Soft tissue sarcomas of the upper extremities: Maximizing treatment opportunities and outcomes (Review)

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Abstract. Soft tissue sarcomas (STS) are rare tumors; they do not even equate to 1% of all malignant tumor cases. One-fifth of all STS occur in the upper extremities, where epithelioid sarcoma, synovial sarcoma, clear cell sarcoma and malignant fibrohistiocytoma are the most frequent subtypes. Surgical resection is the cornerstone of treatment. However, accomplishment of optimal oncological and functional results of STS of the upper extremities may represent a challenge for hand surgeons, due to the complex anatomy. In several cases, preoperative therapies are needed to facilitate tumor resection and improve the oncological outcome. Oligometastatic disease may also be a challenging scenario as curative strategies can be applied. Radiotherapy and chemotherapy are commonly used for this purpose albeit with conflicting evidence. Novel drug combinations have also been approved in the metastatic setting, further improving the quality of life and survival of eligible patients. Thus, prior to any approach, every case should be individually discussed in sarcoma centers with specialized multidisciplinary tumor boards. The aim of the present review was to gather the multidisciplinary experiences of the available therapeutic strategies for STS of the upper extremities.

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1. Introduction

Soft tissue sarcomas (STS) constitute a heterogenic group of tumors that accounts for only 1% of the overall human burden of malignant tumors, with an annual incidence of approximately 4.5/100,000 in Europe (1). Around 70 to 80% of patients are diagnosed in a local or locally advanced stage of the disease. The median age at diagnosis is 58 years, and the STS-related death around 65 years (2).

Extremities are the most frequent location for STS, accounting for 21.7% of all STS. However, a lower incidence (25-30% of STS of the extremities) and an earlier median age of diagnosis (38 years, ranging from 4 to 77 years) have been reported for the upper extremities compared to lower extremities’ STS (1). Around 50% of STS of upper extremities arise in the shoulder-upper arm region, 30 to 40% in the elbow-forearm and only a 10 to 20% in the wrist-hand (3,4). However, when the upper extremity is subdivided in proximal and distal, a distribution of 50% in each site has been observed (5) (Table I).

2. Histological examination

Over 50 different histological and molecular subtypes have been described, occurring ubiquitously throughout the human body (6). As every subtype of sarcoma has a particular biological behavior and response profile to systemic therapy, histologic diagnosis is a crucial criterion when selecting the appropriate therapy. Distribution of these subtypes varies between registries, due to evolution of classification of STS as a result of histological and molecular biology advances (6).

Potentially, all histologic subtypes can arise on the upper extremities, but a higher incidence of malignant histiocytoblastoma/undifferentiated pleomorphic sarcoma (UPS) and synovial sarcoma (SS) has been reported, reaching 50-65% (7).

Of note is that epithelioid sarcoma (ES) arises almost exclusively on the extremities, while clear cell sarcoma (CCS) is...
considered a specific subtype of the hand and the wrist (2). Histological, immunohistochemical and cytotumoral characteristics of the specific subtypes of STS of the upper extremities are described in Table II.

Accurate pathologic characterization of STS requires adequate and representative tumoral tissue, such as that harvested by core needle biopsy (CNB), which attains a specificity of 70 to 98% (8). The most accepted grading system for STS is the FNCLCC (Fédération Nationale des Centres de Lutte Contre le Cancer) system, based on three scores: Differentiation, mitotic count and necrosis. STS of the upper extremities are categorized as high grade in 45-70% of cases, due to the high score attributed to each of the four most frequent but aggressive histologic subtypes (3,9).

3. Diagnostic approach

Most STS of the hand and upper extremities present as a painless, slow growing and movable mass. In rare cases, the mass may cause nerve compression and present clinically as a nerve compression neuropathy. Thus, malignant lesions are painless, slow growing and moveable mass. In rare cases, the frequent but aggressive histologic subtypes (3,9). cases, due to the high score attributed to each of the four most frequent but aggressive histologic subtypes (3,9).

Magnetic resonance imaging (MRI) is the method of choice for the radiological evaluation of suspicious lumps, informing on the anatomical relations with the surrounding tissues for optimal surgical planning (14). Despite established imaging criteria, precise diagnosis can be made on the basis of MRI in only 24% of cases (15). Gadolinium contrast administration provides important information on tumor heterogeneity, guiding biopsy to the most vascular, non-necrotic part of the lesion (16).

A percutaneous core 14G-needle biopsy is frequently performed under local anesthesia. Through a skin stub 3-4 tissue cylinders are harvested. This procedure allows to harvest adequate tissue volume to make the diagnosis in over 90% of cases, with a sensitivity of 95% for malignancy and 88% for grade (17). Nevertheless, major diagnostic errors associated with the use of CNB can be drawn due to tumor heterogeneity, and low cellularity in cases of lipomatous, hemorrhagic or myxoid tumors. More specifically, in differentiated sarcomas, low-grade and high-grade components coexist in the same mass and a biopsy taken from the low-grade part may therefore result in understimation of the true tumor grade. In order to increase the harvesting of representative tissue, the careful consideration of the MRI features and the accomplishment of CNB under CT or U/S guidance are recommended.

When CNB is repeatedly non-diagnostic, an open biopsy should be performed, as it has a diagnostic accuracy of 94-100%. Open biopsies should be carefully planned and performed. The incision line should be part of the final surgical approach. We avoid transverse incisions as they usually create a soft tissue defect difficult to reconstruct after final resection of the tumor with the biopsy tract. The surgeon should be aware of the complex nerve and vessel anatomy of the upper extremity. The biopsy tract must not violate more than one anatomic compartment and avoid exposure of the neurovascular bundles to the tumor cells. The pseudocapsule of the tumor should be closed with sutures after tissue harvesting. Adequate hemostasis should be performed in order to avoid hematoma formation and when a draining tube is placed it should be in line and close to the incision.

Despite its higher diagnostic value, open biopsy is kept as the last resource as it is expensive, carry a complication rate of up to 16% and may cause contamination of the incisional path. At the final surgery for tumor resection, the surgical path of the biopsy (including 1-3 cm of the skin around the incision and subcutaneous tissues) should be excised en block with the final tumor specimen (18).

In case of small superficial lesions, well planned excisional biopsies with negative margins can be performed (19). An absolute prerequisite to decide for an excisional biopsy is the ability to resect the mass with negative histology margins. The surgeon performing an excisional biopsy should have measured in detail on pre-op MRI the size of the lesion and the relation to surrounding tissue. An exception to this concept, is the benign giant cell tumor of tendon sheath located in the fingers, where frequently a marginal resection is performed that may result in a higher local recurrence rate (20). In contrast to well-planned excisional biopsy, the ‘unplanned excision’ is defined as the gross removal of tumor without pre-operative staging or consideration for the need for removal of normal tissue around the tumor (21). Unplanned excisional biopsies of the upper extremity, frequently leave microscopic residual disease requiring a more aggressive and debilitating subsequent treatment in up to 45% of cases (22).

Once the diagnosis of malignancy is suspected or established, staging is preferably performed by computed tomography (CT) of the thorax and abdomen whereas positron-emission tomography (PET) is reserved for selected cases (23).

4. General considerations on therapeutic approach

Compared to tumors of the lower limb, upper extremity tumors tend to be smaller, more often superficial (3) and more likely to undergo unplanned excision (24). Oncological surgery represents a major challenge in most cases. The complex and intimate surgical anatomy of tendons, vessels and nerves jeopardizes both the success of appropriate surgical margins and the postoperative loss of function (25). An unplanned excision of upper-limb tumors tends to have a higher rate of positive surgical margins, along with a significant higher rate of local recurrence when compared to lower-limb tumors (3).

The addition of adjuvant chemotherapy (26) and radiotherapy (27) has been reported to improve the outcomes of surgery in unselected patients with STS. As a result,
5. Treatment of localized disease

Surgery. Until late 1970s, amputation was regularly performed for localized STS of the extremities, on account of higher rates of local recurrence (30). Enneking et al (31), described four types of resection margins: Intralesional, marginal within the reactive zone, wide with a cuff of normal tissue and radical resection involving excision of the entire anatomical compartment. Through decades, we moved from the surgical principle of resection of the involved anatomic compartment, to tumor free resection margins as a minimum of acceptable resection. Although a wide soft tissue envelop in all directions around the tumor is desirable, the feasible goal is resection to negative margins (1 mm from the inked resection margin) (32). Still, for STS of the hand, amputation of a digit may be necessary to obtain clear margins.

Currently more than 90% of STS can be treated with local resection and limb salvage. However, a primary amputation should be considered when the tumor cannot be excised to clear margins, based on the pre-op imaging. Extensive soft tissue infiltration and/or involvement of a major neurovascular bundle frequently result in amputation. A primary amputation should be considered when tumor resection results in significant loss of soft tissue with severe function impairment, that cannot be reconstructed with available surgical techniques, or the expected complication rate will be high.

Suboptimal biopsies and positive resection margins are associated with local and distant disease recurrence in patients with hand STS (22). Pradhan et al (33), reported on 63 patients with hand STS. Six patients underwent below
elbow amputation and 12 patients had partial amputation. All the amputated patients had clear margins, while 42% out of patients with local tumor excision had involved margins (33). Single ray amputations (excluding thumb) for hand tumors have a low local recurrence rate and high functional scores (34). However, ray transposition should not be performed with ray amputation for tumor excision. In order to achieve negative margins, wider resection may be needed. Double ray amputation results in worse functional outcome than single ray. Good key, tip and tripod pinch can nonetheless be maintained when the deep motor branch of the ulnar nerve is preserved, and this hand can still assist in bimanual hand activities (35).

Preoperative radiation therapy (RT) is useful in cases where the tumor mass is in contact to nerves and vessels, as it may facilitate negative margin resection by inciting a thicker reactive fibrous tumor pseudocapsule, which can be dissected from the neurovascular bundle (36). Clarkson et al (37), concluded that meticulous sharp epineural dissection of the ischial nerve combined with RT is a safe technique and nerve preservation can be attempted when the tumor does not encase the nerve trunk. Although there are no randomized studies available, RT and epineural dissection is the common practice for STS of the upper extremity abutting on major nerves.

When the tumor mass surrounds important vessels, limb-sparing surgery can be performed as an en bloc resection of the sarcoma and vessels with vascular reconstruction. For large diameter vessel reconstruction, the great saphenous vein is usually harvested, reversed and anastomosed proximally and distally to restore anatomic continuity and circulation (38).

Contact of the mass with the bone is a common finding in MRI of large, deep STS and invasion of the bone cortex can be found in cases of SS and UPS. Cortical and medullary signal intensity changes and cortical destruction observed on T1 and T2-weighted MR images are highly sensitive and specific signs of osseous invasion by STS (39). A study from Mount Sinai reported that true bone invasion occurs in a 5.5% and it is associated with increased metastatic disease at presentation and decreased overall survival (OS) (40). Preoperative RT enables resection of the mass with periosteum serving as the deep margin, without expecting increased recurrence rate (41). For STS invading the bone, or when negative margins cannot be achieved using periosteum as a deep margin, en bloc resection of the soft tissue mass with the affected bone should be performed. For segmental bone defects, reconstruction can be done with either avascular bone autograft, allograft, a vascularized fibula graft or a hybrid reconstruction of an allograft combined with a free vascularized fibula graft.

Flap reconstruction is an essential part of STS surgical treatment (42). Tensionless primary wound closure is important to avoid wound healing complications, especially if preoperative RT has been administered. For small size soft tissue defects, wound closure can be achieved either by simple sutures or muscle approximation and split thickness skin grafting. Flap coverage is essential in the case of exposed vessels, nerves or bone. Flap usage is also important in the prevention or management of wound healing complications (43). Frequently used flaps are the lateral arm flap, the radial forearm, anterior-lateral thigh and latissimus dorsi flap (42). Vasileios et al (16) reported on 57 patients with soft tissue malignant fibrous histiocytoma. A rotational or free flap was eventually needed in 28 patients. A major wound complication occurred in 17% of patients. All complications were related to preoperative RT and 90% involved the lower limbs. Wound breakdown was associated with infection in 50% of cases (16).

Prognostic factors. Metastatic relapse after complete surgery occurs in around 40% of patients, leading to death from the disease within the first 8 years after initial diagnosis (44). Several prognostic factors have been identified to assess the probability of recurrence after surgery. High histological grade, size >5 cm, deep location and positive surgical margin status have been characterized as the most important poor prognostic factors (45). Of them, histological grade has been pointed out as the factor with the heaviest prognostic impact for systemic control after surgery (46), while surgical margin status has been described as the most important factor for local control (47). In order to reduce the high probability of relapse, complementary radiation therapy and chemotherapy may be applied.

Radiation therapy. Radiation therapy (RT) is a crucial adjunct to surgery for STS of the extremities. The most important outcome by the use of RT is the local control of the disease, but this is not associated with a significant reduction in distant metastasis or improvement in disease-specific survival (48). There are several RT modalities applied such as external beam radiation therapy (EBRT), intensity-modulated radiation therapy (IMRT), intraoperative radiation therapy (IORT) and brachytherapy.

External beam radiation therapy. The administration of preoperative or adjuvant EBRT in order to avoid amputation is supported by evidence reported from several clinical trials. Selected randomized prospective clinical trials are shown in Table III. The addition of EBRT after LSS attains similar results as amputation and significantly reduces the local-recurrence rate of LSS alone (27). The benefit of adjuvant EBRT seems higher for STS with poor prognostic factors, and data suggest that it might be omitted in patients with completely resected low-risk STS (49,50). Significant differences in toxicity have been reported with the use of postoperative EBRT compared to LSS alone with respect to edema, limb strength and range of motion. Preoperative EBRT, is significantly related to greater acute toxicity and major wound complications than postoperative EBRT, without differences in local-relapse rates and long-term OS (51).

As complex trade-off issues are involved in the sequencing of LSS and RT for patients with localized STS of the extremities, it seems important to define subsets of patients who might be adequately treated by surgery alone and the optimal sequence of surgery and EBRT for patients who require both types of local therapy (52). An attempt has been made to develop a nomogram to quantify the 3- and 5-year risk of local recurrence after LSS without postoperative EBRT that includes age, size, margin status, grade of tumor and histology (53).

Intensity-modulated radiation therapy. The main advantage of IMRT is its ability to deliver high dose RT to the tumor minimizing the dose of RT to the surrounding normal structures. Such a tight margin might compromise tumor coverage and result in a higher rate of local recurrences. A retrospective
after treatment with function-conserving surgery and postoperative IMRT (34 patients), reported encouraging data on acute and late toxicity. The 5-year local control rate was 94%, which compares favorably with that of historical controls ranging from 82% in negative margins to 51% in positive margins (54).

A retrospective comparative study including 319 patients with STS of the extremities treated with postoperative EBRT (154 patients) or IMRT (165 patients) with similar dosing schedules indicated that IMRT was associated with significantly reduced local recurrence compared with conventional EBRT (55).

A following prospective phase II study included 80 patients with localized STS of extremities (51 patients) and trunk wall (29 patients). After treatment with function-conserving surgery and postoperative IMRT, an excellent local control was assessed, with low IMRT-associated toxicity such as edema and joint stiffness (56). As a result, IMRT is a promising RT approach in STS of the extremity as it provides excellent local control in a group of patients with high-risk features with a beneficial effect in sparing the surrounding normal tissue.

**Intraoperative radiation therapy.** Intraoperative RT consists of a single large dose of RT, administered during LSS after tumor removal and prior to wound suturing. As a result, the tumor bed can be irradiated directly sparing the surrounding normal tissue. IORT is usually combined with postoperative RT, but special hospital infrastructures are required. IORT used as a boost to EBRT seems to provide excellent local control with only mild acute side effects, as indicated by a retrospective study of 17 patients with STS of upper or lower extremities (57).

A more recent retrospective analysis of 61 patients with upper-extremity STS treated with surgery, IORT (12.50 Gy) and EBRT (45-50 Gy), associated this strategy with excellent local control, limb preservation and survival, even in patients with positive margins. Only 4 patients developed RT-associated toxicity (58).

**Brachytherapy.** Brachytherapy is the direct application of radioactive sources into the tumor bed through catheters, which allows a high dose of RT to the tumor in a more conformal way compared to EBRT. This is translated in shorter treatment periods, fewer side effects and a faster recovery. It permits evaluation at the time of surgery and complications can be avoided by sparing the surrounding tissues (59).

A prospective trial randomized 164 patients to receive either adjuvant brachytherapy or no further RT after complete resection of STS of the extremity (60). The RT was administered by iridium-192 implants, which delivered 42 to 45 Gy over 4 to 6 days. The results indicated that in patients with high-grade histologies brachytherapy provides convenient means to complete RT within a short period with no long-term functional sequelae and with a local control benefit comparable to that obtained with more protracted courses of EBRT. These data suggest that brachytherapy is an effective alternative to EBRT.

**Adjuvant and preoperative chemotherapy.** Chemotherapy for resectable STS of the extremities has been evaluated in both the adjuvant and the preoperative settings.

**Adjuvant chemotherapy.** The data of nearly twenty clinical trials, carried out from 1980's to 2008, have been gathered in two different meta-analyses (61,62). Tierney et al (61) showed that adjuvant chemotherapy for unselected patients results in an absolute benefit at 10 years of 4%. When subgroup analysis was carried out, sarcoma of the extremities had an absolute benefit at 10 years of 7% (p<0.029) (61). In a more recent meta-analysis, Pervaiz et al (62) evaluated data from 1,953 patients receiving...
postoperative chemotherapy. According to this meta-analysis, administration of adjuvant anthracyclines and ifosfamide leads to a significant reduction in the risk of recurrence and death of 10 and 11%, respectively (62).

Conversely, the largest, phase III, placebo controlled, clinical trial for adjuvant chemotherapy in STS, which randomized 351 patients to receive or not 5 cycles of Adriamycin and ifosfamide after surgery, found no significant differences, either in relapse free survival or in OS. However, a clear advantage of chemotherapy can be inferred from the forest plot for extremities, grade 3 and greater size (63).

Preoperative chemotherapy. The role of preoperative chemotherapy was first evaluated retrospectively, showing a positive effect mainly for patients with deep, high-grade chemotherapy was first evaluated retrospectively, showing a positive effect mainly for patients with deep, high-grade tumors over 10 cm (64). Prospectively, a large phase III clinical trial randomized 328 high-risk patients to receive neoadjuvant epirubicin 120 mg/m² and ifosfamide 9 g/m² for 3 or 5 cycles (65). It was confirmed that 3 cycles do not worsen survival rates (5-year OS of 68% with 3 cycles versus 70% with 5 cycles), and they are comparable to those seen with the same combination in the adjuvant setting (66). The addition of RT was permitted and subgroup analysis of patients with affected surgical margins showed a local relapse rate of 17% if RT was administered after surgery versus 0% when it was performed before (67).

The unslected populations of the previously mentioned clinical trials constitutes a major hindrance when deciding the best therapeutic option for sarcoma patients. Evidence indicates that chemotherapy is more useful for sarcomas of the extremities and the trunk wall, but no differences between upper and lower limbs have been reported. In the absence of clinical trials with selected populations, chemotherapy for STS of the upper limb should be administered for chemosensitive histologic subtypes when poor prognostic factors are present.

Despite the preoperative treatments described hitherto, some cases are not amenable but with amputation of the extremity. These cases led to the exploration of other methods in an attempt to improve the LSS. The isolated limb perfusion (ILP) consists in the administration of chemotherapy after separating the circulation of a limb from that of the rest of the organism. The introduction of TNF-α in combination with melphalan after some frustrating results of ILP with doxorubicin (68) allowed the LSS in 76% of patients who, otherwise, would have required amputation (69).

Follow-up and recurrence. As for STS of other locations, follow-up of STS of the upper extremities has the objective of controlling the sequelae from the administered treatments as well as the detection of local or metastatic relapse. Thus, follow-up is also an important part of the multidisciplinary approach, since early and late complications from surgery, radiotherapy and chemotherapy have to be diagnosed and treated promptly.

Surveillance for uncovering local and metastatic relapse is a controversial subject. Local recurrence of the upper extremity is easily detected by physical examination and even by self-examination, which calls into question the routine use of image tests of the limb (70). Surveillance periodicity varies between clinical guidelines and it depends on several factors, such as histologic grade and subtype or even the experience of every sarcoma center. Despite the tendency to use high-definition image tests, it seems that chest CT does not add a benefit to the use of simple X-ray in the detection of resectable pulmonary metastases (71). However, no prospective trials have determined to date the best surveillance strategy for the detection of both local and metastatic relapse of STS.

Local relapse of a previously treated STS of the upper extremities may not be amenable to re-excision, but the option of radiation and even re-radiation could be possible in selected cases (72).

Similarly, the therapeutic strategy for metastatic disease depends on the number and the site of metastases, potentially managed with local treatments, and on the histologic subtype, potentially sensitive to systemic therapy. The lung is the most common site of metastases, as up to 80% of metastatic STS present with lung metastases (73). Although prospective and randomized studies are lacking, pulmonary metastasectomy with complete resection of all disease burden may be considered for selected patients. According to different reported series, pulmonary metastasectomy attains a 5-year survival rate of 15 to 50.9% (74). The complete resection of all metastases is the most important prognostic factor, as it duplicates the survival compared to that of incomplete resection (75). The number of metastases has also been indicated as an important prognostic factor, though the maximum number of metastases contraindicating the surgery has not been established (74).

6. Chemotherapy and targeted therapy for metastatic STS of the upper extremities

After multidisciplinary curative treatment, the risk of metastatic relapse within the next 2 years after surgery is as high as 46% (29). Thus, improvement in the treatment of metastatic disease is imperative.

If not amenable to salvage surgical procedures, treatment of metastatic sarcoma is still based primarily on chemotherapy. Despite the intrinsic heterogeneity, clinical trials have often been designed for all histological subtypes taken together. Since the decade of the 80 s, anthracyclines have been the drug class of choice for the first line treatment, attaining a limited response rate of under 25% at conventional dose of 60-75 mg/m² (76). In order to optimize the effectiveness of treatment, combinations of doxorubicin and ifosfamide have been investigated at different doses. Effectiveness of the combination has been reported superior at conventional doses (doxorubicin 60-75 mg/m² and ifosfamide up to 9 g/m2) at the expense of a greater toxicity, without significant improvement in OS (77,78). High doses of doxorubicin and ifosfamide have also been tested, resulting in a better clinical benefit and progression-free survival (PFS) with the counterpoint of a mainly hematological greater toxicity (79).

Combination of doxorubicin and olaratumab, showed a significant improvement in OS that could not be confirmed in the recently reported, phase III clinical trial ANNOUNCE (NCT02451943), which did not meet its primary endpoint of OS (80).
Following failure on preoperative, adjuvant or first-line chemotherapy with anthracyclines, therapeutic options for STS include other cytotoxic agents, tyrosine-kinase inhibitors (TKI) and immunotherapy (81). Monotherapy with trabectedin at 1, 5 mg/m² in a continuous 24-hour infusion every 3 weeks is an approved second line option after anthracyclines, achieving a median PFS of 4.2 months (82). Combinations of the antimeabolite gemcitabine, with docetaxel (83) or dacarbazine (84) achieve a median PFS of 6.2 and 4.2 months, respectively. The TKI pazopanib, targeting the vascular endothelial growth factors (VEGF) 1‑3, the PDGFR A and B and mast/stem cell growth factor receptor KIT, was tested in a randomized phase III clinical trial and a median PFS of 4.6 months was found (81).

As several reports have indicated differential responses to the available drugs, the choice of specific therapeutics may vary according to histologic subtypes (Table IV) and toxicity (Table V).

Undifferentiated pleomorphic sarcoma. The UPS subtype has been included in almost all clinical trials that led to the approval of second line therapies. Anthracyclines, ifosfamide, gemcitabine and trabectedin are active drugs in this histology (86). Based on a phase II study, the combination of gemcitabine with docetaxel was considered to be active in UPS (83), but later a larger phase III study confirmed that the combination of epirubicin with ifosfamide

Table IV. First and second line options for the predominant histological subtypes of STS of the upper extremities.

| Sarcoma type | First line | Second and further lines | Drugs under investigation |
|--------------|------------|--------------------------|---------------------------|
| UPS          | Doxorubicin ± Ifosfamide<br>Gemcitabine-Docetaxel<br>Trabectedin<br>Pazopanib | Pembrolizumab | |
| SS           | Doxorubicin ± Ifosfamide<br>Trabectedin<br>Pazopanib | Tazemetostat | |
| ES           | Doxorubicin ± Ifosfamide<br>Gemcitabine-Docetaxel<br>Pazopanib | Tazemetostat | |
| CCS          | - | - | Caffeine-potentiated doxorubicin<br>Sorafenib<br>Sunitinib<br>Tinvatinib |

*Table IV. First and second line options for the predominant histological subtypes of STS of the upper extremities.*

Table V. Selection of the toxicities of principal available drugs for soft tissue sarcomas.

| Drug               | Very common and common                           | Uncommon                                      | Rare and very rare                                       |
|--------------------|-------------------------------------------------|-----------------------------------------------|----------------------------------------------------------|
| Doxorubicin        | Myelosuppression, Cardiotoxicity                | Dehydration                                  | Tissue necrosis                                          |
| Ifosfamide         | Myelosuppression, Hepatotoxicity, Hemorrhagic cystitis, Acute renal failure | Peripheral neuropathy, Stomatitis | CNS toxicity, Dermatitis                                 |
| Gemcitabine        | Myelosuppression, Elevation of liver transaminases, Allergic skin rash, Influenza-like symptoms | Interstitial pneumonitis | Anaphylactoid reaction, PRES, Capillary leak syndrome |
| Docetaxel          | Myelosuppression, Hypersensitivity, Peripheral neuropathy, Fluid retention | Arthralgia, Elevation of liver transaminases | Cardiotoxicity                                            |
| Trabectedin        | Myelosuppression, Elevation of liver transaminases | Capillary leak syndrome, Pulmonary edema | Hepatic failure                                           |
| Pazopanib          | Hypothyroidism, Hypertension, Hair color change, Elevation of liver transaminases, Diarrhea | Hypomagnesaemia, Retinal detachment, Cardiotoxicity, Intestine perforation | Thrombotic microangiopathy, Posterior reversible, encephalopathy, Pneumonitis |

The following was utilized for the classification of frequency: Very common ≥1/10; common ≥1/100 to <1/10; uncommon ≥1/1,000 to <1/100; rare ≥1/10,000 to <1/1,000; and very rare <1/10,000. CNS, Central nervous system; PRES, Posterior reversible; encephalopathy syndrome; PPEDS, Palmar-plantar erythrodysesthesia syndrome.
was more effective (87). Despite the meager research of immunotherapy in STS, a small phase II study testing the anti-PD1 antibody pembrolizumab in UPS showed 40% objective response rate (88), placing pembrolizumab a potential therapeutic option for the future.

**Synovial sarcoma.** Although SS is particularly sensitive to ifosfamide (89), high dose of ifosfamide is not superior to the combination of anthracycline and ifosfamide in the first line (87). After first-line chemotherapy, trabectedin monotherapy showed better responses for SS compared to other STS subtypes (90), as well as the TKI pazopanib (85).

Research on new targeted agents has led to the exploration of potential molecular targets. In SS, the specific gene fusion SYT-SXX leads to the hyperexpression of intracellular pathways involved in survival and metastases, highlighting a number of potential targets (91). The oncogenic fusion SYT-SXX results in SMARCB1/INI1 proteolytic degradation, boosting the action of EZH2 on heterochromatin (92). Tazemetostat, an EZH2-inhibitor, has shown activity in preliminary results of a phase 1/2 clinical trial (NCT02601950) (93) and a phase 2 trial (NCT02601950) is currently ongoing.

**Epithelioid sarcoma.** Due to its rarity, prospective data on effectiveness of chemotherapeutic agents in ES are scarce. Retrospective analyses showed that gemcitabine-based and anthracycline-based chemotherapy regimens are active in metastatic ES (94). The combination gemcitabine/docetaxel is thus a second-line option for these patients. Trabectedin has been reported ineffective in ES (95), and pazopanib showed an inferior PFS and OS when compared to anthracyclines and to gemcitabine in a retrospective study (81).

ES is marked by SMARCB1/INI1 deficiency in a 90% of cases, and patients with this histology are included in the previously detailed clinical trials with tazemetostat (NCT02601950; NCT02601950).

**Clear cell sarcoma.** The CCS is a rare entity without prospective clinical trials and it is considered a primarily chemo-resistant sarcoma (81).

No objective responses have been reported with pazopanib, and only a small case series reporting partial responses with sorafenib and sunitinib has been reported (96).

Tivantinib, a MET-inhibitor, has been tested in a phase II trial with 11 cases where a clinical benefit rate of 36% and a median PFS of 1.9 months were documented (97). The MET/ALK-inhibitor crizotinib and several immunotherapeutic options have also been tested in CSS patients without objective responses (98,99).

7. Discussion

STS of the upper extremity represent less than 10% of all STS and affect young patients, with a mean age at diagnosis of 38 years. Surgery remains the cornerstone of the treatment of STS of the upper extremities, which tend to be small and...
Figure 2. Therapeutic algorithm of the localized STS of the upper extremities with poor prognostic factors. The poor prognostic factors included, high histological grade, size >5 cm, deep location and positive surgical margin status. STS, soft tissue sarcomas.

Figure 3. Therapeutic algorithm of the metastatic STS of the upper extremities. STS, soft tissue sarcomas.
superficial, leading to a higher number of unplanned resections. This may be the cause of a higher rate of relapse of STS of this location. Besides, the anatomical particularities represent a surgical challenge, as wide excision may worsen the functional outcomes and debilitate this young patient subgroup. Although amputation could still be an obligatory option for selected cases, preservation of as many anatomical structures as possible performing a limb-sparing surgery (LSS) is desirable and represents the standard of care, using different reconstruction techniques to safeguard member's functionality.

However, about half of the patients, most commonly those with poor prognostic factors, who undergo surgery will relapse within the first 5 years following the intervention.

Adjuvant external-beam radiation therapy (RT) achieves similar outcomes as amputation when combined with LSS. Nowadays, different RT methods are available with less toxicity and similar efficacy but restricted to centers with expertise in the field. Whether administration of RT is preferable before or after surgery is an unanswered question. There are no great differences in terms of effectiveness and preoperative RT is associated with a higher rate of acute toxicity, but it may be preferred when downsizing of the tumor is required for increasing the probabilities of a successful LSS.

Adjuvant chemotherapy improves the survival of selected patients with STS of the extremities. However, clinical trials for adjuvant chemotherapy lack of specificity as their design have not considered the intrinsic heterogeneity of the disease and the prognostic factors due to the histological rarity and molecular heterogeneity of the subtypes. Given the greater incidence of high-grade STS in the upper extremities, it is expected that most patients with this diagnosis will undergo adjuvant chemotherapy.

Metastatic disease must be radically treated with surgery or radiotherapy whenever it is possible, as this practice may achieve an improvement in overall survival. Systemic therapy improves survival when radical therapy cannot be accomplished. The understanding of the underlying biology of each subtype of sarcoma is essential for the selection of one or another drug in each line of treatment.

With the information gathered above, the authors propose a therapeutic algorithm for the treatment of STS of the upper extremities according to the absence (Fig. 1) or presence of poor prognostic factors (Fig. 2) in the early disease stage or metastatic disease (Fig. 3). However, each individual patient should be discussed in the context of a specialized multidisciplinary meeting at the earliest possible stage of diagnosis, in order to establish a radical therapeutic plan, maximizing the profitability of the procedures and shortening the time between interventions. Indeed, the presence of a dedicated tumor board has been associated with an improvement of about 5% in the 2-year disease-free survival, and its absence has been defined as a new poor-prognostic factor for STS (29).

STS of the upper extremities represent a challenge due to anatomical and histopathological particularities, as well as to their low incidence. Although multidisciplinary treatments have increased the functional outcomes and the survival of these patients, a need of improvement of treatment for metastatic disease is of utmost importance, urging for multinational cooperation for the recruitment of patients in multicenter clinical trials.

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AK and VK conceived and designed the review. JDM performed the literature review. All authors were involved in the preparation and revision of the manuscript.

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Competing interests
The authors declare that they have no competing interests.

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