Radiation-induced meningioma

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The long-term or delayed side effects of irradiation on neural tissue are now known to include the induction of new central nervous system neoplasms. However, during the first half of the 20th century, human neural tissue was generally considered relatively resistant to the carcinogenic and other ill effects of ionizing radiation. As a result, exposure to relatively high doses of x-rays from diagnostic examinations and therapeutic treatment was common.

In the present article the authors review the literature relating to radiation-induced meningiomas (RIMs). Emphasis is placed on meningiomas resulting from childhood treatment for primary brain tumor or tinea capitis, exposure to dental x-rays, and exposure to atomic explosions in Hiroshima and Nagasaki. The incidence and natural history of RIMs following exposure to high- and low-dose radiation is presented, including latency, multiplicity, histopathological features, and recurrence rates. The authors review the typical presentation of patients with RIMs and discuss unique aspects of the surgical management of these tumors compared with sporadic meningioma, based on their clinical experience in treating these lesions. (DOI: 10.3171/FOC/2008/24/5/E7)

KEY WORDS • ionizing radiation • radiation-induced meningioma • tinea capitis

Abbreviations used in this paper: CT = computed tomography; MR = magnetic resonance; RIM = radiation-induced meningioma; SM = spontaneous meningioma; WHO = World Health Organization.
1967), a founder of 20th century genetics, published evidence in *Science* that x-ray exposure produced a 150-fold increase in the natural mutation rate of fruit flies and that the rate of mutations was proportional to the dose. He showed that x-rays broke apart and rearranged the genes. In 1946, this research was recognized with the Nobel Prize in Medicine. Muller spoke against needless exposure to x-rays in medicine and urged protection for individuals routinely exposed to ionizing radiation.

Neurological tissue was long considered relatively resistant to damage from x-rays, and some authors suggested that higher doses allowed longer survival in patients with glioblastoma multiforme. In 1938, Davidoff et al. described the clinical and histological effects of delivering 10–50 Gy in a single exposure, or 48–72 Gy in 2 fractions, applied directly to the brains and spinal cords in monkeys. The authors observed profound physiological and morphological changes, which were especially marked in the glia and nerve tissues. The intensity of histological change was determined primarily by x-ray dose and, to a lesser extent, by the time lapsed between irradiation and autopsy. Clinical and histological changes were progressive. Other authors also documented pronounced degenerative effects on nerve tissue from exposure to ionizing radiation.

However, only after atomic bombs were dropped on Nagasaki and Hiroshima, and atomic tests were performed during the 1950s, did the general public and much of the medical community became aware of risks from exposure to ionizing radiation. In 1956, the Committee on Biological Effects of Atomic Radiation of the National Academy of Sciences introduced the concept of regulating the overall dose to the general population, suggesting a 0.1-Gy limit during the reproductive years, a span averaging 30 years.

Today we know that exposure to ionizing radiation overcomes the binding energy of electrons orbiting atoms and molecules. In biological material, hydroxyl radicals formed by x-ray interaction with water molecules cause strand breaks or base damage in nearby DNA. The x-rays may also ionize DNA directly. While most radiation-induced damage to DNA is repaired by mechanisms within the cell, double-strand breaks are more difficult to repair, and occasional misrepair can result in point mutations, chromosomal translocations, and gene fusions, all of which have the potential for induction of cancer.

Radiation-induced changes to neural tissue are now known to include visual deterioration, hearing loss, hormonal disturbances, vasculopathy, brain and bone necrosis, atrophy, demyelination, calcification, fatty replacement of bone marrow, and induction of central nervous system neoplasms, and are increasingly evident radiologically. A dose effect has been documented for many changes, including tumor induction.

Both x-rays and gamma rays are forms of ionizing radiation, which is defined as radiation that has sufficient ener-
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...gy to displace electrons from molecules. Free electrons can damage human cells. At very high doses, the damage inflicted on tissues is evident within days of exposure. Late-onset effects such as cancer and other neoplasms, however, may develop even with exposure to relatively low doses.

**Radiation-Induced Meningioma**

Radiation-induced meningioma is the most common brain neoplasm known to be caused by ionizing radiation. Harrison et al. grouped RIMs into 3 categories: those due to high-dose (> 20 Gy), intermediate-dose (10–20 Gy), and low-dose (< 10 Gy) radiation. Other authors define doses > 10 Gy as high.

### High-Dose RIMs

In 1953, Mann et al. described a meningioma in a 4-year-old girl who had undergone radiation treatment for optic nerve glioma. The child’s meningioma eventually became malignant. Several authors have summarized up to 126 cases of high-dose RIMs reported in the literature between 1953 and 2002. The radiation doses ranged from 22 to 87 Gy, and the majority of patients had undergone radiotherapy as children. The mean latency from irradiation to diagnosis of the meningioma was ~ 19 years, with a tendency for shorter latency in patients treated with higher doses and those who had undergone radiotherapy at younger ages.

### Low-Dose RIMs

The increased incidence of meningiomas following exposure to low-dose radiation has been reported in patients who, as children, had undergone radiation treatment for tinea capitis, those whose heads and necks were exposed to medical and dental x-rays at a young age, and survivors of the atomic explosions in Hiroshima and Nagasaki.

**Irradiation for Tinea Capitis.** Prior to the introduction of griseofulvin in 1960, the world standard for treatment of tinea capitis was scalp irradiation. Treatment for tinea capitis was administered using the Adamson–Keinbock technique, which was designed to irradiate the entire scalp as uniformly as possible through exposure to 5 overlapping treatment areas. Phantom studies using this technique showed that the radiation dose to the scalp was 5–8 Gy, the surface dose to the scalp was 5–8 Gy, the surface dose to the brain was 1.4–1.5 Gy, and the skull base received an average dose of 0.7 Gy. In 1966, Albert et al. reported a substantial increase in cancer, mental disease, and permanent damage to scalp hair in 1908 patients in whom radiotherapy was performed for tinea capitis at the New York University Medical Center between 1940 and 1958, compared with 1801 patients treated for tinea capitis without radiotherapy during the same years. Other reports followed.

In 1974, Modan and colleagues at the Sheba Medical Center in Tel Aviv published a retrospective cohort study showing a significantly higher risk of malignant and benign head and neck tumors among ~ 11,000 Israeli adults treated for tinea capitis as children. This major epidemiological work is widely credited as proving the causal role of radiation in the development of meningiomas in some patients.

Follow-up studies of the Israeli cohort showed that a radiation dose of only 1–2 Gy administered during childhood led to a 9.5-fold increase in meningioma incidence. Continuing follow-up shows that elevated risk of brain tumor, including meningioma, is positively associated with dose, with excess relative risk for benign meningioma rising to 18.82 for doses > 2.6 Gy. These studies showed a higher prevalence of calvarial tumors and multiple meningiomas, and higher recurrence rates in RIMs than SMs. The mean latency was ~ 36 years.

**Low-Dose RIM Due to Dental Radiographic Examination.** In 1953, Nolan alone and with Patterson published studies in which significant blood changes occurred in patients exposed to 1.15–2.8 Gy during full-mouth radiographic examinations. Nolan observed that full-mouth series delivered converging lines of radiation, with high points near the meninges. In 1980, nearly 30 years later, Preston-Martin et al. noted a higher incidence of meningioma in women with a history of full-mouth dental x-rays. The risk in this cohort study was higher in patients who had undergone radiographic examinations as children or teenagers and in those in whom radiographic examinations were conducted before 1945, when doses were higher. The majority of tumors were located in the tentorial or subtentorial region. The authors of more recent case-control studies in Sweden and the US have found that dental radiographic examinations performed during the adult years may also increase the risk of meningioma.

**Radiation-Induced Meningiomas in Survivors of Atomic Bombs Dropped on Hiroshima and Nagasaki.** The increased incidence of meningiomas among survivors of the 1945 atomic explosions in Japan was shown only in 1994, when Shibata et al. demonstrated a higher incidence of meningiomas in survivors of the bombing in Nagasaki. In 1997 Shintani et al. published similar findings after studying data obtained in Hiroshima survivors. Due to the relatively low-dose exposure to ionizing radiation among these survivors compared with those undergoing radiotherapy for tinea capitis, the average latency was greater in the Japanese studies. The risk of meningioma induction was shown to increase with closer proximity to the bombs’ epicenters and in those exposed during childhood.

**Clinical Presentation**

Radiation-induced meningiomas typically display female preponderance, although the difference between sexes may be less distinct compared with SMs. Strojan et al. reported female predominance of 1:3.3:1 in a survey of 126 cases of high-dose RIMs. In a review of the incidence of central nervous system tumors among survivors of the atomic bombs in Hiroshima and Nagasaki, Preston et al. found that excess relative risk for RIMs is higher in men than in women.

Scanty hair, or alopecia, and an atrophic scalp are hallmarks of irradiation. We are also of the impression that microcephaly is increased in patients with low-dose RIMs compared with the general population, although no evidence has been reported in the literature. Microcephaly most likely results from premature closure of the skull sutures when radiation is administered to the head in young children.
The location of the RIM is primarily related to the site of exposure. The authors of several large studies conducted in Israel have documented a primarily calvarial location in patients treated for tinea capitis.\(^5^9,6^4,7^5\) Ghim et al.\(^1^7\) found that 11 of 13 pediatric patients with high-dose RIMs had calvarial meningiomas, whereas skull base meningiomas were found in 4–19% of patients with RIM in whom there was a history of high-dose irradiation for primary brain tumors.\(^3^9,5^9,6^7,7^6\) Epidemiological studies of patients with RIM linked to full-mouth dental x-rays have shown that skull base meningiomas are more common following this type of radiation exposure.\(^5^4\)

The latency period between exposure and clinical diagnosis of meningioma varies with radiation dose and age at initial treatment. Sadetzki et al.\(^6^4\) reported a 36.3-year average latency (range 12–49 years) in the Israeli low-dose cohort study. In a thorough review of the literature, Harrison et al.\(^2^4\) calculated that a mean of 35.2 years elapsed between low-dose irradiation (\(<\ 10\ Gy\)) and diagnosis, whereas a mean of 26.1 and 19.5 years, respectively, elapsed in patients receiving moderate (10–20 Gy) and high-dose (\(>\ 20\ Gy\)) treatment. In their survey of reported cases of RIM in patients exposed to high-dose radiation, Strojan et al.\(^7^8\) observed average latency of 18.7 years and also noted shorter latency in patients in whom radiation treatments were administered at younger ages. Ghim et al.\(^1^7\) found a mean latency of only 10.8 years (range 5–15.5) in a study of 13 pediatric high-dose RIM patients with mean age at diagnosis 13 years (range 6–18). These patients had been treated for brain tumors at an average age of 2.5 years. A 14-month latency period was reported in an 11-year-old boy following high-dose irradiation for a tumor in the posterior fossa.\(^1^2\)

Radiation-induced meningioma differs from SM in patient age at presentation and in the multiplicity, aggressiveness, and rate of tumor recurrence. Mean age at presentation is reported as 29.2–37.9 years in patients exposed to high-dose radiation\(^2^1,1^4,2^4,4^1,6^6,6^7,7^5\) and 45–58 years in those who received low-dose treatment,\(^2^1,2^4,3^2,3^9,7^6\) whereas SM generally arises in the 5th or 6th decade of life.

The incidence of multiple lesions is elevated in both patients with low- and high-dose RIMs, and it ranges from 4.6 to 18.7% of cases.\(^5^9,7^6\) Multiplicity has been reported in 15.8% of patients with RIMs, compared with 2.4% of those with SMs in the Israeli cohort study.\(^5^4\)

Protocols for MR and CT imaging of RIM and SM are comparable. The appearance of RIM on MR and CT images is comparable with SM; thus, the differential diagnosis cannot be based on imaging studies. The presence of multiple meningiomas in a patient with a history of head irradiation, as well as typical skin changes, raises the suspicion of RIM (Fig. 3).

Meningotheliomatous, transitional, and fibroblastic histological subtypes are the most common in RIM.\(^2^7,7^6\) Histological features are distinctive compared with SM. Soffer et al.\(^7^6\) have noted high cellularity, nuclear pleomorphism, an increased mitotic rate, focal necrosis, bone invasion, and tumor infiltration of the brain in a series of 42 patients with low-dose RIM. Rubinstein et al.\(^2^9\) reported several findings: a high degree of cellularity; pleomorphic nuclei with great variation in nuclear size, shape, and chromatin density; numerous multinucleated and giant cells; and nuclei with vacuolated inclusions. They also noted frequent mitoses, psammoma bodies, foam cells, and thickened blood vessels.

![Fig. 3. Multiplicity is elevated in RIM.](image1)

![Fig. 4. Photograph of a patient with an RIM and related alopecia and scalp atrophy.](image2)
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that did not stain for amyloid. Louis et al.20 found that RIMs were more often atypical, or aggressive and multifocal, with higher proliferation indices than SMs. Musa et al.42 reported a 23% incidence of atypia or malignancy among 79 cases of high-dose RIM.

Radiation-induced meningiomas may also exhibit more aggressive clinical behavior than SMs,20,24,52,59,67,76 including high recurrence rates following surgery and radiotherapy. Rubinstein et al.59 noted a 25.6% recurrence rate in 43 patients with RIM, and 11.6% of this group experienced multiple recurrences, compared with an 11.4% recurrence rate in 258 patients with SMs in their series. Both groups were followed for only 4 years. Higher recurrence rates in patients with RIMs than those with SMs have also been reported elsewhere.64,76

In a preliminary investigation, we found evidence of increased recurrence rates similar to those seen in other studies, but there was a significantly higher rate of multiplicity and much higher frequency of atypical (WHO Grade II) and anaplastic (WHO Grade III) histopathology in patients with RIMs compared with SMs treated in our institution (F. Umansky et al., unpublished data).

Management of RIM

Surgical Intervention

Surgical removal is the treatment of choice for most cases of RIM, although complete and safe resection may not be possible due to the lesion’s frequent multiplicity, involvement of osseous structures and vessels, and aggressive nature. Paradoxically, stereotactic radiosurgery or fractionated stereotactic radiosurgery may be appropriate adjuncts to surgery or may be performed in lieu of surgery in some patients, despite the radiation-related origins of RIMs. In some patients, angiography may be appropriate for visualization of the tumor’s vascular anatomy and preoperative embolization.

The first consideration when planning the surgical approach for resection of RIM is often scalp atrophy, followed by multiplicity and the high rate of recurrence. The scalp may be slightly or severely atrophic and poorly vascularized, depending on the radiation dose received. In some patients, the thickness of the entire galeal/skin flap may reach only 1.5–2 mm, necessitating careful planning and meticulous technique to avoid cerebrospinal fluid leakage and skin flap dehiscence. The risk of these complications increases in patients with an atrophic scalp who require multiple craniotomies, which is a common scenario in the RIM population (Fig. 4). In such cases, it may be appropriate to include a plastic surgeon on the team.

A large percentage of RIMs are located at the calvaria, and in cases of parasagittal or falce meningiomas preoperative assessment of the superior sagittal sinus is critical in planning surgery (Fig. 5). We prefer the use of MR imaging with MR venography for assessment of patency, although angiography may be required in rare cases. In patients with highly vascular tumors, preoperative embolization may be beneficial.

If healthy skin is available at the periphery of the irradiated, atrophic scalp, it should be used.42 In our experience, skin damage is often less severe in the low forehead, low temporal, or occipital areas compared with the convexity and vertex. When skin in these basal areas is indeed healthier and better vascularized, incisions may be made more inferiorly than usual. Use of hemostatic clamps or coagulation is not advised as they may further damage this fragile skin and may cause drying or shrinkage of the skin edges. Straight or slightly curved incisions are preferred to horseshoe contours, and flap size should be held to the minimum.

Given the aggressive nature and high recurrence rates of RIMs, wide resection margins in the involved dura mater are vital.8,9,26,74,80 In cases in which the tumor involves under- or overlying bone, the osseous portion of the tumor should be radically removed.24,59,76,77,80 because osseous invasion has been linked to higher rates of tumor recurrence.26,74 When we find bone flap invasion or suspect it has

Fig. 5. Studies obtained in a 56-year-old woman who had undergone radiotherapy for tinea capitis at age 6 years. A: In recent years she had detected a growing lump on her scalp and had recently developed difficulty walking. B and C: Axial and sagittal CT scans revealed a large parasagittal meningioma. Magnetic resonance imaging had also shown an additional small, parasellar meningioma (data not shown). Note the presence of alopecia, a hallmark of radiation to the scalp.
occurred, we autoclave the flap or replace it with acrylic graft. Recurrence rates are higher among patients with tumors involving the skull base or a major cranial sinus, where wide resection margins are often impossible to achieve.\textsuperscript{3,10,14}

Fragile skin conditions frequently necessitate special considerations when closing the wound. Multilayer closure may be impossible, especially in convexity meningiomas, since the atrophic scalp may not accept additional sutures. In patients with a very thin scalp, we close the skin and galea aponeurosis with a single layer of running 3-0 sutures, with or without locking, using nylon, not Vicryl. This closure should be performed carefully to provide a single, watertight layer, capable of preventing cerebrospinal fluid leakage; however, if the stitches are closed too tightly, skin necrosis may result and cause potentially serious consequences. Staples are not suitable for closure in these patients. They provide poor approximation, without hermetic closure, and can easily damage a fragile, atrophic scalp.

\textit{Stereotactic Radiosurgery and Fractionated Stereotactic Radiotherapy}

Surgery may be followed by radiotherapy in patients with RIMs if radical excision cannot be achieved. When patients have already received the maximum tolerable dose of radiation, stereotactic radiotherapy or stereotactic radiosurgery may be preferable to conventional radiotherapy. Adjuvant conventional conformal radiation is generally recommended following surgery or radiosurgery to treat viable tumor cells that remain along the dura and in brain parenchyma in patients with atypical (WHO Grade II), anaplastic (WHO Grade III), or malignant (WHO Grade IV) meningiomas.\textsuperscript{5,50} Careful consideration should be given to original doses and treatment fields when planning new therapeutic irradiation.

\textit{Other Therapies}

Systemic treatment with tamoxifen\textsuperscript{21} and hydroxyurea\textsuperscript{35,47,49,58,60,99} has been attempted as an adjuvant therapy in patients with recurrent, unresectable SMs. In the past, we have used both of these agents in selected RIM patients with recurrent tumors when surgery and radiotherapy failed, with no measurable response. In our experience, these agents did not provide measurable clinical benefit for the management of meningioma.

\textit{Conclusions}

Exposure to ionizing radiation has been shown to significantly increase the risk of meningioma, a solid tumor arising from cells of the meningeal coverings of the brain. The risk of meningioma formation increases, and the latency period between exposure and tumor development decreases, with higher doses of radiation; however, exposure to even low radiation doses has been shown to significantly increase the risk of meningioma.\textsuperscript{51,62,69-72}

The Israeli experience has shown that even low-dose radiation in children leads to a high and significant increase in the risk of developing meningioma. This risk should be considered when decisions are made to use radiation in routine imaging examinations, such as CT scanning,\textsuperscript{16} or to administer higher doses of radiation for treatment of benign tumors during childhood, adolescence, and young adulthood.\textsuperscript{2-4,17,42,78,81}

Radiation-induced meningiomas are characterized by marked changes to the scalp, including alopecia, atrophy, and poor vascularization. Patients with RIMs frequently present with multiple tumors. Compared with SMs, a higher proportion of RIMs are atypical or anaplastic, and recurrence rates are higher. Poor skin condition may necessitate adjustment of the surgical approach and skin flap placement. Paradoxically, stereotactic radiotherapy may be used to treat patients with unresectable or residual/recurrent tumors.

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