CASE REPORT

Leiomyosarcoma of the oropharynx and neurogenic tumors in a young patient with Turner’s syndrome

ANNAROSARIA DE CHIARA, GAETANO APICE, GIUSTINO SILVESTRO, SIMONA LOSITO, GERARDO BOTTI, FRANCESCO IONNA, VINCENZO DE ROSA, ANNAMARIA BORGHESE* & VITO NINFO*

Istituto Nazionale dei Tumori di Napoli “G. Pascale”, Facoltà di Medicina e Chirurgia di Napoli Federico II*, Facoltà di Medicina e Chirurgia di Padova*

Abstract

Patient: A case of Turner’s syndrome developing a leiomyosarcoma of the oropharynx and metachronous neurogenic tumors (mediastinal ‘ganglioneuroblastoma intermixed’, subcutaneous neurilemoma) is described.

Discussion: To our knowledge, this case is the second reported leiomyosarcoma in a patient with Turner’s syndrome. Also the site of involvement (palate and oropharynx) is particularly unusual for the already rare leiomyosarcomas in the young age.

Key words: Turner’s syndrome, adolescents, neurogenic tumors, sarcoma, leiomyosarcoma, oropharynx.

Introduction

The occurrence of gonadal tumors in gonadal dysgenesis is a well known phenomenon.1–3 A higher incidence of extragonadal neoplasms with a preponderance of neurogenic tumors is also reported, suggesting that developmental neural defects, hamartomatous growths, and susceptibility to neurogenic tumors are related to one another as an expression of Turner’s phenotype.4 A few example of soft tissue sarcomas are also reported5–8 but to our knowledge only one leiomyosarcoma (LMS) has already been described in this setting.9 On the other hand, leiomyosarcoma is a rare tumor in children and adolescents.10,11 The visceral ones are often related to genetic12 or acquired immunodeficiency disorders,13–15 or immunosuppressive drugs16 and arise in the hepatobiliary, gastrointestinal, and tracheopulmonary systems. In most cases they were shown to be associated with Epstein-Barr virus (EBV).17

We describe a leiomyosarcoma occurring in a patient with Turner’s syndrome and metachronous neurogenic tumors (mediastinal ‘ganglioneuroblastoma intermixed’, subcutaneous neurilemoma). Also the site of involvement (palate and oropharynx) is particularly unusual for the already rare leiomyosarcomas at a young age.

Clinical history

The patient was diagnosed with Turner’s syndrome a few months after birth. For this reason, she had estrogen replacement therapy for the last 7 years. She underwent to thoracotomy because of a mediastinal ‘ganglioneuroblastoma intermixed’ (Shimada: schwannian stroma-rich) (Fig. 1) when she was 4-year-old. She was treated with radiotherapy and she was well and free of disease for 18 years.

Fig. 1. Mediastinal ‘ganglioneuroblastoma intermixed’. In a stromal-rich background, microscopic foci of differentiating neuroblasts and more or less mature ganglion cells are interspersed. (H&E ×10)
At the age of 22, she was referred to our hospital with a huge tumor, more than 10 cm. in largest diameter, infiltrating the whole soft palate, the ventral side of the posterior hard palate, the right tonsillar pillar and deeply into the parapharyngeal space and retro-molar trigone on the right side (Fig. 2). An incisional biopsy showed spindle and rounded cells with often vacuolar or clear cell change in the cytoplasm (Fig. 3), strongly positive to muscle specific actin (MSA) (Fig. 4) and to desmin (Fig. 5), with the diagnosis of epithelioid leiomyosarcoma G3. At the same time, two cutaneous nodules (left and right thigh) of 1.5 cm. in largest diameter and a subcutaneous lesion of 2 cm. (in the right axilla) were noted and excised. The cutaneous nodules proved to be sclerotic dermatofibromas with some large atypical mesenchymal cells; the subcutaneous one was a neurilemoma. Serological tests for HBV, HCV, HIV and EBV were negative.

The cytogenetic analysis performed on tissue from the neurilemoma by standard cytogenetic method showed the expected monosomy of chromosome X and chromosome aberrations with a complex karyotype (Fig. 6A). FISH analysis revealed that the most

![Fig. 2. CT scan sections through the oral cavity show a large mass in the oropharynx infiltrating soft palate with ventral invasion of the posterior edge of the hard palate.](image)

![Fig. 3. Epithelioid leiomyosarcoma of the oropharynx. Pleomorphic round to spindle cells with clear or vacuolar cytoplasm and with atypical mitoses are present beneath the oral mucosae. (H&E ×20)](image)

![Fig. 4. MSA is strongly positive in the neoplastic cells. (immunoperoxidase ×20)](image)

![Fig. 5. Desmin is strongly positive in the neoplastic cells. (immunoperoxidase ×20)](image)

![Fig. 6. A) G-banded karyotype of a cell from the neurilemoma shows multiple abnormalities and structural rearrangements such as del(1)(p), del(6)(q22), del(8)(p), der(12)dup(q13–22), loss of a chromosome 13, 14, 19 and X, other unknown rearrangements defined as markers which probably are derived from chromosome 3, 7, 8, 12 and 18. B) FISH with Chromophobe Multiprobe (Cy 3). Chromosome 6 hybridization: the small spot indicates a rearranged chromosome 6.](image)
frequent abnormalities were loss of chromosomes 13, 14 and 22, deletions [such as del(1)(p), del (6)(q22)] and rearrangements [such as der(12)dup(q13–22), t(7;9)(q11;q11), t(6;22)] (Fig. 6B). FISH revealed also the presence of part of the Y chromosome translocated on a chromosome of the D group in 10% of the metaphases.

Because of the site and size of the lesion, the patient was treated by chemotherapy (epirubicin 80 mg/m², ifosfamide 7.5 gr/m², DTIC 900 mg/m²), but after two courses she developed a 2.5 cm. right retromandibular lymph node metastases while the oropharyngeal mass was unmodified by CT scan. Then she went to a full course of radiation therapy. The Target Volume included the whole palate, the oropharynx and the lymph nodes of the left neck. A total dose of 66 Gy (2 Gy/fraction) was delivered in 64 days using two lateral portals with 6 MV photons to gross tumor volume (GTV) and bilateral neck lymph nodes, and two opposed antero-posterior fields for supraclavicular nodes for a total dose of 50 Gy. Radiation therapy was stopped on two occasions (a week each time) because of severe laryngeal mucositis. Seventy-five days from the end of radiotherapy, the MRI showed a significant reduction in the tumor mass with fairly good clinical improvement. But a month later, she developed local progression and then lung metastases. She died after a month from respiratory failure.

Discussion

Leiomyosarcoma is a rare tumor in children and adolescents. However, at least some cases reported as LMS, especially those diagnosed in neonates, may represent examples of congenital-infantile myofibromatosis with an excellent prognosis. Some authors also propose that some tumors present in the bronchopulmonary tree, particularly when in endobronchial location, are more often misdiagnosed as LMS while in fact are better regarded as primary bronchopulmonary fibrosarcoma. These histotypes have a relatively good outlook, representing the bronchopulmonary counterpart of the congenital-infantile myofibromatosis of the soft tissue. A recent review of 20 childhood LMSs shows that they have a relatively good prognosis in the majority of the cases and the occurrence of this neoplasm in the oral cavity is exceedingly rare, as already reported. While leiomyosarcomas account for about 1% of all soft tissue tumors in childhood (18, also personal experience of one of us, V.N.), they are the second most frequent malignancy in children with acquired immunodeficiency syndrome (AIDS). In this cohort they are often located in unusual sites especially visceral ones such as hepatobiliary, gastrointestinal, and tracheobronchial systems. Based on the reported cases, EBV seems to be involved in the pathogenesis of these tumors in immunosuppressed patients, while there is no evidence of its presence in the LMS in immunocompetent patients. Thus the oncogenesis of these neoplasms seems to be different in the two subgroups of patients.

Individuals with a history of childhood cancer have been estimated to have 10 to 20 times the lifetime risk of a second cancer compared with age-matched controls. Within the first 20 years after the initial diagnosis, the incidence appears to be on the order of 3% to 12%. In most cases, the second malignancy is related to ionizing radiation or aggressive chemotherapy, especially alkylating agents. The two main criteria to classify a sarcoma as radiation induced are: 1) the patient should have been irradiated for another malignancy at least 3 years prior to sarcoma diagnosis, and 2) the sarcoma should have developed within the field of radiation. In our case, the patient received radiation for a mediastinal tumor, but the oropharyngeal sarcoma she developed after 18 years was clearly outside of the prior radiation field. She had not received any chemotherapy for the ganglioneuroblastoma. Therefore, we concluded that the second tumor had no links to the therapy.

Alternatively, it is postulated that the occurrence of second cancers may be related to the presence either of an underlying systemic disease (such as von Recklinghausen’s neurofibromatosis) or of chromosomal abnormalities shared by multiple organs in which they are tumorigenic such as occurs in some congenital tumors or in Li-Fraumeni syndrome. Thus the simultaneous or sequential appearance of different tumors within an individual may be the clue to their shared genetic etiology. It appears that the majority of hereditary cancer predisposition is attributable to germline mutation in tumor suppressor genes now known to be critically important in growth control of both sporadic and hereditary tumors. Mapping and cloning of the retinoblastoma gene proved Knudson ‘two-hits’ hypothesis and elucidated the nature of tumor suppressor genes.

The increased risk for malignant transformation within dysgenetic gonads has been documented in numerous reports, most of which occurred in Turner’s syndrome with 45,X karyotype or in a phenotypic female with Y chromosome. The most frequent types of tumors in this setting are gonadoblastoma and dysgerminoma. Non gonadal malignancies are also reported with a preponderance of melanocytic nevi and peripheral neurogenic neoplasms, both neural crest derivatives, among children and young adults. In these cases, it has been postulated that these syndromes may have a mechanism of tumor susceptibility such as occurs in the neurocutaneous disorders, particularly neurofibromatosis type I, in which the NF1 gene is homozygously inactivated in both benign and malignant tumors related to it, and nevoid basal cell carcinoma syndrome (NBCC). So Turner’s syndrome may be added to those congenital disorders in which aberrant develop-
ment of neural-crest derivatives predisposes to neoplasia.\textsuperscript{4} We have to underline that our patient was affected by Turner's syndrome, cytogenetically diagnosed a few months after birth. She presented a mediastinal 'ganglioneuroblastoma intermixed' (Shimada: schwannian stroma-rich) at the age of 4 years, and later in life a subcutaneous neurilemoma in the right axilla and two sclerotic dermatofibromas in the left and the right thigh.

Some childhood solid tumors occur in close association with hereditary syndromes. For example, soft tissue sarcomas especially rhabdomyosarcoma (RMS), osteosarcoma, brain tumors are frequent in families affected by the Li-Fraumeni syndrome. Survivors of hereditary retinoblastoma, are prone to develop later in life osteosarcomas of bone and soft tissue sarcomas, mainly RMS and fibrosarcoma. In addition, children with neurofibromatosis type I are at increased risk to develop soft tissue malignancies such as malignant peripheral nerve sheath tumors (MPNST), RMS and angiosarcoma.\textsuperscript{22} Instead, the occurrence of sarcomas in Turner's syndrome is a very rare event with only few published cases.\textsuperscript{4–9} For the endometrial sarcomas, as already postulated for endometrial carcinomas,\textsuperscript{23,24} it has been suggested that the risk may be increased by estrogen replacement therapy or endogenous disorders that lead to unopposed estrogenic stimulation of the uterus.\textsuperscript{6,7} This mechanism has also been recalled for RMS of the uterus\textsuperscript{25} and oral cavity\textsuperscript{26} of patients without Turner's syndrome. Experimental studies showed that treatment with testosterone propionate (TP) and dyethylstilbestrol (DES) or TP and estradiol (E2) for 8–9 months causes RMS in vas deferens or uterus of Golden Syrian hamsters at a frequency of 100%. The authors hypothesize that treatment with TP and E2 causes a loss of the protective enzyme hGSTYBX leaving the cells vulnerable to the genotoxic effects of estrogen or estrogenic metabolites.\textsuperscript{27} LMS with clonal abnormalities display highly complex karyotypic changes and extensive heterogeneity, suggesting that the karyotypic profile is more dependent on site of origin than on microscopic features.\textsuperscript{28} In our case, the cytogenetic analysis was done on tissue from the subcutaneous neurilemoma that showed a very complex,...
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