PILOCYTIC ASTROCYTOMA FOLLOWING RADIOTHERAPY FOR CRANIOPHARYNGIOMA

CASE REPORT

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ABSTRACT - Administration of fractionated doses of irradiation is part of the adjutant therapy for CNS tumours such as craniopharyngiomas and pituitary adenomas. It can maximise cure rates or expand symptom-free period. Among the adverse effects of radiotherapy, the induction of a new tumour within the irradiated field has been frequently described. The precise clinical features that correlate irradiation and oncogenesis are not completely defined, but some authors have suggested that tumors are radiation induced when they are histologically different from the treated ones, arise in greater frequency in irradiated patients than among normal population and tend to occur in younger people with an unusual aggressiveness. In this article, we report a case of a papillary astrocytoma arising in a rather unusual latency period following radiotherapy for craniopharyngioma.

KEY WORDS: pilocytic astrocytoma, radiotherapy adverse-effects, oncogenesis.

Astrocitoma pilocítico pós-radioterapia para tratamento de craniofaringioma: relato de caso

RESUMO - A administração de doses fracionadas de radiação é parte do tratamento adjuvante para tumores do sistema nervoso central como craniofaringiomas ou adenomas hipofisários. A radioterapia pode aumentar as chances de cura ou expandir o período livre de sintomas. Dentre seus efeitos adversos, é conhecida a possibilidade de aparecimento de novo tumor na região irradiada. Os critérios que definem se um novo tumor é de fato induzido por radiação ainda não são completamente definidos. Admite-se, no entanto, que se o novo tumor é histologicamente diferente do primeiro, aparece em indivíduos mais jovens e é mais agressivo que o habitual, provavelmente sua origem é devida à oncogênese da radiação. Descrevemos um caso de astrocitoma pilocítico possivelmente induzido por radioterapia, com surgiimento em tempo de latência inesperadamente curto.

PALAVRAS-CHAVE: astrocitoma pilocítico, efeitos adversos de radioterapia, oncogênese.

Conventional radiation therapy is commonly applied as adjutant treatment for central nervous system (CNS) tumours. Lesions that continue to growth despite surgical ablation or are not amenable to complete resection tend to recur lately when submitted to irradiation. Among benign CNS tumours, pituitary adenomas and craniopharyngiomas are those for which standard fractionated radiation or stereotatic radiosurgery is usually performed. For these lesions adjunctive radiation has a clear-cut positive effect maximising cure rates and adjourning the reappearance of symptoms. Radiation therapy can produce side effects such as tissue damage and neoplastic transformation in the irradiated field. Tumour development after radiotherapy is constantly described as an unusual but invariably serious medium term complication of irradiation. For patients undergoing sellar irradiation, gliomas are the most frequent tumours, usually presented with marked aggressive behaviour and young age of onset.


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In this article we report a case of radiation induced pilocytic astrocytoma with clinical aggressive behaviour developed in a rather unusual period (6 months) after radiotherapy for a craniopharyngioma.

CASE

A 36-year-old previously healthy woman was referred in 1996 to a neurosurgical consult by her endocrinologist with a 18-month history of symptoms of panhypopituitarism and amenorrhoea. A skull roentgen showed a normal sellar region, while an MRI depicted a cystic suprasellar lesion. In July of 1996, the patient underwent a right temporo-fronto-sphenoidal craniotomy with complete resection of the tumour. A histological examination of the specimen showed a suprasellar papillary squamous epithelium (Fig 1). The definitive diagnosis was papillary craniopharyngioma due to the complexity of the epithelium and the absence of skin adnexae and keratin, expected in dermoid and epidermoid cysts. One month after surgery the patient started radiation therapy with 5860cGy, with daily fractions of 180cGy. Six months after the beginning of radiotherapy, the patient presented a progressive bitemporal hemianopia and a new MRI presented a 1.0 x 0.9 cm supra sellar lesion infiltrating the optic chiasm. In June of 1997 the patient underwent a new surgical procedure with partial tumour resection. Histological analysis showed a lesion with conspicuous elongated and citologically bland cells intermixed with Rosenthal fibers (Fig 2). Due to its marked cellularity and the presence of the Rosenthal fibers, pylocitic astrocytoma was the definitive diagnosis, opposed to gliosis. The patient quickly developed complete blindness and diabetes insipidus after surgery and died 5 months later, reinforcing the diagnosis of a second tumour.

DISCUSSION

Tissue damage in the irradiated field yield side effects that in the brain and CNS may include optic neuritis, vasculopathy, cognitive and psychological disturbances and hypothalamic insufficiency. Moreover, due to base pair disruption and deoxyribonucleic acid (DNA) strand alteration, irradiation can produce neoplastic transformation. Tumour development within the irradiated field constitutes a known consequence with very detrimental outcome.

The most common intracranial nonglial CNS tumours induced by radiation are meningiomas and sarcomas. After sellar irradiation gliomas are the most frequent glial tumours. For patients undergoing sellar irradiation, the relative risk for developing a glioma is approximately 8 bigger.

Fig 1. Cystic craniopharyngioma: rounded complex squamous epithelium with loose center. H&E., 400x.
than in the normal. Tumour development after radiotherapy is considered radiation induced when some criteria are met. These include: the new lesion arises within the field of radiation; it is histologically different from the first tumour; it appears after a latency period that reasonably permits a secondary tumour development and there are not other conditions predisposing tumorigenesis.

Tumours evolving after sellar irradiation are generally reported by sporadical case descriptions. Despite the relative abundance of tumour neogenesis examples, the appearance of a second and unrelated CNS neoplasm after brain irradiation for a primary lesion is not the rule. However, the increased risk of developing a new tumour after irradiation and the peculiar settings presented by these second lesions argue against a simple coincident association. In a cohort study performed in 1992 with 334 patients undergoing 4500 cGy sellar irradiation for treatment of pituitary lesions, the relative risk of a new tumour development was significantly increased (7.92 times) compared to the one bore by the normal non-irradiated population. Large series have shown that skull low dose irradiation (350-425cGy) for pediatric treatment of Tinea capitis increases the relative risk for developing a brain tumour by 2.6 times. Likewise, malignant CNS tumours have been witnessed to evolve in animal models submitted to brain irradiation with proton or x-ray in a much higher frequency than in non-irradiated animals.

The use of fractionated doses of radiotherapy associated with spatial strategies to avoid excessive exposure of single brain territories or paths have been applied in order to minimise side effects and neoplastic transformation originated by radiation. It is not yet known whether a clear-cut dose response curve may be used regarding new tumour developing, but it seems that fractionated radiation is responsible to lessen the appearance of side effects. The use of 50 to 55 cGy in 25 sessions is capable of controlling primary tumour growth with a somewhat small level of complications.

Tumours appearing after irradiation usually differ from their “spontaneous” counterparts mainly because of the age of onset and aggressiveness observed. The average range for glioma appearance in the normal population is the sixth decade. Gliomas appearing before or in the lower extreme of this range are generally expected to have a benign course.
Comprehensive analysis of new cases of glioma after radiation for craniopharyngioma depict that its average age of onset is roughly 27 yr., much younger than the age of onset of glial tumours in the normal population or even after radiotherapy for pituitary adenomas, what is explained by the younger presentation of craniopharyngiomas. Opposed to what should be expected in young patients also shows a rather different histological behaviour, with increased frequency of more aggressive patterns, comprising mainly grade III and IV gliomas.

We observed that in 8 cases of gliomas after radiotherapy for craniopharyngiomas (Table), the average latency period between irradiation and symptoms due to new tumour was 12.1 yr., with the smallest latency period reported of 5 years.

**Conclusion**

In this article we report an unusual case with unfavourable clinical outcome of a patient with craniopharyngioma who underwent surgical and irradiation treatment. As confirmed by post-surgical histological analysis, there was not recurrence of the first tumour but appearance of what could have been considered post-surgical gliosis or another tumour. Since the patient was submitted to radiation therapy and had a fast unfavourable clinical course, we accepted that it was indeed a totally new lesion with very aggressive behaviour. We also considered that the histological feature of the second lesion was apparently benign, corresponding to a low-grade glioma. However, due to the rapid clinical deterioration, we considered it to be in fact a dynamic and rapid changing tumour, evolving quickly to aggressiveness. This hypothesis, unfortunately, could not have been confirmed by autopsy, which was not desired by the patient’s relatives. Nonetheless, the data presented above strong suggests that the new lesion had a very aggressive and rapid changing status, based upon clinical settings.

The new tumour reported fulfils every variable suggested by the criteria above in order to be considered induced by radiation. However, it arised in a rather unusual latency period after radiotherapy (6 months). We therefore want to draw attention to irradiated induced tumours that may be formed in different settings, considering that it could have been originated by radiation, despite its rapid appearance.

**REFERENCES**


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**Table. Review of case reports of glioma after radiotherapy for craniopharyngiomas:**

<table>
<thead>
<tr>
<th>Reference</th>
<th>Age of diagnosis</th>
<th>Radiation dose</th>
<th>Latency period</th>
<th>Glioma type (WHO*)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ushio et al. (1997)</td>
<td>6 years</td>
<td>5460 cGy</td>
<td>5 years</td>
<td>IV</td>
</tr>
<tr>
<td>Sogg et al. (1978)</td>
<td>9 years</td>
<td>6007 cGy</td>
<td>6 years</td>
<td>IV</td>
</tr>
<tr>
<td>Komaki et al. (1977)</td>
<td>22 years</td>
<td>5400 cGy</td>
<td>6 years</td>
<td>IV</td>
</tr>
<tr>
<td>Gutjahr et al. (1979)</td>
<td>4 years</td>
<td>6000 cGy</td>
<td>8 years</td>
<td>IV</td>
</tr>
<tr>
<td>Maat-Schieman et al. (1985)</td>
<td>5 years</td>
<td>6000 cGy</td>
<td>14 years</td>
<td>III</td>
</tr>
<tr>
<td>Shimizu et al. (1994)</td>
<td>20 years</td>
<td>6000 cGy</td>
<td>16 years</td>
<td>III</td>
</tr>
<tr>
<td>Kitanaka et al. (1989)</td>
<td>7 years</td>
<td>6000 cGy</td>
<td>17 years</td>
<td>III</td>
</tr>
<tr>
<td>Liwnicz et al. (1985)</td>
<td>11 years</td>
<td>5900 cGy</td>
<td>25 years</td>
<td>IV</td>
</tr>
</tbody>
</table>

*WHO, World Health Organization.