Pituitary adenomas are known to have a propensity to invade surrounding structures despite their benign histological appearance. Aggressive behavior by such tumors, defined by rapid growth, radiological or intraoperative evidence of gross invasion of parasellar structures, and/or histopathological evidence of dural invasion, may preclude complete surgical resection. Radiotherapy has been used to treat both invasive pituitary tumors and incompletely resected or recurrent tumors. Subsequent development of a sellar or parasellar sarcoma is a rare event.\textsuperscript{1-15} In this report we describe a case of sarcomatous change within a locally aggressive growth hormone (GH)-secreting adenoma four years after irradiation of the sellar region. The patient also received hormonal treatment with pegvisomant, an inactive analogue of GH and antagonist of its receptor.

CLINICAL HISTORY

The patient’s symptoms began at age 11 when he began to grow faster than his identical twin brother. Over the next several years he gained substantially in height. In 1995 at age 20, he was 6 feet 8 inches tall compared to his twin who was 5 feet 10 inches tall. On presentation at that time, he had the classical features of acromegaly with a broad forehead and nose, prominent chin and brow, and large hands and feet. He denied any visual symptoms. His initial magnetic resonance imaging (MRI) showed a large sellar and suprasellar enhancing mass with involvement of both cavernous sinuses, with invasion of the right foramen ovale, and extension into the infratemporal fossa (Figure 1). His serum GH level was 1033 ng/mL (normal, < 2.3 ng/mL). After three weeks of treatment with somatostatin analogue (octreotide), the GH level had fallen to 126 ng/mL. The level of insulin-like growth factor I (IGF-I) fell from 9060 ng/mL (normal, 114-492 ng/mL) before treatment to 850 ng/mL during the same interval. Repeat MRI six months after starting octreotide showed a 20% reduction in tumor size. One year after the clinical diagnosis was made and medical treatment initiated, the patient underwent a right transfacial transmaxillary approach to the sella, sphenoid sinus, and infratemporal fossa with a substantial but subtotal removal of tumor. Although a full panel of anti-hormonal primary antibodies was used, this tumor was positive only for GH. Prior to surgery, his levels of GH (54 ng/mL) and IGF-I (720 ng/mL) were both elevated although lower than at the onset of octreotide therapy. He made a good postoperative recovery and his IGF-I levels decreased further, to 250 ng/mL. He continued on octreotide postoperatively.

Because of the residual mass and the aggressive nature of the tumor, he next underwent radiation treatment to a field covering the sphenoid sinus, clivus, both cavernous sinuses, both posterior maxillary sinuses, and the floor of each middle fossa to the foramen ovale, with a 1-cm margin. A total of 50 Gy was given at 2 Gy fractions over five weeks in late 1996 using lateral opposing portals with high-energy photons. Prior to radiotherapy his GH levels had risen again to > 1000 ng/mL. During this time, his thyroid function tests and testosterone levels were low and the patient was started on supplemental thyrotropin and testosterone.

Because of persistent clinical symptoms of acromegaly, the patient was taken off octreotide and started on pegvisomant three years after his first operation. His symptoms and clinical features of acromegaly improved thereafter, but his GH levels never normalized (Figure 2). He stopped taking pegvisomant a year later after supplies of this experimental drug ran short, and resumed octreotide injections.

Five years after completing radiotherapy, the patient presented with blurred vision and worsening headaches. On examination he still had the stigmata of acromegaly along with a partial left abducens palsy. Magnetic resonance imaging showed tumor regrowth mainly within the sella and suprasellar region and infiltrating the left cavernous sinus (Figure 3). He underwent a transsphenoidal resection of this recurrent tumor in November 2001. The tumor was much firmer than the soft tumor removed during his first surgery, was grossly invasive through dura, and was excised with some difficulty from the sella and from the suprasellar cistern.

After surgery the patient completed five cycles of treatment with adriamycin and ifosfamide over seven months. His symptoms markedly improved and follow-up MRI has shown a significant reduction in tumor size (Figure 4). However, dose-limiting toxicity from the chemotherapy forced its cessation and the tumor then grew into the adjacent brainstem, leading to the patient’s demise 11 months after diagnosis of the sarcoma.
MATERIALS AND METHODS

Tumor specimens were fixed in 10% neutral buffered formalin followed by paraffin embedding and sectioning for H&E staining. Immunohistochemistry was performed using commercially available anti-human antibodies for growth hormone (Zymed), chromogranin (Zymed), vimentin (clone V9-1, Dako), smooth muscle actin (clone 1A4, Sigma), desmin (clone D33, Dako), CD34 (clone MY10, Becton-Dickinson), and S-100 (rabbit polyclonal 15E2E2, Ventana Labs). Staining for the IGF-I receptor was performed using a rabbit polyclonal antibody directed towards the beta subunit (Santa Cruz Biotechnology, Santa Cruz, CA). Following antigen retrieval (microwaving in citrate buffer) this primary antibody was used at a dilution of 1:200. Appropriate HRP-conjugated secondary antibodies (Vector Labs) were applied according the manufacturer’s instructions and a diaminobenzidine substrate was used as a chromagen.

RESULTS

Histology of the patient’s initial lesion revealed a typical pituitary adenoma, containing cells with uniform round nuclei and “salt-and-pepper” chromatin arranged in a trabecular pattern (Figure 5A). Immunohistochemistry for GH revealed positivity in the cytoplasm of most tumor cells (Figure 5B). In contrast, the specimen from the patient’s second procedure showed a biphasic tumor, with a predominant spindle cell component consistent with sarcoma. The spindle cells demonstrated significant nuclear atypia and a mitotic count of 22 per 10 high power fields (400x), features diagnostic of high-grade sarcoma. Islands of neuroendocrine cells with features similar to the patient’s original lesion were also identified (Figure 5C). Reticulin and GH staining were performed on adjacent sections and showed positive reticulin staining in only the spindled areas (Figure 5D), and positive GH staining in only the neuroendocrine tumor cells (Figure 5E). A pattern similar to the GH staining was seen using antibody against chromogranin (not shown). In order to characterize the sarcomatous component, staining for vimentin, smooth muscle actin, desmin, CD34, and S-100 was performed, revealing positivity only for vimentin. The overall histological appearance and staining profile were considered most consistent with fibrosarcoma. In addition, strong cytoplasmic staining within sarcoma cells was demonstrated for the IGF-I receptor (Figure 5F).
Figure 5: Histology and immunohistochemistry of pituitary lesion. (A) Representative area of initial pituitary adenoma with a benign appearance (hematoxylin and eosin, X200). (B) Immunohistochemistry of the original tumor using a specific anti-GH antibody (X400). (C) Representative section of patient’s second tumor showing spindle cells consistent with undifferentiated sarcoma (X200). Staining of the same area of tumor from adjacent sections (X400) showing (D) reticulin and (E) GH. Staining for (F) IGF-I receptor (X400) reveals positive cytoplasmic staining in tumor cells and suggests interaction between the patient’s acromegaly and growth of the sarcoma.
DISCUSSION

 Postsurgical treatments for persistent acromegaly include radiotherapy, dopamine agonists (bromocriptine and cabergoline), and somatostatin analogues (octreotide, long-acting octreotide, and lanreotide). More recently pegvisomant, a growth hormone receptor antagonist, has also been used in clinical trials to lower significantly circulating levels of IGF-I, the principal mediator of GH action.16-18 The patient reported here was exposed to irradiation and to both octreotide and pegvisomant. As neither of those drugs is known to stimulate malignant transformation, radiotherapy probably induced this fibrosarcoma.

 Although spontaneous sarcomas of the parasellar region are occasionally seen including cases of osteogenic sarcoma,19 alveolar soft-part sarcoma,20 rhabdomyosarcoma,21 and chondrosarcoma,22 radiation-induced sarcomas are rare within the sella. Terry, et al13 were the first to document the occurrence of these tumors following radiotherapy for pituitary adenomas. Two nonfunctional adenomas have been described following radiotherapy of pre-existing pituitary adenomas.1-5 Although 60% have arisen from clinically nonfunctional adenomas, immunohistochemical analysis is lacking in most of these cases. Another 30% have been reported in patients with clinical acromegaly but only one case other than ours has been shown to stain positive for GH.2 Two fibrosarcomas have also been described which occurred spontaneously in the sella,23,24 and another after treatment of a prolactinoma with bromocriptine with no history of sellar irradiation.25

 The time from onset of symptoms to the diagnosis of postirradiation sarcoma varies from two to 20 years.11,26 Several types of radiotherapy including orthovoltage radiation and more recently cobalt-60, betatron, linear accelerator therapy, and heavy particle cyclotron radiotherapy have been associated with the development of fibrosarcomas.27,28 The sarcomas described are mostly of the spindle cell variety but may show mixed patterns of differentiation: recently an osteosarcoma of the pituitary was described in a pre-existing fibrosarcoma.4 Single-fraction stereotactic delivery of doses up to 35 Gy has recently been used to treat newly diagnosed and residual pituitary tumors,27 and at least one case of sarcoma following stereotactic radiotherapy (in a vestibular schwannoma) has been reported.29 However, long-term follow-up of these patients is lacking and the true incidence of late-onset sarcomas cannot yet be stated with that modality.

 The cellular origin of this relatively undifferentiated sarcoma cannot be determined with certainty. Although osteosarcoma has been induced in parasellar bone,29 this tumor’s identity as a fibrosarcoma might suggest that it arose from the dura lining the sella, or from the pituitary gland itself. However, the interlacing of the sarcomatous and adenomatous components implies that the sarcomatous component was derived from the pre-existing pituitary adenoma. Sarcomatous change has also been reported in other irradiated intracranial tumors, including gliomas.30

 This patient’s sarcoma was strongly positive for the IGF-I receptor (Figure 5F). If these receptors are functional, then his ongoing GH and/or IGF-I excess may have encouraged the growth of the malignancy. Others have reported sarcoma development in acromegalic patients. Wright’s31 series of 194 patients with acromegaly contains one with a co-existing fibrosarcoma, as does Nabarro’s32 series of 265 patients with GH excess. In addition, Barzilay et al33 reported one patient with an unclassified sarcoma among 87 with acromegaly. Thus, the incidence of sarcoma among these 537 cases of acromegaly was 0.56%, a relatively high number that supports an association between sarcomagenesis and excess circulating GH and/or IGF-I.

 IGF-I induces cell proliferation in many (but not all) sarcoma cell lines.34 When bound by the appropriate ligand, the IGF-I receptor activates ERK 1 and 2, STATs, and Akt, which govern the phenotypic characteristics of both normal and neoplastic cell lines.35 It also activates the downstream molecule p38, which rescues tumor cells from otherwise lethal DNA damage by modifying degradation of p53.36 This allows damaged cells to continue to proliferate, and may thereby promote malignant transformation as seen in our case.

 Following the diagnosis of fibrosarcoma the patient was started on ifosfamide and adriamycin with good results. There was, at first, symptomatic improvement along with reduction in tumor size (Figure 3A). However, after five cycles the patient developed dose-limiting toxicity and medical therapy had to be stopped. Although adjuvant chemotherapy is palliative rather than curative, it has occasionally been successful in treating radiation-induced sarcomas and offers an additional avenue for managing this difficult problem.36 However, the positive effects of such treatment may diminish in the face of ongoing IGF-I excess, a limitation that might be overcome by administering pegvisomant to such patients while they continue a chemotherapeutic regimen.

 This case highlights the possibility of developing a sarcoma after irradiation of a pituitary adenoma. As this event is disproportionately reported in acromegalic patients and, as the patient reported here had a sarcoma positive for IGF-I receptor, we speculate that overactivity of the GH/IGF-I axis encourages such sarcoma formation.

REFERENCES