Malignant Triton Tumor of the Sciatic Nerve as a Secondary Malignancy after Extended Field Radiotherapy and Chemotherapy of Hodgkin’s Disease

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Key Words
Triton tumor · Radiotherapy · Second malignancy · Hodgkin’s disease

Abstract
Late effects of therapy for Hodgkin’s disease include secondary malignancies like leukemia, lymphoma or solid tumors developing after long periods of latency. Ionizing radiation often causes the last group. The highest risks have been described for induced breast and lung cancers. We are the first to report a malignant triton tumor (MTT) as a secondary malignancy after radiotherapy and chemotherapy for Hodgkin’s lymphoma. MTT is a very rare subtype of malignant peripheral nerve sheath tumors with rhabdomyoblastic differentiation and an aggressive course of disease.

Case Presentation
At the first diagnosis of Hodgkin’s lymphoma (CSIIIA), the patient was 15 years of age (fig. 1). He received total lymphoid irradiation with a cobalt-60 unit. The field configuration consisted of a mantle field (including the major lymph node regions above the diaphragm)
up to 33.4 Gy and an inverted Y field (including the retroperitoneal, iliac and inguinal lymph nodes) up to 30 Gy with an additional paravertebral/para-aortal (L2/L3) boost of 6 Gy to the tumor region. Afterwards, 2 cycles of COPP chemotherapy (cyclophosphamide, vincristine, procarbazine, prednisolone) were applied.

Twenty-one years later, the patient developed progressive pain in the left gluteal muscle. An MRI scan showed a tumorous mass of 3 × 4.4 cm in the left gluteal region expanding to the greater trochanter. A microsurgical operation revealed an isolated tumor of the left sciatic nerve, unfortunately with an R2 resection status.

Pathology

The pathological specimen consisted of multiple, gray-white pieces of hard consistency measuring about 13 × 8 × 4 cm in diameter. Microscopically, the tumor tissue was streaked by ‘geographic’ zones of necrosis and surrounded by a pseudocapsule of dense collagenous tissue. The density of the tumor cells varied. In areas of high-cell density, the tumor cells consisted of plump spindle cells, sometimes with wavy nuclei, which showed a high chromatin density. The mitotic rate was high (31 mitoses/10 HPF). The tumor cells lay concentrically around delicate vessels, in which they sometimes herniated. In areas of lower-cell density, the tissue was sometimes of a myxoid quality. Rarely, the tumor cells grew in fascicles or concentric eddies. Scattered, large, polygonal, eosinophilic cells were crowded with eosinophilic filaments (rhabdomyoblasts) and some elongated cells with cross-striations were seen. There were foci of chondroid differentiation (fig. 2, fig. 3).

Clinical Course of the Disease

Two months after the microsurgical operation, the tumor had regrown up to 9.9 × 4.7 × 5.9 cm (fig. 4). A second microsurgical operation was undertaken with neurolysis and decompression of the left sciatic nerve, again resulting in R2 resection margins. This time, the patient refused a complete resection of the sciatic nerve as he feared the unavoidable loss of function. Yet another 2 months later, during the planning phase of local radiotherapy, further imaging showed a fulminant progress of the disease. Accordingly, the treatment plans were changed, and 1 cycle of VIDE chemotherapy (vincristine, ifosfamide, doxorubicin, etoposide) without doxorubicin and 1 cycle of I²VA (ifosfamide, vincristine, actinomycin-D) chemotherapy were applied in a pseudoneoadjuvant attempt. At restaging, progressive disease was again diagnosed, and another 2 cycles of I²VA were applied, ultimately resulting in a stable disease.

In a curative attempt, the tumor including the sciatic nerve was resected during a third operation with a narrow R0 resection status (1 mm). Another 2 months later, an MRI scan showed 7 satellite metastases distributed in the initial tumor region. Hyperfractionated radiotherapy was applied to the whole region with 1.2 Gy twice daily up to 50.4 Gy in a curative attempt. The loco-regional metastases received an additional boost dose of 14.4 Gy, resulting in a cumulative dose of 64.8 Gy (fig. 5). Despite intensive treatment, the disease progressed rapidly, leading to an exulcerating local tumor. Ultimately, the patient underwent left hemipelvectomy with a narrow R0 resection status. Only half a year later, the patient experienced extensive local progress and pulmonary metastases. He died of uncontrollable bleeding despite embolization.
Discussion

Generally, the traditional Cahan criteria are used to define radiation-induced malignancy [1], i.e. (1) the disease must have arisen in the irradiated field with (2) a latency period of >4 years between irradiation and induced malignancy, (3) the induced tumor must have undergone biopsy and (4) the tissue of the induced tumor must have been normal prior to radiation exposure. All 4 points are fulfilled in the reported case. Secondary solid tumors in Hodgkin’s disease are mostly related to radiotherapy and occur with a long latency period of more than 7 years. There is a 5.6-fold increased risk for secondary female breast cancer (57 cases/10,000 persons/year) and an increased risk for secondary lung cancer after doses as low as 5 Gy, especially when patients continue to smoke after therapy [2, 3]. The relative risk of developing a secondary malignancy after combined modality treatment for Hodgkin’s disease is 2.3–2.9 with an absolute risk of 44.5 cases/10,000 patients/year, including leukemia, lymphoma and solid tumors [4, 5].

A malignant peripheral nerve sheath tumor with rhabdomyosarcomatous differentiation was first described by Masson in 1932 [6]. This tumor is defined by histological and immunohistochemical features of a malignant peripheral nerve sheath tumor with focal rhabdomyosarcomatous elements or as a rhabdomyosarcomatous tumor with focal nerve sheath elements, which occur in a peripheral nerve or in neurofibromatosis type 1 (NF-1) [7, 8]. Histology and immunohistochemistry are the mainstay of diagnosis: The tumor cells express S100, desmin, muscle-specific actin, myosin, vimentin, myoglobin and, as in the present case, MyoD1 and Myf4.

Malignant triton tumor (MTT) is associated with NF-1 in 50–70% of all cases, where it is diagnosed at a younger age and predominantly in males. Patients without NF-1 are diagnosed at a mean age of 32 years with an equal sex distribution. The overall survival rate is low with only 26% in both groups. The local recurrence rate is about 40%, metastases occur in about 50% [9, 10]. Survival depends on the location of the disease, as patients with MTT of the head and neck or extremities generally have a longer life expectancy than those with MTT of the gluteal region, trunk or retroperitoneum [6, 9, 11–13]. The extent of the excision is another factor with a significant impact on the local outcome and survival, as wide resection margins usually result in a better response to adjuvant treatment and better outcome [9, 11, 14]. Therefore, the treatment of MTT must compulsorily start with a complete resection and large resection margins whenever possible.

In our case, the tumor was localized in the left gluteal region surrounding the left sciatic nerve. The patient’s initial wish was to preserve the function of the sciatic nerve with minimal surgery. For this reason, the necessary radical approach with adequate margins was omitted. However, soon after the third operation – eventually with a complete resection of the tumor with an R0 resection status at close margins – local recurrence developed during the planning phase of adjuvant radiotherapy, so that ultimately chemotherapy and hemipelvectomy had to be performed.

Up to date, the effectiveness of adjuvant therapy in MTT is still undefined, as no prospective randomized trials are available. In most of the reported cases, the disease progressed rapidly despite adjuvant chemotherapy and radiotherapy [8, 11, 13]. However, it has been suggested that adjuvant therapies may prolong survival after the radical resection of the MTT [7, 15–17].
References

Fig. 1. Total lymphoid irradiation with a cobalt-60 unit. The subdiaphragmal field configuration consisted of an inverted Y field including the retroperitoneal, iliac and inguinal lymph nodes. Treatment planning included a lymphography with lipiodol. At the left hilum of the kidney, pathological lymph nodes were visible; they received an additional boost dose.

Fig. 2. Myxoid tumor tissue with cell whorls herniating rarely into the vascular lumina (HE, original magnification: ×100).
Nitsche et al.: Malignant Triton Tumor of the Sciatic Nerve as a Secondary Malignancy after Extended Field Radiotherapy and Chemotherapy of Hodgkin’s Disease

**Fig. 3.** a Tumor tissue consisting of plump spindle cells with many mitoses marked by arrows (HE). b Cytoplasm-rich polygonal cells (so-called rhabdomyoblasts) and elongated multinucleated strap cells (HE). c Tumor tissue made up of spindle cells and a chondroid area with a hyaline matrix and lacunar cells (HE). d S100-positive tumor cells in a perivascular location (anti-S100, ABC). Original magnification: ×200 in a and b, ×100 in c and d.

**Fig. 4.** Coronal T1-weighted MRI scan revealing the recurrent disease. An extensive tumor (10 × 5 × 6 cm) expanding from the left sciatic nerve to the gluteal region.
Fig. 5. The dose distribution of the radiotherapy treatment planning: yellow indicates the 95% isodose of the prescribed dose of 50.4 Gy, red indicates the 95% isodose of the prescribed boost dose of 64.8 Gy – representing an additional boost to the region of the recurrent tumor.