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Sarcoma European and Latin American Network (SELNET) Recommendations on Prioritization in Sarcoma Care During the COVID-19 Pandemic

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Disclosures of potential conflicts of interest may be found at the end of this article.

Key Words. COVID-19 • Sarcoma • Guidelines • Patient care • Multidisciplinary

ABSTRACT

Background. The COVID-19 outbreak has resulted in collision between patients infected with SARS-CoV-2 and those with cancer on different fronts. Patients with cancer have been impacted by deferral, modification, and even cessation of therapy. Adaptive measures to minimize hospital exposure, following the precautionary principle, have been proposed for cancer care during COVID-19 era. We present here a consensus on prioritizing recommendations across the continuum of sarcoma patient care.

Material and Methods. A total of 125 recommendations were proposed in soft-tissue, bone, and visceral sarcoma care. Recommendations were assigned as higher or lower priority if they cannot or can be postponed at least 2–3 months, respectively. The

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consensus level for each recommendation was classified as “strongly recommended” (SR) if more than 90% of experts agreed, “recommended” (R) if 75%–90% of experts agreed and “no consensus” (NC) if fewer than 75% agreed. Sarcoma experts from 11 countries within the Sarcoma European-Latin American Network (SELNET) consortium participated, including countries in the Americas and Europe. The European Society for Medical Oncology-Magnitude of clinical benefit scale was applied to systemic-treatment recommendations to support prioritization.

Results. There were 80 SRs, 35 Rs, and 10 NCs among the 125 recommendations issued and completed by 31 multidisciplinary sarcoma experts. The consensus was higher among the 75 higher-priority recommendations (85%, 12%, and 3% for SR, R, and NC, respectively) than in the 50 lower-priority recommendations (32%, 52%, and 16% for SR, R, and NC, respectively).

Conclusion. The consensus on 115 of 125 recommendations indicates a high-level of convergence among experts. The SELNET consensus provides a tool for sarcoma multidisciplinary treatment committees during the COVID-19 outbreak. The Oncologist 2020;25:e1562–e1573

Implications for Practice: The Sarcoma European-Latin American Network (SELNET) consensus on sarcoma prioritization care during the COVID-19 era issued 125 pragmatical recommendations distributed as higher or lower priority to protect critical decisions on sarcoma care during the COVID-19 pandemic. A multidisciplinary team from 11 countries reached consensus on 115 recommendations. The consensus was lower among lower-priority recommendations, which shows reticence to postpone actions even in indolent tumors. The European Society for Medical Oncology-Magnitude of Clinical Benefit scale was applied as support for prioritizing systemic treatment. Consensus on 115 of 125 recommendations indicates a high level of convergence among experts. The SELNET consensus provides a practice tool for guidance in the decisions of sarcoma multidisciplinary treatment committees during the COVID-19 outbreak.

Introduction

Deferral, modification, and even cessation of therapy are hindrances affecting patients with cancer during the current COVID-19 outbreak. The balance between adapting measures to minimize hospital exposure, following the precautionary principle, and offering a more relaxed diagnostic or therapeutic management while preserving the patient’s survival options is not easy.

More than 80 reports, often short reports and letters, have addressed the intersection of cancer and COVID-19 care. Although the risk of morbidity and mortality is almost always higher in patients with cancer than in patients infected with COVID-19 (i.e., risk of death for pancreatic cancer is greater than 90%, whereas for COVID-19-infected patients, it is usually lower than 5%) [1], the truth is that the precautionary principle has prevailed over cancer care continuation in many cases [2]. Elective surgeries were largely suspended, in some instances, because of shortage of ventilators, and thus the operating theaters became extemporaneous intensive care units. Some reports have focused on the real impact (direct and indirect) of the COVID-19 outbreak in patients with cancer. In some locales where the admissions for SARS-CoV-2 far exceeded the hospital capacity, it was to be expected that cancer care would be compromised in a variety of ways.

Patients with cancer constitute a large population with a spectrum of risk with respect to immunosuppression. Cancer immunodeciting, which represents the interplay between tumor and immune system, ultimately leads to changes in immune cells, immune modulators, cytokines, and molecules expression toward the escape phase and the development of an immunosuppressive tumor microenvironment [3]. Both specific diagnoses and their treatment with chemotherapy, surgical resection, and newer treatments variably compromise immune function and render some patients with cancer more susceptible to infections [4].

As with other infections, the innate and adaptive arms of the immune system play different key roles in the COVID-19 infection and evolution. SARS-CoV-2 causes an overwhelming persistent innate-induced inflammation that can lead to a cytokine storm, cytokine-associated lung injury, and diffuse organ involvement [5]. Alterations of the CD4+ and CD8+ T cells subsets have been observed, with loss of functional diversity in CD4+ T cells and increased expression of regulatory molecules in CD8+ T cells [6, 7]. Hence, it can be assumed that the systemic immunosuppressive state of patients with cancer might result in an increased risk of COVID-19 infection and poorer prognosis for this group of patients.

Epidemiological statistics of the cases in China suggested that patients with cancer were at greater risk than the general population, data which appear to be borne out more for hematologic malignancies than solid tumors, although there still appears to be greater risk even for patients with solid tumor for fatal outcomes from SARS-CoV-2 [8]. Noteworthy, it has been reported that COVID-19–related deaths were strongly associated with male sex, older age, and deprivation; severe asthma; uncontrolled diabetes; cancer; and several other previous, clinical conditions [9].

Importantly, a recent epidemiological report from the U.K. based on 800 patients with cancer with COVID-19 infection observed a mortality rate of 28%, with older age, male gender, and comorbidities, such as cardiovascular disease, significant prognostic factors related to higher mortality. Remarkably, chemotherapy or other systemic therapy administered within 4 weeks of testing positive for COVID-19 did not have significant effect on mortality from COVID-19 disease [10].
Taken together, rapidly evolving data indicate that oncologic patients, such as patients with sarcoma, constitute a high-risk group more likely to suffer higher mortality and morbidities than in the general population if infected by SARS-CoV-2.

Additionally, during the COVID-19 outbreak, the major risk for patients with cancer is the inability to receive necessary medical services. Decisions on whether or not to postpone cancer treatment and clinical trials need to be made on a patient-by-patient basis and according to the inherent tumor risk in each patient and the prevailing situation, because delays could lead to tumor progression and, ultimately, poorer outcomes [11]. An article on patients’ perspectives reported that up to 30% of oncologic patients have had changes in their treatment or follow-up care during the COVID-19 pandemic [12]. Beyond the increased risks, including requiring mechanical ventilation and death, that should be prospectively analyzed in oncologic patients with COVID-19 infection, it is also relevant to take into account the impact of the precautionary principle that is implemented in our patients [13]. The latter implies that patients with cancer have suffered delays or cancellation of diagnostic tests, surgeries, radiation therapies, or systemic treatments.

The aim of this article is to build a consensus on prioritization aspects in sarcoma care during the current COVID-19 outbreak or future outbreaks, which could appropriately balance the precautionary principle while preserving the highest survival probabilities in sarcoma patients.

This consensus has been developed within the Sarcoma European-Latin American Network (SELNET). This is a consortium granted by the European Commission within the Horizon 2020 Call, within the program H2020-SC1-BHC-2018-2020 (better health and care, economic growth, and sustainable health systems). The aim of the SELNET consortium is to improve clinical outcome in sarcoma care, through a network of reference centers in sarcoma [14].

These guidelines are intended to add value over and above other clinical practice guidelines reported for oncologic patients in the context of COVID-19 outbreak. Thus, our guidance focuses specifically on patients with sarcoma and thoroughly provides precise details on prioritization of 125 clinical sarcoma situations, whereas the National Institute for Health and Care Excellence (NICE) offers a general recommendation for any oncologic patient [15], and the European Society for Medical Oncology (ESMO) offers a more general recommendation of prioritization for patients with sarcoma [16].

**Materials and Methods**

The recommendations have been divided into two scenarios, higher and lower priority, to make them simple and replicable. Higher priority is defined as an undelayable procedure, because the inherent risk of the tumor, affecting survival or morbidity, would likely exceed the risk of COVID-19 infection if this procedure had not been performed. By contrast, lower priority is defined as a delayable procedure (at least 2–3 months), because the inherent risk of the tumor, affecting survival or morbidity, would be low enough that the patient could minimize or avoid hospital frequentation and, consequently, reduce the risk of infection by COVID-19. A total of 125 recommendations have been proposed in soft tissue, bone, and visceral sarcomas to thoroughly cover details for diagnosis, surgery, radiation therapy, systemic treatments, follow-up, and clinical research.

These recommendations were proposed by the SELNET coordinator team and shared with the official partners of SELNET network. Associated partners were not involved in this consensus. Each point derived from multidisciplinary (MDT) expertise discussion from the SELNET coordination aimed to be pragmatic, detailed, and patient-oriented for each recommendation.

Official partners had the competence to give their own opinion if this was clear enough for them or to give the opinion of their MDT on certain recommendations.

In this consensus, MDT experts participated from AC Camargo Cancer Center (Sao Paulo, Brazil), Instituto Alexander Fleming (Buenos Aires, Argentina), Instituto Nacional de Enfermedades Neoplasicas (Lima, Peru), Instituto Nacional de Cancerología (Mexico DF, Mexico), Hospital Calderon Guardia (San José, Costa Rica), Centre Leon Berard (Lyon, France), Istituto Nazionale dei Tumori (Milan, Italy), IRCCS Istituto Ortopedico Rizzoli (Bologna, Italy), Abramson Cancer Center (Philadelphia, Pennsylvania), University Hospital Lund (Lund, Sweden), Mannheim University Medical Center (Mannheim, Germany), MD Anderson Cancer Center (Houston, Texas), Spanish group for Research on Sarcomas, and University Hospital Virgen del Rocio (Sevilla, Spain). Besides that, a European patient advocacy group (Sarcoma Patients EuroNet) was also involved in this consensus, as well as the external scientific and ethics committees of SELNET.

Among 31 participating experts in the consensus, there were 18 oncologists (15 for adults and 3 for pediatrics), 4 orthopedic surgeons, 4 surgical oncologists, 2 radiation oncologists, 1 pathologist, and 1 radiologist. As SELNET mainly focuses on adult-type sarcomas, no exclusive pediatric oncologist participated in the consensus. Fourteen Latin American sarcoma experts participated in the consensus: nine oncologists, four surgeons, and one radiation therapist. Correlations between Latin American and European Union (E.U.-U.S.) expert recommendations were evaluated using Pearson’s \( \chi^2 \) test for bivariate options (agreement or disagreement) and Mann-Whitney \( U \) test for multivariate options (strongly agree, agree, disagree, and strongly disagree).

For each recommendation, every participant chose from the following options: strongly agree, agree, disagree, and strongly disagree. The neutral option was not offered to avoid ambiguity.

Those recommendations with \( \geq 90\% \) of consensus (at least strongly agree plus agree obtained in \( \geq 90\% \) of participants) were considered as strongly recommended. Those with \( \geq 75\% \) but \( < 90\% \) were considered as recommended, and the remaining recommendations were considered as “no consensus was obtained.” The cutoff of 75% has been the median threshold to define consensus in Delphi studies [17], whereas the demanding 90% cut-off was arbitrarily chosen to indicate the highest consensus range if at least this percentage was reached.
To mitigate the subjectiveness, the ESMO-Magnitude of Clinical Benefit Scale (MCBS) V1.1 was applied to recommendations involving systemic treatments to support this prioritization strategy (supplemental online Appendix 1) [18, 19]. Although MCBS has not been formally evaluated for sarcoma therapies, MCBS methodology was followed for the estimation of the benefit obtained with different sarcoma systemic treatments (supplemental online Appendix 2).

RESULTS
From a total of 31 fulfilled questionnaires analyzed, the recommendations were distributed as strongly recommended (SR) in 80, recommended (R) in 35, and no consensus (NC) in 10. The consensus was higher among the 75 higher-priority recommendations (85%, 12%, and 3% for SR, R, and NC, respectively) than in the 50 lower-priority recommendations (32%, 52%, and 16% for SR, R, and NC, respectively).

Table 1 describes the consensus level for each recommendation and indicates the MCBS in most contexts of systemic treatments. The statistical distribution for each recommendation is presented in supplemental online Appendix 1.

Of note, statistically significant differences between Latin American and E.U.-U.S. expert recommendations were only detected by Pearson’s \( \chi^2 \) and Mann-Whitney \( U \) test in 3 of 125 recommendations (Table 2). E.U.-U.S. experts proved to be more conservative for the advice of adjuvant chemotherapy in high grade chondrosarcoma, the advice of adjuvant radiotherapy in low-grade, deep soft-tissue sarcoma (STS) larger than 5 cm, and for the advice of adjuvant radiotherapy in superficial STS larger than 5 cm.

DISCUSSION
Because sarcoma multidisciplinary committees are so critical to patient outcomes [20–27], we emphasize the message that these meetings should continue, at least in tele-committee format, in the COVID-19 era. Telecommittees should be scheduled on a regular basis (e.g., weekly), and the following specialists should participate: pathologists, radiologists, surgeons, radiation oncologists, and medical oncologists.

In this article, a prioritization of diagnostic, care, and follow-up procedures across different sarcoma contexts has been reached by consensus among sarcoma experts from different disciplines and from 11 countries among Latin American, North America, and Europe. This consensus has been developed within the SELNET consortium as a guide to protect cancer care in the complex and heterogeneous field of sarcoma during the COVID-19 outbreak. This consensus is intended to offer precise advice on different clinical sarcoma scenarios that the oncology community could face, with the aim of prioritizing or postponing different clinical decisions in patients with sarcoma. The general oncology community should network with expert centers from sarcoma suspicion, and these guidelines can be used to determine which actions can and cannot be postponed in the management of patients with sarcoma in the COVID-19 era.

Several oncology societies and health care providers issued recommendations, most often on their websites, regarding adaptive strategies for emergent standards of care in patients with cancer during the COVID-19 pandemic [28]. The Cancer Core Europe, which encompasses seven cancer centers and oncologic institutes in seven European countries, published a general consensus on the adaptive measures to minimize the number of hospital visits and to prevent anticancer treatment that could induce complications of COVID-19 infections. The methodology for these recommendations was not defined, however [29].

The NICE issued prioritization guidelines on the use of systemic and radiation treatments for cancer. This guide established six prioritization levels, from the first for curative treatments offering more than 50% chance of success and adjuvant or neoadjuvant treatment that adds at least 50% chance of cure compared with surgery or radiation therapy alone, to the last level, which is defined as a non-curative regimen with a 15%–50% chance of palliation or temporary tumor control and less than 1 year expected extension of life [30]. Similarly, a radiotherapy prioritization guide was issued with five levels. The general modifications are based on postponing or avoiding radiation therapy in cases of little added value or changing treatment plans to shorten the number of visits to a health care facility [15]. ESMO issued several guidelines, by tumor type, for prioritizing recommendations involving systemic treatments to support adaptive strategies for emergent standards of care during the COVID-19 pandemic. The intermediate profile is exemplified by a patient with a noncritical situation but one in which delay beyond 6 weeks could impair clinical outcomes [16]. The Society of Surgical Oncology has joined individual sarcoma expert opinions and has issued six recommendation points based on prioritizing actions considering several sarcoma contexts [31].

Certainly, this prioritization does not focus only on individual patient risk but also highlights the community risks and benefit: the reduction of people in transit and the isolation of patients in general decreases contagion risk in the community, leaving more resources to treat the impact of COVID-19 pandemic. However, there are consequences for other group of patients, such as patients with cancer. In The Netherlands, a remarkable drop in cancer diagnosis was noticed from pandemic initiation according to a nationwide cancer registry [32].

There are three principal approaches to conducting consensus methodology research: the consensus development panel, the nominal group process, and the Delphi technique [33]. The latter two require at least two sessions or rounds, being more complex. The consensus development panel is the most common approach used in health care research. This approach consists of organized meetings of experts in a given field and usually requires experts in different disciplines to make a multidisciplinary consensus. Usually, this approach is supported by the literature evidence, and there are some face-to-face discussions. In sarcoma oncology, ESMO guidelines follow this consensus development panel
Table 1. Prioritizing recommendations classified by higher and lower priority with consensus results for each recommendation (percentage agreeing with the recommendation)

| Higher Priority | Multidisciplinary Sarcoma Telecommittees | Lower Priority |
|-----------------|-----------------------------------------|----------------|
| **Soft-tissue, bone, or visceral sarcoma: Localized and advanced disease** | | |
| 1. Every new suspicion of sarcoma (soft tissue, bone, or visceral) diagnosis. The only exceptions being the cases likely to be well-differentiated liposarcoma or low-risk gastric GIST [Strongly recommended, 97%]. | | |
| 2. Therapeutic plan for patients previously diagnosed with any sarcoma with metastatic life-threatening lesions; Ewing sarcoma; high-grade recurrent resectable tumors; high-risk localized sarcoma; osteosarcoma; rhabdomyosarcoma (embryonal, alveolar, and sclerosing) [Strongly recommended, 100%]. | | |
| 3. Therapeutic plan for recurrent or progressing sarcomas [Strongly recommended, 97%] | | |
| **Bone sarcoma: Localized disease** | | |
| 17. Any new tumoral lesion with suspicion of malignancy [Strongly recommended, 100%]. | | |
| 18. Bone tumors without suspicion of malignancy but with risk of fracture [Strongly recommended, 100%]. | | |
| 19. Osteochondromas or GCTB with suspicion of malignant transformation [Strongly recommended, 97%]. | | |
| 20. Any suspicion of local recurrence [Strongly recommended, 100%]. | | |
| **Bone sarcoma: Advanced disease** | | |
| 21. Any new bone tumor with metastatic spread [Strongly recommended, 97%]. | | |
| 22. Any new metastatic recurrence with unexpected behavior for the tumor context [Strongly recommended, 94%]. | | |
| **GIST or other visceral sarcomas: Localized disease** | | |
| 26. Clinically evident intramural gastrointestinal lesion [Strongly recommended, 97%]. | | |
| 27. Clinical and radiological suspicion of uterine leiomyosarcoma (subserosal mass, with recent symptoms) [Strongly recommended, 97%]. | | |
| **GIST or other visceral sarcomas: Advanced disease** | | |
| 28. Any appearance of new nodules suspected of metastatic spread (except for micrometastasis or indolent). Biopsy of metastatic nodules will not be necessary in the context of consistent natural history [Recommended, 84%]. | | |
| **Soft-tissue sarcoma: Localized disease** | | |
| 32. High-risk (>40% risk for death, assessed by Sarculator) localized STS of limbs/trunk wall (after neoadjuvant treatment if indicated) [Strongly recommended, 100%]. | | |
| 33. Intermediate-risk STS (20%-40% risk for death, assessed by Sarculator) [Strongly recommended, 97%]. | | |
| **Soft-tissue sarcoma: Localized disease** | | |
| 12. Lesions not fitting with the points 1 or 2 of the higher priority [Strongly recommended, 97%]. | | |
| 13. Retroperitoneal mass with the appearance of well-differentiated liposarcoma [No consensus, only 74% agreed]. | | |
| 14. Lesions with appearance of desmoid tumors or oligosymptomatic lesions with appearance of TGCT [No consensus, 74%]. | | |
| **Soft-tissue sarcoma: Advanced disease** | | |
| 15. Lung micrometastasis (<1 cm) [Recommended, 81%]. | | |
| 16. Appearance of metastatic spread in the context of indolent tumors (i.e., EMCh, or ASPS) [No consensus, only 74% agreed]. | | |
| **Bone sarcoma: Localized disease** | | |
| 23. Bone lesions likely to be benign without symptoms or complication risks [Recommended, 87%]. | | |
| **Bone sarcoma: Advanced disease** | | |
| 24. Lung micrometastasis (<1 cm) [Recommended, 77%]. | | |
| 25. Indolent metastatic disease (i.e., adamantinoma, periosteal osteosarcoma) [Recommended, 87%]. | | |
| **GIST or other visceral sarcomas: Localized disease** | | |
| 29. Intramural lesions <1-2 cm in the gastrointestinal tract [Strongly recommended, 90%]. | | |
| 30. Uterine mass predominantly intramural, no recent history of symptoms or signs, more likely to be myofibromas [Recommended, 87%]. | | |
| **GIST or other visceral sarcomas: Advanced disease** | | |
| 31. Appearance of metastatic spread in the context of indolent GIST (PDGFRA mutant, KIT/PDGFRA wild type) [Recommended, 81%]. | | |
| **Surgery** | | |
| 40. Low-risk tumors (<20% risk for death) [well-differentiated liposarcoma; atypical liposarcoma; low-risk SFT] [Recommended, 81%]. | | |
| 41. Local recurrence of low-risk tumor [Recommended, 87%]. | | |
### Table 1. (continued)

| Higher Priority | Lower Priority |
|-----------------|----------------|
| **Soft-tissue sarcoma: Advanced disease** |
| 34. Any local recurrence of grade 2 or 3 STS of limbs/trunk wall [Strongly recommended, 97%] |
| 35. Local therapy for extraskeletal Ewing sarcoma or rhabdomyosarcoma (embryonal, alveolar, and sclerosing) [Strongly recommended, 97%] |
| 36. Any surgical complication entailing risk for the patient [Recommended, 87%] |
| 37. Retroperitoneal sarcoma (dedifferentiated liposarcoma, leiomyosarcoma, other high/intermediate grade sarcomas) [Strongly recommended, 100%] |
| 38. Metastasectomies in oligometastatic patients with an adequate time interval without progression [Strongly recommended, 94%] |
| 39. Any life-threatening resectable metastatic spread in adequate MDT discussion [Strongly recommended, 97%] |
| **Bone sarcoma: Localized disease** |
| 40. High-grade conventional osteosarcoma and skeletal Ewing sarcoma after neoadjuvant chemotherapy [Strongly recommended, 97%] |
| 41. Mesenchymal chondrosarcoma [Strongly recommended, 100%] |
| 42. High-grade or intermediate-grade osteosarcoma in local recurrence [Recommended, 87%] |
| 43. Indicated metastasectomy but due to indolent behavior of STS, this can be postponed (i.e., EMCh, ASPS) [Recommended, 87%] |
| 44. Pulmonary micronodules with uncertain malignant nature [Recommended, 84%] |
| **Bone sarcoma: Advanced disease** |
| 45. Indicated metastasectomy but due to indolent behavior, this can be postponed (i.e., adamantinoma, low-grade bone tumors) [Strongly recommended, 90%] |
| 46. In the context of unique or oligometastatic responding patients for whom extending 3-mo imatinib could be difficult [Strongly recommended, 90%] |
| 47. Multicentric GIST or with nodal involvement (i.e. KIT/PDGFRA wild type GIST) [Strongly recommended, 94%] |
| **GIST or other visceral sarcomas: Localized disease** |
| 48. Clinically evident GIST (consider neoadjuvant imatinib for gastroesophageal junction or gastric antrum or duodenal or rectal GIST or any bulky GIST to facilitate surgery) [Strongly recommended, 100%] |
| 49. Symptomatic GIST not suitable for neoadjuvant imatinib [i.e., imatinib intolerant or genotype resistant to imatinib] [Strongly recommended, 97%] |
| 50. Any other visceral grade 2 or grade 3 sarcomas or symptomatic low-grade [Strongly recommended, 100%] |
| **GIST or other visceral sarcomas: Advanced disease** |
| 51. Multicentric GIST or with nodal involvement (i.e. KIT/PDGFRA wild type GIST) [Strongly recommended, 94%] |
| 52. Any metastatic lesion causing symptomatic or other serious effect not amenable with imatinib [Strongly recommended, 97%] |

### Medical Oncology

### Soft-tissue sarcoma: Localized disease

71. Any low-grade visceral sarcoma without relevant symptoms or other complications [Recommended, 87%]

### Soft-tissue sarcoma: Advanced disease

82. Perioperative chemotherapy in localized STS of limbs/trunk wall larger than 5 cm, grade 3, and deep tumors but with risk of death less than 40% based on Sarculator would be postponed or not recommended [Strongly recommended, 90%]
Table 1. (continued)

| Higher Priority                                                                 | Lower Priority                                                                 |
|---------------------------------------------------------------------------------|--------------------------------------------------------------------------------|
| 75. Perioperative chemotherapy in high-risk STS of limbs/trunk wall (>40% death-risk based on Sarculator) [3 cycles of epirubicin + ifosfamide in selected patients] [Strongly recommended, 97%] | Soft-tissue sarcoma: Advanced disease |
| 76. Potentially resectable localized STS [No consensus, only 74% agreed]          | 83. Newly diagnosed metastatic disease with micronodules [Recommended, 84%]     |
| **Soft-tissue sarcoma: Advanced disease**                                       | 84. Indolent SSTS or slow progressive STS with barely or no symptomatic impact [Strongly recommended, 90%] |
| 77. Overtly disease progression [Strongly recommended, 94%]                     | 85. Progressive disease beyond second with low probability of clinical benefit [Recommended, 84%] |
| 78. Newly diagnosed metastatic disease > 100% or doubtfull micronodules         | **Bone sarcoma: Localized disease**                                             |
| 79. Symptomatic patients in relation to their tumor volume [Strongly recommended, 94%] | 92. Perioperative chemotherapy in high-grade bone sarcoma (i.e. Undifferentiated high-grade pleomorphic bone sarcoma) [No consensus, only 68% agreed] |
| 80. Patients with already initiated chemotherapy, or other systemic treatment with clinical or radiological benefit [Strongly recommended, 100%] | 93. Perioperative chemotherapy in high-grade chondrosarcoma [Recommended, 81%] |
| 81. Potentially resectable advanced STS [Strongly recommended, 90%]             | **Bone sarcoma: Advanced disease**                                             |
| **Bone sarcoma: Localized disease**                                            | 94. Systemic treatment beyond second line of metastatic disease in osteosarcoma or beyond third line in Ewing sarcoma [Recommended, 81%] |
| 86. Neoadjuvant chemotherapy in osteosarcoma or Ewing skeletal sarcoma          | 95. Any chemotherapy line in chondrosarcoma [Recommended, 87%]                 |
| [Strongly recommended, 94%]                                                     | 96. Imatinib in advanced/recurrent asymptomatic chordoma [Strongly recommended, 94%] |
| 87. Neoadjuvant chemotherapy in mesenchymal chondrosarcoma potentially resectable. [Recommended, 87%] | **GIST and other visceral sarcomas: Localized disease**                         |
| **Bone sarcoma: Advanced disease**                                            | 101. Adjuvant imatinib with <40% of recurrence risk (heat maps) [Recommended, 81%] |
| 88. Upfront chemotherapy in metastatic osteosarcoma or Ewing sarcoma [Recommended, 100%] | **GIST and other visceral sarcomas: Advanced disease**                         |
| [Strongly recommended, 94%]                                                     | 102. Metastatic indolent disease (i.e., PDGFRA mutant) [Recommended, 87%]      |
| 89. Chemotherapy in recurrent advanced osteosarcoma not suitable for surgery [Strongly recommended, 100%] | 103. Radiological progression without clinical impact [No consensus, 88%] |
| 90. Chemotherapy in recurrent advanced Ewing sarcoma [Strongly recommended, 100%] | **GIST and other visceral sarcomas: Advanced disease**                         |
| 91. Metastatic undifferentiated high-grade bone sarcoma [Strongly recommended, 94%] | **Bone sarcoma: Localized disease**                                            |
| **GIST and other visceral sarcomas: Localized disease**                         | 92. Perioperative chemotherapy in high-grade bone sarcoma (i.e. Undifferentiated high-grade pleomorphic bone sarcoma) [No consensus, only 68% agreed] |
| 97. Neoadjuvant imatinib in locally advanced GIST with sensitive genotype (to facilitate posterior surgery) [Strongly recommended, 94%] | 93. Perioperative chemotherapy in high-grade chondrosarcoma [Recommended, 81%] |
| 98. Neoadjuvant imatinib in gastro-esophageal junction, or gastric antrum or rectal GIST (to minimize morbidity) with sensitive genotype [Recommended, 100%] | **Bone sarcoma: Advanced disease**                                            |
| 99. Adjuvant imatinib in patients with >40% of recurrence risk (heat maps) [Strongly recommended, 100%] | 94. Systemic treatment beyond second line of metastatic disease in osteosarcoma or beyond third line in Ewing sarcoma [Recommended, 81%] |
| **GIST and other visceral sarcomas: Advanced disease**                         | 95. Any chemotherapy line in chondrosarcoma [Recommended, 87%]                 |
| 100. TKI for first, second, and third line in metastatic disease with imatinib, sunitinib, and regorafenib, respectively [Strongly recommended, 94%] | 96. Imatinib in advanced/recurrent asymptomatic chordoma [Strongly recommended, 94%] |
| **Soft-tissue sarcoma: Localized disease**                                     | **GIST and other visceral sarcomas: Advanced disease**                         |
| 104. Perioperative radiation therapy (preferably postoperative during COVID-19 outbreak) in grade 2–3, >5 cm, and deep STS of limbs/trunk wall [Strongly recommended, 97%] | **Bone sarcoma: Localized disease**                                            |
| 105. Perioperative radiation therapy (preferably postoperative during COVID-19 outbreak) in >5 cm and superficial STS of any grade in limbs/trunk wall [Recommended, 77%] | 92. Perioperative chemotherapy in high-grade bone sarcoma (i.e. Undifferentiated high-grade pleomorphic bone sarcoma) [No consensus, only 68% agreed] |
| 106. Perioperative radiation therapy (preferably postoperative during COVID-19 outbreak) in <5 cm and deep STS of any grade in limbs/trunk wall [No consensus, only 61% agreed] | 93. Perioperative chemotherapy in high-grade chondrosarcoma [Recommended, 81%] |
| 107. Radiation therapy in cases of head and neck sarcomas [Strongly recommended, 90%] | **Bone sarcoma: Advanced disease**                                            |
| 108. Radiation therapy in rhabdomyosarcomas (embryonal, alveolar, and sclerosing) and extraskeletal Ewing sarcoma [Strongly recommended, 100%] | 94. Systemic treatment beyond second line of metastatic disease in osteosarcoma or beyond third line in Ewing sarcoma [Recommended, 81%] |
| **Soft-tissue sarcoma: Advanced disease**                                       | 95. Any chemotherapy line in chondrosarcoma [Recommended, 87%]                 |
| 109. Any symptomatic metastatic lesion that could be alleviated with radiation therapy, balancing risk/benefit [Strongly recommended, 97%] | 96. Imatinib in advanced/recurrent asymptomatic chordoma [Strongly recommended, 94%] |
| **Soft-tissue sarcoma: Localized disease**                                     | **GIST and other visceral sarcomas: Localized disease**                         |
| 111. Postoperative radiation therapy in low grade, >5 cm, and deep STS of limbs or trunk wall [No consensus, 71%] | **GIST and other visceral sarcomas: Advanced disease**                         |
| **Soft-tissue sarcoma: Advanced disease**                                       | 112. Radiation therapy for local control of asymptomatic primary tumor in the context of metastatic STS [Recommended, 84%] |
Table 1. (continued)

| Higher Priority                                                                 | Lower Priority                                                                 |
|---------------------------------------------------------------------------------|-------------------------------------------------------------------------------|
| 110. Radiation therapy for symptomatic primary tumor in the context of metastasis [Strongly recommended, 100%] |                                                                                  |
| **Bone sarcoma: Localized disease**                                              |                                                                                  |
| 113. Radiation therapy is skeletal Ewing sarcoma according to CPG [Strongly recommended, 100%] |                                                                                  |
| 114. Unresectable osteosarcoma after induction chemotherapy [Strongly recommended, 94%] |                                                                                  |
| 115. Definitive radiation therapy for grade 3 chondrosarcoma [Recommended, 87%] |                                                                                  |
| **Bone sarcoma: Advanced disease**                                              |                                                                                  |
| 116. Any symptomatic metastatic lesion that could be alleviated with radiation therapy, balancing risk/benefit [Strongly recommended, 100%] |                                                                                  |
| **GIST and other visceral sarcomas: Localized disease**                          |                                                                                  |
| 117. Radiation therapy for unresectable breast or uterine sarcoma [Strongly recommended, 97%] |                                                                                  |
| 118. Any symptomatic metastatic lesion that could be alleviated with radiation therapy, balancing risk/benefit [Strongly recommended, 100%] |                                                                                  |
| **GIST and other visceral sarcomas: Advanced disease**                          |                                                                                  |
| 119. Postoperative radiation therapy in breast sarcoma larger than 5 cm or high grade [No consensus, only 65% agreed] |                                                                                  |
| 120. Radiation therapy for unresectable breast or uterine sarcoma with metastatic spread [No consensus, only 74% agreed] |                                                                                  |
| **Follow-Up**                                                                   |                                                                                  |
| 121. The follow-up recommendation for high-risk STS, conventional osteosarcoma, Ewing sarcoma, rhabdomyosarcoma (embryonal, alveolar, or sclerosing), high-risk GIST after completion of adjuvant imatinib is to schedule CT scans or chest x-ray (in some cases): every 3–4 mo for the first 3 years; every 6 mo in fourth or fifth years; every year after the fifth year [Strongly recommended, 97%] | Consider, in the appropriate context, to prolong the interval 2–3 mo if the patient is beyond the first 3 years of follow-up. The objective is to minimize patient contact with the hospital during the COVID-19 outbreak. As much as possible, teleconsultations should be used during follow-up. |
| **Clinical Trials**                                                             |                                                                                  |
| 123. To new enrollment: therapies likely to improve clinical outcome (drugs with strong preclinical rationale, drugs showing promising results in previous clinical trials, drugs with robust predictive biomarker, therapy targeting additive signaling pathway in some tumors, therapy targeting relevant signaling in orphan diseases) [Strongly recommended, 97%] |                                                                                  |
| 124. To maintain the treatment under trial: patients with clinical or radiological benefit, patients still not assessed for efficacy without relevant toxicity [Strongly recommended, 100%] |                                                                                  |

The benefit from alternating cycles of VDC (vincristine, doxorubicin, cyclophosphamide) and IE (ifosfamide and etoposide) over vincristine, doxorubicin, cyclophosphamide chemotherapy in Ewing sarcoma [39] shows level A of recommendation when applying the European Society for Medical Oncology (ESMO)-Magnitude of Clinical Benefit Scale (MCBS) V1-1: Form 1.

The benefit from VDC/IE over vincristine, ifosfamide, doxorubicin, etoposide chemotherapy in Ewing sarcoma [40] shows level B of recommendation when applying the ESMO-MCBS V1-1: Form 1.

The benefit from ifosfamide, vincristine, actinomycin D when compared with other regimens with more drugs (IVA + Carbopeto-epidexametrex in and IVA (ifosfamide, vincristine, actinomycin D) + doxorubicin) in rhabdomyosarcoma [41, 42] shows level B of recommendation when applying the ESMO-MCBS V1-1: Form 1 (IVA is the recommended option as resulted in less toxicity with the same survival outcome).

The benefit from three cycles of epirubicin and ifosfamide in the perioperative setting in high-risk patients with STS [43–45] shows level A of recommendation when applying the ESMO-MCBS V1-1: Form 1. (Advanced disease)

The benefit from eribulin in liposarcoma [46] shows level 4 of recommendation when applying the ESMO-MCBS V1-1: Form 2A (benefit in overall survival OS). The benefit from eribulin in L-sarcoma (liposarcoma and leiomyosarcoma) [46] shows level 3 of recommendation when applying the ESMO-MCBS V1-1: Form 2A. The benefit from trabectedin in L-sarcoma (liposarcoma and leiomyosarcoma) [47] shows level 3 of recommendation when applying the ESMO-MCBS V1-1: Form 2B. The benefit from trabectedin in translocation-related sarcoma [48] shows level 4 of recommendation when applying the ESMO-MCBS V1-1: Form 2A (benefit in OS). The benefit from pazopanib in pretreated STS excluding liposarcoma [49] shows level 3 of recommendation when applying the ESMO-MCBS V1-1: Form 2B.

The benefit from the combination of gemcitabine and dacarbazine in pretreated STS [50] shows level 4 of recommendation when applying the ESMO-MCBS V1-1: Form 2A (benefit in OS). The benefit from...
The combination of gemcitabine and docetaxel in advanced STS shows level 1 of recommendation when applying the ESMO-MCBS V1.1: Form 2C. The benefit from high-dose ifosfamide in advanced STS shows level 4 of recommendation when applying the ESMO-MCBS V1.1: Form 2B.

The benefit from cisplatin in advanced osteosarcoma shows level 4 of recommendation when applying the ESMO-MCBS V1.1: Form C.

The benefit from sorafenib plus everolimus and regorafenib in advanced osteosarcoma shows level 2 of recommendation when applying the ESMO-MCBS V1.1: Form C.

There are no comparative studies analyzing, for instance, the delay of some treatments in patients with sarcoma. In addition, no formal face-to-face meeting was organized to establish the consensus. Despite the proposed prioritizing approach [25, 27, 34]. The consensus we present here has two main differences in comparison with the consensus development panel. Despite the proposed prioritizing approach, there are no comparative studies analyzing, for instance, the delay of some treatments in patients with sarcoma. In addition, no formal face-to-face meeting was organized because of inherent confinement constraints. In contrast, the fact that all expert participants provided their own opinion on each recommendation ensures more independence and the quality of the conclusions, avoiding inferential differences that could arise in a face-to-face discussion.

In contrast, with the aim to minimize the subjectivity in the recommendation process, the ESMO-MCBS was fulfilled, at least for systemic therapies applied in sarcoma. We tried to adopt the grade 3 or higher and grades A or B scores, in the systemic treatment, as the cut-off for preserving the use of such treatments in the COVID-19 pandemic era. The ESMO-MCBS v1.1 is a validated scale measuring the magnitude of clinical benefit and adds value with just the statistical significance focus [18]. Although there is not yet an ESMO-MCBS for sarcoma, our exercise has followed the rules of the scale and is an additional support for this consensus.

The fact that 80 of 125 recommendations were “strongly recommended” and only 10 were “no consensus” indicates a high grade of accord among sarcoma experts in this consensus on prioritization. The fact that lower consensus was obtained in the low priority group might indicate the reticence of sarcoma experts in postponing treatment, even in indolent or low-risk cases.

This consensus has been mainly addressed keeping in mind Latin-American communities, and thus it has pursued simplicity and concision, taking into consideration that there are many health care providers in each country. Hence, assigning higher or lower priority to those undelayable or delayable treatments, respectively, facilitates the decision-making process in patients with sarcoma. Additionally, this SELNET consensus issues 125 recommendations, which indicates a high level of thoroughness, covering a substantial spectrum of sarcoma care.

Follow-up recommendations in the context of sarcoma remains largely unstudied [35] and usually are based on conventions; thus, in high-risk patients, for example, the imaging tests are performed every 3–4 months for the first 3 years, then every 6 months for the 4th and 5th years, and so on, once per year. This strategy is based on the higher risk of recurrence observed in the first 3 years after treatment for localized disease and the fact that the asymptomatic recurrence, detected by computed tomography scan for instance, could potentially have higher curative options. There are some reports addressing the relevance of smaller size as independent prognostic relevance, at the time of metastatic disease, for longer survival [36–38]. Nevertheless, the truth is that convincing evidence that determined strategy is related with better survival is lacking, and a lead-time bias can occur in highly interventionist follow-up.
Summary

This SELNET consensus provides a tool for multidisciplinary sarcoma committees during the COVID-19 outbreak. The detail of different recommendations and the distinction between the two levels of prioritization enables a practical approach for Latin-American health care providers and sarcoma expert centers.

Table 2. Mann-Whitney U and Pearson’s χ² test for those recommendations with statistical differences between LATAM and E.U.-U.S. experts

| Recommendations | Mann-Whitney test | Pearson’s χ² test |
|-----------------|-------------------|------------------|
|                 | LATAM, % | E.U.-U.S., % | p value | LATAM, % | E.U.-U.S., % | p value |
| Recommendation no. 93 |                      |                 | .001 |                      |                |
| Strongly agree   | 38       | 7             |       | Agree       | 57       | 100   | .009 |
| Agree           | 31       | 64            |       | Disagree    | 43       | 0     |       |
| Disagree        | 31       | 22            |       | Strongly disagree | 0    | 7     |       |
| Recommendation no. 105 |                      |                 | .008 |                      |                |
| Strongly agree   | 25       | 64            |       | Agree       | 100      | 63    |       |
| Agree           | 38       | 36            |       | Disagree    | 0        | 37    |       |
| Disagree        | 31       | 0             |       | Strongly disagree | 0    | 0     |       |
| Recommendation no. 111 |                      |                 | .021 |                      |                |
| Strongly agree   | 41       | 21            |       | Agree       | 43       | 94    | .002 |
| Agree           | 53       | 21            |       | Disagree    | 57       | 6     |       |
| Disagree        | 6        | 58            |       | Strongly disagree | 0    | 0     |       |

Conclusion

This SELNET consensus provides a tool for multidisciplinary sarcoma committees during the COVID-19 outbreak. The detail of different recommendations and the distinction between the two levels of prioritization enables a practical approach for Latin-American health care providers and sarcoma expert centers.

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REFERENCES

1. Lazzerini M, Putoto G. COVID-19 in Italy: Momentous decisions and many uncertainties. Lancet Glob Health 2020;8:e641–e642.
2. Moujaes E, Kourie HR, Ghosn M. Cancer patients and research during COVID-19 pandemic: A systematic review of current evidence. Crit Rev Oncol Hematol 2020;150:102972.
3. Mittal D, Gubin MM, Schreiber RD et al. New insights into cancer immunodecting and its three component phases—Elimination, equilibrium and escape. Curr Opin Immunol 2014;27:16–25.
4. Kamboj M, Sepkowitz KA. Nosocomial infections in patients with cancer. Lancet Oncol 2009; 10:S89–S97.
5. Mehta P, McAuley DF, Brown M et al. COVID-19: Consider cytokine storm syndromes and immunosuppression. Lancet 2020;395:1033–1034.
6. Huang C, Wang Y, Li X et al. Clinical features of patients infected with 2019 novel coronavirus in Wuhan, China. Lancet 2020;395:497–506.
7. Zheng HY, Zhang M, Yang CK et al. Elevated exhaustion levels and reduced functional diversity of T cells in peripheral blood may predict severe progression in COVID-19 patients. Cell Mol Immunol 2020;17:541–543.
8. Liang W, Guan W, Chen R et al. Cancer patients in SARS-CoV-2 infection: A nationwide analysis in China. Lancet Oncol 2020;21:335–337.
9. Williamson E, Walker AJ, Bhaskaran KJ et al. OpenSAFELY: Factors associated with COVID-19-related hospital death in the linked electronic health records of 17 million adult NHS patients. medRxiv Preprint posted online May 7, 2020. doi: https://doi.org/10.1101/2020.05.06.20092999
10. Lee LYW, Cazier JB, Angelis V et al. COVID-19 mortality in patients with cancer: chemotherapy or other anticancer treatments: A nationwide electronic health records of 17 million adult NHS patients. medRxiv Preprint posted online May 7, 2020. doi: https://doi.org/10.1101/2020.05.06.20092999
11. Al-Shamsi HO, Alhazzani W, Alhuraiji A et al. A practical approach to the management of cancer patients during the novel coronavirus disease 2019 (COVID-19) pandemic: An international collaborative group. The Oncologist 2020;25: e936–e945.
12. de Joode K, Dumoulin DW, Engelen V et al. Impact of the coronavirus disease 2019 pandemic on cancer treatment: The patients’ perspective. Eur J Cancer 2020;136:132–139.
13. Hanna TP, Evans GA, Booth CM. Cancer, COVID-19 and the precautionary principle: Prioritizing treatment during a global pandemic. Nat Rev Clin Oncol 2020;17:268–270.
14. Selnet. Available at https://selnet-h2020.org/. Accessed May 12, 2020.
15. COVID-19 rapid guideline: Delivery of radiotherapy. National Institute for Health and Care Excellence. Available at https://www.nice.org.uk/guidance/ng162. Accessed May 12, 2020.
16. ESMO-EURACAN Management and Treatment Adapted Recommendations in the COVID-19 Era: Sarcomas. Available at https://www.esmo.org/guidelines/cancer-patient-management-during-the-covid-19-pandemic/sarcomas-in-the-covid-19-era. Accessed May 12, 2020.
17. Diamond IR, Grant RC, Feldman BM et al. Defining consensus: A systematic review recommends methodologic criteria for reporting of Delphi studies. J Clin Epidemiol 2014;67:401–409.
18. Cherry NJ, Dafni U, Bogaerts J et al. ESMO-Magnitude of Clinical Benefit Scale version 1.1. Ann Oncol 2017;28:2340–2366.
19. Cherry NJ, Sullivan R, Dafni U et al. A standardised, generic, validated approach to stratify the magnitude of clinical benefit that can be anticipated from anti-cancer therapies: The European Society for Medical Oncology Magnitude of Clinical Benefit Scale (ESMO-MCBS). Ann Oncol 2015;26:1547–1573.
20. Bauer HC, Trovik CS, Alvegråd TA et al. Monitoring referral and treatment in soft tissue sarcoma: Study based on 1,851 patients from the Scandinavian Sarcoma Group Register. Acta Orthop Scand 2001;72:150–159.
21. Mathoulin-Pélissier S, Chevreau C, Bellera C et al. Adherence to consensus-based diagnosis and treatment guidelines in adult soft-tissue sarcoma patients: A French prospective population-based study. Ann Oncol 2014;25:225–231.
22. Ray-Coquard I, Thiesse P, Ranchère-Vincen D et al. Conformity to clinical practice guidelines, multidisciplinary management and outcome of treatment for soft tissue sarcomas. Ann Oncol 2004;15:307–315.
23. Hoekstra HI, Haas RLM, Verhoef C et al. Adherence to Guidelines for Adult (Non-GIST) soft tissue sarcoma in The Netherlands: A plea for dedicated sarcoma centers. Ann Surg Oncol 2017;24:3279–3288.
24. Martin-Brito J, Hindi N, Cruz J et al. Relevance of reference centers in sarcoma care and quality item evaluation: Results from the prospective registry of the Spanish Group for Research in Sarcoma (GEIS). The Oncologist 2019;24:e338–e346.
25. Casali PG, Bielack S, Abeccasis N et al. Bone sarcomas: ESMO-PaedCancer-EURACAN Clinical Practice Guidelines for diagnosis, treatment and follow-up. Ann Oncol 2018;29:iv79–iv95.
26. Casali PG, Abeccasis N, Aro HT et al. Soft tissue and visceral sarcomas: ESMO-EURACAN Clinical Practice Guidelines for diagnosis, treatment and follow-up. Ann Oncol 2018;29:iv68–iv78.
27. Casali PG, Abeccasis N, Aro HT et al. Gastrointestinal stromal tumours: ESMO-EURACAN Clinical Practice Guidelines for diagnosis, treatment and follow-up. Ann Oncol 2018;29(Suppl 4):iv68–iv78.
28. Burki TK. Cancer guidelines during the COVID-19 epidemic. The Oncologist 2020;25:701–706.
29. van de Haar J, Hoes LR, Coles CE et al. Car- ing for patients with cancer in the COVID-19 era. Nat Med 2020;26:665–671.
30. COVID-19 rapid guideline: Delivery of systemic anticancer treatments. National Institute for Health and Care Excellence. Available at https://www.nice.org.uk/guidance/ng161. Accessed May 14, 2020.
31. Resource for Management Options of Sarcoma During COVID-19. Society of Surgical Oncology. Available at https://www.surgonc.org/wp-content/uploads/2020/03/Sarcoma-Resource-during-COVID-19-3.30.20.pdf. Accessed May 14, 2020.
32. Dimmohamed AG, Visser O, Verhoeven RHA et al. Fewer cancer diagnoses during the COVID-19 epidemic in The Netherlands. Lancet Oncol 2020;21:750–751.
33. Waggoner J, Carlile JD, Durning SJ. Is there a consensus on consensus methodology? Descriptions and recommendations for future consensus research. Acad Med 2016;91:663–668.
34. Casali PG, Abeccasis N, Bauer S et al. Soft tissue and visceral sarcomas: ESMO-EURACAN Clinical Practice Guidelines for diagnosis, treatment and follow-up. Ann Oncol 2018;29(suppl 4):iv51–iv67.
35. Rothermundt C, Whelan JS, Dilop P et al. What is the role of routine follow-up for localised limb soft tissue sarcomas? A retrospective analysis of 174 patients. Br J Cancer 2014;110:2260–2262.
36. Lindner LH, Litière S, Sleijfer S et al. Prognostic factors for soft tissue sarcoma patients with lung metastases only who are receiving first-line chemotherapy: An exploratory, retrospective analysis of the European Organization for Research and Treatment of Cancer-Soft Tissue and Bone Sarcoma Group (EORTC-STBSG). Int J Cancer 2018;142:2610–2620.
37. Hompland I, Bruland JS, Holmebak T et al. Prediction of long-term survival in patients with metastatic gastrointestinal stromal tumor: Analysis of a large, single-institution cohort. Acta Oncol 2017;56:1317–1323.
38. Rutkowski P, Andrzejuk J, Blyja E et al. What are the current outcomes of advanced gastrointestinal stromal tumors: Who are the long-term survivors treated initially with imatinib? Med Oncol 2013;30:765.
39. Grier HE, Krailo MD, Tarbell NJ et al. Addition of ifosfamide and etoposide to standard chemotherapy for Ewing’s sarcoma and primitive neuroectodermal tumor of bone. N Engl J Med 2003;348:694–701.
40. Wheatley K, Moroz V, Marec-Berard P et al. First results of the EURO EWING 2012 (EE2012) trial comparing two chemotherapy regimens in newly diagnosed Ewing Sarcoma (ES). Poster presented at CTOS Annual meeting, Tokyo, Japan: November 13–16, 2019:48.
41. Oberlin O, Rey A, Sanchez de Toledo J et al. Randomized comparison of intensified six-drug
versus standard three-drug chemotherapy for high-risk nonmetastatic rhabdomyosarcoma and other chemotherapy-sensitive childhood soft tissue sarcomas: Long-term results from the International Society of Pediatric Oncology MMT95 study. J Clin Oncol 2012;30:2457–2465.

42. Bisogno G, Jenney M, Bergeron C et al. Addition of dose-intensified doxorubicin to standard chemotherapy for rhabdomyosarcoma (EpSSG RMS 2005): A multicentre, open-label, randomised controlled, phase 3 trial. Lancet Oncol 2018;19:1061–1071.

43. Frustaci S, Gherlinzoni F, De Paoli A et al. Adjuvant chemotherapy for adult soft tissue sarcomas of the extremities and girdles: Results of the Italian randomized cooperative trial. J Clin Oncol 2001;19:1238–1247.

44. Gronchi A, Ferrari S, Quagliullo V et al. Histotype-tailored neoadjuvant chemotherapy versus standard chemotherapy in patients with high-risk soft-tissue sarcomas (ISG-STS 1001): An international, open-label, randomised, controlled, phase 3, multicentre trial. Lancet Oncol 2017;18:812–822.

45. Gronchi A, Palmerini E, Quagliullo V et al. Neoadjuvant chemotherapy in high-risk soft tissue sarcomas: Final results of a randomized clinical trial from the Italian (ISG), Spanish (GEIS), French (FSG), and Polish (PSG) Sarcoma Groups. J Clin Oncol 2020;38:2178–2186.

46. Schöffski P, Chwalba S, Maki RG et al. Eribulin versus dacarbazine in previously treated patients with advanced liposarcoma or leiomyosarcoma: A randomised, open-label, multicentre, phase 3 trial. Lancet 2016;387:1629–1637.

47. Demetri GD, von Mehren M, Jones RL et al. Efficacy and safety of trabectedin or dacarbazine for metastatic liposarcoma or leiomyosarcoma after failure of conventional chemotherapy: Results of a phase III randomized multicentre clinical trial. J Clin Oncol 2016;34:786–793.

48. Kawai A, Araki N, Sugiiura H et al. Trabectedin monotherapy after standard chemotherapy versus best supportive care in patients with advanced, translocation-related sarcoma: A randomised, open-label, phase 2 study. Lancet Oncol 2015;16:406–416.

49. van der Graaf WT, Blay JY, Chwalba SP et al. Pazopanib for metastatic soft-tissue sarcoma (PALETTE): A randomised, double-blind, placebo-controlled phase 3 trial. Lancet 2012;379:1879–1886.

50. García-Del-Muro X, López-Pousa A, Maurel J et al. Randomized phase II study comparing gemcitabine plus dacarbazine versus dacarbazine alone in patients with previously treated soft tissue sarcoma: A Spanish Group for Research on Sarcomas study. J Clin Oncol 2011;29:2528–2533.

51. Maki RG, Watthen JX, Patel SR et al. Randomized phase II study of gemcitabine and docetaxel compared with gemcitabine alone in patients with metastatic soft tissue sarcomas: Results of sarcoma alliance for research through collaboration study 002 [corrected]. J Clin Oncol 2007;25:2755–2763.

52. Pautier P, Floquet A, Penel N et al. Randomized multicenter and stratified phase II study of gemcitabine alone versus gemcitabine and docetaxel in patients with metastatic or relapsed leiomyosarcomas: A Federation Nationale des Centres de lutte contre le Cancer (FNCLCC) French Sarcoma Group Study (TAXOvEM study). The Oncologist 2012;17:1213–1220.

53. Buesa JM, López-Pousa A, Martín J et al. Phase II trial of first-line high-dose ifosfamide in advanced soft tissue sarcomas of the adult: A study of the Spanish Group for Research on Sarcomas (GEIS). Ann Oncol 1998;9:871–876.

54. Benjamin RS, Wiernik PH, Bachur NR. Adriamycin: A new effective agent in the therapy of disseminated sarcomas. Med Pediatr Oncol 1975;1:63–76.

55. Eilber F, Giuliani A, Eckardt J et al. Adjuvant chemotherapy for osteosarcoma: A randomized prospective trial. J Clin Oncol 1987;5:21–26.

56. Bernthal NM, Federman N, Eilber FR et al. Long-term results (>25 years) of a randomized, prospective clinical trial evaluating chemotherapy in patients with high-grade, operable osteosarcoma. Cancer 2012;118:5988–5993.

57. Meyers PA, Schwartz CL, Krakó MD et al. Osteosarcoma: The addition of muramyl tripeptide to chemotherapy improves overall survival—a report from the Children’s Oncology Group. J Clin Oncol 2008;26:633–638.

58. Ochs JJ, Freeman AI, Douglass HO, Jr et al. cis-Dichlorodiammineplatinum (II) in advanced osteogenic sarcoma. Cancer Treat Rep 1978;62:239–245.

59. Voute PA, Souhami RL, Nooj M et al. A phase II study of cisplatin, ifosfamide and doxorubicin in operable primary, axial skeletal and metastatic osteosarcoma. European Osteosarcoma Intergroup (EOI). Ann Oncol 1999;10:1211–1218.

60. Gentet JC, Brunat-Mentigny M, Demeaille MC et al. Ifosfamide and etoposide in childhood osteosarcoma. A phase II study of the French Society of Paediatric Oncology. Eur J Cancer 1997;33:232–237.

61. Berrak SG, Pearson M, Berberoglu S et al. High-dose ifosfamide in relapsed pediatric osteo- sarcoma: Therapeutic effects and renal toxicity. Pediatr Blood Cancer 2005;44:215–219.

62. Grignani G, Palmerini E, Ferrari V et al. Sorafenib and everolimus for patients with unresectable high-grade osteosarcoma progressing after standard treatment: a non-randomised phase 2 clinical trial. Lancet Oncol 2015;16:98–107.

63. Duffaud F, Mir O, Boudou-Rouquette P et al. Efficacy and safety of regorafenib in adult patients with metastatic osteosarcoma: A non-comparative, randomised, double-blind, placebo-controlled, phase 2 study. Lancet Oncol 2019;20:120–133.

64. McCabe MG, Moroz V, Khan M et al. Results of the first interim assessment of rEECuR, an international randomized controlled trial of chemotherapy for the treatment of recurrent and primary refractory Ewing sarcoma. J Clin Oncol 2019;37:11007a.

65. Mora J, Castañeda A, Perez-Jaume S et al. GEIS-21: A multicentric phase II study of intensive chemotherapy including sorafenib and docetaxel for the treatment of Ewing sarcoma of children and adults: A report from the Spanish sarcoma group (GEIS). Br J Cancer 2017;117:767–774.

66. Meazza C, Casanova M, Luksh R et al. Prolonged 14-day continuous infusion of high-dose ifosfamide with an external portable pump: Feasibility and efficacy in refractory pediatric sarcoma. Pediatr Blood Cancer 2010;55:617–620.

67. Joensuu H, Eriksson M, Sundby Hall K et al. One vs three years of adjuvant imatinib for operable gastrointestinal stromal tumour: A randomised trial. JAMA 2012;307:1265–1272.

68. van Oosterom AT, Judson I, Verweij J et al. Safety and efficacy of imatinib (STI571) in metastatic gastrointestinal stromal tumours: A phase I study. Lancet 2001;358:1421–1423.

69. Demetri GD, van Oosterom AT, Garrett CR et al. Efficacy and safety of sunitinib in patients with advanced gastrointestinal stromal tumour after failure of imatinib: A randomised controlled trial. Lancet 2006;368:1329–1338.

70. Demetri GD, Garrett CR, Schöffski P et al. Complete longitudinal analyses of the randomized, placebo-controlled, phase III trial of sunitinib in patients with gastrointestinal stromal tumour following imatinib failure. Clin Cancer Res 2012;18:3170–3179.

71. Demetri GD, Reichardt P, Kang YK et al. Efficacy and safety of regorafenib for advanced gastrointestinal stromal tumours after failure of imatinib and sunitinib (GRID): An international, multicentre, randomised, placebo-controlled, phase 3 trial. Lancet 2013;381:295–302.

72. von Mehren M, Serrano C, Bauer S et al. INVICTUS: A phase 3, interventional, double-blind, placebo-controlled study to assess the safety and efficacy of ripretinib as 4th-line therapy in patients with advanced gastrointestinal stromal tumours (GIST) who have received treatment with prior anticancer therapies (nct03357535). Ann Oncol 2019;30(suppl 5):v851–v934.