Glioblastoma of the Conus Medullaris Following Treatment of Hodgkin’s Lymphoma: History of a Case and Literature Review

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Abstract

Purpose: Reports of a glioblastoma arising in a previously irradiated field are rare in the literature, even more so in the conus medullaris of the spinal cord.

Method: Case report and review of the literature reporting other radiation-induced intra-medullary glioblastomas.

Results: We report a case of glioblastoma of the conus, which subsequently metastasized to the brain, arising in a 45 years old man, nine years after treatment for Hodgkin’s lymphoma, which had included the administration of 41 Gy to the brain and spinal cord.

Conclusion: Despite a well-conducted treatment by several lines of chemotherapy, the prognosis of radiation-induced glioblastoma is poor especially since these patients often cannot benefit from a re-irradiation due to the maximum dose supported by the spinal cord.

Implications of cancer survivors: Associations are now estabilished between therapeutic exposures and specific complications but considerable inter-individual variability is observed for a given therapeutic exposure. In this context, identification of genetic susceptibilities for specific treatment-associated late effect is a very promising future approach. Improvement in our knowledge on genetic susceptibilities may help define more personalized primary therapies that weigh treatment efficacy with the risk of late complications.

Keywords: Glioblastoma; Spinal cord; Hodgkin’s lymphoma; Radiotherapy; Radiation-induced; Second cancer

Clinical Practice Points

Glioblastomas of the conus medullaris is a very rare grade 4 glial tumor that can appear in irradiated zone on the terminal medullary cone and may be in this case a serious complication after lymphoma.

Such secondary cancer and others emphasize the needs of improvement in our knowledge on genetic susceptibilities, which may help define more personalized primary therapies that weigh treatment efficacy with the risk of late complications.

Introduction

Glioblastomas, also known as grade IV astrocytomas, are extremely aggressive malignant tumors of the glial cells of the central nervous system. They arise more frequently in the brain than the spinal cord. Their estimated incidence in all sites is 1-5/100,000 in European and North American populations. The prognosis remains grave, whatever the initial site, and despite any surgical, chemotherapeutic or radiological treatment that may be tried.

Glioblastomas may be classified as primary or secondary. Among the secondary tumors of the central nervous system, a few cases of intracerebral glioblastomas in previously irradiated tissue have been reported [1-3], though they are considerably less frequent than secondary meningiomas [4,5]. A tumor must fulfil several criteria to be considered of radiation-induced etiology [2,6,7]: It must develop in a previous radiation field with an interval of several years between the tumors clinical presentation and the original radiotherapy; and be confirmed as histologically distinct from the tumor for which the radiotherapy was originally given.

The goal of this article is to emphasize with the case of a severe adverse effect after radiotherapy for Hodgkin's lymphoma the needs of Improvement in our knowledge on genetic susceptibilities which may help define more personalized primary therapies that weigh treatment efficacy with the risk of late complications.
Case Report

A medullary glioblastoma was diagnosed in a 45-year-old Caucasian male. Relevant medical history included a melanoma in situ on the skin of the back treated by excision, and a stage IIISB nodular sclerosing Hodgkin’s lymphoma (Ann Arbor classification), both presenting at the age of 35. No relevant family history of malignancy was noticed.

History of the Hodgkin’s lymphoma

Six months after his melanoma had been treated, the patient complained of enlarged glands in the region of the right clavicle, associated with sweats. A node of the left transverse cervical chain enlarged to 3 cm, was entirely filling the left supraclavicular fossa. Open biopsy and frozen section of the lesion suggested a nodular sclerosing Hodgkin’s Lymphoma (HL). The full work-up confirmed the HL, stage IIIB of Ann Arbor classification with bilateral subclavicular, mediastinal involvement, and splenomegaly. The patient responded only partially to the current protocol of chemotherapy ABVD (Adriamycin-bleomycin-vinblastine-dacarbazine), so a course of vindesin-doxorubicin-carmustine-etoposide-methylprednisolone then followed. The chemotherapy was supplemented by radiotherapy over ten weeks. X-ray photons (15 MV) from a linear accelerator were administered in two courses of treatment: to a supra-diaphragmatic mantle field encompassing the mediastinum, and the subclavicular and axillary nodes on each side, receiving 40 Gy in 20 fractions administered daily over four weeks; and a sub-diaphragmatic block field that included the aortic, coeliac and splenic nodes, receiving 30 Gy in 15 fractions over six weeks. The maximum dose received by the spinal cord was 41 Gy. The treatment was reasonably well tolerated, apart from some episodes of nausea and epigastric pain. This treatment resulted in complete remission.

History of the Glioblastoma

Nine years after radiotherapy, the patient presented with progressive difficulty in walking and urinary urgency. On examination, the patient was found to be ataxic with paraparesis. There was a motor deficit of the dorsiflexors of the feet, absent patellar reflexes, and loss of sensation to both touch and point discrimination anteriorly on both the thigh and leg bilaterally.

Lumbar puncture and analysis of the CSF revealed high protein (1.85 g/L), significant lymphocytosis with no B-lymphocytes, normal LDH, and normal immuno-electrophoresis; an lumbar and sacral MRI showed an inflammatory hypersignal with increased cord volume as high as T8/9 with a 5 cm area of multiple homogenous contrast uptake in the cord and some nerve roots (Figure 1).

Figure 1: MRI of lower thorax and lumbar region. a) Gadolinium-enhanced T1 sagittal slices show contrast uptake in a multi-lobed tumor of the conus medullaris with spontaneous T1 hypersignal of the vertebral bodies secondary to previous radiotherapy. b) FLAIR T2-weighted sagittal slices showing diffusely enlarged conus medullaris extending throughout the lower part of the cord. c) T1 transverse slice with axis centered on the lesion (Magnevist).

Positron-emission tomography with 18-fluorodeoxyglucose (18-FDG) showed cord involvement at T11-T12 with bony involvement of T10.

A decompressive and diagnostic surgical procedure was performed to exclude a relapse of the lymphoma. The neurological symptoms showed no improvement on high dose corticosteroids. Pathology reported a high grade astrocytoma of the spinal glial cells of glioblastoma type. Immuno-markers showed a heterogenous expression of GFAP, absent CD20 B-lymphocytes; some CD3 marking of reactive T-lymphocytes, p53 expression in 15% of cell nuclei, and no expression of EGF-receptors or to a modified anti-IDH1 antibody. The proportion of proliferating cells expressing MIB-1 was 15% (Figure 2a and 2b).
As the patient had already received 41 Gy to his spinal cord, the maximum dose being 45 Gy, further radiotherapy was contraindicated. Despite several courses of treatment including first line chemotherapy with monthly temozolomide alone, second line treatment with irinotecan and bevacizumab, third line with fotemustine, and finally a trial of carboplatin and etoposide (VP16), the tumor progressed, both locally and metastasizing to the brain, causing the patient’s death 22 months after diagnosis.

Discussion

Secondary malignancy following curative treatment for a Hodgkin’s lymphoma is well-recognized in the literature [8,9]. We have found nine cases of spinal cord glioblastomas which were possibly secondary to radiotherapy [3,7,10-16], of which the radiotherapy had been indicated for lymphatic disease in five [7,10,11,13,14]. All patients were male, aged less than 50 years old [7,10,11,13,14] with a presentation of lymphatic disease in nine cases reported in the world literature [3,7,10-16] with a poor prognosis of around six months [27]. Taking best known practice in cerebral glioblastoma as a guide, the optimum treatment would seem to be complete macroscopic surgical resection followed by a combination of radiotherapy and chemotherapy [28]. However, this course is rarely feasible given the range of possible therapeutic choices, such a decision would generally be reached in consultation with a multidisciplinary team. Radiotherapy should be avoided in malignancy associated with p53 mutations or the Li-Fraumeni syndrome.

Radiation-induced glioblastoma of the spinal cord remains a rare condition with nine cases reported in the world literature [3,7,10-16] and the threshold dose to the spinal cord likely to cause a secondary tumor, has not, to our knowledge, been established. In the six cases reported of giall tumors arising in the irradiated field following treatment for Hodgkin’s lymphoma, significant doses were received by the spinal cord, of between 37.8 and 49.69 Gy (Table 1). The radiation-induced cerebral glioblastomas were more likely to occur in the physical penumbra at the margins of the treatment field than at the center, where the maximum dose had been received [3,20].

Distant metastasis arising from spinal cord glioblastomas has already been reported [7,21-27], with a poor prognosis of around six months [27]. Taking best known practice in cerebral glioblastoma as a guide, the optimum treatment would seem to be complete macroscopic surgical resection followed by a combination of radiotherapy and chemotherapy [28]. However, this course is rarely feasible given the previous radiotherapy, the unresectable nature of the tumor, and the patient’s poor general condition. Further administration of radiotherapy is usually compromised by the dose already received by the spinal cord, at risk for radiation myelitis, though three centers have nevertheless attempted additional radiotherapy [7,13,14]. Where surgery seems contraindicated, stereotactic radiotherapy may be contemplated, with the goal of effective decompression of the cord and pain relief. Palliative chemotherapy often seems the only therapeutic option. We were unable to find any report of a patient with a radiation-induced spinal cord glioblastoma surviving longer than 24 months. This grave prognosis seems impervious to all therapeutic manoeuvres attempted to date (Table 1).

In any case, a family history for genetic predisposition should be sought, and referral to the onco-genetic service may be advisable to inform judgment in the consideration of the risks and benefits of any therapeutic course proposed. Given the range of possible therapeutic choices, such a decision would generally be reached in consultation with a multidisciplinary team. Radiotherapy should be avoided in malignancy associated with p53 mutations or the Li-Fraumeni syndrome.

Figure 2: Spinal cord histology. a) HES (x20): Astrocytic fibrillary and gemistocytic tumor cells with marked nuclear pleomorphism. b) GFAP (x20): cytoplasmic immunostaining of tumor cells.
| Number of cases | Sex | Initial disease | Age at first radiotherapy (years) | Radiation dose (Gy) | Maximum dose to spinal cord (Gy) | Radiation field | Interval between presentations (years) | Glial tumor | Tumor site | Follow-up duration | Secondary tumor treatments |
|----------------|-----|-----------------|---------------------------------|-------------------|---------------------------------|----------------|----------------------------------------|-------------|------------|-----------------|-----------------------------|
| 1              | Male| Hodgkin's lymphoma | 21                              | 50/30             | 49.69                           | Mantle +cervical +para-aortic | 6                        | 6 | Cervico-thoracic C4-T2 | 10 weeks                   | Laminectomy               |
| 1              | Male| Hodgkin's lymphoma | 19                              | 40                | Not reported                     | Mantle, Waldeyer' s ring, para-aortic, splenic | 7                        | 7 | Astrocytoma Grade II-III | Cervical C6 | 6 months | Laminectomy, chemotherapy |
| 1              | Male| Hodgkin's lymphoma | 30                              | 40                | Not reported                     | Mediastinal         | 9                        | 9 | Anaplastic glioma | Cervico-thoracic C6-T2 | 11 months | Laminectomy, etoposide, cisplatin, fotemustine +radiotherapy (54 Gy) |
| 1              | Male| Hodgkin's lymphoma | 26                              | 30.6              | Not reported                     | Residual lymphnodeopathy in the anterior mediastinum | 3                        | 3 | Cervico-thoracic stoma | Thoracic | A few weeks | Radiotherapy (50 Gy) +temozolomide |
| 1              | Male| Lymphoblastic T-cell lymphoma | 21                              | 30                | 37.8                            | Residual lymphnodeopathy in the anterior mediastinum | 9                        | 9 | Cervico-thoracic C6-T6 with craniospinal dissemination | Cervico-thoracic C6-T6 with craniospinal dissemination | 9 months | Cranio-spinal radiotherapy (30 Gy)+peri-ventricular boost (30 Gy) +cervico-thoracic boost (22.5 Gy) +simultaneous temozolomide plus maintenance |
| 1              | Male| Hodgkin's lymphoma | 35                              | 40/30             | Not calculated and reported in the file | Mantle plus lumbar-aortic and splenic block | 9                        | 9 | Cervico-thoracic stoma | Terminal cone extending rostrally to T8/T9 then disseminating craniospinally | 24 months | Laminectomy, temozolomide, irinotecan +bevacizumab, fotemustine, carboplatine +etoposide |

Table 1: Malignant glial tumors of the spinal cord following radiotherapy for lymphoma.

Progress remains to be made on the evaluation of a patient’s radiosensitivity with the future goal of tailoring radiation doses to the patient’s genetic characteristics. Identification of genetic susceptibilities for specific treatment-associated late effect is a very promising future approach. Improvement in our knowledge on genetic susceptibilities may help define more personalized primary therapies that weigh treatment efficacy with the risk of late complications. For this patient, another treatment may have been chosen if physicians knew he had a specific radio-sensitivity. Tumors arising in irradiated fields must be histologically distinguished from a late relapse of the original malignancy. A radiation-induced spinal cord glioblastoma following Hodgkin’s lymphoma remains an exceptionally rare occurrence. To date, because of the very small number of cases, no standard therapy can be rationally suggested. The prognosis remains grim with- or without cerebro-spinal metastasis, and no treatment has been shown to be effective.

Compliance with Ethical Standards

Informed consent

Written informed consent was obtained from the patient's wife for publication of this report and the accompanying images.

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Declaration of interested conflicts

The authors declare no interested conflicts with this article.
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