3) the volume of tissue irradiated, and (4) the specific organ irradiated. Patients can experience acute radiation syndrome and severe bone marrow effects of radiation even though only a portion of the body is irradiated. For example, patients with ovarian cancer are sometimes treated with abdominal-pelvic radiation fields which cover the marrow of the lumbar-sacral (LS) spine and pelvis. Typical doses are 3000 cGy (rad) over six weeks time. Nausea, vomiting, and diarrhea are frequently seen side effects, in addition to bone marrow suppression. The most common limiting acute toxicity is blood-count depression (white blood cells platelets), which is due to bone-marrow injury. Patient A received approximately 2000 cGy (rad) minimum dose to the bone marrow of the LS spine region over a very short time (4 days). Because of this relatively short time, the effects on the bone marrow would be much greater than if Patient A would have received 3000 cGy (rad) over 6 weeks.

Bowel intolerance is another major dose-limiting factor in the treatment of tumors of the abdomen and pelvis. Acute small bowel reactions result in nausea, vomiting, painful abdominal cramps, and diarrhea. In the human, it is known that the small bowel mucosa denudes between 3-7 days following a large radiation dose. Superficial ulceration of bowel with denuding of the villi allow a pathway for bacterial invasion. With sufficient dose and volume of bowel irradiated, death due to the gastrointestinal (GI) syndrome can occur within 3-10 days with significant diarrhea leading to dehydration, with electrolyte loss. Patient A had already been irradiated to normal small bowel tolerance in 1991 when she had to be hospitalized several times due to severe chemo-radiation GI toxicity and related secondary infections.

In addition to the effects on the bone marrow, the intestine, and related infectious complications, the radiation effects on the rectum, the bladder, and the kidney are important in this case. Doses in the range of a half million to a million cSv (rem) should cause extensive necrosis in a few days and possible rectal perforation. If rectal perforation had not occurred by the time Patient A died in 5 days, it would have occurred later. Similar conclusions could be drawn on the effects on the bladder and perhaps also on the kidneys, which received lower doses.

The local HDR treatment that Patient A received on November 16, 1992, was an attempt to control the local cancer without causing additional irradiation to the small bowel and other normal tissues away from the anal tumor. Unfortunately, the HDR overdose was so massive, normal tissues higher and outside the pelvis were irradiated beyond normal tissue tolerance levels.

Within a few hours of Patient A's return to the SHM nursing home, Patient A's condition suddenly and dramatically deteriorated, starting with repeated episodes of nausea, vomiting,

and increased rectal pain. Observations after several days included a warm, reddened, swollen rectal area, diarrhea, tachycardia, fever, poor skin turgor suggesting dehydration, and finally a piece of grey-black tissue stuck to the catheter (which presumably contained the source) that fell out after 4 days. NRC's Medical Consultant concluded that the clinical deterioration of the patient coincided with the radiation accident. The sequence of symptoms and observations were consistent with a severe acute radiation syndrome.

Analysis of Patient A's medical records by NRC's Medical Consultant indicated that the massive radiation overdose was either a probable cause of death, or a probable contributing cause of death in Patient A. Some life-threatening effects including bone marrow suppression had not yet reached their peak. In addition, if Patient A had lived longer, she would have suffered within weeks, or months, with other severe, acute, or delayed effects, including new or additional episodes of bowel obstruction, bowel perforation, fistulae, strictures, and tissue necrosis along with delayed vital organ failure owing to the radiation dose having exceeded the expected normal tissue tolerance for that organ system. This radiation accident would probably have resulted in a fatality in a younger, healthier patient within days or weeks of an equivalent exposure. Based on analysis of the clinical records, the approximate location of the iridium-192 source in the pelvis, and the massive overexposure Patient A received, the effect of the radiation could contribute to her death in a number of ways, alone or in combination, within a 5-day period. Analysis of medical records and physical dosimetry would suggest that one likely mechanism of death from radiation for the range of possibilities would be acute GI injury with secondary sepsis, dehydration, and electrolyte imbalance.

6.2.3 Postmortem Examination

A postmortem examination was conducted on December 21, 1992, at the request of the Indiana County Coroner on Patient A. In order to better understand the radiation effects and to pinpoint the exact location and distance of various critical organs from the source location in the pelvis, the Indiana County Coroner invited NRC's Medical Consultant to observe the autopsy. At the request of the Coroner, radiological contamination surveys were performed by NRC's Medical Consultant. No radioactive contamination externally or internally was found.

From the direct measurements taken from the body, the actual perineal needle puncture sites were found to be in general agreement with the diagrams provided by the IRCC. In addition, NRC's Medical Consultant stated that there appeared to be gross changes and tissue destruction consistent with radiation effect. In a press release dated January 26, 1993, the Indiana County Coroner stated that the cause of death listed in the official autopsy report was "Acute Radiational Exposure and Consequences Therof."

**Table 6.1(a) Normal Tissue Distances from
Possible Source Location in Patient A**

	Source location in centimeters				
	A	B	C	D	E
1. Rectum	1.5	1.5	1.5	1.5	1.5
2. Bladder (closest point)	4.7	6.0	3.0	2.2	2.2
3. Bladder (median point)	9.1	8.6	6.7	5.7	6.2
4. Small bowel (closest point)	8.5	9.3	7.3	6.0	6.3
5. Small bowel (median point)	15.5	15.2	12.7	11.4	11.4
6. Small bowel (furthest point)	23.9	23.7	22.5	22.1	22.2
7. LS spine/pelvis bone marrow (furthest point)	25.4	25.3	24.9	25.7	25.4
8. Right kidney (median point)	19.3	20.7	21.2	22.3	22.3
9. Left kidney (median point)	20.6	19.2	18.7	20.3	19.8
10. Liver (median point)	25.1	25.4	24.6	25.4	25.2
11. Right lung (median point)	38.6	39.2	39.0	39.8	39.5
12. Left lung (median point)	39.2	38.5	37.8	38.7	38.1
13. Heart (closest point)	30.4	29.7	28.8	29.6	29.1
14. Heart (left ventricle)	34.1	33.5	32.8	33.6	33.1
15. Brain	71.0	71.0	71.0	71.0	71.0

**Table 6.1(b) Normal Tissue Dose (cGy) Versus
Possible Source Location in Patient A**

	Source location				
	A	B	C	D	E
1. Rectum	776,956	776,956	776,956	776,956	776,956
2. Bladder (closest point)	78,221	47,605	193,483	360,663	360,663
3. Bladder (median point)	20,142	22,667	37,979	52,857	44,520
4. Small bowel (closest point)	23,226	19,244	31,836	47,605	43,086
5. Small bowel (median point)	6,369	6,655	9,897	12,497	12,497
6. Small bowel (furthest point)	2,283	2,332	2,653	2,772	2,742
7. LS spine/pelvis bone marrow (furthest point)	1,956	1,976	2,058	1,898	1,956
8. Right kidney (median point)	3,842	3,250	3,068	2,712	2,712
9. Left kidney (median point)	3,289	3,889	4,139	3,406	3,616
10. Liver (median point)	2,017	1,956	2,122	1,956	1,996
11. Right lung (median point)	620	593	602	567	580
12. Left lung (median point)	593	625	660	616	645
13. Heart (closest point)	1,219	1,298	1,409	1,310	1,371
14. Heart (left ventricle)	887	932	989	925	964
15. Brain	86	86	86	86	86

Note: 1 cGy = 1 rad.

Table 6.2 Tolerance of Different Tissues to Radiation Exposure*

Organ	Type of damage	TD 5/5 (Gy)	TD 50/5 (Gy)	Total or partial organ irradiation (Field size or length)
Bone marrow	Aplasia, pancytopenia	2-5/30	4-5/40	Total/partial
Liver	Acute and chronic hepatitis	25/15	40/20	Partial/total
Bladder†	Contracture, fistula	-	80	Whole
Intestine	Ulcer, perforation, hemorrhage	45/50	55/65	400 cm ² /100 cm ²
Brain	Infarction, necrosis	50	60	Whole
Rectum†	Proctitis, ulcer, stricture, fistula	-	80	100 cm ²
Heart	Pericarditis and pancarditis	45/70	55/80	50%/25%
Lung	Acute and chronic pneumonitis	30/15	50/25	100 cm ² /total
Kidney	Acute and chronic nephrosclerosis	15/20	20/25	Total/partial

TD: tolerance dose (5/5 and 50/5% and 50% of severe complications at 5 years) The alternatives in the last column refer to the alternative values of dose and to total or partial organ irradiation. For example, 2-5 Gy to the whole bone marrow results in 5% severe complications and 4-5 Gy gives 50%. For partial irradiation the doses are 30 Gy and 40 Gy. For purposes of this report,

1 Gy = 1 Sv.

Note: 1 Gy = 100 rads

* Source: "Introduction to Radiobiology," by Maurice Tubiana, Jean Dutreix, and Andre Wambersie, Taylor & Francis, p. 138, London, England, 1990

† Source: "Radiation Biology and Radiation Pathology Syllabus," American College of Radiology, 1975

6.3 Evaluation of Radiation Doses for Occupational Workers at the Indiana Regional Cancer Center

Employees at IRCC were exposed to the iridium-192 source during patient treatment, catheter manipulation, implanted catheter removal, and source retrieval. Employees at IRCC who may have received occupational doses of radiation were —

- Registered Therapy Technician (RTT)-A
- RTT-B
- Registered Technologist Radiographer (RTR)
- Nurse A
- Medical Physicist A
- Physician A

The evaluation of radiation exposure for each of these employees is based on the dosimeter reports and interviews with the IRCC staff. The results are presented in Table 6.3 and discussed in the following sections.

6.3.1 Dose for RTT-A

When the staff experienced difficulty inserting the source wire in the last of five catheters and when subsequent dummy wire insertions were unsuccessful, RTT-A entered the treatment room several times to attempt to straighten the connecting catheter. After several error messages appeared on the HDR afterloader computer screen, Physician A instructed RTT-A to disconnect the catheters from the patient and move the patient to the stretcher room with the assistance of the RTR. After moving the patient to the stretcher room, RTT-A returned to accelerator patient treatment duties. The team used RTT-A's dosimeter reading of 8.20 mSv (0.820 rem) for November 5 to December 4, 1992, as the dose received from exposure to the source. RTT-A's year-to-date exposure was 8.20 mSv (0.820 rem).

6.3.2 Dose for RTT-B

During the failures in dummy wire insertions previously mentioned, RTT-B entered the treatment room twice with RTT-A to attempt to straighten the connecting catheter. RTT-B then returned to paperwork duties.

The team used RTT-B's dosimeter reading of 1.10 mSv (0.110 rem) for November 5 to December 4, 1992, as the dose received from exposure to the source. RTT-B's year-to-date exposure was 1.10 mSv (0.110 rem).

6.3.3 Dose for RTR

The RTR entered the treatment room three times to see to the patient's needs and to assist RTT-A. After several error messages appeared on the computer screen, the RTR was

instructed to assist RTT-A in removing Patient A from the treatment room to the stretcher room. RTR then returned to accelerator patient treatment duties. The team used RTR's dosimeter reading of 1.40 mSv (0.140 rem) for November 5 to December 4, 1992, as the dose received from the exposure to the source. The RTR's year-to-date exposure was 1.50 mSv (0.150 rem).

6.3.4 Dose for Nurse A

After Patient A was removed from the treatment room and placed in the stretcher room, Nurse A and Physician A removed the implanted catheter found to have loose stitches. The team used Nurse A's dosimeter reading of 6.30 mSv (0.630 rem) for November 5 to December 4, 1992, as the dose received from the exposure to the source. Nurse A's year-to-date exposure was 6.30 mSv (0.630 rem).

6.3.5 Dose for Medical Physicist A

Medical Physicist A's only exposure to the source was during source retrieval. Medical Physicist A and Physician A retrieved the source from BFI-Carnegie. The team used Medical Physicist A's dosimeter reading of 1.20 mSv (0.120 rem) for November 5 to December 4, 1992, as the dose received from the exposure to the source. Medical Physicist A's year-to-date exposure was 1.90 mSv (0.190 rem).

6.3.6 Estimated Dose for Physician A

As described in the previous sections, Physician A was exposed to the source on three occasions:

1. When he entered the treatment room after another set of error messages with RTT-A and RTT-B to attempt to straighten the catheter. Physician A was estimated to have been 0.5 to 1.0 meters (20 to 39 inches) away from the source for 1 minute while in the treatment room.
2. When he removed the implanted catheter with Nurse A. Physician A was estimated to have been 0.5 to 1.0 meter (20 to 39 inches) away from the source for 3 minutes while in the stretcher room.
3. When he retrieved the source with Medical Physicist A. Physician A was estimated to have been 0.5 to 1.0 meter (20 to 39 inches) from the source for 1.2 minutes while at BFI-Carnegie.

Physician A stated that he did not wear his dosimeter during any of these possible exposures. The calculated doses from the above time and distance data give an estimated exposure from 1.82 to 7.21 mSv (182 to 721 mrem). The dose can also be estimated by summing the film badge doses received by individuals that participated with Physician A in each of the above

activities. This total would have been 8.60 mSv (0.860 rem). Therefore, the team is estimating the dose to Physician A to be the sum of each dose received by RTT-B, Nurse A, and Medical Physicist A, totaling 8.60 mSv (0.860 rem). Even with this more conservative approach, Physician A's dose is less than the NRC quarterly limit.

Table 6.3 Summary of Radiation Doses for Indiana Regional Cancer Center Occupational Workers

Exposed individuals	Activity causing radiation exposure	Range of distance from source (m)	Dose rate range* (mSv/hr)	Estimated exposure time (Minutes)	Total dose range (mSv)	Total dose range (mrem)
Physician A	Catheter manipulation and catheter removal on 11-16-92	0.5-1.0	21.0-83.2	4.0	1.82-7.21 (8.60) ‡	182-721 (860) ‡
	Source retrieval on 12-01-92	0.5-1.0	21.0-83.2	1.2		
RTT-A	Catheter manipulation and patient removal from treatment room	-	-	-	8.20 †	820 †
RTT-B	Catheter manipulation	-	-	-	1.10 †	110 †
RTR	Patient care and patient removal from treatment room	-	-	-	1.40 †	140 †
Medical Physicist A	Source retrieval	-	-	-	1.20 †	120 †
Nurse A	Catheter removal	-	-	-	6.30 †	630 †
					Person-Sv	Person-rem
TOTALS:					0.027	2.7

* Rate from unshielded iridium-192 source containing $1.579 \text{ E}+11 \text{ Bq}$ (4.267 Ci) on November 16, 1992, using Microshield 3.13 with point-source geometry and Berger build-up factor. $1 \text{ mSv} = 100 \text{ mrem}$

† Total dose reported from dosimeter reports

‡ Total dose reported from dosimeter reports for RTT-B + Nurse A + Medical Physicist A

6.4 The Public

When the team arrived in Indiana, Pennsylvania, on December 2, 1992, NRC's medical consultant worked with the SHM nursing home staff to determine the number of employees who may have been exposed to the iridium-192 source. In addition, residents and visitors of SHM nursing home may have been exposed to the source while it remained at the nursing home from November 16 through 25, 1992.

BFI employees at both the Carnegie, Pennsylvania, and Warren, Ohio (BFI-Warren), sites were also exposed to radiation from the iridium-192 source while it remained at these two facilities from November 25 to December 1, 1992. These exposures, however, were much less than those encountered at SHM nursing home because of two major factors: (1) the distance from the source to the persons exposed was, on the average, much greater at the BFI facilities, and (2) the time spent near or around the source was relatively short, in most cases, less than a few hours.

In some isolated cases, however, an individual may have been exposed to the iridium-192 source any number of ways from November 16 to December 1, 1992, for example: (1) while visiting a resident other than Patient A at the nursing home; (2) while driving by or parking near the rear entrance of SHM nursing home when the source was in the outside storage area; (3) by driving past one of the BFI trucks as it carried the source during transit; or (4) possibly, standing near one of the BFI trucks as it drove by carrying the source. Even in the most conservative estimates for an individual inadvertently exposed to the source in these specific scenarios, the potential calculated dose to this person would be less than the allowable exposure to a member of the general public according to NRC regulations (10 CFR Part 20).

The estimated potential and actual radiation exposures are described in the following sections.

6.4.1 Nonoccupational Radiation Doses for Workers and Visitors to Indiana Regional Cancer Center

The team estimated potential doses of nonoccupational radiation exposure for employees who were not involved with treating Patient A and visitors to the IRCC who were exposed to the iridium-192 source on November 16, 1992.

During the treatment, Patient A was in a shielded treatment room. After the IRCC staff experienced difficulty using the HDR afterloader to insert the source wire into the last catheter, Patient A was moved from the shielded room to the stretcher room (see Figure 6.1). Patient A remained in this room for 10 minutes awaiting the arrival of the ambulance service employees to transport Patient A to SHM nursing home. For this 10-minute period, all nonoccupational workers and visitors to IRCC were potentially exposed to the source. Table 6.4 depicts their range of distances from the source location in the stretcher room and,

hence, the range of potential doses received by those individuals who received nonoccupational exposures at IRCC. Depending on the estimated distance from the source, the calculated doses to these individuals ranged from as low as 0.04 mSv (4.0 mrem) to as high as 13.9 mSv (1.39 rem). Figure 6.1 shows isodose curves at the IRCC on November 16, 1992.

Table 6.4 Summary of Radiation Doses for Indiana Regional Cancer Center Nonoccupational Workers and Patients						
Exposed individuals	Activity causing radiation exposure	Range of distance from source (m)	Dose rate range* (mSv/hr)	Estimated exposure time (Minutes)	Total dose range* (mSv)	Total dose range* (mrem)
Patient C	Waiting-GWA	6.2- 6.5	0.55- 0.60	10	0.09 - 0.10	9-10
Patient D	Waiting-GWA	6.2- 6.5	0.55- 0.60	10	0.09 - 0.10	9-10
Patient E	Waiting-GWA	6.2- 6.5	0.55- 0.60	10	0.09 - 0.10	9-10
Patient F	Waiting-GWA	6.2- 6.5	0.55- 0.60	10	0.09 - 0.10	9-10
Patient G	Waiting-GWA	6.2- 6.5	0.55- 0.60	10	0.09 - 0.10	9-10
Patient H	Waiting-GWA	6.2- 6.5	0.55- 0.60	10	0.09 - 0.10	9-10
Patient I	Waiting-GWA	6.2- 6.5	0.55- 0.60	10	0.09 - 0.10	9-10
Patient J	Receiving care in exam room adjacent to S.R.	1.9- 4.7	1.02- 5.92	10	0.17 - 0.99	17-99
Phlebotomist	Stood beside patient in T.R.	0.5- 1.0	21.0 -83.2	10	3.5 -13.9	350-1390
Office Manager	Working in secretary office	6.4-10.0	0.24- 0.56	10	0.04 - 0.09	4-9
Medical Secretary	Working in secretary office	6.4-10.0	0.24- 0.56	10	0.04 - 0.09	4-9
Tumor Registrar	Working in secretary office	6.4-10.0	0.24- 0.56	10	0.04 - 0.09	4-9
GWA = Gowned waiting area near stretcher room S.R. = Stretcher room T.R. = Treatment room					Person-Sv	Person-rem
TOTALS:					0.004-0.016	0.442-1.586

* Rate from unshielded iridium-192 source containing 1.579 E+11 Bq (4.267 Ci) on November 16, 1992, using Microshield 3.13 with point-source geometry and Berger build-up factor.

1 mSv = 100 mrem

6.4.2 Estimates of Possible Radiation Doses for Employees and Visitors to the Doctor's Office Adjacent to the IRCC Stretcher Room

The team found that three employees of the adjacent doctor's office (ADO) and two patients visiting the ADO were potentially exposed to the source on November 16, 1992. The ADO is located southwest and adjacent to the IRCC stretcher room (see Figure 6.1). Using conservative estimates, the team estimated the exposure time for these individuals to be 10 minutes (the time Patient A spent in the IRCC stretcher room). The team estimated the distance the ADO employees and visitors spent near the source from the office's layout plan, which was supplied by the staff in the ADO, and the IRCC's blueprint of the building. As shown in Table 6.5, the maximum estimated exposure that these persons received was 4.3 mSv (430 mrem).

Table 6.5 Summary of Radiation Doses for Adjacent Physician's Personnel and Patients

Exposed individuals	Activity causing radiation exposure	Range of distance from source (m)	Dose rate range* (mSv/hr)	Estimated exposure time (minutes)	Total dose range (mSv)	Total dose range (mrem)
Patient K	Waiting for care adjacent to IRCC stretcher room	0.90-1.20	14.64-25.88	10	2.4 - 4.3	240 - 430
Patient L	Waiting for care adjacent to IRCC stretcher room	0.90-1.20	14.64-25.88	10	2.4 - 4.3	240 - 430
Administrative Aide A	Working in office near IRCC stretcher room	2.50-6.50	0.55- 3.46	10	0.09 - 0.58	9 - 58
Administrative Aide B	Working in office near IRCC stretcher Room	2.50-6.50	0.55- 3.46	10	0.09 - 0.58	9 - 58
Laboratory Employee	Working in lab near IRCC stretcher room	4.00-7.00	0.48- 1.39	10	0.08 - 0.23	8 - 23
					Person-Sv	Person-rem
				TOTALS:	0.005-0.010	0.51 - 1.00

* Dose rate from unshielded iridium-192 source containing $1.579 \text{ E} + 11 \text{ Bq}$ (4.267 Ci) on November 16, 1992, using Microshield 3.13 with point-source geometry and Berger build-up factor.
 1mSv = 100 mrem

6.4.3 Estimates of Possible Radiation Doses for Citizens Ambulance Service Staff

Two ambulance service employees were exposed to the iridium-192 source during transport of Patient A after Patient A received HDR treatment on November 16, 1992. The team measured probable distances that the ambulance service employees maintained from the source in Patient A and estimated the time spent at each distance based on the ambulance service's dispatch log. The team estimates that the ambulance aide received a dose in the range of 13.3 mSv (1.3 rem) to 25.7 mSv (2.6 rem), while the ambulance driver received from 4.8 mSv (0.48 rem) to 8.4 mSv (0.84 rem) (Table 6.6). The estimated dose range for the ambulance aide is higher because he spent the time in transit to SHM nursing home sitting across from Patient A approximately 58 cm (22.8 inches) away (Figure 6.2).

Table 6.6 Summary of Radiation Doses for Ambulance Service Employees

Exposed individual	Activity causing radiation exposure	Range of distance from source (m)	Dose rate range* (mSv/hr)	Estimated exposure time (Minutes)	Total dose range (mSv)	Total dose range (mrem)
Ambulance Driver	Loaded patient	0.80-1.10	17.39-32.69	6	4.8-8.4	480-840
	Drove ambulance	2.40-2.60	3.20- 3.75	13		
	Unloaded patient	0.80-1.10	17.39-32.69	8		
Ambulance Aide	Loaded patient	0.80-1.10	17.39-32.69	6	13.3-25.7	1330-2570
	Sat beside-transit	0.50-0.70	42.60-83.17	13		
	Unloaded patient	0.80-1.10	17.39-32.69	8		
					Person-Sv	Person-rem
TOTAL:					0.02-0.03	1.81-3.41

* Dose Rate from unshielded iridium-192 source containing 1.579 E+11 Bq (4.267 Ci) on November 16, 1992, using Microshield 3.13 with point-source geometry and Berger build-up factor. 1mSv = 100 mrem.

6.4.4 Nursing Home

The SHM nursing home is a State-licensed, 58-bed nursing home offering skilled, intermediate, and respite care to the community of Indiana, Pennsylvania, located approximately 104.6 km (65 miles) east-northeast of Pittsburgh, Pennsylvania. On average, there are 45 occupants residing in the home at any given time.

The home employs three working shifts: 7 a.m. to 3 p.m.; 3 p.m. to 11 p.m.; and 11 p.m. to 7 a.m. For the first two shifts, three staff members are assigned to each resident: a registered nurse (RN) for treatment, a licensed practical nurse (LPN) for administering medication to residents, and a certified nurse assistant (CNA) as an aide. In those instances where a graduate practical nurse (GPN) was employed on a shift, an RN was still required to provide any medical treatment that a resident would need. For the evening shift (11 p.m. to 7 a.m.), only two staff members were assigned to a resident: an LPN and a CNA. Table 6.7 identifies the SHM nursing home staff assigned to Patient A the week of November 16 through 20, 1992.

**Table 6.7 Schedule for Nursing Staff Assigned to Patient A
Week of November 16 - 20, 1992**

Date	Shift	Assigned employee	Estimated hours with Patient A
11/16/92	7:00 a.m. - 3:00 p.m.	RN A LPN A CNA A	1 1.3 2
	3:00 p.m. - 11:00 P.M.	RN B GPN A CNA B	1 1.3 2
	11:00 p.m. - 7:00 a.m.	LPN B CNA C	1.3 2
11/17/92	7:00 p.m. - 3:00 p.m.	RN B LPN C CNA D	1 1.3 2
	3:00 p.m. - 11:00 p.m.	RN C LPN A CNA E	1 1.3 2
	11:00 p.m. - 7:00 a.m.	LPN B CNA C	1.3 2
11/18/92	7:00 a.m. - 3:00 p.m.	RN A GPN A CNA A	1 1.3 2
	3:00 p.m. - 11:00 p.m.	RN C LPN D RN B (CNA duties)	1 1.3 2
	11:00 p.m. - 7:00 a.m.	LPN B CNA F	1.3 2
11/19/92	7:00 a.m. - 3:00 p.m.	RN B LPN C CNA G	1 1.3 2
	3:00 p.m. - 11:00 p.m.	RN C GPN A CNA E	1 1.3 2
	11:00 p.m. - 7:00 a.m.	LPN B CNA C	1.3 2
11/20/92*	7:00 a.m. - 3:00 p.m.	RN D GPN A CNA H	0.2 0.2 0.2

* These time estimates assume that each of the three employees on duty for Patient A spent 10 minutes near the stored red-bag waste (0.8-1.0 meters) inside the soiled utility room where the source was kept until approximately 8:30 a.m. on November 20, 1992.

For purposes of estimating distances from the iridium-192 source for each of the three types of nursing staff (RNs, LPNs, and CNAs) as well as other SHM nursing home employees, the following information was based on interviews with the nursing staff and individuals involved with providing care to Patient A and nurse's notes about this resident from November 16 through 20, 1992:

- **Registered Nurses** These individuals were on the average between 1 to 1.3 meters (39.4 to 51.2 inches) away from Patient A when providing medication to Patient A. Some of those interviewed stated that the RNs at SHM nursing home spent approximately 10 to 15 minutes with Patient A every 2 hours ($15 \text{ min}/2 \text{ hr} \times 8 \text{ hr/day} = 1 \text{ hr}$).
- **Licensed Practical Nurses** were on the average between 0.8 to 1.0 meters (31.5 to 39.4 inches) away when providing medication and checking vital signs of Patient A. Some of those interviewed stated that the GPN and LPNs at SHM nursing home spent approximately 20 minutes every 2 hours ($20 \text{ min}/2 \text{ hr} \times 8 \text{ hr/day} = 1.3 \text{ hr}$) with Patient A.
- **Certified Nurse Assistants** were on the average between 0.75 to 0.9 meters (29.6 to 35.5 inches) away from Patient A while performing their daily duties. These duties include bed baths, perineum care, assistance with feeding, assistance with bed pan, and any other assistance necessary that Patient A requested. Some of those interviewed stated that the CNAs at the SHM nursing home spent approximately 15 to 30 minutes every 2 hours ($30 \text{ min}/2 \text{ hr} \times 8 \text{ hr/day} = 2 \text{ hr}$), or as Patient A required assistance.

Because of the specific care that the CNA's provided to Patient A, these individuals were most likely to have received the highest extremity dose of any of the SHM nursing staff owing to either having their hand at close proximity to the source or by actually handling the catheter in which the source was dislodged. The radiation dose to the hand is usually limited by the magnitude of dose received by the skin because the hand does not include vital tissues such as blood-forming organs (i.e., active bone marrow). The maximum calculated dose to the hand (deep tissue) of an individual was 1.6 Gy (160 rad) (see Table 6.9). Because there was no beta contribution to these doses, it is reasonable to assume that the maximum dose to the skin of the hand was also 1.6 Gy (160 rad). In addition, Table 6.14 provides extremity dose estimates for BFI employees involved in this incident. This maximum dose was found to be below the threshold for early erythematous reactions of 2 Gy (200 rad). No skin damage is anticipated because the maximum dose was also well below the moist skin desquamation induced dose of approximately 11 Gy (1100 rad).⁹

- **Activities Director** was estimated to have been 4.1 to 9.1 meters (13.5 to 29.8 feet) away from the outside waste storage area (where the source was stored before shipping) when working in the Recreation Room. On the basis of discussions with

the Activities Director, the team determined that this individual spent approximately 5 hours per day in this room on November 20, 23, and 24. On November 25, when the medical waste containing the source was removed from the SHM nursing home, the team estimated from an interview with the Activities Director that the Director spent 1.5 hours at her desk that morning, which was approximately 9.1 meters (29.8 feet) away from the outside waste storage area where the source was stored November 20 through 25, 1992.

- **Dietician** was, on the average, 1.5 to 2.0 meters (59 to 78.7 inches) away when providing meals (or other information regarding the resident's diet) to Patient A. The team estimated that the dietician at SHM nursing home spent approximately 10 minutes each day with Patient A.

Figure 6.3 is a diagram of the dose rates at different distances for individuals who may have been in contact with Patient A from November 16 through 20, 1992.

- **Maintenance Man** was estimated to have been 0.4 to 0.5 meters (15.8 to 19.7 inches) away from the source when moving the red-bag waste from the inside soiled utility room (next to Room 9) to the outside waste storage room on November 20, 1992. Figure 6.4 shows the soiled utility room illustrating isodose curves from the iridium-192 source on November 20, 1992. The amount of time estimated to walk from the inside soiled utility room to the outside waste storage area, and then to place the red-bag waste in a BFI carton was approximately 5 minutes.

The amount of time the maintenance man spent thereafter in the outside waste storage room is 2 minutes each for the 7:30 a.m. and 3:30 p.m. pickup of red-bag waste. The estimated distance from the source during this period was between 0.5 to 0.75 meters (19.7 to 29.6 inches). The total time that the maintenance man spent inside the outside waste storage room after November 20 was [4 min/day x 5 days (11/21-11/25)] 20 minutes.

The team assessed the doses to two individuals known to be pregnant at the time of the incident on November 16, 1992. One of these individuals, CNA-I, was not assigned to Patient A during the week of November 16 through 20, 1993, but helped to assist other CNA's who were assigned to Patient A during this period. Blood samples were taken from this individual on two separate occasions during the incident investigation to verify the team's initial dose calculations for this individual. In addition, time-motion studies were performed to calculate an estimated dose. CNA I stated that she assisted other CNA's on November 17, but had spent most of her time with Patient A on November 20, 1992. Because the source was no longer in Patient A's body on November 20 and because the time she spent with Patient A before this date was relatively short, the team estimates that CNA-I received on average a dose of about 5 mSv (500 mrem).

In addition to CNA I, the Dietician was also known to be pregnant at the time of the incident. As described in Table 6.8, the Dietician received an average dose of about 5 mSv (500 mrem) for this time period. Current guidance provides a limit of 5 mSv (500 mrem) for the embryo-fetus.

Table 6.8 gives a summary of whole body dose estimates for SHM nursing staff during the week of November 16 through 20, 1992. Table 6.9 gives a summary of dose estimates for extremity (i.e., hand and forearm) exposures for SHM nursing staff the week of November 16 through 20, 1992.

Table 6.8 Summary of Whole Body Dose Estimates for SHM Nursing Home Staff

Exposed individuals	Activity causing radiation exposure	Range of distance from source (m)	Dose rate range* (mSv/hr)	Estimated exposure time (hours)	Total dose range (mSv)	Total dose range [rem]
RN A	Treatment	1.0 - 1.3	21.0-12.5	2	42 - 25	4.2 - 2.5
RN B	Treatment	1.0 - 1.3	21.0-12.5	3	137 - 89 (Total)	13.7 - 8.9 (Total)
	Aide (11/18)	0.75 - 0.9	37.1-25.8	2		
RN C	Treatment	1.0 - 1.3	21.0-12.5	3	63 - 38	6.3 - 3.8
RN D**	Near Utility room	0.8 - 1.0	32.6-21.0	0.17	5.5 - 3.6	0.55 - 0.36
GPN A	Medication	0.8 - 1.0	32.6-21.0	4.17	136 - 88	13.6 - 8.8
LPN A	Medication	0.8 - 1.0	32.6-21.0	2.67	87 - 56	8.7 - 5.6
LPN B	Medication	0.8 - 1.0	32.6-21.0	5.33	174 - 112	17.4 - 11.2
LPN C	Medication	0.8 - 1.0	32.6-21.0	2.67	87 - 56	8.7 - 5.6
LPN D	Medication	0.8 - 1.0	32.6-21.0	1.33	43 - 28	4.3 - 2.8
CNA A	Aide	0.75 - 0.9	37.1-25.8	4	148 - 103	14.8 - 10.3
CNA B	Aide	0.75 - 0.9	37.1-25.8	2	74 - 52	7.4 - 5.2
CNA C	Aide	0.75 - 0.9	37.1-25.8	6	223 - 155	22.3 - 15.5
CNA D	Aide	0.75 - 0.9	37.1-25.8	2	74 - 52	7.4 - 5.2
CNA E	Aide	0.75 - 0.9	37.1-25.8	4	148 - 103	14.8 - 10.3
CNA F	Aide	0.75 - 0.9	37.1-25.8	2	74 - 52	7.4 - 5.2
CNA G	Aide	0.75 - 0.9	37.1-25.8	2	74 - 52	7.4 - 5.2
CNA H**	Near Utility room	0.8 - 1.0	32.6-21.0	0.17	5.5 - 3.6	0.55 - 0.36
CNA I	Assisting CNAs 11/17/92	0.75 - 0.9	37.1-25.8	0.17	6.2 - 4.3	0.62 - 0.43
Maintenance Man A	Initial transfer of red-bag waste on 11/20/92	0.4 - 0.5	130-83.1	0.08	38 - 19 (Total)	3.8 - 1.9 (Total)
	Subsequent transfer of daily waste	0.5 - 0.75	83.1-37.1	0.33		
Dietician	Visits, residents rooms	1.5 - 2.0	9.4 - 5.3	0.67	6.3 - 3.6	0.63 - 0.36
Activities Director	Develops and participates in SHM activities 11/20, 11/23-24	4.1 - 9.1	1.32- 0.29	15	22 - 4.8 (Total)	2.2 - 0.48 (Total)
	Develops activities at desk 11/25	9.0 - 9.1	0.296-0.290	1.5		
TOTALS					Person-Sv 1.67 - 1.10	Person-rem 167 - 110

* Dose rate from an unshielded iridium-192 source containing 1.579 E+11 Bq [4.267 Ci] on November 16, 1992, using MICROSIELD 3.13 with point-source geometry and Berger build-up factor.

** These individuals spent approximately 10 minutes near (0.8 - 1.0 meters) the soiled utility room where the source was inadvertently placed on November 20, 1992.
Note: 1 mSv = 100 mrem

Table 6.9 Summary of Extremity Dose Estimates for Selected SHM Nursing Home Staff

Exposed individuals	Activity causing radiation exposure	Range of distance from source [†] (m)	Dose rate range [*] (Sv/hr)	Total estimated exposure time (hours)	Total dose range (Sv)	Total dose range (rem)
LPN B ^{**}	Assisting CNA C (11/20)	0.10 - 0.20	2.1 - 0.51	0.17	0.36 - 0.09	36 - 8.7
CNA A	Aide	0.20 - 0.30	0.51 - 0.23	2	1.02 - 0.46	102 - 46
CNA B	Aide	0.20 - 0.30	0.51 - 0.23	1	0.51 - 0.23	51 - 23
CNA C ^{**}	Aide	0.20 - 0.30	0.51 - 0.23	2.83	1.6 - 0.74	160 - 74
	Aide (11/20)	0.15 - 0.20	0.92 - 0.51	0.17	(Total)	(Total)
CNA D	Aide	0.20 - 0.30	0.51 - 0.23	1	0.51 - 0.23	51 - 23
CNA E	Aide	0.20 - 0.30	0.51 - 0.23	2	1.02 - 0.46	102 - 46
CNA F	Aide	0.20 - 0.30	0.51 - 0.23	1	0.51 - 0.23	51 - 23
CNA G	Aide	0.20 - 0.30	0.51 - 0.23	1	0.51 - 0.23	51 - 23

[†] Ranges of distances for perineal care estimated from time-motion studies and personal interviews. Distances are approximated.

^{*} Dose rate from an unshielded iridium-192 source containing 1.579 E+11 Bq (4.267 Ci) on November 16, 1992, using MICROSHIELD 3.13 with point-source geometry and Berger build-up factor.

^{**} These individuals found the dislodged iridium-192 source in Patient A's bed on November 16, 1992.

Note: 1 Sv = 100 rem

In addition to the number of staff who were exposed to the radiation source during November 16 through 20, 1992, Patient A received seven different visitors (relatives and friends). Because there was no visitor log maintained at the entrance of the facility, the team cannot determine the exact number of visitors that may have entered the facility and traveled past Room 4B. Even if several visitors were to have been near this room, or in the hallway near Room 4B, the resulting exposures would have been relatively low due to the short period of time spent at these locations. Distances and estimated times of the hours visited were established through interviews with relatives, friends, and the SHM nursing home staff. In addition, the daily nurse's notes provided valuable information as to when visitors arrived and departed. Interviews with Relative A and Friend A gave the team specific information about the time each spent with Patient A and at what distance. Relative A stated that she spent approximately one-third of her time with Patient A at close proximity [between 0.5 to 1 meter (1.6 to 3.3 feet)]. Friend A stated that she spent approximately 2.5 hours with Patient A [estimated distance 0.75 to 1.5 meters (29.5 to 59 inches)]. Table 6.10 provides detailed information as to the dates, times of visitation, estimated distance from Patient A, and calculated doses while they were at the SHM nursing home.

The team estimated doses for several of the residents that were known to have been exposed to the source for the entire period of November 16, 1992, 11 a.m., through November 20, 1992, 6:15 a.m., (91.25 hours total) while the source remained inside Patient A. Isodose curves for the rooms adjacent to Patient A are shown in Figure 6.5. These estimated doses are summarized in Table 6.11. All of the residents listed in Table 6.11 except Residents K and M participated in several activities in the Recreation Room during November 20 through 25, 1992.

The team assumed that the residents in nearby rooms were exposed to the source while it remained inside Patient A for 91.25 hours. The highest of these doses, those for Residents B, C, and M, were verified independently by cytogenetic analysis (see Section 6.5.2).

**Table 6.10 Summary of Radiation Doses for Relatives and Friends
During November 16 - 20, 1992**

Exposed individuals	Date visited	Range of distance from source (m)	Dose rate range* (mSv/hr)	Estimated exposure time (hours)	Total dose range (mSv)	Total dose range [rem]
Relative A	11/17/92	0.5 - 1.0	83.1-21.0	1.25	166 - 54.4 (Total)	16.6-5.4 (Total)
		1.0 - 1.5	21.0- 9.4	3.0		
Relative B	11/16/92	1.0 - 1.2	21.0-14.6	2.0	42.0-29.2	4.2-2.9
Relative C	11/16/92	0.9 - 1.2	25.8-14.6	0.5	64.5-36.5 (Total)	6.4-3.6 (Total)
	11/18/92			2.0		
Relative D	11/16/92	1.2 - 1.5	14.6- 9.4	0.5	36.5-23.5 (Total)	3.6-2.4 (Total)
	11/18/92			2.0		
Relative E	11/18/92	1.0 - 1.2	21.0-14.6	0.75	31.5-21.9 (Total)	3.1-2.2 (Total)
	11/19/92			0.75		
Relative F	11/18/92	1.0 - 1.2	21.0-14.6	0.75	31.5-21.9 (Total)	3.1-2.2 (Total)
	11/19/92			0.75		
Friend A	11/19/92	0.75- 1.5	37.1-9.4	2.5	92.8-23.5	9.3-2.4
					Person-Sv	Person-rem
TOTALS					0.46-0.21	46.3-21.1

* Dose rate from an unshielded iridium-192 source containing $1.579 \text{ E}+11 \text{ Bq}$ (4.267 Ci) on November 16, 1992, using MICROSHIELD 3.13 with point-source geometry and Berger build-up factor.

Table 6.11 Summary of Radiation Doses for SHM Residents During November 16 - 20, 1992

Exposed Individuals	Activity causing radiation exposure	Range of distance from source (m)	Dose rate range* (mSv/hr)	Estimated exposure time (hours)	Total dose range (mSv)	Total dose range (rem)
Resident B	In adjacent room	2.6 - 4.9	3.2 - 0.94	40	128 -37.6	12.8-3.8
Resident C	In adjacent room	2.7 - 4.9	2.9 - 0.94	68	197 -63.9	19.7-6.4
Resident D	Room down the hall	4.3 - 8.5	1.2 - 0.33	70	84.0-23.1	8.4-2.3
Resident E	Room down the hall	10.6 - 12.4	0.22 - 0.16	70	15.4-11.2	1.6-1.1
Resident F	Room across hall	6.7 - 7.2	0.52 - 0.45	61	31.7-27.5	3.2-2.8
Resident G	Room across hall	8.1 - 9.0	0.36 - 0.30	61	22.0-18.3	2.2-1.8
Resident H	Room across hall	6.7 - 7.2	0.52 - 0.45	76	39.5-34.2	4.0-3.4
Resident I	Room across hall	8.1 - 9.0	0.36 - 0.30	72	25.9-21.6	2.6-2.2
Resident J	Room across hall	8.7 - 11.4	0.32 - 0.19	76	24.3-14.4	2.4-1.4
Resident K	Room across hall	9.8 - 11.4	0.25 - 0.19	91.25	22.8-17.3	2.3-1.7
Resident L	In adjacent room	7.2 - 11.1	0.45 - 0.20	61	27.5-12.2	2.8-1.2
Resident M	Room near soiled utility room	11.4 - 15.5 (Distance from Room 4B)	0.19 - 0.11	91.25	90.9-57.3 (Total)	9.1-5.7 (Total)
		0.8 - 1.0 (Distance from soiled utility room)	32.7 - 21.0	2.25		
Resident N	Room near soiled utility room	11.4 - 16.1 (Distance from Room 4B)	0.19 - 0.10	60	23.4-13.8 (Total)	2.3-1.4 (Total)
		2.0 - 2.5 (Distance from soiled utility room)	5.35 - 3.46	2.25		
					Person-Sv	Person-rem
TOTALS					0.73-0.35	73.2-35.2

* Dose rate from an unshielded iridium-192 source containing 1.579 E+11 Bq [4.267 Ci] on November 16, 1992, using MICROSHIELD 3.13 with point-source geometry and Berger build-up factor.

The SHM nursing home provides medical care and, in addition, a recreation room that offers residents a variety of diversions from which to choose; therefore, many of the residents in the home occupy the recreation room daily. On the basis of an interview with the Activities Director, who was responsible for developing and coordinating all activities at the SHM nursing home, and a review of the activities calendar for November 1992; the team developed Table 6.12 that describes the number of activities, the approximate number of residents in the Recreation Room for each activity, the estimated distance the residents were from the outside waste storage room (where the source was later stored), estimated total dose range for each resident that participated in each activity, and the total collective dose for each activity in the Recreation Room. Figure 6.6 shows the isodose curves for the Recreation Room for the iridium-192 source while located in the outside storage room from November 20 through 25, 1992.

Table 6.12 shows that the total collective dose for *each* activity ranged from 0.003 to 0.030 person-Sv (0.26 to 3.0 person-rem). Because there are no records of the residents who were participating in each activity, only the collective dose for each activity can be calculated. If it is assumed, however, that the same residents participated in all of the activities at the SHM nursing home during November 20 through 25, 1992, then the total dose for each of the residents would be between 8.32 to 3.36 mSv (832 to 336 mrem). The collective dose for the recreation room activities is the sum of the dose for each activity, or 0.15 to 0.06 person-Sv (15.0 to 5.9 person-rem).

The catheter containing the source remained in SHM's outside waste storage room until 9:25 a.m. on Wednesday, November 25, 1992, when BFI Driver A picked up all the red-bag waste (3 boxes) generated by the SHM nursing home.

**Table 6.12 Summary of Radiation Doses for SHM Nursing Home Residents While in Recreation Room
During November 20 - 25, 1992**

Date	Time for activity (hr)	Activity	Estimated no. of residents	Ranges of distance from source (m)	Dose rate range ^a (mSv/hr)	Dose ^b range for each resident (mSv)	Total collective dose ^c for each activity (person-Sv)	Total collective dose ^c for each activity (person-rem)
11/20/92	0.5	Reading	15	4.1-8.4	1.32-0.34	0.66-0.17	0.010-0.003	0.99-0.26
11/20/92	1.0	Games	20	4.1-8.4	1.32-0.34	1.32-0.34	0.026-0.007	2.6 -0.68
11/20/92	1.0	Church	20	4.1-6.1	1.32-0.62	1.32-0.62	0.026-0.012	2.6 -1.2
11/22/92	1.5	Music	15	4.1-6.1	1.32-0.62	1.98-0.93	0.030-0.014	3.0 -1.4
11/23/92	1.0	Exercise	20	4.1-6.1	1.32-0.62	1.32-0.62	0.026-0.012	2.6 -1.2
11/23/92	1.0	Music	15	7.5-8.4	0.40-0.34	0.40-0.34	0.006-0.005	0.60-0.51
11/24/92	1.0	Birthdays	20	4.1-8.4	1.32-0.34	1.32-0.34	0.026-0.007	2.6 -0.68
11/25/92	1.0	Comedy ^d	N/A	N/A	N/A	N/A	N/A	N/A
TOTALS						mSv ^e	Person-Sv	Person-rem
						8.32-3.36	0.15-0.06	15.0-5.9

^a Dose rate from an unshielded iridium-192 source containing 1.521 E+11 Bq [4.110 Ci] on November 20, 1992, using MICROSHIELD 3.13 with point-source geometry and Berger build-up factor.

^b Time participating in activity (hours) multiplied by the dose rate (mSv/hour).

^c The collective dose equals the total dose (in mSv) multiplied by the number of residents participating in each activity.

^d Waste containing source was removed from SHM at 9:25 a.m. Wednesday, November 25, 1992, prior to comedy activity.

^e Total dose for each resident assuming that each resident went to all the activities listed above.

Note: 1 mSv = 100 mrem

6.4.5 Browning-Ferris Industries Transit and Storage

BFI is a waste management company that had two separate facilities involved in the transit and storage of the iridium-192 source. At the time of the incident, BFI had a firm policy that prohibits the acceptance of specific levels of radioactive material in biomedical waste. The policy set for the screening levels of medical waste is 0.5 μSv (50 μrem) per hour. BFI has an informal agreement with the Ohio Environmental Protection Agency to report all received shipments above 0.5 μSv (50 μrem) per hour.

For comparison purposes, the naturally occurring background radiation level at each of the BFI facilities—BFI-Carnegie and BFI-Warren—varies between 0.07 to 0.15 μSv (7 to 15 μrem) per hour. BFI-Warren places a portable survey meter in each vehicle and fixed radiation monitors at each processing facility to sound an alarm at readings at or above 0.2 μSv (20 μrem) per hour. Neither BFI-Carnegie nor BFI-Warren were authorized or permitted to handle radioactive material at any of the medical waste facilities and, thereby, established a strict policy to prohibit the entrance of any radioactive waste.

BFI employees are trained in the use and operation of portable survey meters. Each BFI-Carnegie straight truck (see Figure 2.9) that is used to pick up medical waste from generators is equipped with a portable survey meter similar to the one illustrated in Figure 6.7. In addition, all BFI-Carnegie straight-truck drivers are also required to wear radiation dosimeters (i.e., film badges) during work hours.

All BFI-Carnegie straight truck drivers are required to perform a radiation survey with their portable survey meters before accepting any waste on the truck. In addition, the drivers must sign a statement on the bottom of the shipping manifest that all packages at each location of pick-up have been surveyed with their portable survey meter. Contrary to BFI's policy, BFI Driver A, who was the driver of the straight truck that picked up waste at the SHM nursing home, stated that he did not use his survey instrument at the SHM nursing home on November 25, 1992.

Although the BFI-Carnegie policy states that (1) each straight-truck driver must survey all waste containers that are placed in the tractor-trailer, (2) survey the tractor-trailer before transportation of the waste, and (3) sign a statement on the manifest form stating that all these surveys have been completed, BFI-Carnegie Driver B was not supplied with a portable survey meter. Therefore, BFI Driver B could not have surveyed the tractor-trailer or boxes because he did not have a survey meter available. As a followup note, BFI has since purchased portable survey meters for each BFI tractor-trailer.

6.4.6 Browning-Ferris Industries Search and Retrieval

For purposes of estimating distances from the radioactive source for each of the BFI employees, the team developed the following information from interviews with employees

and time-motion studies at BFI-Carnegie regarding isolation of the box that contained the radioactive source:

- **BFI Driver A** drove the BFI-Carnegie straight truck on November 25, 1992. Because the SHM nursing home was his 12th stop of the day, the team estimated that the source was located approximately midway in the truck, or about 4.6 meters (15 feet) from the driver. The team estimated that for each stop the driver made after the SHM nursing home, he would be between 4.5 to 4.6 meters away from the box containing the source. For a brief period while he unloaded the truck (approximately 1 minute), he was estimated to be between 2 to 2.5 meters away from the box containing the radioactive source (10 stops x 1 min/stop = 10 minutes between 2 to 2.5 m). The total dose estimated for BFI Driver A is between 7.3 to 6.4 mSv (0.73 to 0.64 rem).

However, on January 15, 1993, BFI Driver A provided a statement to the team which stated that he was wearing a film badge on November 25, 1992. BFI Driver A's film badge report indicated that BFI Driver A received 1.7 mSv (0.17 rem) for this period. The team used the dosimetry report value of 1.7 mSv (0.17 rem) for the assigned dose to this individual.

Although BFI Driver A stated that he did not survey for radioactive material at the SHM nursing home, he did state that he surveyed for waste at 10 additional stops *after* picking up the waste at the home. Because of the high radioactivity of the source, the team finds it unlikely that the source was not detected, and that his film badge reading was not higher than 1.7 mSv (0.17 rem).

- **BFI Driver B** drove the tractor-trailer from BFI-Carnegie to BFI-Warren on November 27, 1992. Because the team determined that the container was the last loaded onto the trailer, it was approximately 12.2 meters (40 feet) from the driver during transit. Except for initially checking the shipment at the beginning of the trip, which took about 15 minutes at a distance of about 1.5 meters (4.9 feet) from the box containing the source, Driver B was between 12.0 to 12.2 meters away from the source for the duration of the trip.
- **BFI Driver C** drove the tractor-trailer from BFI-Warren to BFI-Carnegie on November 27, 1992 (see Figure 2.13). Because it had been determined that the trailer contained radioactive material, the driver stayed at the maximum distance away from this material while driving it back to BFI-Carnegie. The driver was estimated from the source to be between 12.0 to 12.2 meters (39 to 40 feet) during transit.
- **Medical Waste Supervisor A** spent approximately 1.5 hours at a distance between 0.75 to 1.5 meters (2.5 to 5 feet) away from the box while trying to determine which of the red bags contained the radioactive source. Once the source was located,

Supervisor A left for the day, and asked Safety Technicians A and B to continue searching through the box for identifying information.

- **Safety Tech A** spent approximately 1.5 hours within a distance of 0.75 to 1.25 meters (2.5 to 4.1 feet) and helped to identify the box containing the radioactive source. An additional 30 minutes was spent after the initial location of the box as Safety Technician A opened up the box and began to search through the red-bag waste [approximate distance during these 30 minutes was 0.5 to 0.75 meter (1.6 to 2.5 feet)]. In addition, this individual helped Medical Physicist A and Physician A when they retrieved the source from BFI-Carnegie on December 1, 1992.
- **Safety Tech B** spent approximately 1.5 hours within a distance of 0.75 to 1.25 meters (2.5 to 4.1 feet) and helped to identify the box containing the radioactive source. After the initial location of the box, Safety Technician B spent an additional 30 minutes at an approximate distance of 0.75 to 1.0 meters (2.5 to 3.3 feet) from the source holding the portable survey meter and searching through the red-bag waste.

Figure 6.8 shows the position of Supervisor A, Safety Technician A, and Safety Technician B around the box [78.7 cm by 53.3 cm by 53.3 cm (31 inches by 21 inches by 21 inches)] showing isodose curves in mSv per hour.

- **Other BFI Employees** (11 employees) worked on Saturday, November 28, 1992: three individuals in the maintenance shop, four in the main office, and four in the recyclery. The closest of these buildings, the container shop, is approximately 107 to 122 meters (350 to 400 feet) from the trailer containing the source. The exposure rate at this distance is between 0.0021 to 0.0016 mSv (0.21 to 0.16 mrem) per hour. Assuming that every worker was working in the container shop, each worker would receive a total dose of 0.017 to 0.013 mSv (1.7 to 1.3 mrem) for this 8-hour day. On November 29, 1992, only two people worked at BFI-Carnegie. These employees worked in the maintenance shop on this day. Assuming an 8-hour work day, at a distance of between 150 to 180 meters (490 to 590 feet), their total dose would have been 0.007 to 0.004 mSv (0.7 to 0.4 mrem). The total collective dose for November 28 and 29, 1992, would be 0.19 to 0.14 person-mSv (0.019 to 0.014 person-rem).

Table 6.13 summarizes the whole body dose estimates for BFI-Carnegie employees for November 25 through December 1, 1992. Table 6.14 summarizes the extremity doses for BFI-Carnegie employees for November 25 through December 1, 1992.

Table 6.13 Summary of Whole Body Radiation Doses for BFI Employees

Exposed individuals	Date & time	Activity causing radiation exposure	Range of distance from source (m)	Dose rate range* (mSv/hr)	Estimate of time for exposure (hours)	Total dose range (mSv)	Total dose range [rem]
Driver A	11/25/92 (9:45a.m.-3:30p.m.)	Loading and unloading straight truck	-	-	-	1.7†	0.170†
Driver B	11/27/92 (6:15a.m.-8:45a.m.)	Initial 15-min safety check	1.0 - 1.5	19.3- 8.7	0.25	5.2 - 2.5 (Total)	0.52 - 0.25 (Total)
		Driver from BFI-Carnegie to BFI-Warren	12.0 -12.2	0.16-0.15	2.5		
Driver C	11/27/92 (2:30p.m.-4:45p.m.)	Driver from BFI-Warren to BFI-Carnegie	12.0 -12.2	0.16-0.15	2.25	0.36- 0.34	0.036-0.034
Supervisor A	11/30/92 - 12/01/92	Unloading boxes	0.75- 1.5	34.2- 8.7	1.5	51.3- 13.0	5.1 -1.3
Safety Technician A	11/30/92	Locating source	0.75 -1.3	34.2-11.5	1.5	89.5- 34.3 (Total)	9.0 -3.4 (Total)
		Searching through red-bag waste	0.5 - 0.75	76.4-34.2	0.5		
Safety Technician B	11/30/92	Locating source	0.75 - 1.3	34.2-11.5	1.5	68.4- 26.9 (Total)	6.8 -2.7 (Total)
		Searching through red-bag waste	0.75 - 1.0	34.2-19.3	0.5		
Other BFI Employees	11/28/92	11 Workers at BFI-Carnegie	107 - 122	0.0021-0.0016	8	0.19-0.15 (Total Person-mSv for 11/28 and 11/29)	0.019-0.015 (Total Person-rem for 11/28 and 11/29)
	11/29/92	2 Workers at BFI-Carnegie	150 - 180	0.00089-0.00052	8		
						Person-Sv	Person-rem
TOTALS						0.22-0.08	0.022-0.008

† Total dose reported from dosimeter reports.

* Dose rate from an unshielded Ir-192 source containing 1.451 E+11 Bq [3.922 Ci] on November 25, 1992, using MICROSIELD 3.13 with point-source geometry and Berger build-up factor.

Note: 1 mSv = 100 mrem

Table 6.14 Summary of Extremity Dose Estimates For Selected BFI Employees

Exposed individuals	Activity causing radiation exposure	Range of distance from source[†] (m)	Dose rate range[*] (Sv/hr)	Total estimated exposure time (hours)	Total dose range (Sv)	Total dose range (rem)
Safety Technician A	Searching through red-bag waste	0.10 - 0.25	1.9 - 0.30	0.5	0.95 - 0.15	95 - 15
Safety Technician B	Searching through red-bag waste	0.20 - 0.25	0.48 - 0.30	0.5	0.24 - 0.15	24 - 15

[†] Ranges of distances for this activity were estimated from time-motion studies and personal interviews. Distances are approximate.

^{*} Dose rate from an unshielded iridium-192 source containing 1.451 E+11 Bq (3.922 Ci) on November 25, 1992, using MICROSHIELD 3.13 with point-source geometry and Berger build-up factor.

Note: 1 Sv = 100 rem

6.5 Blood Studies

When the NRC was notified of the incident on December 1, 1992, the NRC immediately consulted a Medical Consultant to help assess the extent of the radiation exposures to individuals at the SHM nursing home. On December 3, 1992, when the team arrived in Indiana, Pennsylvania, the NRC Medical Consultant contacted the SHM nursing home administrator to review all the medical records of Patient A. In addition, the NRC Medical Consultant worked with the Medical Director of the SHM nursing home to--

- interview most, if not all, the SHM nursing home employees, especially those in close contact with the patient, to see if they had been ill since the iridium-192 source was inside the facility (November 16 through 20, 1992);
- assess the SHM nursing home residents in the vicinity of Patient A's room (Room 4B) and any visitors who may have spent time in Room 4B between November 16 to 20, 1992; and
- ask all SHM nursing home employees, selected SHM nursing home residents, and relevant visitors to have blood counts done as a group during the following day at the SHM nursing home.

6.5.1 Blood Counts

NRC's Medical Consultant contacted REAC/TS in Oak Ridge, Tennessee, to request special prepackaged blood kits by overnight mail for blood studies of SHM nursing home employees. This allowed blood samples to be drawn and platelets to be analyzed at a local hospital and chromosome analysis of cultivated lymphocytes to be analyzed by REAC/TS.

On December 3, 1992, NRC's Medical Consultant supervised the drawing of blood samples at SHM nursing home between 2 p.m. and 4 p.m. (39 of 51 employees had blood samples taken). These samples were drawn by two laboratory technologists from a local hospital. Twelve of the SHM nursing home employees did not have their blood taken for analysis because they (1) had infrequent contact with Patient A at the SHM nursing home; (2) had not worked near Patient A for extended periods of time; or (3) had not been on duty the week of November 16 through 20, 1992.

Samples for possible cytogenetic studies were drawn at the same time and were carefully labelled, handled, and packaged. The NRC's Medical Consultant explained to the SHM nursing home Medical Director and RN B that the blood tests were precautionary only and they were asked to reassure their staff. The analogy was given of similar precautionary blood tests familiar to all health care workers, such as those tests that are taken to rule out tuberculosis or hepatitis exposures from patient care.

The results of the above tests, as stated by NRC's Medical Consultant, were as follows:

- No SHM nursing home employees or residents located near Patient A were known to be ill between November 16 through 20, 1992, or to have experienced any effects consistent with acute radiation syndrome;
- blood work consisting of complete blood count, differential count, and platelet counts were not consistent with a significant exposure to cause their depression.¹⁰ Since the blood was drawn on December 3, 1992, and the exposure period was from November 16 through 20, 1992, sufficient time had elapsed to allow for depression of not only the lymphocytes, but also the neutrophils and platelets (see Reference 10).

All blood samples taken from the SHM nursing home staff were sent to REAC/TS for processing, but not for further analysis until further instructions from either NRC or NRC's Medical Consultant were given. All the initial 39 blood samples were hand-carried by an NRC contractor to the REAC/TS facility on December 3, 1992. This was done so that if any new information became available, such as depressed lymphocyte, neutrophil, or platelet counts, any of the blood samples could be analyzed for cytogenetics without losing time. In addition, as more time-motion studies and interviews were performed, the team selected additional persons to be judged at risk of higher exposures. In fact, an additional two individuals were later evaluated for cytogenetic analysis because of calculated high exposures. Of the 40 blood samples (total) taken from the SHM nursing home, eight (five employees and three residents) were sent to REAC/TS in Oak Ridge, Tennessee, for cytogenetic and independent analysis of dose distributions. An additional 10 blood samples were taken from relatives and visitors to Patient A.

In addition to the SHM nursing home employees, visitors, and friends who may have been near the source during this period, six BFI employees' blood samples were also taken for initial screening and analysis. The results of the time-motion studies and blood tests indicated that none of the BFI employees were exposed to an amount of radiation greater than the minimum detectable limit needed [approximately 20 cGy (rad)] for cytogenetic evaluation.

On December 23, 1992, a member of the public was identified and contacted about the amount of time spent at the SHM nursing home November 16 through 20, 1992. This individual stated that she had spent approximately 2.5 hours visiting with Patient A at very close proximity [between 0.75 to 1.5 meters (29.5 to 59 inches)] the evening of November 19, 1992. Because of this individual's close proximity to the resident, additional blood samples were taken from this person (Friend A) on January 7, 1993, and sent directly to REAC/TS for cytogenetic analysis. The results of these tests are provided in Table 6.15.

After the team became aware of this previously unidentified visitor, NRC placed an advertisement in the local newspaper on December 2, 1992, informing any person who was concerned about having coming into contact with radiation from the source and who had not

been interviewed to contact the NRC. A collect telephone number was provided. This lead to two additional contacts, the results of which are discussed in Table 6.10.

6.5.2 Cytogenetics

To confirm the range of estimated radiation exposure to the nine individuals identified as having the highest potential for exposure, the team arranged for cytogenetic evaluation of these individuals. Cytogenetic analysis is used to determine if any of the persons sustained any radiation-induced chromosome aberrations, a biological indicator of significant radiation exposure. Cytogenetic dosimetry is based on the premise that ionizing radiation induces identifiable chromosome aberrations in lymphocytes and that this induced frequency is strictly dependent on cellular dose.¹¹ The cytogenetic evaluation involved the examination of a randomly selected set of lymphocytes (white blood cells) to determine how many in each set exhibited radiation-induced chromosome aberrations.

The team arranged with the Oak Ridge Institute for Science and Education (ORISE), REAC/TS, Cytogenetics Laboratory in Oak Ridge, Tennessee, to supply "cytogenetics kits" for collecting two blood specimens for each person that was evaluated. For all of the individuals tested, the contracting physician arranged for blood specimens to be collected at the SHM nursing home. Of the 41 blood samples sent to REAC/TS, the team requested that nine of the blood samples be evaluated for cytogenetic analysis based on the team's initial dose calculations. Although the samples were collected at different times, REAC/TS stated that as long as the blood samples were taken within the first 6 weeks of the incident involving radiation exposure, no difference in the cytogenetic evaluation would be noted.

REAC/TS's procedure for cytogenetic evaluation involves sampling 500 first-division metaphases from blood lymphocyte cultures. REAC/TS reported to the team that the observed frequencies of dicentric chromosomes ranged from 1 to 4 dicentrics in the 500 metaphase samples from these cultures.¹² In persons who have had no radiation exposure other than background, an average of 1 dicentric per 500 cells scored is routinely observed in lymphocyte cultures. As an example, for a 90-percent confidence interval, an average of 5 dicentrics per 500 cells would indicate a cytogenetic dose estimate of approximately 20 cGy (rad), similarly, an average of 3 dicentrics per 500 cells would indicate a cytogenetic dose estimate of approximately 10 cGy (rad) (Reference 12).

The results of the cytogenetic analysis showed a total of 21 dicentric chromosomes in the 4,500 metaphases examined from these 9 persons (see Reference 12). The average frequency of dicentric chromosomes in >25,000 metaphases from 131 nonirradiated control persons examined at REAC/TS is 1.6 per thousand cells, which is virtually identical to the 1.8 per thousand reported by a recent REAC/TS report, "Chromosome Aberrations in Relation to Radiation Dose following Partial-Body Exposures in Three Populations."¹³

On the basis of these findings in a nonirradiated control subject, the team expected that about 1 dicentric per 500 metaphases would have been observed in persons having no radiation

exposures other than incidental chest x-rays or natural background exposures. Thus, in a sample of 4,500 metaphases, one would expect to see on average about 9 dicentric chromosomes. However, the team observed a total of 21 dicentrics, or 12 dicentrics in excess to those expected due to background. Thus, the data provides strong evidence that the group as a whole did indeed receive an exposure in excess of background with a cytogenetic dose estimate for the group of 7 rad, with 90 percent confidence intervals of 5 to 11 rad (see Reference 12).

The results from the cytogenetic studies (with 90% confidence intervals for the dose estimates) are shown in Table 6.15. The individuals identified in the table known to have been actually exposed to the iridium-192 source exhibited up to 4 dicentrics per 500 metaphases scored, which was in good agreement with the estimated radiation exposures from time-motion studies the team conducted for each person.

Table 6.15 Comparison of Calculated Whole Body Doses with Cytogenetic Evaluations

Individual	Metaphases scored	No. dicentrics observed	Dose estimate †		90% Confidence interval (cGy)
			Based on time-motion studies (cGy)	Based on cytogenetic evaluation (cGy)	
SHM Dietician	500	1	0.6 - 0.4	0	<1-12
SHM Resident B	500	2	13 - 4	~6	<1-20
SHM Resident C	500	3	20 - 6	~10	3-25
SHM Resident M	500	3	9 - 6	~10	3-25
SHM CNA C	500	4	22-16	~16	6-29
SHM CNA E	500	2	15 - 10	~6	<3-20
SHM LPN B	500	3	17 - 11	~10	3-25
SHM Maintenance Man A	500	2	4 - 2	~6	<1-20
Friend A	500	1	9 - 2	0	<1-12

† Estimate of "equivalent" dose to whole body for iridium-192 gamma rays. For purposes of this report, 1 cGy = 1 cSv.

Note: 1 cGy = 1 rad

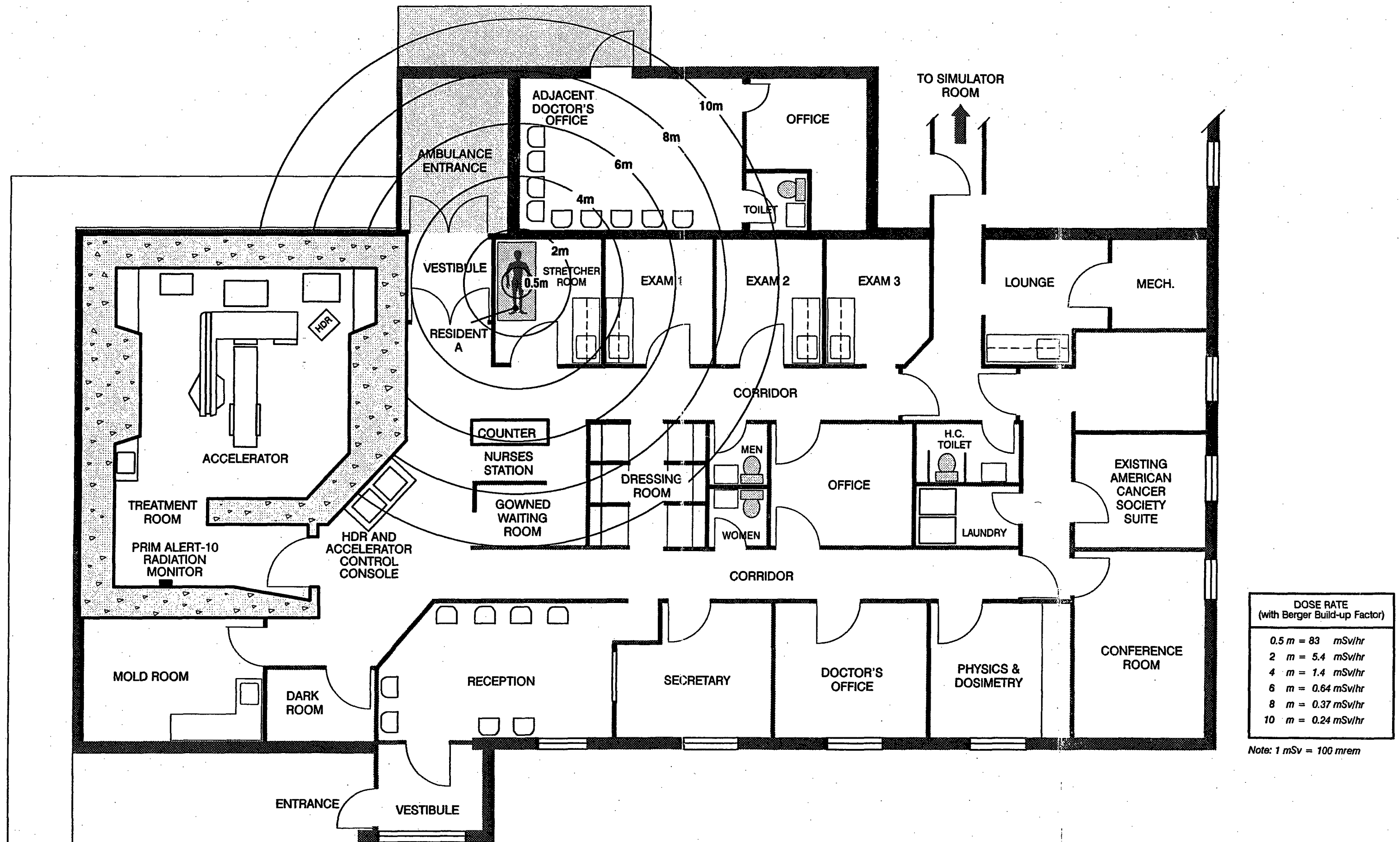


Figure 6.1 Diagram of the Indiana Regional Cancer Center Showing Isodose Curves from the Iridium-192 Source on November 16, 1992.

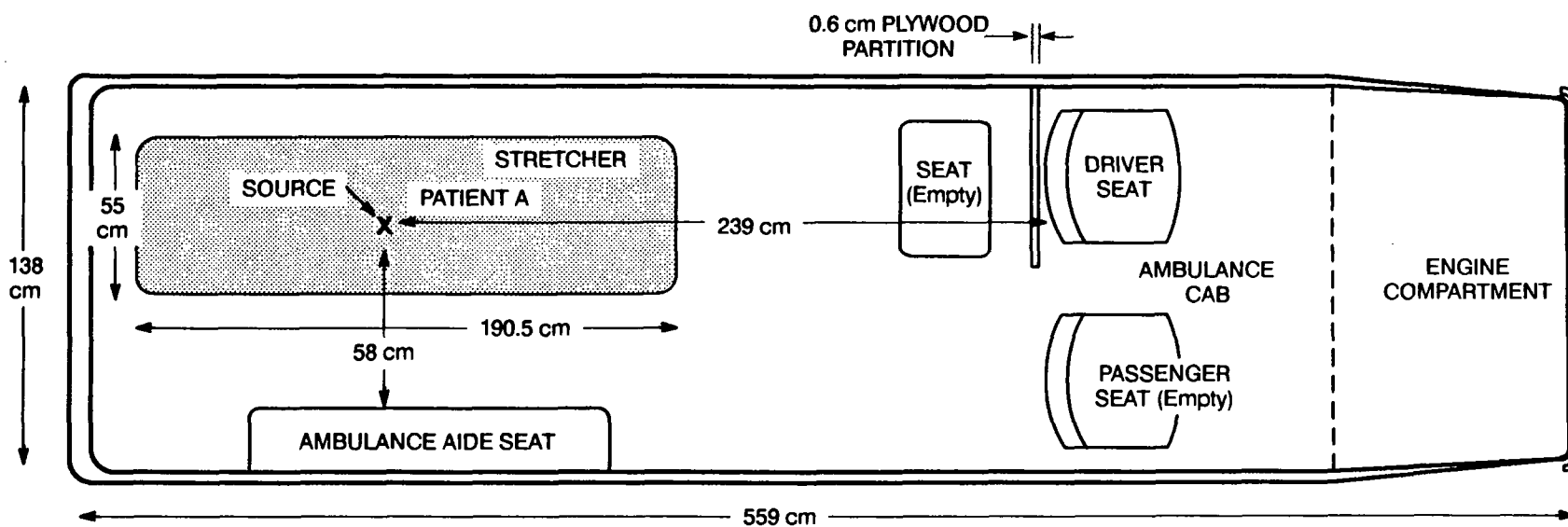
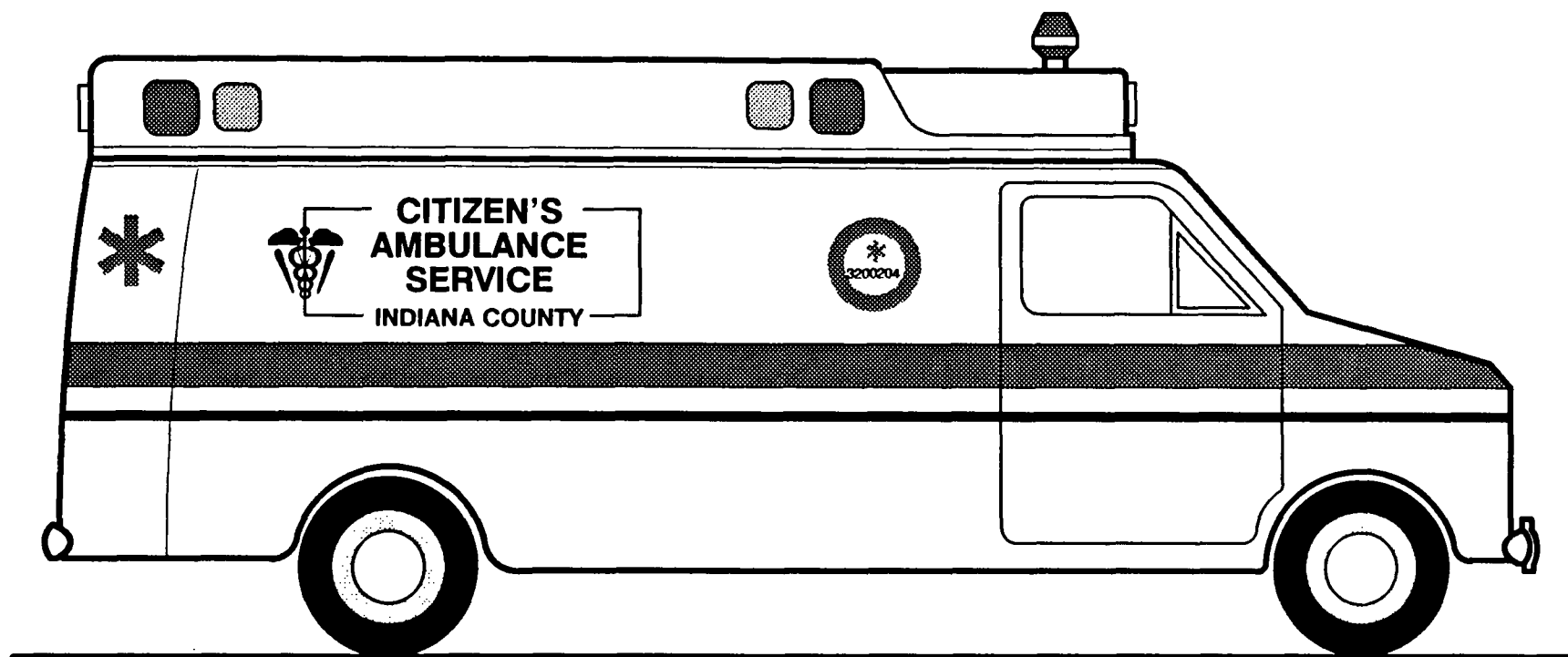


Figure 6.2 Location of Ambulance Staff and Patient A in Citizen's Ambulance

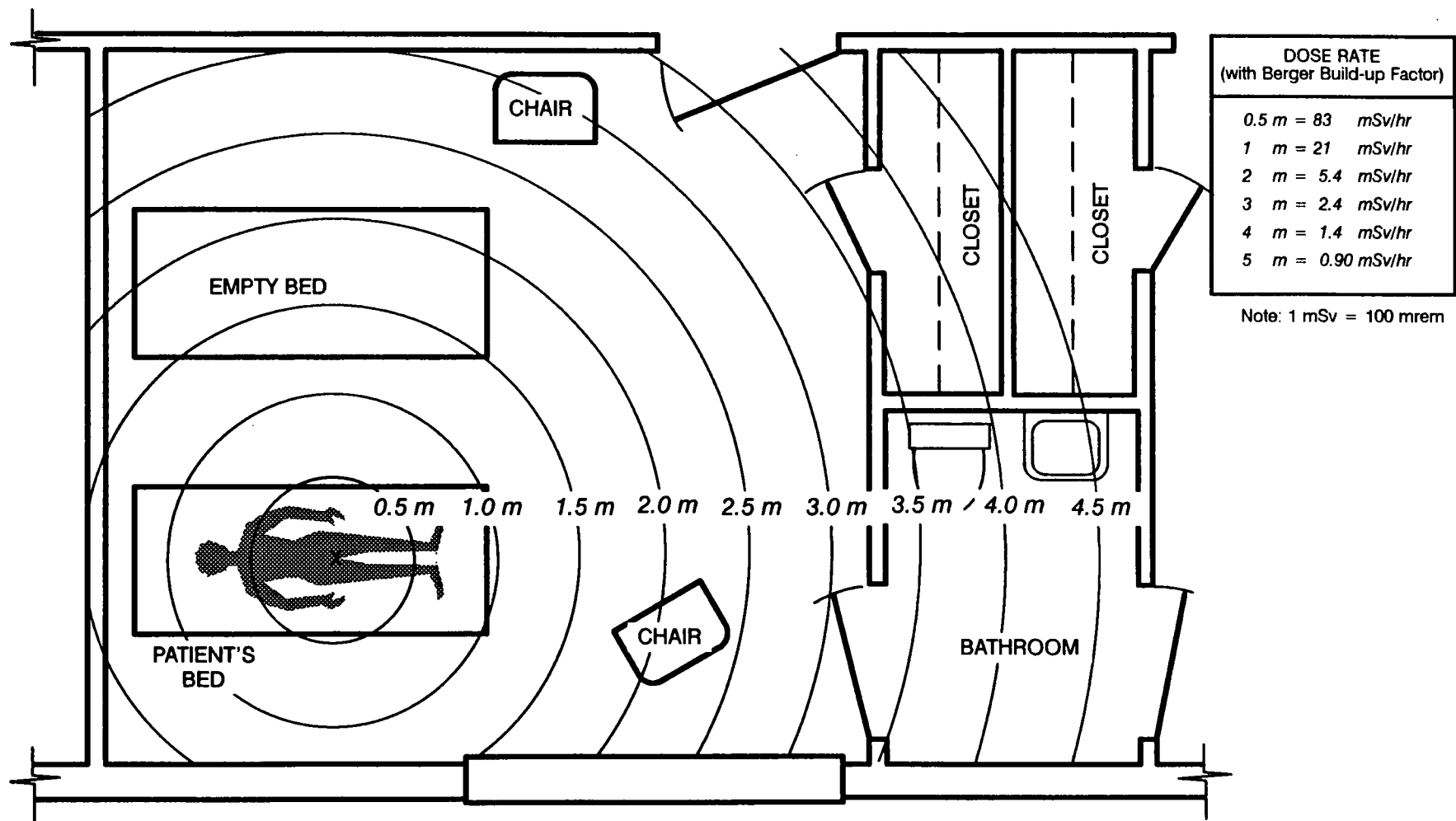


Figure 6.3 Diagram of Room 4B at the Scenery Hill Manor Nursing Home Showing Isodose Curves from the Iridium-192 Source on November 16, 1992

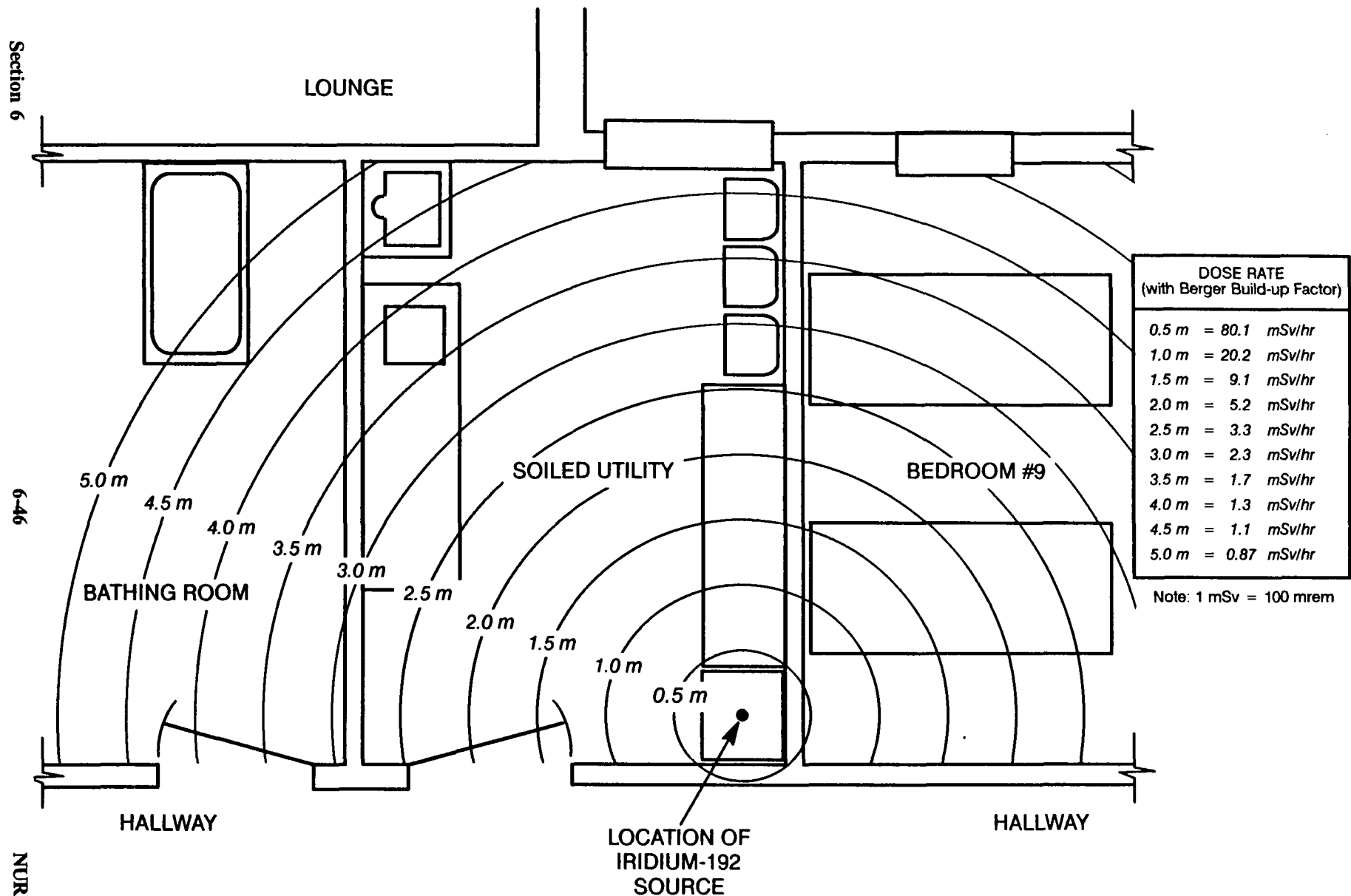


Figure 6.4 Diagram of the SHM Nursing Home Soiled Utility Room Showing Isodose Curves from the Iridium-192 Source on November 20, 1992

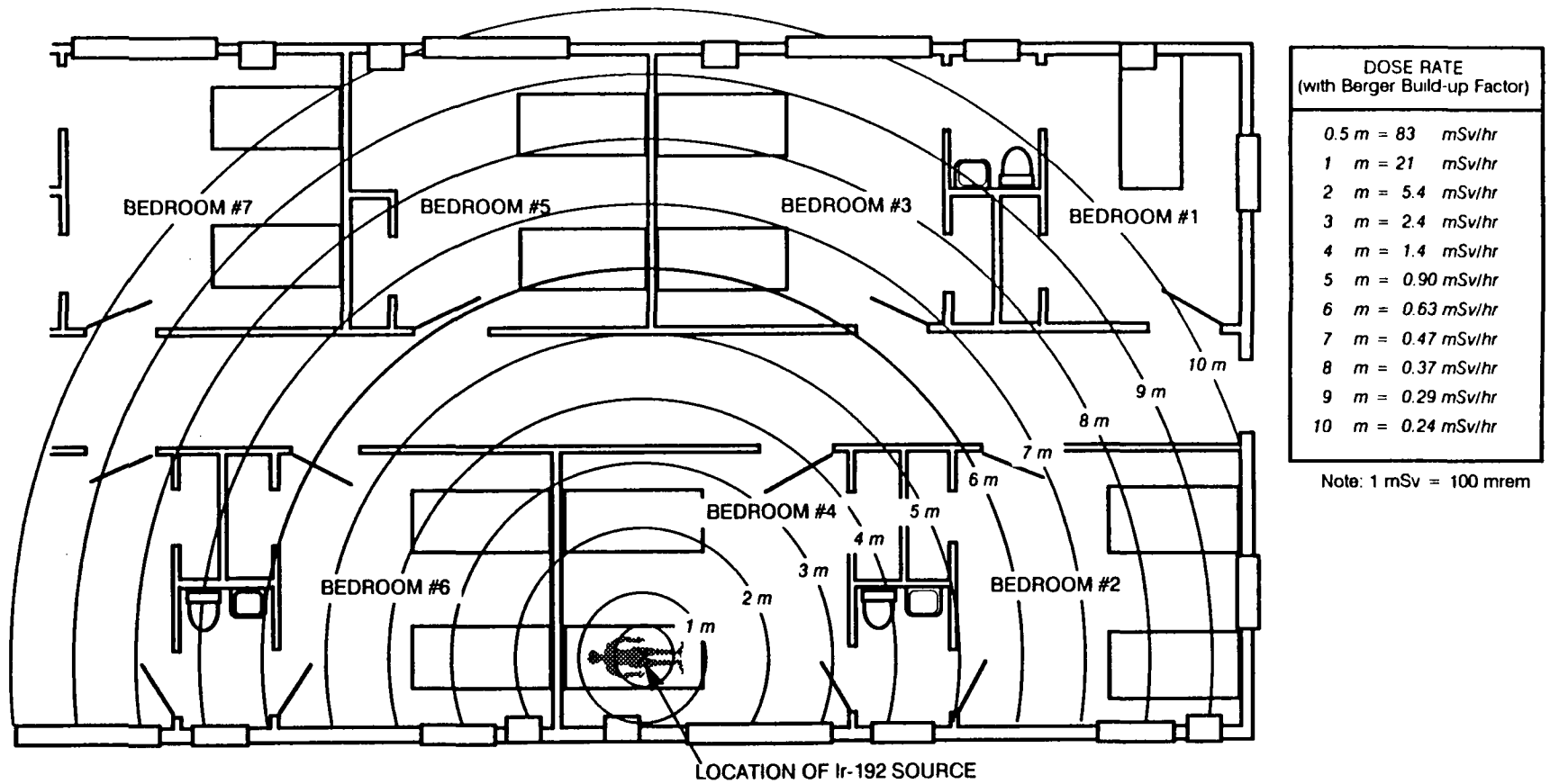


Figure 6.5 Diagram of Rooms Adjacent to Patient Room 4B Showing 1-Meter Isodose Curves from the Iridium-192 Source on November 16, 1992

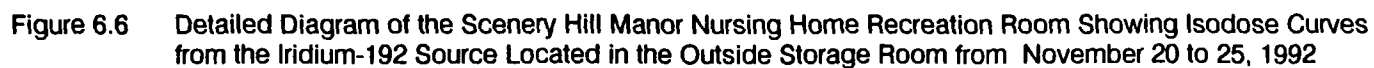




Figure 6.7 Portable Radiation Survey Meter from BFI-Carnegie Straight Truck

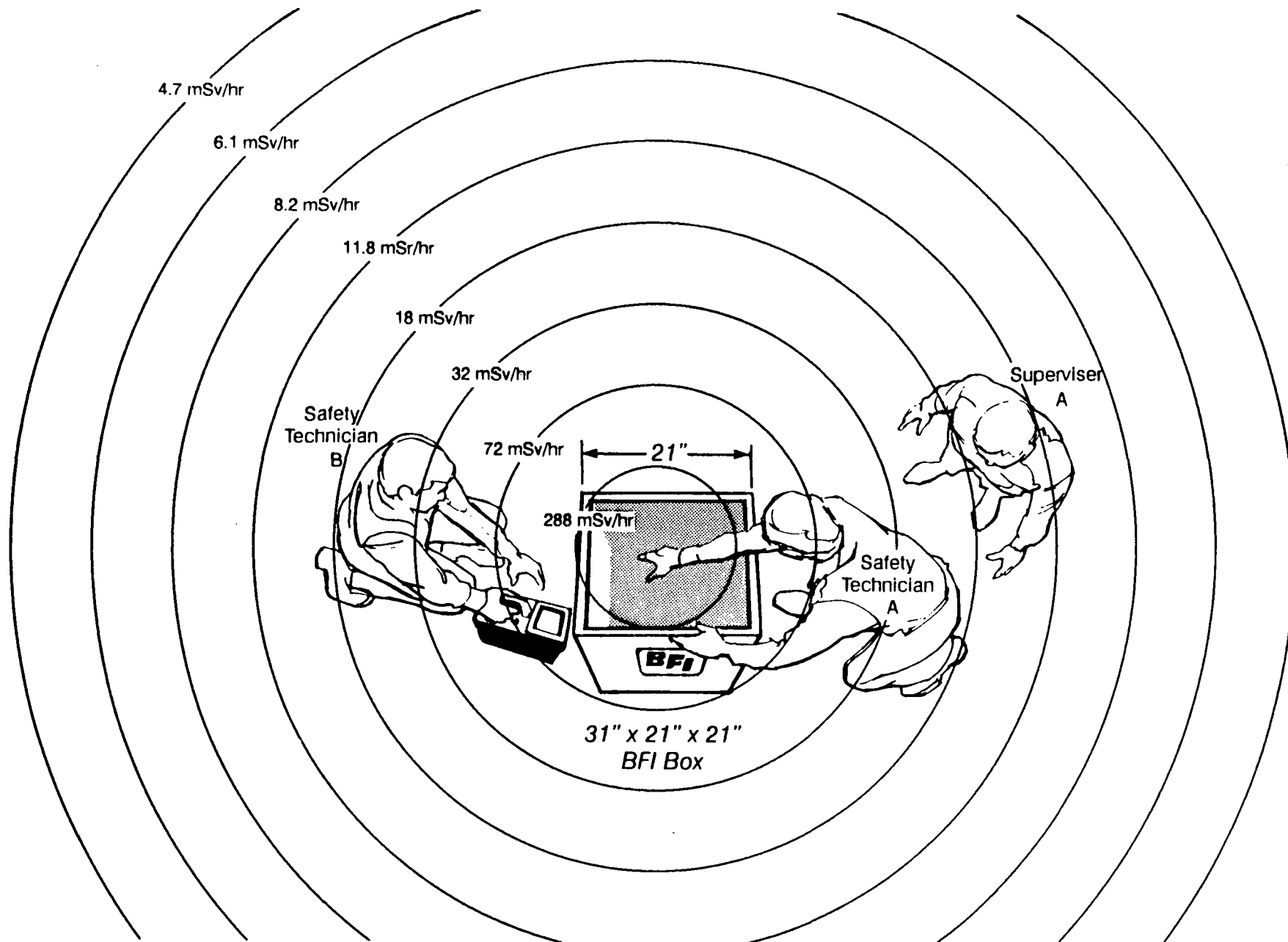


Figure 6.8 Aerial View of the Positions of Supervisor A, Safety Technician A, and Safety Technician B Around the Box Containing the Iridium-192 Source on December 1, 1992, Showing Isodose Curves in mSv per Hour

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7 LICENSEE AND VENDOR RESPONSE TO INCIDENT

The following section briefly summarizes the licensee's actions related to the November 16, 1992, misadministration notification, reports to the NRC, licensee assessment of personnel radiation exposure, and assessment of patient's dose. This section also describes the licensee's initial corrective actions and the vendor's actions.

7.1 Misadministration

As defined in NRC regulations Part 35 of Title 10, "Energy," U.S. Code of Federal Regulations (10 CFR Part 35), a brachytherapy misadministration would occur when the calculated administered dose differs from the prescribed dose by more than 20 percent of the prescribed dose. This incident far exceeded this criteria.

In this incident, Patient A received a severe medical misadministration. The physician intended to deliver a total dose of 1800 cGy (rad) to the tumor, delivered in three 600 cGy (rad) fractions. By the licensee's own calculations, the delivered dose to the same treatment point was in the order of 1,600,000 cGy (rad).

The NRC's medical consultant concluded that Patient A experienced a severe acute radiation syndrome from an overdose of radiation, which was a probable contributing cause of death.

7.2 Licensee Actions

7.2.1 Initial Event Notification

The Medical Physicist A initially notified NRC's Region I Office on December 1, 1992, of a radiological incident involving the loss of a $1.37 \text{ E}+11 \text{ Bq}$ (3.7-Ci) iridium-192 sealed source from the facility (Indiana Regional Cancer Center (IRCC)) in Indiana, Pennsylvania. The licensee reported that Browning-Ferris Industries (BFI) (a nonradioactive waste removal company) notified a local nursing home of the discovery of radioactive material in waste that BFI-Carnegie, a BFI facility in Carnegie, Pennsylvania, previously removed from the nursing home. Suspecting that the radioactive material may have originated from a nursing home resident that had previously undergone radiation therapy at the IRCC, the nursing home immediately notified the IRCC that BFI personnel had discovered radioactive material.

Knowing that a Scenery Hill Manor (SHM) nursing home resident had undergone high dose rate (HDR) treatment at the IRCC, Medical Physicist A performed radiological surveys, observed the PrimAlert-10, and performed an x-ray film autoradiograph evaluation of the HDR afterloader source wire to ascertain the location of the source. The licensee reported that the PrimAlert-10 did not alarm when the HDR afterloader was activated and the radiological surveys and the autoradiograph indicated the HDR afterloader iridium-192 source was missing.

Initial investigation of the incident and the licensee reports to the NRC indicated that the HDR afterloader was last used for patient treatment on November 16, 1992, when the patient was brought to the IRCC by local ambulance from the SHM nursing home for HDR treatment at the IRCC.

The licensee stated that the patient was returned to the SHM nursing home after HDR treatment at the IRCC and, apparently, the iridium-192 source became dislodged from the HDR afterloader and moved with the patient to the SHM nursing home. The licensee's initial report did not indicate how the iridium-192 source was dislodged, whether the source was still implanted in the patient or lost in the patient bedding/clothing material, or how the source was found in the waste at BFI-Carnegie. The licensee reported that the iridium-192 source was secured inside an enclosure at BFI-Carnegie. The licensee informed the NRC that plans were completed, with all appropriate radiological safety precautions, to retrieve the iridium-192 source from the BFI-Carnegie facility and return it to the IRCC. At approximately 4 p.m. on December 1, 1992, the licensee, IRCC, reported to the NRC that the iridium-192 source had been successfully retrieved and would be secured later at the IRCC.

Medical Physicist A stated that the source retrieval was completed in about 70 seconds. Forceps and a cylindrical container (similar to a 5-gallon drum) with a lead shield were used during source retrieval and transportation. The source container was surveyed by Medical Physicist A. The container was placed in the back of Medical Physicist A's truck for transport to IRCC. Dose rates on contact with the container's side and at 1 meter (3 feet) were recorded by Medical Physicist A as 0.35 mSv/hr (35 mrem/hr) and 0.03 mSv/hr (3 mrem/hr), respectively.

The NRC's independent survey of the source container at IRCC on December 2, 1992, confirmed the medical physicist's measured dose rates at contact and at 1 meter (3 feet) from the side of the container. However, the dose rate at contact with the top of the container was approximately 2 cSv (rem) per hour. Medical Physicist A resurveyed the top of the source container and measured approximately 2 cSv (rem) per hour. Medical Physicist A stated that he recalled inserting the source in its shielded location within the source container when the source was transported from BFI-Carnegie to the IRCC on December 1, 1992. The Medical Physicist added that it was possible that the source may have moved within the shielded container and that this would have contributed to the elevated dose rate on the container's top. Medical Physicist A stated that no radioactive material placarding was used on the source transport truck; however, the source container was well shielded and braced during transportation.

An Omnitron International, Inc. (Omnitron) representative opened the source container and pushed the source further into its shielded location. This reduced the dose rate at the top of the source container to 0.35 mSv/hr (35 mrem/hr). Medical Physicist A stated that the source container was placed in the HDR treatment room on the top of a table at the corner of

the room far from the accelerator treatment area with the top of the container facing up. Therefore, the dose to personnel or patients would have been minimal.

7.2.2 IRCC Incident Reports

On December 10, 1992, Physician A submitted a written report on the incident to the NRC Incident Investigation Team (the team) Leader. The report described the incident and the subsequent events, including (1) the BFI notification to the nursing home of radioactive material that was found, (2) the IRCC's discovery of the lost HDR afterloader source, (3) the IRCC's notification to NRC of the incident, (4) source retrieval activities by IRCC personnel, (5) radiation exposure to personnel, (6) patient exposure, and (7) IRCC actions taken to prevent recurrence of similar incidents.

The December report stated that on November 13, 1992, an 82-year-old female patient with anal canal carcinoma was surgically implanted with five catheters at a local hospital. Three days later, on November 16, 1992, the patient was brought by ambulance to IRCC. After being seen by Physician A, the patient was taken to the HDR treatment room and the patient's five implanted catheters were connected to the HDR afterloader under the supervision of Medical Physicist A. The licensee reported that no difficulty was encountered during the treatment using the first four catheters and no error messages were observed on the HDR computer monitor screen until the fifth and final catheter was entered by the source wire, at which point the treatment was interrupted and a message on the HDR computer monitor screen indicated that an obstruction had prevented entrance of the source wire into the fifth catheter. After retri al to treat the fifth catheter had failed, Physician A ordered the procedure to be terminated.

Although the December 10 report indicated that, at some point in the incident, one technologist mentioned that the PrimAlert-10 area radiation monitor light had been flashing red, Physician A did not observe, to his recollection, any flashing of the PrimAlert-10 monitor. However, Physician A informed the NRC staff on two separate occasions on December 2 and 3, 1992, that he noticed the PrimAlert-10 flashing red light. A licensee's consultant audit report, dated January 5, 1993, stated that Physician A, RTT-A, and RTT-B entered the treatment room and did not notice that the PrimAlert-10 light was flashing red, indicating that radiation levels were still elevated in the room. The licensee's consultant audit report stated that RTR did notice the PrimAlert-10 light flashing red and called it to the attention of the other three individuals. The consultant's report stated that the other three individuals chose to disregard it since it had flashed previously when there was no problem. This report adds that they unplugged the PrimAlert-10 and reconnected it later after the patient had been taken to an examination room.

The team learned from interviews with IRCC staff that they had informed Physician A before he had entered the HDR treatment room of the flashing of the PrimAlert-10 red light that they had observed when they first encountered difficulties inserting the source wire into the fifth catheter.

In the December 10, 1992, report, the IRCC stated that after termination of the patient treatment, Physician A and Nurse A examined the patient. Physician A stated, "This examination revealed that the fifth, most posterior catheter, was extruded approximately 2 cm from its desired position. Its suture was cut and the catheter and suture were removed."

The team review indicated that the treatment plan for Patient A required placement of the five implanted catheters as shown in Figure 7.1. Physician A and Medical Physicist A identified the implanted catheter placements as being the same as those shown in Figure 7.1. Physician A and Nurse A identified the location of the implanted catheter removed on November 16, 1992, as being the implanted catheter connected to Channel 5 in Figure 7.1. However, RTT-A, who had connected the implanted and connecting catheters to the HDR afterloader's turret head, identified the locations of the implanted catheters as that shown in Figure 7.2, which is the reverse of the treatment plan. This reversal of the connections made to the HDR afterloader's turret head places the implanted catheter connected to Channel 5 on top (see Figure 7.2). The SHM nursing home staff identified the top location as being the location of the implanted catheter that had extruded from Patient A on November 20, 1992, and had been subsequently removed and placed in SHM's nursing home outside waste storage room for BFI waste pickup.

Therefore, given RTT-A's description of the implanted catheter placement, and the SHM staff's recollection of the extruded implanted catheter location, the source was left in the implanted catheter attached to Channel 5 (top location in Figure 7.2), which remained in the patient until November 20, 1992, and was eventually recovered on December 1, 1992.

7.2.3 Licensee Dose Assessment Reports

On December 23 and 24, 1992, IRCC and Oncology Services Corporation (OSC) submitted two reports to the NRC. The reports described the incident, the dose received by potentially exposed personnel, the patient dose, the licensee's corrective action, and the results of audits of the HDR therapy program at OSC centers. In a facsimile dated January 7, 1993, entitled "Supplement to Part 20 and 30 Reporting Requirements," the licensee provided additional information relevant to the chronology of events, the effect on patient and patient dosimetry, effects on other individuals exposed, action to prevent recurrence, and a preliminary "independent audit" report on the incident.

The licensee conducted interviews of personnel potentially exposed to radiation from the iridium-192 source. Also, blood samples of selected individuals were taken and blood counts were performed to identify personnel with potential high levels of radiation exposures (see Section 6). Section 6 also presents the team's calculations of doses for those that may have been or were exposed to the source.

IRCC's December 10, 1992, report indicated that dosimetric evaluations were done on potentially exposed personnel. IRCC reported that 16 SHM nursing home staff members had received radiation doses in the range of 1.5 to 102 mSv (0.15 to 10.2 rem). However, the

team's examination of the SHM nursing home staff exposure indicated that 21 identified workers had received doses in the range of between 4.0 to possibly as high as 223 mSv (0.4 to 22.3 rem). All calculated doses are given as a range of values.

IRCC reported that six persons visited the patient at the nursing home and that their estimated exposure ranged from 23 to 68 mSv (2.3 to 6.8 rem). The team's assessment of these individual exposures indicated that one person whose dose was estimated at 34 mSv (3.4 rem) by the licensee did not receive any exposure since their visit occurred after the catheter containing the source was removed. Also, the NRC's assessment of potential exposures of patient visitors at the nursing home indicated that seven visitors had received exposures in the range of 22 to 166 mSv (2.2 to 16.6 rem). The licensee's consultant who performed the dose calculation stated he assumed 14 cm or 2 half-value layers of tissue in the path of the radiation. The team's calculations show that the use of narrow beam attenuation factors without buildup is inappropriate (see Appendix B).

The licensee's reports did not address potential exposures to the nursing home residents. The NRC's assessment of potential exposure of the nursing home residents identified 13 individuals who received calculated doses in the range of 11 to 197 mSv (1.1 to 19.7 rem).

The licensee reported that the two ambulance service attendants' who transported the patient to the nursing home had a total exposure of 1.6 mSv (0.16 rem) each. The team's assessment of the two ambulance service attendants exposure indicated that the two individuals may have received exposures in the range of 4.8 to 25.7 mSv (480 to 2570 millirem). OSC or IRCC did not consider the location of the second attendant sitting near, about one-half meter, from the patient.

The licensee's report did not address potential exposure to individuals at an office adjacent to the IRCC. The team's assessment of the potential exposure of individuals in the adjacent office identified five individuals who may have received doses in the range of 0.08 to 4.3 mSv (8 to 430 mrem).

The licensee reported that six BFI individuals may have received doses in the range of 0.3 to 28 mSv (0.03 to 2.8 rem). The team's assessment indicated that these six individuals may have received doses in the range of 0.34 to 90 mSv (0.034 to 9.0 rem).

The licensee reported no occupational exposure that exceeded the NRC regulatory limit [12.5 mSv (1.25 rem) in a calendar quarter]. The dosimeter results indicated that exposures for the calendar quarter (up to December 1) for technologists RTT-A, RTT-B, and RTR were 8.2, 1.1, and 1.4 mSv (820, 110, and 140 mrem), respectively. The licensee estimated Nurse A's exposure to be 4.0 mSv (0.4 rem). The team noted Nurse A's film badge showed 6.3 mSv (0.63 rem). The licensee estimated the exposure of a laboratory technician, who spoke to the patient for about 10 minutes, to be 3.0 mSv (0.3 rem). The NRC's exposure estimate of 3.5 to 13.9 mSv (0.35 - 1.39 rem) for the laboratory technician (phlebotomist)

was based on an estimated time of 10 minutes at a distance of 0.5 to 1.0 meters (20 to 39 inches) from the patient.

The licensee estimated that Physician A's exposure as a result of both Patient A's examination on November 16 and the source retrieval on December 1, 1992, was 5.2 mSv (520 mrem). Physician A received an estimated 1.2 mSv (120 mrem) based on Medical Physicist A's whole-body dosimeter reading during source retrieval because they had worked together on December 1, 1992. Physician A did not use the assigned whole-body dosimeter either during Patient A's examination on November 16, 1992, at IRCC or during retrieval of the source on December 1, 1992, at the BFI-Carnegie facility. The team's assessment of Physician A's exposure of 8.60 mSv (860 mrem) was based on summing the exposure for Nurse A, RTT-B, and Medical Physicist A as indicated by their film badges.

7.2.4 The Licensee's Assessment of Patient's Dose

In the December 23, 1992, addendum to the December 10, 1992, report, the licensee stated that the patient dosimetry calculation indicated a minimum total body dose of 1.6 Gy (158 rad), a dose of 6.3 Gy (630 rad) to surface of a sphere with a 50-cm radius, and a minimum dose of 158 Gy (15759 rad) to surface of a sphere with a 10-cm radius. The licensee added that the mean or median total body dose probably lies between the 1.6-Gy (158-rad) and 6.3-Gy (632-rad) values, and may approach 4 Gy (400 rad).

Physician A stated in his December 10 report that although the 4-Gy (400-rad) dose, "is below the published LD50 dose (lethal dose to cause mortality in 50% of exposed population) of 5 Gy (500 rad), it is of sufficient magnitude that I believe that it is highly probable that the radiation exposure was at least a contributing factor to the patient's subsequent death." Physician A, disagreeing with NRC Bulletin 92-03, stated that it is not consistent with the facts as known in this case to state "that the patient either died as a result of exposure to radiation or that radiation exposure was a major contributor to her death," as was stated in NRC Bulletin 92-03, dated December 8, 1992. Physician A did not evaluate the doses to significant internal organs or the potential consequences of such doses.

The December 23, 1992, report was supplemented by information the team received from OSC on January 7, 1993. In a January 7, 1993, OSC facsimile, the licensee stated that one of the effects on the patient was significant local tissue damage. Possible significant tissue damage to organs outside the treatment area would depend on the progression of radiation damage over time before the patient expired. The prescribed dose at 1 cm was 18 Gy (1800 rad) to be delivered in three treatments. The licensee reported that the source remained in the patient for 92.7 hours resulting in the following doses based on a $1.4 \text{ E}+11 \text{ Bq}$ (3.8-Ci) source and including the effects of tissue absorption and scatter. At 1 cm, the dose was $1.6 \text{ E}+04 \text{ Gy}$ ($1.6 \text{ E}+06 \text{ rad}$). The licensee reported that at 10 cm, the dose was 140 Gy ($1.4 \text{ E}+04 \text{ rad}$). At 20 cm, the dose was 1.9 Gy (190 rad).

The team's independent dose assessment for patient organs, provided by the NRC's Medical Consultant, is described in Section 6.2.

7.2.5 Corrective Actions

The team reviewed the licensee's ongoing actions to prevent recurrence of similar incidents. Corrective actions were initiated by Physician A and by the Radiation Safety Officer (RSO).

In a written policy statement dated December 9, 1992, Physician A directed the staff to follow additional precautions for HDR brachytherapy treatments. These proposed precautions included additional continuous radiological measurements to be performed during HDR patient treatment by placing a diode detector probe near or over the center of the treatment location. Further, this detector will be used to verify that no radioactive material remains in the patient after each treatment. Also, immediately after each treatment, each patient will be surveyed again with a portable survey meter for radiation before leaving the clinic. Physician A added that all personnel involved in HDR treatment will be initially trained on appropriate radiation safety practices and trained semiannually thereafter. Improvements in the documentation of radiological survey, quality control verifications, and training will also be completed.

The RSO stated and reported to the NRC that in addition to the audit conducted by an independent contractor, he was performing an internal audit. The RSO also stated OSC had taken "extraordinary precautions" by suspending HDR treatments at many selected OSC centers. The HDR treatments will be conducted at certain OSC centers under the supervision of a physicist. In addition, the authorized physician will be required to be present at the HDR console while patient treatment is in progress. The RSO also stated that radiological safety training, including emergency drills and practical factors, will be provided to all personnel before initiating any additional HDR treatment at centers where HDR treatments have been suspended. Additional resources, including qualified physicists, will be hired to ensure that radiation safety training is provided and verified to be effective. In addition, radiological safety training and quality control verifications will be documented. The RSO stated that the center is devoting significant effort to ensuring that high quality patient care is provided and that operations are conducted safely by well-trained personnel.

The team's assessment of the OSC's corrective actions indicated that full implementation and verification of the effectiveness of these actions by the licensee may significantly contribute to enhancement of radiological safety at OSC Centers. These actions may provide greater assurance that HDR treatment would be conducted in a manner that precludes radiological consequences of potential radiological occurrences.

7.3 Vendor Actions

This section summarizes the actions taken by Omnitron the manufacturer of the Omnitron 2000 brachytherapy afterloader system associated with the incident that occurred at the IRCC on November 16, 1992.

OSC notified the NRC on December 1, 1992, about the event at its IRCC facility involving the Omnitron 2000 HDR afterloader. They also notified Omnitron that the radioactive source that had previously broken from the source wire had been secured and stored in a lead storage cask. The RSO and the Vice-President, Product Development, from Omnitron, arrived at IRCC's site in Indiana, Pennsylvania, on December 2, 1992. An NRC Region I inspection team also arrived at the IRCC facility on December 2, 1992. After obtaining approval of the NRC inspection team, the RSO surveyed the afterloader and storage cask for radioactive contamination. The RSO found trace levels of radioactive contamination. The HDR afterloader was then secured in a locked room in the clinic.

On the following day, December 3, 1992, Omnitron's Vice-President, Product Development, was allowed temporary access to the afterloader and the treatment console to retrieve and print out all information, including error messages, recorded from November 16, 1992, to December 3, 1992. The NRC team continuously monitored the Omnitron representatives' areas of access. Omnitron then supplied this information to the NRC team, and transmitted a copy to the Omnitron software developer. This information was transmitted to the software developer to preliminarily identify what caused the source to dislodge itself from the source wire and why this condition was not communicated to the technologists at the IRCC.

Physician E at the Greater Pittsburgh Cancer Center (GPCC) informed the team that he was not called by OSC but that a doctor from Texas, presumed to be from Omnitron, notified him of the event at IRCC before the GPCC incident on December 7, 1992.

On December 7 and 8, 1992, the vendor notified all Omnitron 2000 user sites by telephone about a second incident involving an active source dislodging from the guide wire. The users were told that Omnitron recommends that they temporarily not use their HDR afterloader systems until further information indicating reasons for the source-wire break is obtained. The users were also told that if they should choose to use the device, they should strictly adhere to all safety precautions indicated in the user manual and that the patients should be surveyed for radioactivity both before and after the HDR afterloader treatment.

During the investigation of the source-wire breaks (failures), initial observations indicated that dilute hydrogen fluoride solutions embrittle the nickel-titanium wire. This solution may have been generated in the teflon liner of the shipping container for the source. Omnitron noticed deterioration of the teflon liner and, as a result, made a design change in late September 1992, changing the teflon liner to a stainless steel liner. Because of this initial observation, Omnitron stated in a memorandum, dated January 8, 1993, that they contacted

customers using the old shipping container by telephone and told them not to use the source wire until their source wire is replaced.

Numbers represent placement of implanted catheters within the patient and correspond to the treatment channel number indicated on the turret head.

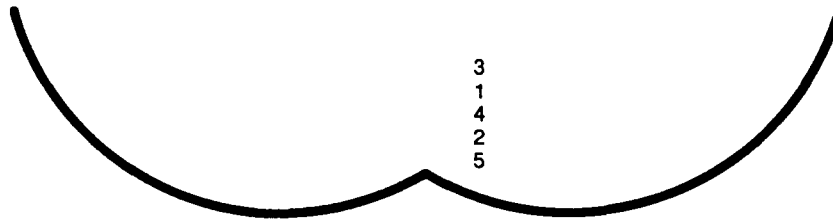


Figure 7.1 Treatment Plan Catheter Location

Numbers represent placement of implanted catheters within the patient and correspond to the treatment channel number indicated on the turret head.



Figure 7.2 Actual Catheter Location

8 REGULATORY OVERSIGHT

8.1 High Dose Rate Afterloader Device Review by the Food and Drug Administration

The team and one of its contractors prepared a description of the regulatory review process followed by the Food and Drug Administration (FDA) for the Omnitron International Inc.'s (Omnitron's) 2000 high dose rate (HDR) afterloader based on FDA's Omnitron file. This section was presented to the FDA to ensure its accuracy; however, the team has not received FDA's comments. Therefore, a description of the FDA's role has not been included in this report. Information on the case has been shared with the FDA who conducted their own inspection. The FDA stated they will furnish the NRC with an accurate description of their role, but it would not be furnished in time to be included in this report.

8.2 Source and Device Registration by the State of Louisiana

Any device or sealed source that contains radioactive material must be evaluated and approved by the NRC or an Agreement State before the device or sealed source can be distributed. The manufacturer or distributor typically provides the information to the licensing authority on behalf of their customers (licensees) so that each customer does not have to provide this information.

In order to approve a sealed source or device for distribution, sufficient information pertaining to the design and construction, labeling, conditions of use, prototype testing, external radiation profiles, and quality assurance (QA) and quality control (QC) program must be submitted so that the reviewer can determine whether the device or sealed source is radiologically safe for use. If the license reviewer determines that the device or sealed source is adequate, then (1) the reviewer issues a certificate summarizing the information submitted, (2) lists the limitations placed on the device or sealed source, and (3) states their conclusion (i.e., a safety analysis) of the sealed source or device. The reviewer signs the document. The document is then independently reviewed by another individual. If this individual agrees with the reviewer's findings then the document is signed by this individual and the sealed source or device is approved for distribution.

Omnitron's office located in Lake Charles, Louisiana, submitted to the State of Louisiana information pertaining to the registration of the nickel-titanium source wire and the HDR afterloader. The nickel-titanium source wire was approved for use on March 9, 1992 (Registration Certificate LA-0760-S-102-S). The HDR afterloader was approved for use on March 26, 1991 (Registration Certificate LA-0760-D-101-S).

8.3 NRC's Regulations and Guidance for Quality Assurance and Quality Control Programs for Sealed Source and Device Vendors

The NRC or an Agreement State must evaluate and approve a device or sealed source that contains radioactive material before the device or sealed source can be distributed. The NRC or Agreement State evaluates the manufacturer's and/or the distributor's QA/QC program as part of the overall evaluation.

The NRC does not have specific regulations or guidance that relate to QA/QC programs for vendors of sealed sources and devices. Section 32.210 of Title 10 of the Code of Federal Regulations (10 CFR 32.210) states that a manufacturer or distributor must include sufficient information pertaining to their QC program to "provide reasonable assurance that the radiation safety properties of the source or device are adequate to protect health and minimize danger to life and property." However, the regulation does not mention what the QC program must entail.

The NRC issued two regulatory guides, Regulatory Guide 10.10, "Guide for the Preparation of Applications for Radiation Safety Evaluation and Registration of Devices Containing Byproduct Material," March 1987, and Regulatory Guide 10.11, "Guide for the Preparation of Applications for Radiation Safety Evaluation and Registration of Sealed Sources Containing Byproduct Material," June 1987. These guides provide assistance to manufacturers and distributors of byproduct sealed sources and devices on how to file an application and on what information and documentation is required in their submittal.

The guidance states that part of the vendor's submittal should describe its "quality control program and procedures to be followed to ensure that each finished device (or sealed source) meets the specifications furnished to the NRC." However, the guidance does not give details on what is necessary for the QA/QC program. The guidance suggests using industry standards. In the team's review of the standards pertaining to the manufacturer of brachytherapy sources and devices (i.e., American National Standard Institute ANSI N542-1977, "Sealed Radioactive Sources, Classification"; ISO 22919-1980 (E), "Sealed Radioactive Sources, Classification"; American National Standard Institute, ANSI N44.1-1973, "Integrity and Test Specifications for Selected Brachytherapy Sources") no specific guidance for QA/QC was found. ISO-2919 was revised in September 1992. This revision mentions a QA program and provides an example of a QA program. The example references ISO-9000, "Quality Management and Quality Assurance Standards Guidelines for Selection and Use," which specifies areas that a QA/QC program should cover.

NRC has evaluated different guidance on QA/QC, such as military standards, ISO-9000, and FDA's GMP, to establish a QA/QC program for vendors of sealed sources and devices. The NRC completed a draft Quality Assurance Manual in October 1991. This draft manual has been used on several visits to vendors of sealed sources and devices in the United States and other countries. The draft QA manual outlines and explains the basic procedures,

policies, and practices needed in a QA/QC program. This manual was used as guidance for the review of Omnitron's QA/QC program.

8.4 NRC Licensing of Oncology Services Corporation

Oncology Services Corporation (OSC) is licensed by the NRC, as prescribed in 10 CFR Part 35 to use several types of HDR brachytherapy remote afterloaders to treat humans. The license was issued by NRC Region I. NRC's Office of Nuclear Materials Safety and Safeguards provides guidance for medical use programs in Regulatory Guide 10.8, "Guide for the Preparation of Applications for Medical Use," Revision 2, August 1987, and specific licensing guidance for brachytherapy remote afterloaders in Policy and Guidance Directive FC 86-4; "Information Required for Licensing Remote Afterloading Devices," issued on February 20, 1986.

Part 35, issued in substantially its current form on October 16, 1986, prescribes requirements for the medical use of byproduct material. Subpart G of Part 35, "Sources for Brachytherapy," provides specific requirements for brachytherapy; however, this subpart is mostly applicable to low dose rate brachytherapy application. Although the latest amendments to Part 35 became effective 6 months after FC 86-4, the regulations do not explicitly recognize HDR brachytherapy, and only some requirements in Subpart G are applicable to HDR brachytherapy. Regulatory Guide 10.8 provides guidance only for low dose rate brachytherapy. Because of the substantial changes to Part 35 in 1986, certain references to an earlier revision of Part 35 in FC 86-4 are inaccurate. These inaccuracies are not significant and appear to have no practical consequence. However, the licensing guidance in FC 86-4 is outdated and is not well integrated with NRC medical regulations or other NRC licensing guides.

The NRC issued original OSC License 37-28540-01 on August 3, 1990. This license authorized the use of one Gamma Med Ili remote afterloading brachytherapy system in Harrisburg, Pennsylvania. On November 2, 1990, the license was amended in its entirety to add additional users and locations and to permit the single Gamma Med system to be shipped from site to site. The IRCC was one of these sites.

License Amendment 2 was issued January 22, 1992. This amendment added several models of HDR afterloader brachytherapy systems to the license for use at several locations, including the IRCC. One of the models authorized for use was the Omnitron 2000 HDR remote afterloader.

Amendment 3 was issued August 19, 1992. This amendment was the version of the license in effect at the time of the incident. It added to the license authorization for a stand-alone HDR brachytherapy shielded facility at the existing Harrisburg facility.

8.5 NRC Inspection Program and Process

NRC Inspection Manual Chapter (MC) 2800 establishes the inspection program for medical licenses, including license priority and inspection frequency. Inspection Procedure (IP)-87100 provides inspection direction for several types of material inspections, including Nuclear Medicine and Medical Teletherapy. The field notes used by inspectors for teletherapy are included in Appendix A to IP-87100 and the field notes for Nuclear Medicine programs, including brachytherapy, in Appendix B to this IP. The section on brachytherapy in the field notes closely follows the requirements in 10 CFR Part 35, Subpart G. None of the guidance explicitly discusses HDR brachytherapy, and FC 86-4 is not referenced, although other NRC licensing guides are referenced in MC-2800. In addition, inspection guidance directs inspectors to review the licensee's license, the license conditions, and referenced applications.

MC-2800 requires an initial inspection within a year of initial licensing of a facility and routine inspections every 3 years for those facilities licensed for medical programs. Facilities licensed for teletherapy programs are to be inspected annually. The frequency for inspecting facilities licensed to use HDR brachytherapy is not explicitly addressed in this MC but is included in its guidance for medical programs. The inspection program guidance is silent for licenses authorizing (1) multiple places of use, (2) use where the Radiation Safety Officer (RSO) can not respond quickly, and (3) license amendments that greatly expand a licensee's scope of operations, including new locations of use after the initial inspection.

The initial inspection of OSC was conducted on September 4, 1991, about 1 year after its initial licensing. The inspection was conducted at the Harrisburg Cancer Center, which is the mailing address of the licensee as well as the routine work location of the RSO. At the time of the inspection, the licensee possessed only one HDR afterloader. The system was transported to other OSC facilities named in the license and stored as well as used in the Harrisburg facility. During this inspection, inspectors identified two Severity Level IV violations (Appendix C to 10 CFR Part 2). Both violations were of modest safety significance and were appropriately categorized. Neither was a precursor of the November 16, 1992, event. A routine reinspection was scheduled for September 1995 in accordance with MC-2800.

The two individuals conducting the inspection in 1991 were experienced and well-qualified inspectors. One was the Chief of the Region I Medical Licensing Section and the other was a senior inspector. Both had work experience in medical radiology in addition to their NRC experience. Their training in brachytherapy was limited to conventional low dose rate brachytherapy; one inspector had extensive work experience in low dose rate brachytherapy.

These inspectors found that the staff involved with the HDR afterloader at the Harrisburg facility had been trained. This finding was based on interviews with six Harrisburg staff members, including the RSO, the physician user, two medical physicists, a driver, and a

technologist. Those questioned were knowledgeable in both operating and emergency procedures. They were also knowledgeable in the areas specified in 10 CFR 19.12. Records of training and retraining were not reviewed and this was stated in the inspector's inspection notes. One inspector stated that, if correctly recalled, they had requested these records. The inspector does not recall why they did not review them. According to the inspector's statement, the inspector may have been distracted. Inspectors did not go to the IRCC during this inspection. The IRCC had no HDR afterloader at this time.

Section 12, Brachytherapy, of the NRC inspector's field notes was marked "HDR only." The inspector stated a belief that the requirements covered by this section (i.e., Subpart G of 10 CFR Part 35) were either not applicable or covered by other sections of the field notes. The inspector noted that operational and emergency procedures were covered by license conditions in lieu of 10 CFR 35.410 "Safety instructions." The inspector correctly notes that 10 CFR 35.415 "Safety precautions," are not applicable to HDR brachytherapy. The inspector believed the requirement in 10 CFR 35.404 to survey the patient immediately after removing the source was met by the area radiation monitor in the treatment room. The inspector stated that this belief was based on the licensee's commitment to comply with the guidance in FC 86-4, which provides for a room monitor to verify the location of a source.

The NRC has no policy on reinspection when a licensee significantly changes or increases the scope of its program by an amendment. At times a licensing visit is performed when such changes are requested.

8.6 NRC Regulatory Activities

The NRC took several actions related to this event before the completion of this report.

1. Bulletin 92-03, issued December 8, informed licensees authorized to use the Omnitron 2000 HDR afterloader of the November 16, 1992, incident and requested that they implement three actions: First, the survey of all patients had to be completed immediately after completing each therapy treatment and before removing the patient from the therapy room. Second, procedures, staff, and equipment had to be available for prompt intervention should the source not retract into the shielded container at the conclusion of the treatment. Third, initial and semiannual training were needed on procedures for both the routine use of the afterloader and in the event of an emergency.
2. Information Notice 92-84, issued December 17, 1992, informed all NRC medical licensees of this incident and reminded them of the requirements for positive assurance that all implanted sources have been removed before patients have been released after treatment. It provided information similar to that in Bulletin 92-03.

3. On January 20, 1993, the NRC suspended the license of Oncology Services Corporation. The action was based on both the findings discussed in this report as well as the results of inspections at two other Oncology Service Corporation facilities that identified training problems similar to those identified by the team at IRCC performed by NRC Region I.

9 PRECURSORS

The following section presents a summary of all hardware-related brachytherapy device precursors that were identified as being similar to the malfunction of the Omnitron International Inc.'s (Omnitron's) 2000 high dose rate (HDR) brachytherapy afterloader unit at the Indiana Regional Cancer Center (IRCC).

9.1 High Dose Rate Remote Afterloader Incidents

For this report, an HDR incident precursor is defined as an event that involves malfunctioning (through breakage, rupture, unintended disconnection) of the source wire or guide tube of a brachytherapy remote afterloader.

From information provided by the Food and Drug Administration, NRC's Office of State Programs, and other NRC Offices, no Omnitron-specific malfunctioning events have been reported to NRC before December 1, 1992. The first Omnitron 2000 system was shipped to a user in February 1991.

To date, the team is aware of one incident involving HDR brachytherapy afterloaders (and sources) manufactured by a company other than Omnitron International, Inc. (Omnitron). However, other similar incidents may have occurred in the United States or in other countries of which the team is not aware.

The precursor, in August 1988, involved a Microselectron HDR afterloader manufactured by Nucletron. In this event, the top of a defective source capsule became detached during a treatment, allowing several high-activity iridium-192 seeds to fall out of the capsule. Several things are important to note. First, when the top of the source capsule separated from the capsule and the body of the capsule returned to the storage safe, the Nucletron device indicated a "safe condition" existed. Second, the PrimAlert-10 monitor in the room was flashing red and the radiation therapy technologists (RTTs) performing the treatment took this signal as an indication that radiation was present in the treatment room. The RTT's actions demonstrated their knowledge of radiation safety and emergency procedures. Third, the RTTs surveyed the patient for radiation after removing the patient from the treatment room.

Subsequent testing and examination performed by the source vendor (Mallinckrodt Diagnostica B.V.) indicated that an incomplete welded joint on the source end cap was the cause of the source failure. To prevent recurrence of this type of event, both the source vendor and Nucletron improved quality control procedures associated with the manufacture of the source. Nucletron has since changed the overall design of the source used in the Microselectron HDR afterloader.

9.2 Omnitron 2000 Performance History

Omnitron was founded in 1988, and its first HDR afterloader system was developed in 1989. Omnitron demonstrated it at an industry show and decided that the system needed some modifications and improvements. The first commercial unit was shipped in February 1991. Currently, 25 units with an active source wire installed are in service in the United States.

Omnitron's President stated that the afterloader's performance history has been excellent. Several individuals who work for Omnitron stated that they were unaware of any production source-wire breaks except for those at the IRCC and the Greater Pittsburgh Cancer Center.

A review of the complaint file provided to the team indicated that 41 complaints have been filed from 15 different customers. The most frequent complaints that required service are categorized as follows:

- Dummy wire fails to retract/park or constricted—7 complaints
- Printer problems—5 complaints
- Planning system problems—4 complaints
- HDR hardware problems—4 complaints
- Computer console hardware problems—2 complaints
- Software problems—2 complaints
- Source wire failed length check—1 complaint
- Contamination problem—1 complaint

The remaining 15 complaints were handled by calling the customer and working through the complaint with them by telephone.

When service calls are placed, or when routine maintenance is required, Omnitron representatives go to the customer's site and perform the required service. A service checklist was provided to the team by Omnitron's service manager. According to this service manager, the checklist is used and discarded after its use. The service checklist shows tests, inspections, and parts that are replaced during service.

The only service records that were saved are the actual service-order documents. These documents list the parts replaced, the service requested, the type of service, and other information. However, in reviewing these records, the team found little or no information explaining why the item was replaced or why the problem occurred.

10 FINDINGS AND CONCLUSIONS

The team has compiled the following list of principal findings and conclusions concerning the incident. Each major conclusion appears as a numbered section.

10.1 Radiological Consequences Were Serious to the Patient and Significant to Many Members of the Public.

Patient A at the Indiana Regional Cancer Center (IRCC) received a severe medical misadministration and subsequently died. The physician intended to deliver a total dose of 1800 cGy (rad) to the tumor delivered in three 600 cGy fractions. By the licensee's own calculations the delivered dose to the same treatment point was in the order of 1,600,000 cGy (rad). Radiation absorbed doses to normal tissues and vital organs were calculated by NRC's Medical Consultant assuming a 92.75-hour exposure from a $1.56 \text{ E}+11 \text{ Bq}$ (4.22-Ci) iridium-192 source, which was the source strength as indicated on the Omnitron computer printout. Doses to the kidneys ranged from 27 to 41 Gy (2,712 to 4,139 rad), doses to the small bowel ranged from 64 Gy (6,369 rad) at the median point to 476 Gy (47,605 rad) to the closest point, and doses to the bladder ranged from 201 to 3,607 Gy (20,142 to 360,663 rad). As stated by NRC's Medical Consultant, doses in the half million to one million cSv (rem) range should cause extensive necrosis in a few days and possible rectal perforation. Therefore, the NRC's Medical Consultant concluded that Patient A experienced a severe acute radiation syndrome from an overdose of radiation, which was a probable contributing cause of death. In a press release dated January 26, 1993, the Indiana County Coroner stated that the cause of death listed in the official autopsy report was "Acute Radiational Exposure and Consequences Thereof."

Forty-nine nonoccupational workers and members of the public received exposures calculated to be in excess of 0.005 Sv (0.5 rem). Of these exposures, 21 were calculated to have exceeded 0.05 Sv (5 rem). The upper limit on exposures is calculated to be 0.15 to 0.25 Sv (15 to 25 rem). The calculations were based on reconstructions of activities and estimates of dwell times that result in uncertainties in individual exposures. All nine cytogenetic evaluations are in agreement with the above upper limit and generally consistent with calculated doses.

The NRC's medical consultant concluded there were no acute effects of radiation seen in the nursing home staff, other residents, visitors, or members of the general public.

Population groups receiving significant exposures included individuals at the IRCC; the adjacent physician's office; the staff, residents, and visitors at the SHM nursing home; the ambulance driver and aide; and BFI workers at Carnegie, Pennsylvania.

The evaluation of radiological doses submitted by the licensee were lower than the team's calculations in some cases by as much as a factor of 3. In addition, the licensee did not

calculate doses for any of the residents of the Scenery Hill Manor (SHM) nursing home, even though in several cases, these residents were known to have been exposed (later confirmed by cytogenetic evaluation) to the radiation source because of their close proximity to Patient A's room.

No occupational worker's exposure exceeded the 0.0125-Sv (1.25-rem) quarterly occupational limit.

The most conservative estimates for individuals not specifically evaluated in the report but inadvertently exposed to the source are less than 1 mSv (100 mrem). This is less than the current allowable exposure limit for a member of the general public.

The source, an iridium-192 wire, released essentially no radioactive material. No contamination was detected at the IRCC, in the ambulance or at the SHM nursing home. Traces, less than the NRC (185 Bq) 5-nanocurie limit, of iridium-192 were detected on the active and dummy source wires.

10.2 Weaknesses in Oncology Services Corporation's Radiation Protection Program Were a Contributing Cause of the Seriousness of the Event and Radiation Exposure Consequences.

OSC's Radiation Protection Program was ineffective and incomplete. The corporation's Radiation Safety Officer (RSO) did not provide sufficient oversight to ensure that all personnel associated with operation of the HDR afterloader were properly trained in its use. The RSO did not ensure that all personnel were properly trained and knowledgeable in basic radiation safety. Further, the RSO was not aware of the IRCC's authorized user's practice of authorizing a technologist, who had not completed the Omnitron International, Inc.'s (Omnitron's) training course, to operate the unit.

Other than the operator's manual provided by Omnitron, approved written policies, procedures, and guidelines were not provided to OSC personnel. Practices followed at the Greater Pittsburgh Cancer Center (GPCC) that limited the consequences of their source-wire-break incident included surveying the patient and the afterloader following patient treatment, and having the medical physicist and the physician present at the console. These practices were not followed at IRCC. The lack of adequate emergency procedures and training contributed to the failure of the IRCC staff to respond to the PrimAlert-10 area radiation monitor alarm. The RSO was not aware of the PrimAlert-10 spurious alarms and the IRCC reaction to the alarms.

Further, owing to the lack of management oversight, guidance, and training, OSC personnel at the IRCC did not adequately respond during the recovery of the source, which resulted in apparent violations of Department of Transportation (DOT) regulations; unnecessary exposure to a nonoccupational worker; and inadequate and incomplete reports to the NRC concerning whole-body and organ doses received by the patient.

10.3 Weaknesses Existed in the Design and Testing of the Omnitron 2000 Remote Afterloader System and Its Source Wire.

The two failures of the source wire in service in the Omnitron 2000 high dose rate (HDR) afterloader may be attributed to weaknesses in design of the source wire and the testing program to validate the design. Specifically, the manufacturer performed no engineering calculations for stress and fatigue of the wire before this event, particularly the source cavity. All breaks during testing and in service have occurred in the bottom section of the cavity. Although they performed prototype testing in accordance with ISO 2919, they performed no additional tests before this event to ensure that environmental conditions would not affect the integrity of the source wire, particularly, tests for environmental effects of radiation, moisture, and the presence of teflon within the shipping container. The material teflon has been known to have poor resistance to radiation in the presence of oxygen.

The HDR afterloader was not designed to positively verify that a source wire has fully retracted if the emergency dc motor is being used to retract the source. If a broken source wire is not initially detected by the HDR afterloader, the HDR afterloader is not designed to detect the broken source wire during subsequent use because it is not designed to measure and record the overall length of the wire.

Personnel performing treatment are briefly notified by an error message displayed on the computer monitor that an emergency dc retraction is occurring (i.e., an error message only appears on the screen for the short duration of the retraction). Once the source wire has retracted, the error message is cleared, and no other design features verify full retraction of the source wire or whether the emergency motor retracted the wire. The manufacturer established no program to recover and examine "spent" source wires to look for evidence of deterioration and confirm the adequacy of the source-wire design.

Although Omnitron's quality assurance and quality control (QA/QC) program was weak, the team identified no other deficiencies that could be directly related to the source-wire breaks.

10.4 Oncology Services Corporation and Indiana Regional Cancer Center Lacked Critical Safety Awareness with Respect to High Dose Rate Brachytherapy.

The safety culture at the IRCC significantly contributed to the incident. A pervasive lack of critical radiation safety awareness existed within the IRCC organization. The authorized user was not wearing a film badge during both encounters with the source. Technologists routinely ignored the PrimAlert-10 alarm and habitually sought to override this alarm by unplugging the unit to reset it. No one at IRCC who operated the HDR afterloader ever verified the operation of safety interlocks associated with the unit before its use. Technologists did not survey patients, the afterloader, or the treatment room following HDR

treatment. The authorized user rarely observed operations and was not present at the console during treatment in this incident. The IRCC technologist who had not completed Omnitron training was allowed to perform unsupervised treatments.

Omnitron personnel believed and apparently had led most OSC personnel to believe that a source-wire break was not possible and, in fact, had not provided emergency procedures adequate to mitigate such an event. Consequently, IRCC personnel did not consider a source-wire break as a credible event.

These assumptions were exacerbated by the facts that the corporation's RSO rarely visited the site and did not audit radiation safety practices and that IRCC personnel were not aware of who was responsible for radiation management and training or who was listed as the RSO on the license.

10.5 Overall Regulatory Oversight Was Weak.

The team identified regulatory oversight weaknesses in several areas although none directly caused this incident or increased the severity of the consequences. The weaknesses were in the areas of regulatory oversight of HDR afterloader users, the licensing and inspection of programs that expand significantly by amendment after initial licensing; and the overall regulation of vendors of devices that use licensed nuclear material.

HDR afterloader license applications for use of HDR afterloaders are reviewed using guidance provided by NMSS Policy and Guidance Directive FC 86-4, "Information Required for Licensing Remote Afterloading Devices." Except for Regulatory Guide 8.33, "Quality Management Program," HDR is not addressed specifically or FC 86-4 referenced in any other NRC regulation, licensing guide, or inspection procedure, most of which were issued after FC 86-4 was issued. Because the current effective 10 CFR Part 35, and its associated medical licensing guide, Regulatory Guide 10.8, Revision 2, were issued after FC 86-4, licensing guidance in FC 86-4 is not totally consistent with the general requirements of 10 CFR Part 35. Most importantly, FC 86-4 does not reference the patient survey requirements in 10 CFR 35.404(a), which require a survey with radiation survey instruments.

Some experienced NRC staff in the Regions were confused over the applicability of some 10 CFR Part 35 requirements to HDR brachytherapy. The NRC staff's medical inspection procedures are consistent with Part 35 requirements and the medical licensing guide but reference neither HDR nor FC 86-4. NMSS Headquarters managers stated they believed that NRC inspectors were not always enforcing the patient survey requirements of 10 CFR 35.404(a). Instead, some NRC inspectors and licensees believed area radiation monitors met the survey requirements of 10 CFR 35.404(a), which was the case at IRCC where patients were not surveyed after treatment.

NRC licensing and inspection guidance did not clearly focus on a program that rapidly expanded, as was the case with OSC, when License Amendment 2 was issued. This

amendment did not require special management controls to ensure the RSO could manage radiation safety training and control at remote facilities. The NRC inspection Manual Chapter 2800, while requiring an early initial inspection, allowed the next inspection to be set 4 years after the initial inspection, regardless of how significantly the scope of licensed activities changed after the initial inspection. Furthermore, even if the initial inspection had been conducted after the program expanded, a performance-based inspection at the Harrisburg facility alone would not have identified the training problem at other OSC facilities.

This investigation also revealed a number of weaknesses in the design and QA and QC program of the vendor. NRC issued two regulatory guides, Regulatory Guide 10.10, "Guide for the Preparation of Applications for Radiation Safety Evaluation and Registration of Devices Containing Byproduct Material," March 1987, and Regulatory Guide 10.11, "Guide for the Preparation of Applications for Radiation Safety Evaluation and Registration of Sealed Sources Containing Byproduct Material," June 1987. These guides provided assistance to manufacturers and distributors of byproduct sealed sources and devices on how to file an application, and what information and documentation were required in their submittal. However, the guidance did not provide details on what the QC program must include. Therefore, the team could find no applicable NRC requirements or guidelines against which details in a QC program could be measured.

The Food and Drug Administration (FDA) has authority over the vendors of medical devices such as the Omnitron 2000 afterloader. The source wires were also licensed as sealed sources by the State of Louisiana. The scope of the IIT did not include an evaluation of the adequacy of FDA or State regulatory activity related to the Omnitron 2000 afterloader. The regulatory interaction between the NRC, FDA, and the involved Agreement States in the regulation and authorization of the Omnitron 2000 HDR afterloader is poorly defined.

10.6 No Regulatory Guidance Exists for Nonradioactive Waste Collectors. In Addition, Browning-Ferris Industries Failed To Follow Their Existing Radiation Control Policies.

No regulatory guidance exists for waste collectors, brokers, and recyclers similar to that developed by the steel industry. Browning-Ferris Industries (BFI) had an aggressive radiation monitoring program to ensure radioactive material was not disposed of inappropriately. Without this program, the source might never have been detected by the BFI-Warren facility, and further significant radiation exposures could have occurred. The radioactivity that BFI workers receive from medical facilities they serve is very low. In this incident, BFI workers encountered a significant radiological hazard for which they were not prepared and for which they had no regulatory guidance that described how to dispose of highly radioactive material.

The failure of the BFI-Carnegie waste pick-up Driver A to survey the waste at the nursing home on November 25, 1992, in accordance with BFI policy, resulted in the exposure of the driver and other BFI employees to significant levels of radiation. If BFI Driver A had surveyed the waste at the nursing home, exposures to all other BFI employees would have been avoided.

The return of the radioactive waste shipment to BFI-Carnegie, coupled with the search for the source of radiation and attempts to trace the material's origin by BFI employees who were neither trained nor equipped to handle a source of this magnitude, should have been avoided. The highest exposed BFI workers were among those who searched the trailer for the source on November 30, 1992. Once BFI-Warren determined that they had a highly radioactive shipment, further shipment violated DOT regulations. In addition, when BFI-Warren located the trailer containing the iridium-192 source, measures should have been taken to isolate the trailer and obtain professional radiological safety help to minimize radiation exposure to others.

GLOSSARY

GLOSSARY

The description of the terms in this glossary do not provide definitions or legal interpretation of these terms. The description of the terms is intended for use in this report.

Absorbed dose means the amount of ionizing radiation absorbed by an object or individual. The special name gray, symbol Gy, has been adopted for the System International (SI) unit of absorbed dose.

Activity means a measure of strength of a radioactive source, measured in units of becquerel (Bq) [curies (Ci)].

Attenuation means the reduction of radiation intensity as it passes through any material, for example, lead shielding.

Autoradiograph means the imaging of radioactive material on x-ray film by exposing the film to the radioactive material. Darkened areas on the film would indicate the presence and distribution of radioactive material.

Background radiation means radiation emitted from naturally occurring radioactive materials in the earth or from cosmic rays. The estimated total effective dose equivalent rate for a member of the public in the United States from various sources of natural background radiation is 3.0 mSv 300 (mrem) per year.

Byproduct material means radioactive material obtained as a byproduct from nuclear reactors. Iridium-192 is byproduct material.

Becquerel means a unit of activity equal to one disintegration per second. (Abbreviation Bq).

Beta particle means a charged particle emitted from the nucleus of an atom with a mass and charge equal in magnitude to that of the electron.

Calibration means a determination of variation from a standard, or accuracy of a measuring instrument to ascertain necessary correction factors.

Chemical analysis means the chemical analysis of bulk samples of a metal part by one or more of several wet chemical techniques to establish the bulk chemical composition of the material for comparison with applicable specifications.

Curie means a unit of activity equal to 3.7×10^{10} disintegrations per second (Abbreviated Ci).

Collective dose means the sum of the individual doses received in a given period of time by a specific population from exposure to a specified source of radiation.

Counting efficiency means a measure of the probability that a count will be recorded when radiation is incident on a detector.

Cytogenetic evaluation means an evaluation of blood cells to determine chromosomal aberrations induced by radiation exposure.

Differential Scanning Calorimetry (DSC) means a thermal analysis technique employed to identify phase transformations in materials as a function of temperature. Small specimens of the material to be analyzed are heated at a constant, predetermined rate in an inert gas environment. Instrumentation records the heat input required to maintain the predetermined heating rate and phase changes (endothermic or exothermic) that result in spikes in a heat input versus temperature plot.

Dose equivalent means the product of D, Q, and N at the point of interest where D is the absorbed dose, Q is the quality factor, and N is the product of all other modifying factors,

$$H = DQN$$

The special name sievert, symbol Sv, has been adopted for the SI unit of dose equivalent in the field of radiation protection. The older conventional unit of dose equivalent is rem ($1 \text{ Sv} = 100 \text{ rem}$). An absorbed dose of 1 Gy (100 rad) from gamma rays results in approximately a dose equivalent of 1 Sv (100 rem).

Dose rate means the absorbed dose delivered per unit time.

Dosimeter means an instrument to detect and measure accumulated radiation exposure (i.e., film badge).

Dwell time means the amount of time a brachytherapy source remains at a specific location within an implant catheter.

Electron capture means a mode of radioactive decay involving the capture of an orbital electron by its nucleus.

Electron volt means a unit of energy equivalent to the energy gained by an electron passing through a potential difference of one volt.

Endobronchial implant means the insertion of brachytherapy source(s) into the bronchial tubes in the patient's lung.

Energy dispersive x-ray spectroscopy (EDX or EDS) means a procedure that provides for identification of the elemental composition of a material. In this procedure, an exciting radiation (electron beam or x-rays) is directed against the specimen in question. Characteristic x-rays are then emitted for each element present and these emitted x-rays are analyzed to identify the individual elements qualitatively or quantitatively.

Exposure means being exposed by ionizing radiation or to radioactive material.

Fractography means the process of examining the fracture surfaces of a failed component or test specimen in detail. Fractographic examinations are carried out at all levels of magnification from the visual level to high magnification (up to 10,000X) in the scanning electron microscope (SEM).

Fracture surface features can identify fracture initiation sites, directions of fracture, and certain mechanical and chronological aspects of the fracture. Fine-scale fracture surface features serve to identify the particular cracking mechanism involved in the fracture.

Gamma ray means short-wave length electromagnetic radiation of nuclear origin emitted from the nucleus.

Geiger-Mueller Counter means a highly sensitive, gas-filled radiation measuring device.

Gray means the SI unit of absorbed dose. One gray is equal to an absorbed dose of one Joule per kilogram (100 rads) (Gy).

Hardness Measurements means measurements made by indenting a material with a standard indenter and a known load. Hardness values are then determined by the depth of penetration or by the projected size of indentation. Hardness is considered as a unique mechanical property but it also provides for relative evaluation of other mechanical properties such as tensile strength.

Hot cell means a shielded box or enclosure for sorting, processing, manufacturing or testing radioactive materials that must be handled remotely.

Interstitial means the insertion of brachytherapy source(s) into spaces within a tissue.

Intracavitary implant means the insertion of brachytherapy source(s) into a body cavity.

Ionization chamber means an instrument designed to measure a quantity of ionizing radiation in terms of the charge of electricity associated with ions produced within a defined volume.

Leak test means a test on sealed sources to ensure that radioactive material is not being released.

Metallography means metallographic examinations that are performed by monitoring a sectioned specimen in a suitable plastic and grinding and polishing the section using successively finer grits (down to 0.05 microns). This specimen is then etched with a suitable reagent to enhance microstructural features. Examinations of the as-polished or polished and etched section in an optical microscope at magnifications up to 1000X provides for evaluation of the microstructure of the material and identification of microstructural defects and abnormalities.

Micro - or milli means those abbreviations commonly used in radiation protection. Several fractions of radiological units are --

millisievert: One-thousandth of a sievert (abbreviated msv)

microsievert: One-millionth of a sievert (abbreviated μSv)

Misadministration, as it applies to brachytherapy, means the administration of a brachytherapy dose that involves the wrong patient, the wrong radioisotope, the wrong treatment site, a leaking source, an administered dose differing by more than 20 percent from the prescribed dose; or when a temporary source is not removed upon completion of the procedure.

Occupational dose means the dose received by an individual in a restricted area or in the course of employment in which the individual's assigned duties involve exposure to radiation and to radioactive material. Occupational dose not involve dose received from background radiation, as a patient from medical practices, from voluntary participation in medical research programs, or as a member of the general public.

Public dose means the dose received by a member of the public from exposure to radiation and to radioactive material released by a licensee, or to another source of radiation either within a licensee's controlled area or in unrestricted areas. It does not include occupational dose or dose received from background radiation, as a patient from medical practices, or from voluntary participation in medical research programs.

Quality assurance (QA) means the planned and systematic actions necessary to provide confidence that a firm or product will perform to establish specifications.

Quality control (QC) means the QC activities, including inspection and testing whose purpose is to provide a means to measure the characteristics of a firm or product to the required specifications.

Quality factor means the modifying factor that is used to derive dose equivalent from absorbed dose. The quality factor for gamma radiation emitted by iridium-192 is one.

Quality management (QM) program means licensee response to the requirements set forth in 10 CFR Part 35.32 that requires medical users of byproduct material to establish a QM program.

Radiation Safety Officer means the individual or the Radiation Safety Officer identified on an NRC licensee responsible for implementing the radiation safety program. The RSO ensures that radiation safety activities are being performed in accordance with approved procedures and regulatory requirements in the licensee's daily operations.

Radioactive decay means the disintegration of the nucleus of an unstable nuclide by spontaneous emission of charged particles and/or photons.

Radioactive half-life means the time required for a radioactive substance to lose 50 percent of its activity by decay.

Radioactive source capsule or source means the small sealed metal capsule containing the radioactive material that emit the gamma rays used in remote afterloading devices.

Red-bag waste means the common term used to describe medical biohazard waste.

Remote afterloader means a radiation therapy device used to insert and retract a radioactive source (or sources) from a patient via remote-controlled mechanism.

Safety culture means the beliefs, perceptions, and expectations that individuals have about the organization in which they work and about the values and consequences that will follow from one course of action or another. Consequently, culture highly influences behavior within the organization.

Sealed source means a radioactive source sealed in an impervious container that has sufficient mechanical strength to prevent contact with and dispersion of the radioactive material under the condition of use and wear for which it was designed.

Sievert means the SI unit of any of the quantities expressed as dose equivalent. The dose equivalent in sieverts is equal to the absorbed dose in grays multiplied by the quality factor ($1 \text{ Sv} = 100 \text{ rems}$).

Time and motion study means an evaluation of the proximity and duration that an individual was near a source of radiation for the purposes of estimating radiation exposure.

Wipe test means an evaluation of removable contamination on a surface or object, wherein an absorbent material such as paper is rubbed across a surface and subsequently analyzed for radioactivity in a counting instrument.

Work traveler means a document that accompanies and records the history of the source wire. Typical work travelers list the steps and operations performed on the wire, the status (pass/fail), who performed inspections on the wire, and the date operations or inspections were performed.

Worst-case scenario means a situation for which it is assumed that no designed or incidental shielding absorbed radiation from the source emitting it.

APPENDIX A

INCIDENT INVESTIGATION TEAM CHARTER



UNITED STATES
NUCLEAR REGULATORY COMMISSION
WASHINGTON, D. C. 20555

December 4, 1992

MEMORANDUM FOR: The Chairman
Commissioner Rogers
Commissioner Curtiss
Commissioner Remick
Commissioner de Planque

FROM: James M. Taylor
Executive Director for Operations

SUBJECT: INVESTIGATION OF A LOSS OF AN IRIDIUM-192 SOURCE AND THERAPY
MISADMINISTRATION AT ONCOLOGY SERVICES CORPORATION, INDIANA
REGIONAL CANCER CENTER, INDIANA, PENNSYLVANIA

On December 1, 1992 at approximately 12:30 P.M. (EST), a medical physics consultant (physicist) with Oncology Services Corporation (OSC) informed Region I that a 3.7 curie, iridium-192 source was missing from the licensee's facility in Indiana, Pennsylvania where it was utilized in a high dose rate remote afterloading (HDR) brachytherapy device. The missing item was found by Browning Ferris Industries (BFI). BFI is a waste removal company. The physicist stated that BFI had contacted OSC to determine whether waste picked up from the nursing home, in the vicinity of OSC's Indiana, Pennsylvania site, may have originated from the OSC facility. Waste picked up in the area set off a radiation monitor alarm at BFI's facility in Warren, Ohio, which initiated the source recovery and followup actions. In this case the source was "found" before it was reported to be "missing."

The licensee's authorized physician user indicated that the HDR brachytherapy device was last used on November 16, 1992, to treat an 82 year old female patient from the Scenery Hill Nursing Home located near OSC's Indiana, Pennsylvania site. Region I was notified on December 1, 1992 that during a brachytherapy treatment of the nursing home patient, the radioactive source had somehow remained in the patient. The patient was subsequently returned to her nursing home where she died a few days later. Therefore, the potential existed for overexposure to the patient, as well as nursing home staff, other individuals and those involved in the inadvertent discovery and retrieval of the source (the dose rate at one meter is of the order of 2R/hour). There are potential generic implications regarding the design and operation of similar devices. Calls by the vendor to other licensees who possess the device, notifying them of the incident, are almost completed. Our current information is that there are 21 such devices in use.

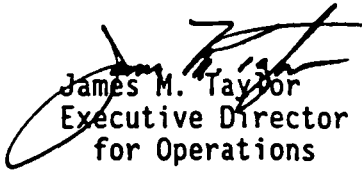
Region I and NMSS dispatched teams to review the licensee's actions and to determine any generic issues. The NRC staff notified the Food and Drug Administration's (FDA's) Center for Devices of the incident. The FDA has primary Federal responsibility for medical devices and indicated its interest to participate in the investigation of the incident. Because of the potential for overexposure for multiple members of the public as well as clinic and

nursing home staff, the complexity of the event, the licensing of the therapy device vendor in an agreement state and the involvement of the FDA in the licensing process, I have requested AEOD to take the necessary actions to upgrade the current investigation activities to an NRC Incident Investigation Team (IIT). The IIT is to: (a) fact-find as to what happened, (b) identify the probable cause as to why it happened, and (c) make appropriate findings and conclusions which would form the basis for any necessary follow-on actions. The IIT charter is enclosed.

The team will report directly to me and is comprised of: Carl Paperiello, (Region III) IIT Leader; Mohamed Shanbaky (Region I), Assistant IIT Leader; Cynthia Jones (NMSS), Thomas Rich (NMSS) and other NRC staff which will be determined later. Contractor support will include Daniel Flynn, MD and additional technical expertise. Because of the limited number of technical experts able to investigate an event of this type, some team members have had previous inspection involvement at the facility. The IIT was selected on the bases of their knowledge and experience in the fields of medical physics, health physics, radiation oncology, human factors, and mechanical engineering. All team members are relieved of all normal duties while assigned to the IIT.

The licensee has agreed to preserve the equipment in an "as-found" state until the licensee and the team has had an opportunity to evaluate the event. The licensee's actions have been confirmed by the Region in a Confirmatory Action Letter which was issued on December 2, 1992.

The IIT report will constitute the single NRC fact-finding investigation report. It is expected that the IIT report will be issued within about 45 days from the time the team exits from the site.


James M. Taylor
Executive Director
for Operations

Enclosure:
Incident Investigation Team
Charter

cc w/encl:
SECY
OGC
ACRS
OPA
OSP
OCA
Regional Administrators

Incident Investigation Team Charter

**LOSS OF AN IRIIDIUM-192 SOURCE AND THERAPY MISADMINISTRATION
AT ONCOLOGY SERVICES CORPORATION,
INDIANA REGIONAL CANCER CENTER, INDIANA, PA**

The scope of the investigation should include: incident chronology; source characterization; analysis of actual and potential dose consequences; human factors aspects; design and experience in operation of the equipment; event reporting and licensee response; and whether the NRC's regulatory process and activities, or the interface with state and other federal jurisdictions preceding the event, contributed to it. Within the framework of this overall scope the IIT should specifically:

With respect to the incident chronology: develop a detailed sequence of events associated with the therapy treatment, transport of the cancer patient, and transport and location of the source from the time it was used at the treatment center until its retrieval from the waste truck.

With respect to the source characterization: determine the manufacture type, radiation types and levels.

With respect to analysis of the actual and potential dose consequences: evaluate the dose received by the cancer patient as a result of the misadministration and the potential consequences; and reconstruct the projected exposure to individuals who were in proximity to the source either during the time it was in the cancer patient, or during storage, waste transport or retrieval.

With respect to the human factors aspects: evaluate training, surveys and supervision associated with the medical personnel's detection and response to the inadvertent retention of the source in the cancer patient.

With respect to the equipment: evaluate (1) the failure mechanism, (2) root causes of the failure, (3) its operating procedures, and (4) generic applicability of the failure or operation to other equipment.

With respect to event reporting and licensee response: evaluate the actions taken by the licensee to report to the NRC the loss of the source and potential overexposure(s) of individual(s); and the actions taken by the licensee to recover the source.

With respect to the NRC's regulatory process and activities: evaluate the regulatory controls concerning this type of event. Also, examine the interface with the involved Agreement States and other Federal jurisdictions.

APPENDIX B
PROPERTIES OF IRIIDIUM-192

Iridium-192*

Atomic number: 77
Atomic weight: 192
Half life: 74.02 days

Betas

	<u>Probability</u> <u>per decay</u>	<u>Maximum</u> <u>(MeV)</u>	<u>Average</u> <u>(MeV)</u>
1.	.056500	.255950	.070800
2.	.414000	.536080	.161200
3.	.483000	.672420	.208900
4.	.004000	.844530	.275900
5.	.000987	.093000	.024933

Electrons

<u>Probability</u> <u>per decay</u>	<u>Energy</u> <u>(MeV)</u>
1. .031078	.006880
2. .001594	.048300
3. .005227	.131920
4. .003649	.192830
5. .001180	.202750
6. .076020	.007240
7. .003505	.051000
8. .001192	.067951
9. .001300	.122470
10. .019237	.217560
11. .017896	.230060
12. .044658	.238110
13. .008792	.282080
14. .002867	.292660
15. .007716	.294580
16. .019470	.302630
17. .002511	.305160
18. .004839	.313210
19. .001483	.315790
20. .010236	.389680
21. .002946	.454190
22. .001509	.526020

Gammas & x-rays

<u>Probability</u> <u>per decay</u>	<u>Energy</u> <u>(MeV)</u>
1. .014625	.008910
2. .011323	.061487
3. .019555	.063001
4. .008399	.071400
5. .004674	.201310
6. .032873	.205800
7. .002615	.283260
8. .007264	.374480
9. .031628	.484580
10. .003985	.489060
11. .000797	.423070
12. .040934	.009440
13. .026350	.065022
14. .045197	.066832
15. .019675	.075700
16. .001806	.136350
17. .290150	.295960
18. .296780	.308460
19. .828530	.316510
20. .006645	.416460
21. .480550	.468070
22. .045735	.588580
23. .082024	.604410
24. .053357	.612460
25. .003016	.884510
26. .000986	.871730

* From Microshield 3.13

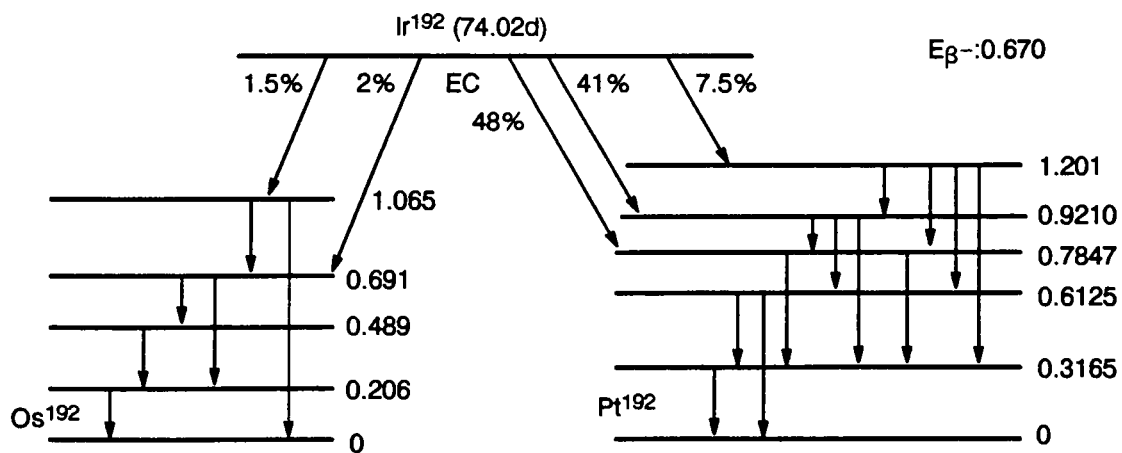
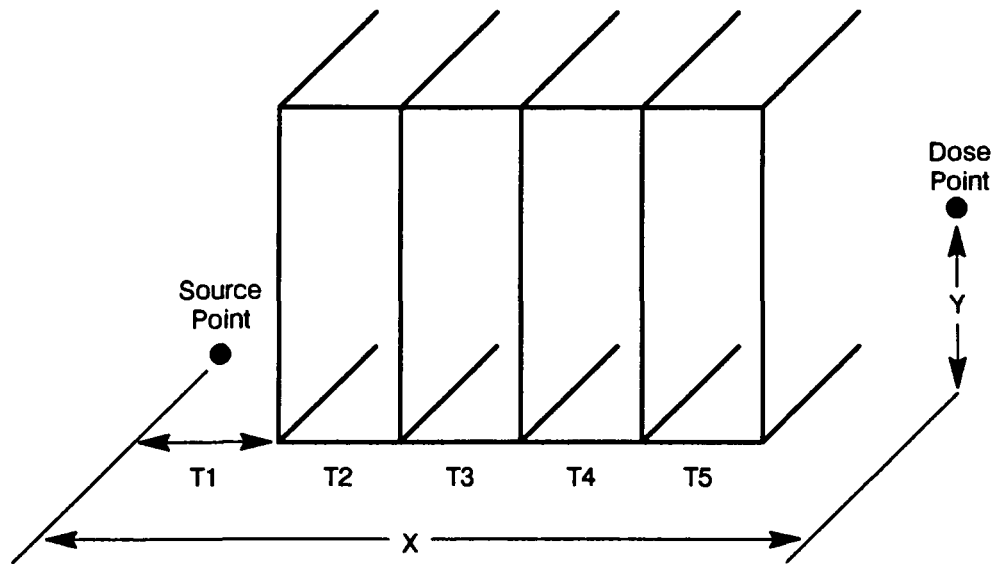


Figure B.1 Simplified Decay Scheme for Iridium-192



- X X is the horizontal distance from the source to the point of interest.
- Y Y is the vertical distance from the source to the point of interest.
- T Six Ts represent shield thicknesses. As many as five shields may be entered by the user. The sixth is reserved for a Microshield assigned air gap.

Figure B.2 Microshield 3.13 Program Showing Point Source and Shields

**Table B.1 Calculated Dose Rates (mSv/hr)* at Selected Distances
for 3.3085 E+11 Bq (8.942 Ci) Iridium-192 Source
Certified August 28, 1992**

Date: Elapsed time (days): Activity (Bq): Activity [Ci]:	11/16/92 79 1.579 E+11 4.267	11/20/92 83 1.521 E+11 4.110	11/25/92 88 1.451 E+11 3.922	11/27/92 90 1.424 E+11 3.850	12/01/92 94 1.372 E+11 3.708
Distance: 0.02 (m) 0.25 0.50 0.75 1.00 1.50 2.00 2.50 3.00 3.50 4.00 4.50 5.00 7.50 10.00 15.00	5.146E+4 3.310E+2 8.317E+1 3.715E+1 2.100E+1 9.427E+0 5.354E+0 3.459E+0 2.424E+0 1.797E+0 1.388E+0 1.106E+0 9.036E-1 4.176E-1 2.432E-1 1.147E-1	4.957E+4 3.188E+2 8.011E+1 3.579E+1 2.023E+1 9.080E+0 5.157E+0 3.332E+0 2.335E+0 1.731E+0 1.337E+0 1.065E+0 8.703E-1 4.022E-1 2.343E-1 1.105E-1	4.730E+4 3.042E+2 7.644E+1 3.415E+1 1.930E+1 8.665E+0 4.921E+0 3.179E+0 2.228E+0 1.652E+0 1.276E+0 1.017E+0 8.305E-1 3.838E-1 2.235E-1 1.054E-1	4.642E+4 2.977E+2 7.502E+1 3.351E+1 1.894E+1 8.504E+0 4.830E+0 3.120E+0 2.187E+0 1.621E+0 1.252E+0 9.979E-1 8.152E-1 3.767E-1 2.194E-1 1.035E-1	4.472E+4 2.876E+2 7.227E+1 3.228E+1 1.825E+1 8.192E+0 4.652E+0 3.006E+0 2.106E+0 1.562E+0 1.206E+0 9.612E-1 7.852E-1 3.629E-1 2.114E-1 9.966E-2

* Dose rate from an unshielded iridium-192 source using Microshield 3.13 with point-source geometry and Berger build-up factor.
1 mSv/hr = 100 mrem/hour

Build-Up Factor

The build-up factor (B) is defined as the ratio of the actual dose rate (dose rate from primary plus scattered and secondary radiation) to the dose rate that is due to the primary radiation alone. The value of B is a function of radiation type and energy, attenuating medium and depth, and geometry (see Reference 1).

From the Microshield 3.13 computer code used in this report to calculate the dose rate to a point of interest, there are four methods available for buildup determination: no buildup, Taylor, geometric progression, and Berger (see Reference 2). The comparison of calculating dose rate with the Berger build-up factor versus no build-up is presented in Table B.2. If a substantial amount of shielding is present, buildup may contribute significantly to the calculated dose rate.

For this report, the Berger build-up factor was used since the Taylor build-up factor is not useful below a gamma ray energy of 2 MeV (see Reference 3). Also, Berger calculations are valid to about 10 mean free paths (10 times $1/\mu$), which is approximately 730 m (2395 feet) of air, 0.9 m (2.95 feet) of water, or 0.4 m (1.31 feet) of concrete (see Reference 4).

**Table B.2 Iridium-192 Dose Rates vs. Distance
With and Without Berger Buildup Factors**

Distance (meters)	Dose Rate in Air		
	With Berger buildup	With no buildup	<u>Dose Rate-Berger</u> <u>Dose Rate-None</u>
	mSv/hr*	mSv/hr*	
0.02	5.146E+4	5.143E+4	1.006
0.25	3.310E+2	3.283E+2	1.008
0.50	8.317E+1	8.182E+1	1.016
0.75	3.715E+1	3.626E+1	1.025
1.00	2.100E+1	2.033E+1	1.033
1.50	9.427E+0	8.984E+0	1.049
2.00	5.354E+0	5.023E+0	1.066
2.50	3.459E+0	3.196E+0	1.082
3.00	2.424E+0	2.206E+0	1.099
3.50	1.797E+0	1.611E+0	1.115
4.00	1.388E+0	1.226E+0	1.132
4.50	1.106E+0	9.632E-1	1.148
5.00	9.036E-1	7.756E-1	1.165
7.50	4.176E-1	3.346E-1	1.248
10.00	2.432E-1	1.827E-1	1.331
15.00	1.147E-1	7.652E-2	1.499

* Dose rate from an unshielded iridium-192 source containing 1.579 E+11 Bq [4.267-Ci] on November 16, 1992, using Microshield 3.13 with point-source geometry.
1 mSv/hr = 100 mrem/hour

Build-up Contribution to Exposure Rate Calculations

As discussed in Section 7 of this report, the team concluded that (1) build-up factors were necessary to include in all exposure rate calculations and (2) there was very little attenuation owing to patient shielding. The following calculations illustrate these conclusions.

Calculation

Material: Muscle

Density (ρ): 1040 kilograms per cubic meter = 1.04 grams per cubic centimeter (g/cm³)
(Reference 5)

Mass Attenuation Coefficient (μ_m) [from interpolation] = 0.1151 square centimeters per gram (cm²/g) for 320 keV gamma ray (Reference 5)

(Note: a 320 keV gamma ray was used since this is the most abundant Ir-192 gamma ray).

$$\text{Linear Attenuation Coefficient}(\mu_l) = \mu_m \times \rho = 0.1151 \text{ cm}^2/\text{g} \times 1.04 \text{ g/cm}^3 \\ = 0.1197 \text{ cm}^{-1}$$

$$\text{Mean free path (mfp)} = 1/\mu_l = 1/0.1197 \text{ cm}^{-1} = 8.354 \text{ cm}$$

$$\frac{X}{X_0} = B \exp(-\mu_l d), \text{ where}$$

X = Final exposure rate

X₀ = Initial exposure rate

B = Build-up factor

d = Distance from the source (cm)

μ_l = Linear attenuation coefficient (cm⁻¹)

a) If d = 10 cm:

$$\text{number of mfp's} = 10 \text{ cm} / 8.354 \text{ cm} = 1.20$$

$$B \text{ (for 1.20 mfp's and a gamma ray energy of 300 keV)} = 3.35 \text{ (Reference 6)}$$

Using the Equation above,

$$\frac{X}{X_0} = 3.35 \exp(-0.1197 \text{ cm}^{-1} \times 10 \text{ cm}) = 1 \text{ (i.e., no attenuation)}$$

b) If $d = 1$ cm:

number of mfp's = $1 \text{ cm} / 8.354 \text{ cm} = 0.12$

B (for 0.10 mfp's and a gamma ray energy of 300 keV) = 1.12 (Reference 6)

$$\frac{X}{X_0} = 1.12 \exp(-0.1197 \text{ cm}^{-1} \times 1 \text{ cm}) = 1 \text{ (i.e., no attenuation)}$$

Sample Hand Calculation (in SI units)

For correcting for radioactive decay of a source,

$$A_t = A_o B e^{-\ln 2 * t / T_{1/2}}$$

- where
- A_t = Corrected source activity (Bq) at time t
 - A_o = Source activity on August 28, 1992 ($3.31 \text{ E}+11 \text{ Bq}$)
 - B = Dose Build-up factor
 - t = Elapsed time in days since August 28, 1992
 - $T_{1/2}$ = Half-life of iridium-192 (74.02 days)

For calculating the exposure rate from a point source at a given distance,

$$\Gamma = 3.65 \times 10^{-9} \sum f_x E_x \frac{(C/kg) m^2}{MBq-hr}$$

- where
- Γ = Exposure rate in coulombs per kilogram (C/kg) per hour at 1 meter from an unshielded iridium-192 source of Activity, A_t
 - f_x = Probability of a gamma ray being emitted
 - E_x = Gamma-ray energy in MeV

For iridium-192:

$$\Gamma = 3.5 \times 10^{-9} \frac{(C/kg) \cdot m^2}{MBq \cdot hr}$$

For calculating the exposure rate from iridium-192 for a specific time period,

$$E_d = \frac{\Gamma T}{d^2}$$

where E_d = Exposure in sievert (Sv) at distance, d
d = Distance in meters from the unshielded iridium-192 source
T = Exposure time in hours
 Γ = Gamma constant

Sample Hand Calculation (in English Units)

For correcting for radioactive decay of a source,

$$A_t = A_o B e^{1n2 \cdot t / T_{1/2}}$$

where A_t = Corrected source activity (Ci) at time t
 A_o = Source activity on August 28, 1992 (8.942 Ci)
B = Dose build-up factor
t = Elapsed time in days since August 28, 1992
 $T_{1/2}$ = Half-life of iridium-192 (74.02 days)

For calculating the exposure rate from a point source at a given distance,

$$\Gamma = 0.5 \sum f_x E_x \frac{R-m^2}{Ci-hr}$$

where Γ = Exposure rate in R per hour at 1 meter from an unshielded iridium-192 source of Activity, A_i

f_x = Probability of a gamma ray being emitted

E_x = Gamma-ray energy in MeV

For iridium-192:

$$\Gamma = 0.48 \frac{R-m^2}{Ci-hr}$$

For calculating the exposure rate from iridium-192 for a specific time period,

$$E_d = \frac{\Gamma T}{d^2}$$

where E_d = Exposure in rem at distance, d

d = Distance in meters from the unshielded iridium-192 source

T = Exposure time in hours

Γ = Gamma constant (0.48 R-m²/Ci-hr)

References

1. Attix, Frank Herbert, "Introduction to Radiological Physics and Radiation Dosimetry", John Wiley & Sons, Inc., 1986.
2. Microshield 3 Manual, Grove Engineering, Inc., Rockville, Maryland, 1987.
3. Taylor, J.J. and F.E. Obershain, USAEC Report WAPD-RM-213, Westinghouse Electric Corp., 1953.
4. Engineering Compendium on Radiation Shielding, Vol. 1, Shielding Fundamentals and Methods, Springer-Verlag, New York, 1968.
5. H.E. Johns and J.R. Cunningham, "The Physics of Radiology," Fourth Edition, Charles C. Thomas, Springfield, Illinois, 1983.
6. M.J. Berger, "Energy Deposition in Water by Photons from Point Isotropic Sources", MIRD/Pamphlet No. 2, J. Nucl. Medicine, Supplement No. 1, Feb. 1968.

APPENDIX C

LOW DOSE RATE AND HIGH DOSE RATE

BRACHYTHERAPY

Appendix C

Low Dose Rate and High Dose Rate Brachytherapy

Brachytherapy is a radiation therapy procedure for the treatment of cancer in which radioactive sources are placed near or in contact with a tumor. The prefix "brachy" means short as opposed to "tele" which means far. Thus, brachytherapy involves the use of a radioactive source at a short distance from the tumor while teletherapy involves the use of a radiation source at a long distance, usually greater than 80 centimeters (cm), from the tumor.

Each treatment method has its own advantages and safety problems. Teletherapy requires the shielding and protection of large sources. Typical sources are $3.7 \text{ E}+14 \text{ Bq}$ (10,000 Ci) of cobalt-60. The output of these devices requires precise measurement. The dose at the treatment site is generally uniform over the volume of the tumor. The rooms in which they are used are heavily shielded with lead or concrete. Treatment times are in the order of minutes. Sources are relatively large sealed sources in large lead or depleted uranium cavities with heavy shields to block the radiation beam when the device is not being used. Problems encountered with these types of devices generally involve errors in output measurement or calculation, errors in treatment time, failure to shield the source at the end of the treatment, and exposure of the wrong part of the body.

One of the original uses of radium was brachytherapy. Today radium is rarely used. Principal sources in use today are cesium-137, iridium-192, cobalt-60, iodine-125, strontium-90, and palladium-103. Original treatment methods with radium did not depend on dose calculations. Instead, they were based on milligrams of radium at certain locations for specific periods of time. Treatment modes were based on physicians' clinical experience. Even today, certain brachytherapy sources are specified in milligram radium equivalents.

Because exposure rates vary with the inverse square of distance and because brachytherapy sources are either very close to or at the treatment site, the dose delivered is relatively nonuniform. When technology advanced, dose could be defined and measured, and systems were developed based on certain source distributions with a dose being specified at certain arbitrary points. Doses were derived from previous clinical practices that proved successful.

Brachytherapy sources are applied with intraluminal, interstitial, intracavitary, or intraoperative techniques. Usually catheters or applicators are placed in the patient to hold the radioactive sources before the sources are placed in the patient. Dummy sources placed in the catheters or applicators are located by x-rays to enable the physician to more accurately place the active sources and more accurately calculate dose. After the physician or technicians identify the proper location for the sources, the dummy sources are replaced by radioactive sources and the treatment begins. Low dose rate brachytherapy generally employs sources similar in strength to the original radium sources used. These sources are generally in the order of $3.7 \text{ E}+09 \text{ Bq}$ (0.1 Ci). Sources are frequently loaded manually,

using forceps. If the physician is skillful and does not handle the sources by hand, the physician's exposure is usually within regulatory limits.

Typically, the sources remain in the patient for several days. The patient's room is isolated and nurses and other support staff are instructed to limit the amount of time spent in the room and, therefore, their exposure. Visitors are likewise limited. At the end of the treatment, the sources are removed, and surveying the patient with a radiation monitor at this point is critical for ensuring that no sources remain in the patient. It is also critical when returning sources to storage after their use to count them and ensure that none are lost. Brachytherapy sources are small and easy to lose. On occasion, a source will fall out of a patient during treatment. Sometimes semiconscious or agitated patients will remove sources inadvertently. Most low dose brachytherapy problems have resulted from sources being left in patients or being lost. NRC regulations require both the patient survey and the source count.

Remote afterloaders are devices for loading active sources into catheters or applicators by machine rather than manually. This protects the physician and other medical personnel from exposure to the source. The sources are stored in an internal lead shield and remotely removed from the shield and guided into the patient through catheters. For low dose rate remote brachytherapy, sources can be retracted when the nursing staff and other medical personnel provide bedside care for the patient. This reduces their exposure and this has significant As-Low-As-Is-Reasonably-Achievable (ALARA) value.

High dose rate brachytherapy combines remote afterloader technology with much higher source strengths in the order of $3.7 \text{ E}+11 \text{ Bq}$ (10 Ci) or about two orders of magnitude greater than low dose rate brachytherapy. The technique is relatively new and it has two principal advantages. First, patients can avoid long hospitalizations in isolation. Treatment times are in the order of minutes. Second, small-diameter high-activity sources allow the treatment of certain sites (e.g., esophagus, bronchus, bile duct, brain) that could not be previously treated by brachytherapy.

However, high dose rate brachytherapy has its own special problems, some of which are similar to teletherapy. The high activity of the source requires the treatment to be conducted in a special shielded room. In addition, the source is not routinely observable. Treatment planning and timing must be much more sophisticated and must strongly depend on computer calculational techniques. Because of the newness of this technique, physicians place more reliance on dose calculations than on previous clinical practice. Emergency response is more difficult because if the source has to be retrieved, the dose rates involved are much higher than those encountered in low dose rate brachytherapy. Nevertheless, high dose rate brachytherapy is a medically important treatment modality.

APPENDIX D
ONCOLOGY SERVICES CORPORATION
MATERIALS LICENSE

MATERIALS LICENSE

Amendment No. 03

Pursuant to the Atomic Energy Act of 1954, as amended, the Energy Reorganization Act of 1974 (Public Law 93-438), and Title 10, Code of Federal Regulations, Chapter I, Parts 30, 31, 32, 33, 34, 35, 39, 40 and 70, and in reliance on statements and representations heretofore made by the licensee, a license is hereby issued authorizing the licensee to receive, acquire, possess, and transfer byproduct, source, and special nuclear material designated below; to use such material for the purpose(s) and at the place(s) designated below; to deliver or transfer such material to persons authorized to receive it in accordance with the regulations of the applicable Part(s). This license shall be deemed to contain the conditions specified in Section 183 of the Atomic Energy Act of 1954, as amended, and is subject to all applicable rules, regulations and orders of the Nuclear Regulatory Commission now or hereafter in effect and to any conditions specified below.

Licensee		In accordance with letter dated June 10, 1992, 3. License number 37-28540-01 is amended in its entirety to read as follows:
1. Oncology Services Corporation		
2. 775 South Arlington Avenue Harrisburg, Pennsylvania 17109		4. Expiration date August 31, 1995
		5. Docket or Reference No. 030-31765
6. Byproduct, source, and/or special nuclear material	7. Chemical and/or physical form	8. Maximum amount that licensee may possess at any one time under this license
A. Iridium 192	A. Sealed sources (Byk Mallinckrodt Model No. GM 252.20-001 or RTS Technology Model No. 722)	A. Not to exceed 11 curies per source and 132 curies total
B. Iridium 192	B. Sealed sources (Byk Mallinckrodt Model No. GM 212.03-000 or RTS Technology Model No. 721)	B. Not to exceed 11 curies per source and 132 curies total
C. Iridium 192	C. Sealed sources (Omnitron International Model SL-777)	C. Not to exceed 11 curies per source and 132 curies total
D. Iridium 192	D. Sealed sources (Byk Mallinckrodt Model No. CI L BV)	D. Not to exceed 11 curies per source and 132 curies total
9. Authorized use		
A. One source per device for use in an Isotopen-Technik Dr. Sauerwein GmbH Gamma Med Iii remote after loading brachytherapy device for the treatment of humans. One source per device in its shipping container as necessary to the replacement of the source in the irradiation device only.		
B. One source per device for use in an Isotopen-Technik Dr. Sauerwein GmbH Gamma Med 12i remote after loading brachytherapy device for the treatment of humans. One source per device in its shipping container as necessary to the replacement of the source in the irradiation device only.		
C. One source per device for use in an Omnitron 2000 remote after loading brachytherapy device for the treatment of humans. One source per device in its shipping container as necessary to the replacement of the source in the irradiation device only.		
D. One source per device for use in a Nucletron MicroSelectron-HDR remote after loading brachytherapy device for the treatment of humans. One source per device in its shipping container as necessary to the replacement of the source in the irradiation device only.		

MATERIALS LICENSE
SUPPLEMENTARY SHEET

License number

37-28540-01

Docket or Reference number

030-31765

Amendment No. 03

CONDITIONS

10. Location of use: Harrisburg Cancer Center, 775 South Arlington Avenue, Harrisburg, Pennsylvania; Life Care Center, R. D. #1, Sandy Lake Road, Stoneboro, Pennsylvania; Greater Pittsburgh Cancer Center, 1145 Bower Hill Road Suite 105, Pittsburgh, Pennsylvania; Exton Cancer Center, 460 Creamery Way, Suite B, Exton Pennsylvania; Indiana Regional Cancer Center 877 Hospital Road, Indiana, Pennsylvania; Mahoning Valley Cancer Center, 800 Mahoning Street, Suite E, Lehighton, Pennsylvania.
11. Radiation Safety Officer: David E. Cunningham, Ph.D.
12. Authorized Users: Material and Use:
- | | |
|----------------------------|---|
| Abdurrahman Unal, M.D. | Iridium-192 in a brachytherapy remote after loader for the treatment of humans. |
| Gilbert Lawrence, M.D. | Iridium-192 in a brachytherapy remote after loader for the treatment of humans. |
| Norman Williams, M.D. | Iridium-192 in a brachytherapy remote after loader for the treatment of humans. |
| Richard M. Yelovich, M.D. | Iridium-192 in a brachytherapy remote after loader for the treatment of humans. |
| James E. Bauer, M.D. | Iridium-192 in a brachytherapy remote after loader for the treatment of humans. |
| David J. Moylan, III, M.D. | Iridium-192 in a brachytherapy remote after loader for the treatment of humans. |
| Bernard R. Rogers, M.D. | Iridium-192 in a brachytherapy remote after loader for the treatment of humans. |
| Roger P. Tokars, M.D. | Iridium-192 in a brachytherapy remote after loader for the treatment of humans. |
13. The following shall be performed only by persons specifically licensed by the Commission or an Agreement State to perform such services:
- Installation, relocation, or removal of high dose after loader units containing sources.
 - Any maintenance or repair operations on a high dose after loader unit involving work on any mechanism that could expose the source, reduce the shielding around the source, or compromise the safety of the unit and result in increased radiation levels.
 - David E. Cunningham, Ph.D., may perform those services listed in Condition 13.A. above, for which he has received specific training from a licensed manufacturer's representative.

APPENDIX E
INTERVIEWS AND MEETINGS
THE INCIDENT INVESTIGATION
TEAM CONDUCTED

Interviews and Meetings the Incident Investigation Team Conducted

Date	Time	Meeting/Interview
12/03/92	3:20 p.m.	Entrance Meeting
12/04/92	1:38 p.m.	Interview of RTR Staff Technologist, Oncology Services Corporation (OSC) Indiana Regional Cancer Center (IRCC)
12/04/92	1:12 p.m.	Group Interview of Physician A, IRCC; Physician D, Director of Brachytherapy, OSC; and RSO, Director of Physics and Bioengineering, OSC, Harrisburg, Pennsylvania
12/04/92	10:20 a.m.	Interview of President, Omnitron International, Inc. (Omnitron)
12/04/92	10:16 a.m.	Interview of RTT-A, Staff Therapist, IRCC
12/04/92	12:08 p.m.	Interview of Medical Physicist A, Consulting Physicist, IRCC
12/04/92		Incident Investigation Team Media Briefing with Dr. Carl Paperiello, NRC, and NRC Medical Consultant
12/04/92	8:20 a.m.	Interview of RTT-B, Registered Radiation Therapy Technician, IRCC
12/04/92	8:25 a.m.	Interview of Nurse A, R.N., IRCC
12/05/92	8:00 a.m.	Interview of Medical Physicist A, Consulting Physicist, was continued on December 5, 1992
12/05/92	8:30 a.m.	Interview of CNA-C, Certified Nurses Aide, Scenery Hill Manor Nursing Home (SHM)
12/05/92	9:34 a.m.	Interview of Physician A, IRCC

Interviews and Meetings the Incident Investigation Team Conducted (continued)

Date	Time	Meeting/Interview
12/05/92	9:55 a.m.	Interview of LPN-B Licensed Practical Nurse, SHM Nursing Home, December 5, 1992
12/05/92	1:00 p.m.	Interview of Administrator, SHM Nursing Home
12/07/92	11:50 a.m.	Interview of Browing-Ferris Industries (BFI) Driver B, Tractor-Trailer Driver, BFI, Carnegie, Pennsylvania
12/07/92	10:30 a.m.	Interview of Supervisor A, Medical Supervisor, BFI, Carnegie, Pennsylvania
12/07/92	10:54 a.m.	Interview of Safety Technician B Assistant District Safety Manager, and Safety Technition A, District Safety Manager, BFI, Carnegie, Pennsylvania
12/08/92	10:17 a.m.	Interview of Medical Physicist B, Senior Radiological Physicist, Oncology Services Corporation, Greater Pittsburgh Cancer Center (GPCC)
12/08/92	12:35 p.m.	Interview of Physician E Medical Director, GPCC and Jefferson Radiation Oncology Center
12/08/92	2:21 p.m.	Interview of CNA-E, Certified Nursing Assistant, SHM Nursing Home
12/08/92	3:05 p.m.	Interview of RTT-C, Radiation Therapist, GPCC
12/08/92	3:07 p.m.	Interview of Maintenance Man A, SHM Nursing Home
12/08/92	10:30 p.m.	Interview of General Manger, Brachytherapy, BFI, Warren, Ohio

**Interviews and Meetings the Incident
Investigation Team Conducted (continued)**

Date	Time	Meeting/Interview
12/09/92	9:30 a.m.	Interview of Physician D Director of Brachytherapy, OSC
12/09/92	11:45 a.m.	Interview of Physician C President and CEO, OSC, State College, Pennsylvania
12/09/92	2:37 p.m.	Interview of Physician B, Radiation Oncologist, Vice President and Medical Director, OSC, State College, Pennsylvania
12/09/92	4:15 p.m.	Interview of RN-D, Registered Nurse, SHM Nursing Home
12/16/92	2:00 p.m.	Group Interview of NRC Headquarters NMSS Staff: Richard E. Cunningham, Director, Division of Industrial & Medical Nuclear Safety; John E. Glenn, Chief, Medical Academic & Commercial Use Safety Branch; and Frederick Combs, Chief, Operations Branch
12/17/92	10:04 a.m.	Interview of RSO Director, Radiation Safety, OSC
12/17/92	1:00 p.m.	Interview of Service Manager, Omnitron International, Inc.
		Interview of Senior V.P., Radiation Science, Omnitron
12/17/92	2:35 p.m.	Interim Exit Interview, Harrisburg Hotel, Harrisburg, Pennsylvania
12/17/92	2:35 p.m.	Interview of V.P., Product Development, Omnitron
12/17/92	4:15 p.m.	Interview of President, Omnitron International, Inc. (Omnitron)

**Interviews and Meetings the Incident
Investigation Team Conducted (continued)**

Date	Time	Meeting/Interview
<hr/>		
12/22/92	10:05 a.m.	Interview (via speakerphone) of Sales Representative for Omnitron
12/31/92	9:45 a.m.	Interview (via speakerphone) of Patient A's best friend
01/05/93	10:10 a.m.	Interview of RSO Director of Physics and Bioengineering, OSC, Harrisburg, Pennsylvania