Injury to the skin and underlying tissues from acute exposure to a large external dose of radiation is referred to as cutaneous radiation injury (CRI). Acute radiation syndrome (ARS)\(^1\) will usually be accompanied by some skin damage; however, CRI can occur without symptoms of ARS. This is especially true with acute exposures to beta radiation or low-energy x-rays, because beta radiation and low-energy x-rays are less penetrating and less likely to damage internal organs than gamma radiation is. CRI can occur with radiation doses as low as 2 Gray (Gy) or 200 rads\(^2\) and the severity of CRI symptoms will increase with increasing doses. Most cases of CRI have occurred when people inadvertently came in contact with unsecured radiation sources from food irradiators, radiotherapy equipment, or well depth gauges. In addition, cases of CRI have occurred in people who were overexposed to x-radiation from fluoroscopy units.

Early signs and symptoms of CRI are itching, tingling, or a transient erythema or edema without a history of exposure to heat or caustic chemicals. Exposure to radiation can damage the basal cell layer of the skin and result in inflammation, erythema, and dry or moist desquamation. In addition, radiation damage to hair follicles can cause epilation. Transient and inconsistent erythema (associated with itching) can occur within a few hours of exposure and be followed by a latent, symptom-free phase lasting from a few days to several weeks. After the latent phase, intense reddening, blistering, and ulceration of the irradiated site are visible. Depending on the radiation dose, a third and even fourth wave of erythema are possible over the ensuing months or possibly years.

In most cases, healing occurs by regenerative means; however, large radiation doses to the skin can cause permanent hair loss, damaged sebaceous and sweat glands, atrophy, fibrosis, decreased or increased skin pigmentation, and ulceration or necrosis of the exposed tissue.

With CRI, it is important to keep the following things in mind:
- The visible skin effects depend on the magnitude of the dose as well as the depth of penetration of the radiation.
- Unlike the skin lesions caused by chemical or thermal damage, the lesions caused by radiation exposures do not appear for hours to days following exposure, and burns and other skin effects tend to appear in cycles.
- The key treatment issues with CRI are infection and pain management.\(^3\)

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2. Both the Gray (Gy) and the rad are units of absorbed dose and reflect the amount of energy deposited in a mass of tissue (1 Gy = 100 rads). In this document, the absorbed dose refers to that dose received by at least 10 cm\(^2\) of the basal cell layer of the skin. The referenced absorbed dose levels in this document are assumed to be from beta, gamma, or x-radiation. Neutron or proton radiation produces many of the health effects described herein at lower absorbed dose levels.
3. On occasion a patient might also be contaminated with radioactive material. To address patient decontamination, please go to the following Web site: http://www.orau.gov/reacts/emergency.htm.
Cutaneous Radiation Injury: Fact Sheet for Physicians
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Stages and Grades of CRI
CRI will progress over time in stages and can be categorized by grade, with characteristics of the stages varying by grade of injury, as shown in Table 1. Appendix A gives a detailed description of the various skin responses to radiation, and Appendix B provides color photographs of examples of some of these responses.

Prodromal stage (within hours of exposure)—This stage is characterized by early erythema (first wave of erythema), heat sensations, and itching that define the exposure area. The duration of this stage is from 1 to 2 days.

Latent stage (1–2 days postexposure)—No injury is evident. Depending on the body part, the larger the dose, the shorter this period will last. The skin of the face, chest, and neck will have a shorter latent stage than will the skin of the palms of the hands or the soles of the feet.

Manifest illness stage (days to weeks postexposure)—The basal layer is repopulated through proliferation of surviving clonogenic cells. This stage begins with main erythema (second wave), a sense of heat, and slight edema, which are often accompanied by increased pigmentation. The symptoms that follow vary from dry desquamation or ulceration to necrosis, depending on the severity of the CRI (see Table 1).

Third wave of erythema (10–16 weeks postexposure, especially after beta exposure)—The exposed person experiences late erythema, injury to blood vessels, edema, and increasing pain. A distinct bluish color of the skin can be observed. Epilation may subside, but new ulcers, dermal necrosis, and dermal atrophy (and thinning of the dermis layer) are possible.

Late effects (months to years postexposure; threshold dose ~10 Gy or 1000 rads)—Symptoms can vary from slight dermal atrophy (or thinning of dermis layer) to constant ulcer recurrence, dermal necrosis, and deformity. Possible effects include occlusion of small blood vessels with subsequent disturbances in the blood supply (telangiectasia); destruction of the lymphatic network; regional lymphostasis; and increasing invasive fibrosis, keratosis, vasculitis, and subcutaneous sclerosis of the connective tissue. Pigmentary changes and pain are often present. Skin cancer is possible in subsequent years.

Recovery (months to years)
### Table 1. Grades of cutaneous radiation injury

<table>
<thead>
<tr>
<th>Grade</th>
<th>Skin dose*</th>
<th>Prodromal stage</th>
<th>Latent stage</th>
<th>Manifest illness stage</th>
<th>Third wave of erythema †</th>
<th>Recovery</th>
<th>Late effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>&gt; 2 Gy (200 rads) ‡</td>
<td>1–2 days postexposure or not seen</td>
<td>no injury evident for 2–5 weeks postexposure</td>
<td>• 2–5 weeks postexposure, lasting 20–30 days: redness of skin, slight edema, possible increased pigmentation • 6–7 weeks postexposure, dry desquamation</td>
<td>not seen</td>
<td>complete healing expected 28–40 days after dry desquamation (3–6 months postexposure)</td>
<td>• possible slight skin atrophy • possible skin cancer decades after exposure</td>
</tr>
<tr>
<td>II</td>
<td>&gt; 15 Gy (1500 rads)</td>
<td>6–24 hours postexposure with immediate sensation of heat lasting 1–2 days</td>
<td>no injury evident for 1–3 weeks postexposure</td>
<td>• 1–3 weeks postexposure; redness of skin, sense of heat, edema, skin may turn brown • 5–6 weeks postexposure, edema of subcutaneous tissues and blisters with moist desquamation • possible epithelialization later</td>
<td>• 10–16 weeks postexposure, injury of blood vessels, edema, and increasing pain • epilation may subside, but new ulcers and necrotic changes are possible</td>
<td>healing depends on size of injury and the possibility of more cycles of erythema</td>
<td>• possible skin atrophy or ulcer recurrence • possible telangiectasia (up to 10 years postexposure) • possible skin cancer decades after exposure</td>
</tr>
<tr>
<td>Grade</td>
<td>Skin dose*</td>
<td>Prodromal stage</td>
<td>Latent stage</td>
<td>Manifest illness stage</td>
<td>Third wave of erythema†</td>
<td>Recovery</td>
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</tr>
<tr>
<td>III</td>
<td>&gt; 40 Gy (4000 rads)</td>
<td>4–24 hours postexposure, with immediate pain or tingling lasting 1–2 days</td>
<td>none or less than 2 weeks</td>
<td>• 1–2 weeks postexposure: redness of skin, blisters, sense of heat, slight edema, possible increased pigmentation • followed by erosions and ulceration as well as severe pain</td>
<td>• 10–16 weeks postexposure: injury of blood vessels, edema, new ulcers, and increasing pain • possible necrosis</td>
<td>can involve ulcers that are extremely difficult to treat and that can require months to years to heal fully</td>
<td>• possible skin atrophy, depigmentation, constant ulcer recurrence, or deformity • possible occlusion of small vessels with subsequent disturbances in the blood supply, destruction of the lymphatic network, regional lymphostasis, and increasing fibrosis and sclerosis of the connective tissue • possible telangiectasia • possible skin cancer decades after exposure</td>
</tr>
</tbody>
</table>
### Cutaneous Radiation Injury: Fact Sheet for Physicians
(continued from previous page)

<table>
<thead>
<tr>
<th>Grade</th>
<th>Skin dose*</th>
<th>Prodromal stage</th>
<th>Latent stage</th>
<th>Manifest illness stage</th>
<th>Third wave of erythema†</th>
<th>Recovery</th>
<th>Late effects</th>
</tr>
</thead>
</table>
| IV    | > 550 Gy (55,000 rads) | occurs minutes to hours postexposure, with immediate pain or tingling, accompanied by swelling | none | • 1–4 days postexposure accompanied by blisters  
• early ischemia (tissue turns white, then dark blue or black with substantial pain) in most severe cases  
• tissue becomes necrotic within 2 weeks following exposure, accompanied by substantial pain | does not occur due to necrosis of skin in the affected area | recovery possible following amputation of severely affected areas and possible skin grafts | • continued plastic surgery may be required over several years  
• possible skin cancer decades after exposure |

*Absorbed dose to at least 10 cm$^2$ of the basal cell layer of the skin  
†Especially with beta exposure  
‡The Gray (Gy) is a unit of absorbed dose and reflects an amount of energy deposited in a mass of tissue (1 Gy = 100 rads).  
§Skin of the face, chest, and neck will have a shorter latent phase than the skin of the palms of the hands and the skin of the feet.
Patient Management

Diagnosis

The signs and symptoms of CRI are as follows:
- Intensely painful burn-like skin injuries (including itching, tingling, erythema, or edema) without a history of exposure to heat or caustic chemicals
  Note: Erythema will not be seen for hours to days following exposure, and its appearance is cyclic.
- Epilation
- A tendency to bleed
- Possible signs and symptoms of ARS

As mentioned previously, local injuries to the skin from acute radiation exposure evolve slowly over time, and symptoms may not manifest for days to weeks after exposure. Consider CRI in the differential diagnosis if the patient presents with a skin lesion without a history of chemical or thermal burn, insect bite, or skin disease or allergy. If the patient gives a history of possible radiation exposure (such as from a radiography source, x-ray device, or accelerator) or a history of finding and handling an unknown metallic object, note the presence of any of the following: erythema, blistering, dry or wet desquamation, epilation, ulceration.

Regarding lesions associated with CRI be aware that,
- days to weeks may pass before lesions appear;
- unless patients are symptomatic, they will not require emergency care; and
- lesions can be debilitating and life threatening after several weeks.

Medical follow-up is essential, and victims should be cautioned to avoid trauma to the involved areas.

Initial Treatment

Localized injuries should be treated symptomatically as they occur, and radiation injury experts should be consulted for detailed information. Such information can be obtained from the Radiation Emergency Assistance Center/Training Site (REAC/TS) at www.orau.gov/reacts/ or (865) 576-1005.

As with ARS, if the patient also has other trauma, wounds should be closed, burns covered, fractures reduced, surgical stabilization performed, and definitive treatment given within the first 48 hours after injury. After 48 hours, surgical interventions should be delayed until hematopoietic recovery has occurred.

A baseline CBC and differential should be taken and repeated in 24 hours. Because cutaneous radiation injury is cyclic, areas of early erythema should be noted and recorded. These areas should also be sketched and photographed, if possible, ensuring that the date and time are recorded. The following should be initiated as indicated:
- Supportive care in a clean environment (a burn unit if one is available)
- Prevention and treatment of infections
- Use of the following:
  - Medications to reduce inflammation, inhibit proteolysis, relieve pain, stimulate regeneration, and improve circulation
  - Anticoagulant agents for widespread and deep injury
- Pain management
- Psychological support
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Recommendations for Treatment by Stage

The following recommendations for treatment by stage of the illness were obtained by summarizing recommendations from Ricks et al. (226) and Gusev et al. (231), but they do not represent official recommendations of CDC.

Prodromal Stage—Use antihistamines and topical antipruriginous preparations, which act against itch and also might prevent or attenuate initiation of the cycle that leads to the manifestation stage. Anti-inflammatory medications such as corticosteroids and topical creams, as well as slight sedatives, may prove useful.

Latent Stage—Continue anti-inflammatory medications and sedatives. At midstage, use proteolysis inhibitors, such as Gordox®.

Manifestation Stage—Use repeated swabs, antibiotic prophylaxis, and anti-inflammatory medications, such as Lioxasol®, to reduce bacterial, fungal, and viral infections
- Apply topical ointments containing corticosteroids along with locally acting antibiotics and vitamins.
- Stimulate regeneration of DNA by using Lioxasol® and later, when regeneration has started, biogenic drugs, such as Actovegin® and Solcoseril®.
- Stimulate blood supply in third or fourth week using Pentoxifylline® (contraindicated for patients with atherosclerotic heart disease).
- Puncture blisters if they are sterile, but do not remove them as long as they are intact.
- Stay alert for wound infection. Antibiotic therapy should be considered according to the individual patient’s condition.
- Treat pain according to the individual patient’s condition. Pain relief is very difficult and is the most demanding part of the therapeutic process.
- Debride areas of necrosis thoroughly but cautiously.

Treatment of Late Effects

After immediate treatment of radiation injury, an often long and painful process of healing will ensue. The most important concerns are the following:

- Pain management
- Fibrosis or late ulcers
  **Note:** Use of medication to stimulate vascularization, inhibit infection, and reduce fibrosis may be effective. Examples include Pentoxifylline®, vitamin E, and interferon gamma. Otherwise, surgery may be required.
- Necrosis
- Plastic/reconstructive surgery
  **Note:** Surgical treatment is common. It is most effective if performed early in the treatment process. Full-thickness graft and microsurgery techniques usually provide the best results.
- Psychological effects, such as posttraumatic stress disorder
- Possibility of increased risk of skin cancer later in life

For More Assistance

Technical assistance can be obtained from the Radiation Emergency Assistance Center/Training Site (REAC/TS) at (865) 576-3131 (M-F, 8 AM to 4:30 PM EST) or (865) 576-1005 (after hours), or at http://www.orau.gov/reacts/, and from the Medical Radiobiology Advisory Team (MRAT) at (301) 295-0316.

Also, more information can be obtained from the CDC Health Alert Network at http://www.bt.cdc.gov or 1-800-311-3435.

June 29, 2005
References


Appendix A: Responses of the Skin to Radiation

**Acute epidermal necrosis** (time of onset: < 10 days postexposure; threshold dose: ~550 Gy or 55,000 rads)—
Interphase death of postmitotic keratinocytes in the upper visible layers of the epidermis (may occur with high-dose, low-energy beta irradiation)

**Acute ulceration** (time of onset: < 14 days postexposure; threshold dose: ~20 Gy or 2000 rads)—
Early loss of the epidermis—and to a varying degree, deeper dermal tissue—that results from the death of fibroblasts and endothelial cells in interphase

**Dermal atrophy** (time of onset: > 26 weeks postexposure; threshold dose: ~10 Gy or 1000 rads)—
Thinning of the dermal tissues associated with the contraction of the previously irradiated area

**Dermal necrosis** (time of onset > 10 weeks postexposure; threshold dose: ~20 Gy or 2000 rads)—
Necrosis of the dermal tissues as a consequence of vascular insufficiency

**Dry desquamation** (time of onset: 3–6 weeks postexposure; threshold dose: ~8 Gy or 800 rads)—
Atypical keratinization of the skin caused by the reduction in the number of clonogenic cells within the basal layer of the epidermis

**Early transient erythema** (time of onset: within hours of exposure; threshold dose: ~2 Gray [Gy] or 200 rads)—
Inflammation of the skin caused by activation of a proteolytic enzyme that increases the permeability of the capillaries

**Epilation** (time of onset: 14–21 days; threshold dose: ~3 Gy or 300 rads)—
Hair loss caused by the depletion of matrix cells in the hair follicles

**Late erythema** (time of onset: 8–20 weeks postexposure; threshold dose: ~20 Gy or 2000 rads)—
Inflammation of the skin caused by injury of blood vessels. Edema and impaired lymphatic clearance precede a measured reduction in blood flow.

**Invasive fibrosis** (time of onset: months to years postexposure; threshold dose: ~20 Gy or 2000 rads)—
Method of healing associated with acute ulceration, secondary ulceration, and dermal necrosis that leads to scar tissue formation

**Main erythema** (time of onset: days to weeks postexposure; threshold dose: ~3 Gy or 300 rads)—
Inflammation of the skin caused by hyperaemia of the basal cells and subsequent epidermal hypoplasia (see photos 1 and 2)

**Moist desquamation** (time of onset: 4–6 weeks postexposure; threshold dose: ~15 Gy or 1500 rads)—
Loss of the epidermis caused by sterilization of a high proportion of clonogenic cells within the basal layer of the epidermis

**Secondary ulceration** (time of onset: > 6 weeks postexposure; threshold dose: ~15 Gy or 1500 rads)—
Secondary damage to the dermis as a consequence of dehydration and infection when moist desquamation is severe and protracted because of reproductive sterilization of the vast majority of the clonogenic cells in the irradiated area
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Telangiectasia (time of onset: > 52 weeks postexposure; threshold dose for moderate severity at 5 years: ~40 Gy or 4000 rads)—
Atypical dilation of the superficial dermal capillaries
Appendix B: Images

Figures 1 & 2. Erythema. These photos display the progression of erythema in a patient involved in an x-ray diffraction accident, 9 days to 96 days postexposure. The day following the exposure (not shown), the patient displayed only mild diffuse swelling and erythema of the fingertips. On day 9, punctuate lesions resembling telangiectasias were noted in the subungal region of the right index finger, and on day 11, blisters began to appear. Desquamation continued for several weeks. The patient developed cellulitis in the right thumb approximately 2 years following exposure. The area of the right fingertip and nail continued to cause the patient great pain when even minor trauma occurred to the fingertip, and he required occasional oral narcotic analgesics to manage this pain. He continued to experience intense pain resulting from minor trauma to the affected areas for as long as 4 years postexposure.

(photos courtesy of Gusev IA and reprinted with permission)
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Figures 3 & 4. Acute ulceration. These photos show acute ulceration in a Peruvian patient who inadvertently placed a 26-Ci (0.962-TBq) irridiun-192 ($^{192}\text{Ir}$) source in his back pocket, 3 days and 10 days postexposure. The source remained in the patient’s pocket for approximately 6.5 hours, at which time he complained to his wife about pain in his posterior right thigh. He sought medical advice and was told he probably had been bitten by an insect. In the meantime, his wife sat on the patient’s pants (her case appears on the next page) while breastfeeding the couple’s 1½-year-old child. The source was recovered several hours later by nuclear regulatory authorities, and the patient was transported to Lima for treatment. This patient exhibited a drastic reduction in lymphocyte count by day 3 postexposure, and a 4-by-4-cm lesion appeared on day 4. Eventually he suffered with a massive ulceration and necrosis of the site with infection, and his right leg was amputated. Grade II and III CRI was also evident on his hands, left leg, and perineum, but he survived and returned to his family.
Figure 5. Moist desquamation. This patient is the wife of the previous case study, 26 days postexposure. She was exposed to the $^{192}$Ir source when she sat on her husband's pants (still containing the source) for approximately 20 minutes after he had changed clothes that evening.

Figure 6. Necrosis, fibrosis, and telangiectasia. Same patient, 2 years following exposure.
(photos courtesy of Ricks RC and reprinted with permission)

For more information, visit [www.bt.cdc.gov/radiation](http://www.bt.cdc.gov/radiation), or call CDC at 800-CDC-INFO (English and Spanish) or 888-232-6348 (TTY).
Acute Radiation Syndrome: A Fact Sheet for Physicians

Acute Radiation Syndrome (ARS) (sometimes known as radiation toxicity or radiation sickness) is an acute illness caused by irradiation of the entire body (or most of the body) by a high dose of penetrating radiation in a very short period of time (usually a matter of minutes). The major cause of this syndrome is depletion of immature parenchymal stem cells in specific tissues. Examples of people who suffered from ARS are the survivors of the Hiroshima and Nagasaki atomic bombs, the firefighters that first responded after the Chernobyl Nuclear Power Plant event in 1986, and some unintentional exposures to sterilization irradiators.

The required conditions for Acute Radiation Syndrome (ARS) are:

- **The radiation dose must be large** (i.e., greater than 0.7 Gray (Gy)\(^1,2\) or 70 rads).
  - Mild symptoms may be observed with doses as low as 0.3 Gy or 30 rads.
- **The dose usually must be external** (i.e., the source of radiation is outside of the patient’s body).
  - Radioactive materials deposited inside the body have produced some ARS effects only in extremely rare cases.
- **The radiation must be penetrating** (i.e., able to reach the internal organs).
  - High energy X-rays, gamma rays, and neutrons are penetrating radiations.
- **The entire body** (or a significant portion of it) must have received the dose.
  - Most radiation injuries are local, frequently involving the hands, and these local injuries seldom cause classical signs of ARS.
- **The dose must have been delivered in a short time** (usually a matter of minutes).
  - Fractionated doses are often used in radiation therapy. These large total doses are delivered in small daily amounts over a period of time. Fractionated doses are less effective at inducing ARS than a single dose of the same magnitude.

The three classic ARS Syndromes are:

- **Bone marrow syndrome** (sometimes referred to as hematopoietic syndrome): the full syndrome will usually occur with a dose greater than approximately 0.7 Gy (70 rads) although mild symptoms may occur as low as 0.3 Gy or 30 rads.\(^4\)
  - The survival rate of patients with this syndrome decreases with increasing dose. The primary cause of death is the destruction of the bone marrow, resulting in infection and hemorrhage.
- **Gastrointestinal (GI) syndrome**: the full syndrome will usually occur with a dose greater than approximately 10 Gy (1000 rads) although some symptoms may occur as low as 6 Gy or 600 rads.

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1 The Gray (Gy) is a unit of absorbed dose and reflects an amount of energy deposited into a mass of tissue (1 Gy = 100 rads). In this document, the referenced absorbed dose is that dose inside the patient’s body (i.e., the dose that is normally measured with personal dosimeters).
2 The referenced absorbed dose levels in this document are assumed to be from beta, gamma, or x radiation. Neutron or proton radiation produces many of the health effects described herein at lower absorbed dose levels.
3 The dose may not be uniform, but a large portion of the body must have received more than 0.7 Gy (70 rads).
4 Note: although the dose ranges provided in this document apply to most healthy adult members of the public, a great deal of variability of radiosensitivity among individuals exists, depending upon the age and condition of health of the individual at the time of exposure. Children and infants are especially sensitive.
Acute Radiation Syndrome: A Fact Sheet for Physicians
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- Survival is extremely unlikely with this syndrome. Destructive and irreparable changes in the GI tract and bone marrow usually cause infection, dehydration, and electrolyte imbalance. Death usually occurs within 2 weeks.
- **Cardiovascular (CV)/ Central Nervous System (CNS) syndrome:** the full syndrome will usually occur with a dose greater than approximately 50 Gy (5000 rads) although some symptoms may occur as low as 20 Gy or 2000 rads.
  - Death occurs within 3 days. Death likely is due to collapse of the circulatory system as well as increased pressure in the confining cranial vault as the result of increased fluid content caused by edema, vasculitis, and meningitis.

The four stages of ARS are:

- **Prodromal stage (N-V-D stage):** The classic symptoms for this stage are nausea, vomiting, as well as anorexia and possibly diarrhea (depending on dose), which occur from minutes to days following exposure. The symptoms may last (episodically) for minutes up to several days.
- **Latent stage:** In this stage, the patient looks and feels generally healthy for a few hours or even up to a few weeks.
- **Manifest illness stage:** In this stage, the symptoms depend on the specific syndrome (see Table 1) and last from hours up to several months.
- **Recovery or death:** Most patients who do not recover will die within several months of exposure. The recovery process lasts from several weeks up to two years.

These stages are described in more detail in [Table 1](#).
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**Table 1. Acute Radiation Syndromes**

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Dose*</th>
<th>Prodromal Stage</th>
<th>Latent Stage</th>
<th>Manifest Illness Stage</th>
<th>Recovery</th>
</tr>
</thead>
</table>
| Hematopoietic (Bone marrow) | > 0.7 Gy (> 70 rads) (mild symptoms may occur as low as 0.3 Gy or 30 rads) | • Symptoms are anorexia, nausea and vomiting.  
  • Onset occurs 1 hour to 2 days after exposure.  
  • Stage lasts for minutes to days. | • Stem cells in bone marrow are dying, although patient may appear and feel well.  
  • Stage lasts 1 to 6 weeks. | • Symptoms are anorexia, fever, and malaise.  
  • Drop in all blood cell counts occurs for several weeks.  
  • Primary cause of death is infection and hemorrhage.  
  • Survival decreases with increasing dose.  
  • Most deaths occur within a few months after exposure. | • In most cases, bone marrow cells will begin to repopulate the marrow.  
  • There should be full recovery for a large percentage of individuals from a few weeks up to two years after exposure.  
  • Death may occur in some individuals at 1.2 Gy (120 rads).  
  • The LD50/60 † is about 2.5 to 5 Gy (250 to 500 rads). |
| Gastrointestinal (GI) | > 10 Gy (> 1000 rads) (some symptoms may occur as low as 6 Gy or 600 rads) | • Symptoms are anorexia, severe nausea, vomiting, cramps, and diarrhea.  
  • Onset occurs within a few hours after exposure.  
  • Stage lasts about 2 days. | • Stem cells in bone marrow and cells lining GI tract are dying, although patient may appear and feel well.  
  • Stage lasts less than 1 week. | • Symptoms are malaise, anorexia, severe diarrhea, fever, dehydration, and electrolyte imbalance.  
  • Death is due to infection, dehydration, and electrolyte imbalance.  
  • Death occurs within 2 weeks of exposure. | • The LD100 ‡ is about 10 Gy (1000 rads). |
| Cardiovascular (CV)/ Central Nervous System (CNS) | > 50 Gy (5000 rads) (some symptoms may occur as low as 20 Gy or 2000 rads) | • Symptoms are extreme nervousness and confusion; severe nausea, vomiting, and watery diarrhea; loss of consciousness; and burning sensations of the skin.  
  • Onset occurs within minutes of exposure.  
  • Stage lasts for minutes to hours. | • Patient may return to partial functionality.  
  • Stage may last for hours but often is less. | • Symptoms are return of watery diarrhea, convulsions, and coma.  
  • Onset occurs 5 to 6 hours after exposure.  
  • Death occurs within 3 days of exposure. | • No recovery is expected. |

* The absorbed doses quoted here are “gamma equivalent” values. Neutrons or protons generally produce the same effects as gamma, beta, or X-rays but at lower doses. If the patient has been exposed to neutrons or protons, consult radiation experts on how to interpret the dose.

† The LD50/60 is the dose necessary to kill 50% of the exposed population in 60 days.

‡ The LD100 is the dose necessary to kill 100% of the exposed population.
Cutaneous Radiation Syndrome (CRS)
The concept of cutaneous radiation syndrome (CRS) was introduced in recent years to describe the complex pathological syndrome that results from acute radiation exposure to the skin.

ARS usually will be accompanied by some skin damage. It is also possible to receive a damaging dose to the skin without symptoms of ARS, especially with acute exposures to beta radiation or X-rays. Sometimes this occurs when radioactive materials contaminate a patient’s skin or clothes.

When the basal cell layer of the skin is damaged by radiation, inflammation, erythema, and dry or moist desquamation can occur. Also, hair follicles may be damaged, causing epilation. Within a few hours after irradiation, a transient and inconsistent erythema (associated with itching) can occur. Then, a latent phase may occur and last from a few days up to several weeks, when intense reddening, blistering, and ulceration of the irradiated site are visible.

In most cases, healing occurs by regenerative means; however, very large skin doses can cause permanent hair loss, damaged sebaceous and sweat glands, atrophy, fibrosis, decreased or increased skin pigmentation, and ulceration or necrosis of the exposed tissue.

Patient Management
Triage: If radiation exposure is suspected:
- Secure ABCs (airway, breathing, circulation) and physiologic monitoring (blood pressure, blood gases, electrolyte and urine output) as appropriate.
- Treat major trauma, burns, and respiratory injury if evident.
- In addition to the blood samples required to address the trauma, obtain blood samples for CBC (complete blood count), with attention to lymphocyte count, and HLA (human leukocyte antigen) typing prior to any initial transfusion and at periodic intervals following transfusion.
- Treat contamination as needed.
- If exposure occurred within 8 to 12 hours, repeat CBC, with attention to lymphocyte count, 2 or 3 more times (approximately every 2 to 3 hours) to assess lymphocyte depletion.

Diagnosis
The diagnosis of ARS can be difficult to make because ARS causes no unique disease. Also, depending on the dose, the prodromal stage may not occur for hours or days after exposure, or the patient may already be in the latent stage by the time they receive treatment, in which case the patient may appear and feel well when first assessed.

If a patient received more than 0.05 Gy (5 rads) and three or four CBCs are taken within 8 to 12 hours of the exposure, a quick estimate of the dose can be made (see Ricks, et. al. for details). If these initial blood counts are not taken, the dose can still be estimated by using CBC results over the first few days. It would be best to have radiation dosimetrists conduct the dose assessment, if possible.

If a patient is known to have been or suspected of having been exposed to a large radiation dose, draw blood for CBC analysis with special attention to the lymphocyte count, every 2 to 3 hours during the first 8 hours after exposure (and every 4 to 6 hours for the next 2 days). Observe the patient during this time for symptoms and consult with radiation experts before ruling out ARS.

If no radiation exposure is initially suspected, you may consider ARS in the differential diagnosis if a history exists of nausea and vomiting that is unexplained by other causes. Other indications are bleeding, epilation, or white blood count (WBC) and platelet counts abnormally low a few days or weeks after unexplained nausea and vomiting. Again, consider CBC and chromosome analysis and consultation with radiation experts to confirm diagnosis.
Initial Treatment and Diagnostic Evaluation
Treat vomiting and repeat CBC analysis with special attention to the lymphocyte count every 2 to 3 hours for the first 8 to 12 hours after exposure (and every 4 to 6 hours for the following 2 or 3 days). Sequential changes in absolute lymphocyte counts over time are demonstrated below in the Andrews Lymphocyte Nomogram (see Figure 1). Precisely record all clinical symptoms, particularly nausea, vomiting, diarrhea, and itching, reddening or blistering of the skin. Be sure to include time of onset.

Figure 1: Andrews Lymphocyte Nomogram


Note and record areas of erythema. If possible, take color photographs of suspected radiation skin damage. Consider tissue, blood typing, and initiating viral prophylaxis. Promptly consult with radiation, hematology, and radiotherapy experts about dosimetry, prognosis, and treatment options. Call the Radiation Emergency Assistance Center/Training Site (REAC/TS) at (865) 576-3131 (M-F, 8 am to 4:30 pm EST) or (865) 576-1005 (after hours) to record the incident in the Radiation Accident Registry System.

After consultation, begin the following treatment (as indicated):
- supportive care in a clean environment (if available, the use of a burn unit may be quite effective)
- prevention and treatment of infections
- stimulation of hematopoiesis by use of growth factors
- stem cell transfusions or platelet transfusions (if platelet count is too low)
- psychological support
- careful observation for erythema (document locations), hair loss, skin injury, mucositis, parotitis, weight loss, or fever
- confirmation of initial dose estimate using chromosome aberration cytogenetic bioassay when possible. Although resource intensive, this is the best method of dose assessment following acute exposures.
- consultation with experts in radiation accident management

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5 Collect vomitus in the first few days for later analysis.
Acute Radiation Syndrome: A Fact Sheet for Physicians
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For More Help
Technical assistance can be obtained from the Radiation Emergency Assistance Center/Training Site (REAC/TSC) at (865) 576-3131 (M-F, 8 am to 4:30 pm EST) or (865) 576-1005 (after hours), or on their web site at www.orau.gov/reacts, and the Medical Radiobiology Advisory Team (MRAT) at (301) 295-0316.

Also, more information can be obtained from the CDC Health Alert Network at www.bt.cdc.gov or by calling (800) 311-3435.

References


