

Multiple Metachronous Malignant Fibrous Histiocytomas of the Upper Limbs – a Case Report

Mnohopočetný metachrónny malígny fibrózny histiocytóm na horných končatinách – kazuistika

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Summary

Background: Soft-tissue sarcomas are rare tumors with the incidence of multiple metachronous or synchronous lesions in the extremities being even more uncommon. In effort to preserve the function of upper extremities, limb-salvage surgery became the treatment of choice for soft-tissue sarcomas. Subsequent adjuvant chemotherapy, as well as radiotherapy, is believed to decrease local recurrence rates, however, their effect on overall survival remains unclear.

Case: We report herein a case of symmetrical bilateral metachronous malignant fibrous histiocytomas of the shoulder. A 19-year-old patient presented with stiffness and pain in the right shoulder. The same symptoms developed 1.5 years later in the other shoulder. The culprit tumors are reported metachronous with regard to the succession in the onset of symptoms. Wide tumor resection was performed in both shoulders, and postoperative radiotherapy was then conducted. Chemotherapy was not indicated after the first surgery; whereas, in the second case it was the patient who refused the recommended adjuvant chemotherapy. **Conclusion:** The phenomenon of either metachronous or synchronous incidence of multiple soft tissue sarcomas is very rare and systematic reporting of every new case in the literature could contribute to further knowledge of tumor's unique behavior.

Key words

malignant fibrous histiocytoma – radiotherapy – upper extremity – neoplasms – multiple primary

Súhrn

Úvod: Sarkómy mäkkých tkanív sú zriedkavé nádory. Medzi nimi mnohopočetné metachrónne alebo synchronné sarkómy mäkkých tkanív končatín sú obzvlášť vzácne. Končatinu zachováajúca chirurgia je terapia voľby u sarkómov mäkkých tkanív, aby sa zachovala funkčnosť horných končatín. Po rozsiahlej resekcii tumoru môže adjuvantná terapia, teda chemoterapia a rádioterapia, znížiť počet lokálnych recidív, ale jej vplyv na celkové prežívanie pacientov zostáva nejasný. **Kazuistika:** V tejto kazuistike sme ukázali prípad mnohopočetných symetrických, metachronných malígnych fibróznych histiocytómov lokalizovaných na oboch ramenách. Prezentovaná je 19-ročná pacientka, ktorá pociťovala napätie a bolesť v pravom ramene. Rovnaké príznaky sa vyvinuli aj v ľavom ramene asi 1,5 roka neskôr. Nádory v ramenách u pacientky boli definované ako metachronné z dôvodu rôznych časových intervalov stanovenia diagnózy. Široká resekcia tumoru bolo vykonaná v oboch ramenách, následne bola aplikovaná pooperačná rádioterapia. Chemoterapia nebola doporučená po prvej operácii; avšak, pacientka odmietla adjuvantnú chemoterapiu po druhej operácii. **Záver:** Fenomén mnohopočetných sarkómov mäkkých tkanív, buď metachronných, alebo synchronných je veľmi zriedkavý. Preto sa domnievame, že každý nový prípad by mal byť opísaný.

Kľúčové slová

malígny fibrózny histiocytóm – rádioterapia – horná končatina – mnohopočetné primárne nádory

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Backgrounds

Soft-tissue sarcoma (STS) is a rare tumor with estimated incidence around 3.0–4.5 per 100,000 individuals [1]. STS accounts for only 1% of all malignancies in adult patients with 60% of STS tumors being located in the extremities. STS constitutes a heterogeneous group of neoplasms of mesenchymal origin, often with a distinct age distribution, site of presentation, natural biological behavior and prognosis [2,3]. The most common types of STS of the upper extremities are epithelioid sarcoma, synovial cell sarcoma, and malignant fibrous histiocytoma (MFH). MFH is a common histologic subtype of this uncommon clinical entity. The origin of the term “malignant fibrous histiocytoma” dates back to the early 1960s. Murray [4] observed that STS cell cultures were characterized by a storiform (i.e. cartwheel-like) growth pattern with pleomorphic and giant tumor cells that displayed amoeba-like movement and phagocytosis. These features were reminiscent of his-

tiocytes (i.e. macrophages of local resident tissue) and gave rise to the term “malignant fibrous histiocytoma” that soon became established in the literature. MFH was further subdivided into five types: 1. storiform-pleomorphic, 2. myxoid (myxofibrosarcoma), 3. giant cell (malignant giant cell tumor of soft parts), 4. inflammatory, and 5. angiomatoid MFH. The inflammatory subtype is the rarest variant constituting 5% of all cases. This tumor presents with a distinctive inflammatory infiltrate comprising neutrophils, lymphocytes, and foamy histiocytes. In 2002 World Health Organization reclassified the inflammatory MFH as “undifferentiated pleomorphic sarcoma with prominent inflammation” [4].

As for the association of STS with other malignancies, this was recorded in patients with genetic disorders such as neurofibromatosis, familial adenomatous polyposis, retinoblastoma, and Li-Fraumeni syndrome. Moreover, several studies have suggested that patients with STS are at an increased

risk of developing a second malignancy, notably breast and kidney cancer, regardless of any underlying genetic disorder present [5–7]. Most cases of STS in adult patients (94%) are not significantly related to common risk factors such as radiation, genetic disease, and chronic lymphedema [1]. Multiple metachronous and synchronous STS of the extremities are very rare. In the study of Murray et al [7], only 4 of 1,201 patients presented with symmetrical bilateral STS of the extremities. The authors encourage physicians in awareness of these sarcomas and their characteristics in order to avoid neglect, since these tumors are typically misdiagnosed and treatment is delayed [7]. Magnetic resonance imaging (MRI) is the imaging modality of choice for the initial staging, as well as further follow-up of sarcomas. Although imaging itself usually cannot reliably predict the histopathological diagnosis or distinguish benign from malignant processes, in combination with clinical history and epidemiologic features of various tumors, it can help to narrow the list of differential diagnoses. In patients with STS, MRI plays a crucial role in outlining the lesion and its relationship to adjacent anatomic structures [8].

The goal of treatment of musculoskeletal sarcomas is to optimize the oncologic outcome without impinging limb's functional status. Surgical resection remains the mainstay of therapy. In patients with musculoskeletal sarcomas of the extremities, limb-sparing resection was proved significantly superior to amputation. Furthermore, wide local excision of the tumor encompassing the muscular compartment, followed by adjuvant chemotherapy and radiation therapy was not associated with an increased risk of recurrence in these patients. Yet, the influence of this treatment on overall survival remains unclear [7,9].

Case

In October 2011, a 19-year-old female patient was referred to the Orthopaedic and Traumatology Clinic of Medical School, Comenius University and Slovak Medical University in Bratislava for

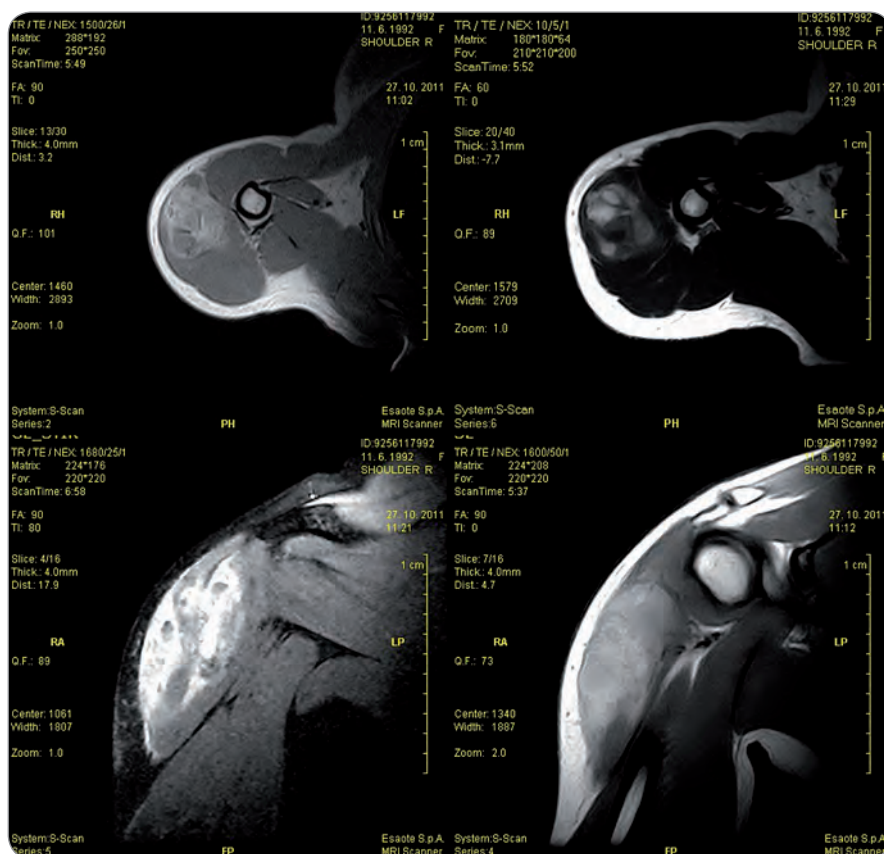


Fig. 1. Magnetic resonance image of right upper limb.

evaluation of stiffness and pain in the right lateral shoulder. She did not report any medical history of previous diseases or family history of genetic disorders. On 27 October 2011, MRI of the right upper limb revealed a well-circumscribed STS located in the deltoid muscle without any signs of shoulder joint involvement or local spread (Fig. 1). A preoperative lung computed tomography (CT) scan performed on 7 November 2011 did not show any lung metastases (Fig. 2).

Hence, wide resection of the tumor was performed on 24 November 2011. The pathology report described a 61-mm inflammatory MFH with 3-mm resection margins.

Adjuvant chemotherapy was not indicated; instead, postoperative radiotherapy was recommended. At the Department of Radiation Oncology of the National Cancer Institute in Bratislava, we conducted external beam radiotherapy on the lateral side of the right shoulder with a total dose of 40 Gy at 2 Gy per day by three fields' technique and a boost to the tumor bed in the second phase, with a total dose of 20 Gy at 2 Gy per day by two fields' technique. In both cases we used a multileaf collimator (80 leaves, 40 × 40) and 6-MV X-ray energy on a linear accelerator (Elekta Synergy; Elekta AB, Stockholm, Sweden). The radiotherapy was planned by co-registration of the preoperative MRI and postoperative planning CT. The patient was subsequently referred to her orthopedic surgeon for further follow-up.

In January 2013, the patient developed the same symptoms in the other shoulder and sought medical care again. On 23 January 2013, MRI of the left upper limb revealed a focal intramuscular lesion in the head of the deltoid muscle. The lesion had unclear boundaries, exhibited infiltrative growth, and measured $15 \times 10 \times 20$ mm. The muscular fascia was intact, and the bones showed normal structure without any focal changes (Fig. 3).

On 8 April 2013, 1.5 years after the first surgery, the patient underwent wide tumor resection on the left shoulder. Histological examination of the lesion revealed a 21-mm MFH with

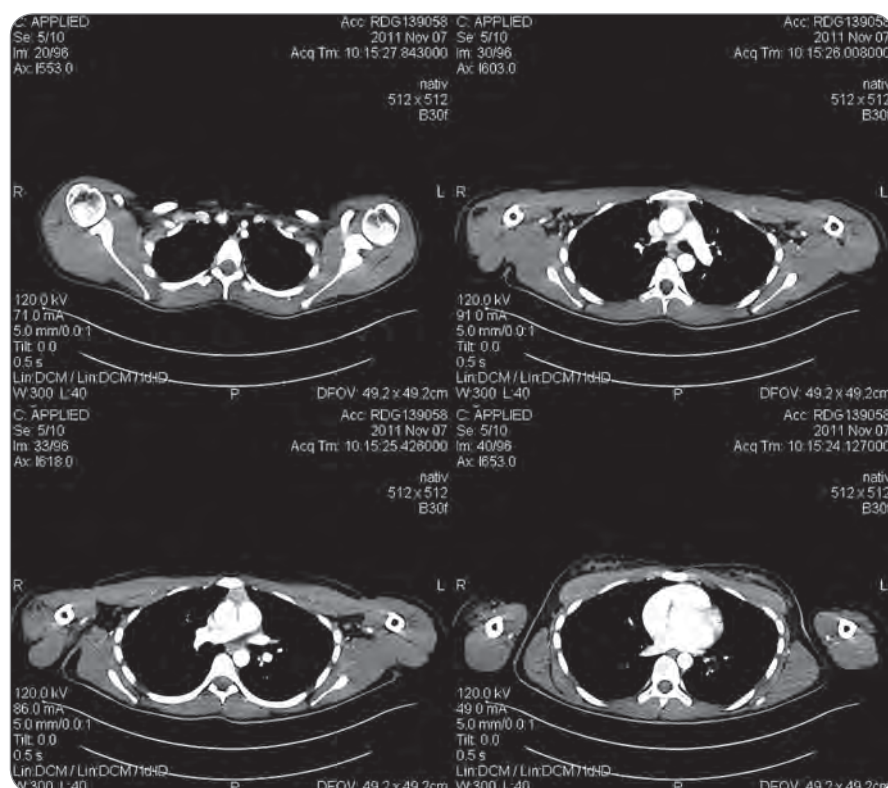


Fig. 2. Computed tomography scan of lung.

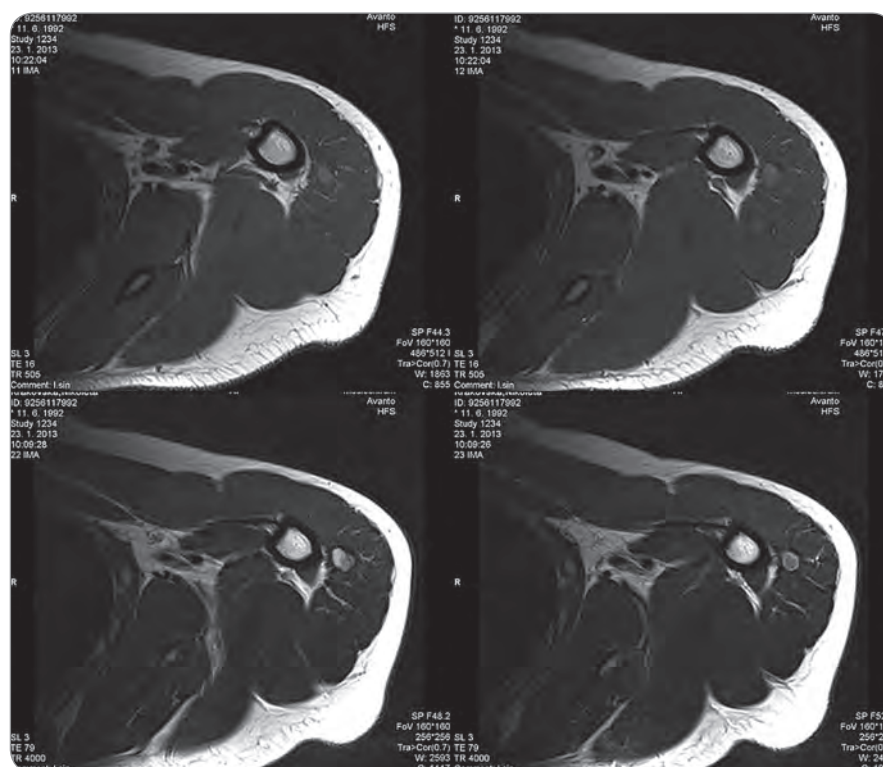


Fig. 3. Magnetic resonance image of left upper limb.

16-mm resection margins. The patient subsequently underwent postoperative radiotherapy on the left lateral shoulder.

Two phases of treatment were performed, involving a total dose of 60 Gy at 2 Gy per day with two fields' technique.

Again, we used a multileaf collimator (80 leaves, 40 × 40) and 6-MV X-ray energy on a linear accelerator (Elekta Synergy; Elekta AB) and planned the radiotherapy by co-registration of the preoperative MRI and postoperative planning CT. The patient refused to undergo adjuvant chemotherapy. She finished radiotherapy on 2 July 2013, and to date, she has exhibited neither signs of local recurrence, nor any distant metastases.

STS is a rare malignancy with an annual incidence of 3 per 100,000 individuals. Although the risk for patients with STS to develop a second malignancy is 12.5 times higher than in individuals with no history of the sarcoma, the occurrence of multiple primary STS in one patient is still uncommon [5,10]. There are many possible explanations for the appearance of multiple primary tumors in the same individual. It was proposed that these patients may have a genetic predisposition to malignant disease, or they may have been exposed to carcinogens [5]. Reports of multiple primary sarcomas unassociated with a predisposing syndrome (e.g. neurofibromatosis, familial adenomatous polyposis, retinoblastoma, or Li-Fraumeni syndrome) are rather anecdotal, particularly those referring to tumors localized in soft tissues of the extremities [6,11,12]. Daigeler et al [6] presented an interesting notion of the subsequent occurrence of a STS being an atypical manifestation of metastatic disease. With respect to the differentiation between a true synchronous, independently developed soft tissue sarcoma and its metastases, this might seem plausible,

especially if soft-tissue metastasis of a primary STS is quite a usual phenomenon (e.g. in cases of myxoid or round-cell liposarcoma) [7,13].

As a matter of fact, according to the new WHO classification from 2013, malignant fibrous histiocytoma was not accepted as a distinct histopathological unit and many of these tumors have now been reclassified into specific sarcoma subtypes. For a long time, malignant fibrous histiocytomas accounted for approximately 50% of sarcoma diagnoses. It took several years to discover that besides several subtypes of pleomorphic sarcomas (leiomyosarcomas, rhabdomyosarcomas, dedifferentiated liposarcoma etc.), this label included unrelated diagnoses such as lymphomas, malignant melanomas and sarcomatoid variants of carcinomas [14].

In this paper, we report a case of multiple metachronous MFH localized symmetrically in both shoulders. The tumors were defined as metachronous because their diagnosis and treatment occurred within a timespan of 1.5 years. This case is also unique because of the symmetrical development of STS bilaterally in the upper limbs. It is noteworthy that our patient had no familial disposition, no history of previous exposure to carcinogenic substances or irradiation.

Conclusion

The phenomenon of either metachronous or synchronous incidence of multiple soft tissue sarcomas is very rare and systematic reporting of every new case in the literature could contribute to further knowledge of tumor's unique behavior.

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