Sarcome-13/OS2016 trial protocol: a multicentre, randomised, open-label, phase II trial of mifamurtide combined with postoperative chemotherapy for patients with newly diagnosed high-risk osteosarcoma

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ABSTRACT

Introduction The controversial results on the mifamurtide efficacy associated with chemotherapy, issued from the American INT-0133-study, in localised osteosarcomas, and the underpowered analysis performed separately in metastatic patients, should be clarified to homogenise international use of this promising drug. The European Commission has granted a marketing authorisation to mifamurtide combined with postoperative chemotherapy in localised osteosarcomas but not in metastatic patients, while the Food and Drug Administration (FDA) has denied this authorisation.

Methods and analysis Sarcome-13/OS2016 trial is a multicentre randomised open-label phase II trial evaluating the survival benefit of mifamurtide administered during 36 weeks in combination with postoperative chemotherapy versus chemotherapy alone, in patients >2 and ≤50 years with newly diagnosed high-risk localised or metastatic osteosarcoma. The main objective is to evaluate the impact on event-free survival (EFS) of mifamurtide on intention-to-treat population. The secondary objectives are to evaluate the impact of mifamurtide on overall survival, to evaluate the feasibility and toxicity of the planned treatment, to correlate biology/immunology with the mifamurtide efficacy/toxicity. With a total of 126 enrolled patients and 51 events, the power is 80% if mifamurtide is associated with an 18% improvement of the 3-year EFS (52% vs 70%, equivalent to an HR=0.55), with a one-sided logrank test alpha=10%. As relevant historical data are available (aggregate treatment effect from the INT-0133 trial and individual data from the control group of the Sarcome-09/OS2006 trial), a Bayesian analysis is also planned.

Strengths and limitations of this study

- Multicentre, randomised, phase II trial design to evaluate the efficacy of mifamurtide combined with postoperative chemotherapy, in children and adults with high-risk osteosarcoma.
- By using ifosfamide-based chemotherapy for all patients, we avoid the possible issue in interpreting the data due to an interaction between mifamurtide and chemotherapy regimen (with or without ifosfamide) as published in the original article of Meyers et al. (J Clin Oncol).
- The relatively small sample size may limit the findings, but it fits with the rare disease setting and a realistic time frame.
- Incorporating historical data (individual control data and aggregate treatment effect) using a power and mixture prior for a Bayesian survival augments the design and analysis of the Sarcome-13/OS2016 trial.

Trial registration number EudraCT 2017-001165-24, NCT03643133

INTRODUCTION

Epidemiology and prognosis of osteosarcoma

Osteosarcoma is the most common primary malignant bone tumour with a peak incidence in adolescents and young adults (AYA). However, in France, this rare disease represents less than 10% of all AYA cancers with 150 new cases/year, including adults. Osteosarcoma treatment relies on preoperative chemotherapy, tumour surgical resection and postoperative chemotherapy. With this multidisciplinary approach, the 3-year event-free survival (EFS) rate reaches 70% in patients with non-metastatic osteosarcoma.
In France, the chemotherapy regimen used for treatment depends on age: children/adolescents are treated with the combination of methotrexate, etoposide and ifosfamide (M-EI), while adults receive a combination of doxorubicin, cisplatin and ifosfamide (API-AI). These regimens formed the backbone of preoperative chemotherapy of the recent Sarcome-09/OS2006 trial. Postoperative chemotherapy is adapted to the metastatic status at diagnosis and, for patients with a localised disease, to the histological response to neoadjuvant chemotherapy according to Huvos' grading.

The main risk factors of relapse are the presence of metastases at diagnosis, a non-operative disease and poor histological response to preoperative chemotherapy, usually defined by ≥10% residual tumour cells. In the Sarcome-09/OS2006 trial, patients with skip metastasis had a poorer outcome than patients with localised disease, close to patients with metastatic disease, and we considered them as metastatic even if the skip metastasis is the unique metastatic lesion. The 3-year EFS in these high-risk patients is around 40%, which has not improved in the last 20 years despite several clinical trials. To address this issue, new treatment strategies for patients with high-risk osteosarcoma are needed.

**Mifamurtide (L-MTP-PE, MEPACT) in osteosarcoma**

Mifamurtide is a fully synthetic lipophilic derivative of the muramyl dipeptide (MDP), encapsulated into liposomes. It binds to extracellular Toll-like receptor-4, activating monocytes and macrophages, and promotes antitumour activity. However, mifamurtide also activates antitumour function of macrophages by intracellular nucleotide-binding oligomerisation domain-2 (NOD) receptor. Mifamurtide enters into macrophages, is degraded into MDP and binds to NOD-2 receptors. It induces NF-κB release through activation of receptor-interacting serine/threonine protein kinases (RICK) signaling pathway and secretion of inflammatory cytokines such as TNFα, IL-6, IL-8 and IL-1. Preclinical studies showed mifamurtide has similar immunostimulatory effects on these cells as the natural MDP, with the additional advantage of a longer half-life in plasma, lower toxicity and better efficacy. The activation of the monocytes/macrophages and dendritic cells may also activate other innate immune cells such as natural killer cells, and may generate an adaptive immune response by T cells. Human monocytes/macrophages, after in vitro activation with mifamurtide, specifically recognised tumour cells and were not cytotoxic to normal cells. In vivo administration of mifamurtide resulted in the inhibition of tumour growth in mouse and rat models of lung metastasis, skin and hepatic cancer, and fibrosarcoma. Interest in mifamurtide for the treatment of osteosarcoma arises from the remarkable results achieved in dogs, in which spontaneous osteosarcoma is common, and similar to the human disease. Randomised studies in dogs of either mifamurtide alone, or in combination with chemotherapy, showed a significantly improved survival in dogs treated with mifamurtide. Results in animal tumour models do not support a significant interaction of most chemotherapeutic drugs on macrophage activation by mifamurtide.

**Trial rationale**

From 1993 to 1997, the randomised, phase III INT-0133 trial, considering patients with newly diagnosed osteosarcoma, and younger than 30, addressed two questions in a factorial design: efficacy of ifosfamide in addition to MAP chemotherapy (methotrexate–doxorubicin–cisplatin); and efficacy of mifamurtide in addition to postoperative chemotherapy. In the 662 patients enrolled with a localised resectable osteosarcoma, the first analysis showed mifamurtide might improve EFS, but a potential interaction between ifosfamide and mifamurtide hampered the results interpretation. A second publication on the same population with a longer follow-up reported a significant benefit of mifamurtide on overall survival (OS) (HR 0.71, 95% CI 0.52 to 0.96, p=0.03) and a 6-year survival improvement rate from 70% to 78%, with no significant ifosfamide/mifamurtide interaction. In addition, 91 patients with synchronous metastatic osteosarcoma recruited in this trial were analysed separately: mifamurtide effect size on OS was similar in this group (HR 0.72, 95% CI 0.40 to 1.30), but not significant (p=0.27).

A retrospective study on metastatic/relapsed osteosarcoma confirmed mifamurtide safety. The most frequent side effects of mifamurtide are chills, fever, fatigue, nausea, tachycardia and headache with mild to moderate grade. Mifamurtide combined with nephrotoxic (cisplatin, ifosfamide) and hepatotoxic (high-dose methotrexate) chemotherapies in the INT-0133 trial did not increase these toxicities.

Based on these results, the European Medicines Agency granted a centralised marketing authorisation on 6 March 2009 for mifamurtide combined with postoperative chemotherapy for patients between 2 and 30 years old and presenting a high-grade non-metastatic osteosarcoma with complete macroscopic resection (EU/1/08/502/001). However, in the USA, no approval has been obtained. In France, the French Transparency Commission required additional investigation before considering mifamurtide as a standard in front-line therapy of osteosarcoma. In several European countries, mifamurtide use is still limited as the results of the INT-0133 trial have been a matter of debate between key-opinion leaders. In addition, European Union authorisation does not include patients over 30, as well as patients with primary metastatic disease for whom outcome improvement is still challenging.

**Aim of the study**

In this context, further investigations of first-line treatment with mifamurtide in osteosarcoma are necessary, especially for patients with a high risk of relapse (metastases at diagnosis or poor histological response to neoadjuvant chemotherapy). As the current postoperative chemotherapy regimens used for the treatment of osteosarcoma...
in France contain ifosfamide, the possible interaction with mifamurtide would not jeopardise the evaluation of the benefit associated with mifamurtide. Sarcome-13/OS2016 trial intends to evaluate the survival benefit of mifamurtide combined with postoperative chemotherapy for treating patients with high-risk osteosarcoma.

EFS will be used as a primary endpoint. We will also study the impact of mifamurtide on OS, and the feasibility of the planned treatment with calculation of cumulative dose and dose intensity of mifamurtide and chemotherapy. Acute and long-term toxicity will be evaluated, as well as biomarkers that could be surrogate markers of mifamurtide pharmacological effect, or predictive factors of efficacy and/or toxicity of mifamurtide. If the efficacy of mifamurtide is confirmed, this may allow the use of a currently promising, original drug, in patients with osteosarcoma with the highest risk of failure.

This article reports the Sarcome-13/OS2016 trial protocol V.2.0 of the 28 May 2018.

METHODS AND ANALYSIS
Study design
Sarcome-13/OS2016 is a French, multicentre, randomised, open-label, phase II trial, with two parallel groups in first-line treatment of high-risk patients with osteosarcoma. This trial is part of a study recruiting all groups in first-line treatment of high-risk patients with randomised, open-label, phase II trial, with two parallel groups: (1) localised disease at diagnosis and poor histological response, (2) metastatic disease at diagnosis (skip or distant metastases) and planned complete removal of lesions and poor histological response of the primary tumour, (3) metastatic disease at diagnosis (skip or distant metastases) and planned complete removal of lesions, but good histological response of the primary tumour, (4) metastatic disease at diagnosis and complete removal of lesions before randomisation and poor histological response of the primary tumour, (5) metastatic disease at diagnosis and planned complete removal of lesions and poor histological response of the primary tumour.

Since osteosarcoma is a rare disease, a large phase III trial with standard alpha level cannot be performed within a reasonable time frame. A randomised phase II trial evaluating the efficacy of mifamurtide at a relaxed alpha level in high-risk patients will produce new independent data, which may be combined with results from the previous INT-0133 trial.

Patient selection criteria
Registration criteria at diagnosis
The following criteria must be met at diagnosis for registration in the study:
1. All newly diagnosed, biopsy-proven, high-grade osteosarcoma, whatever the initial extension of the disease.
2. Age 2–50.
3. Normal haematological, renal, cardiac and hepatic functions.
4. Planned neoadjuvant chemotherapy as follows:
   a. M-EI regimen for patients ≤25 years old.
   b. API-AI regimen for patients 26–50 years old.
5. Written informed consent from patients and/or their parents/guardians before enrolment and any study-related procedure.
6. Affiliation to a social insurance regimen.

Inclusion criteria for randomisation
Patients must meet the following criteria:
1. Histologically proven, confirmed by expert pathologists panel, high-grade osteosarcoma.
2. Registered at diagnosis into the study.
3. Primary tumour resected after preoperative chemotherapy.
4. Osteosarcoma classified as high risk because of at least one risk factor:
   a. Presence of distant metastases or skip metastases at diagnosis.
   b. Poor histological response to preoperative chemotherapy (>10% residual viable cells on the analysis of the primary tumour surgical specimen).
5. Preoperative chemotherapy combining
   a. M-EI regimen for patients ≤25 years old.
   b. API-AI regimen for patients 26–50 years old.
6. Screening laboratory values must meet the following criteria (using Common Terminology Criteria for Adverse Events [CTCAE], V.5) and should be obtained within 7 days prior to randomisation:
   a. Absolute neutrophil count ≥1×10⁹/L.
   b. Platelets ≥100×10⁹/L.
   c. Haemoglobin ≥8g/dL.
   d. Alanine aminotransferase (ALT) and Aspartate aminotransferase (AST) ≤2.5×ULN in the absence of liver metastases or ≤5×ULN in the presence of liver metastases.
Non-inclusion criteria for randomisation

Patients with any of the following conditions are not included in the study:

1. Low-grade osteosarcoma, parosteal or periosteal osteosarcoma.
2. Prior history of other malignancies other than study disease unless the patient has been free of the disease for at least 3 years.
3. Osteosarcoma with multiple metastases for whom complete removal is not expected to be feasible even after shrinkage with chemotherapy.
4. Progressive disease at any site during initial chemotherapy, confirmed before randomisation time and not totally resected during surgery.
5. Any medical condition precluding treatment with protocol chemotherapy.
6. Fractional Shortening <28% or Left Ventricular Ejection Fraction (LVEF) <50% before treatment (only for API postoperative chemotherapy) by echo-cardiogram or multiple-gated acquisition (MUGA) scan.
7. Pregnancy or breast feeding
8. Hypersensitivity to the active substance or to any of the excipients.
9. Concurrent use of immunodepressive treatment such as cyclosporine, tacrolimus or other calcineurin inhibitors.
10. Concurrent use with high-dose non-steroidal anti-inflammatory drugs.
11. Inflammatory or autoimmune disease, allergy or asthma requiring a chronic use of steroid treatment that cannot be stopped.
12. Patients with positive test for HIV or known AIDS.
13. Patients with positive tests for hepatitis-B virus surface antigen or hepatitis-C virus RNA indicating active or chronic infection.

Study description

Intervention

After preoperative chemotherapy and surgery of the primary tumour, patients presenting high-risk osteosarcoma will be randomised to receive the treatment allocated, either:

- Arm-A: Postoperative chemotherapy alone, according to age group.
- Arm-B: Postoperative chemotherapy, according to age group + mifamurtide.

Postoperative chemotherapy is part of standard of care for high-risk osteosarcoma. Chemotherapy should be administered as per local practice at each investigator site. The choice of the chemotherapy regimen is based on experience from Sarcome-09/OS2006 trial, and treatments are determined according to patient’s age (figure 1):
and biological tests before each treatment course, every 3 weeks. Radiological, and echocardiography or cardiac MUGA scintigraphy, and audiogram will also be performed. During the mifamurtide period without chemotherapy, patients will be seen in clinic every week by the physician (before each mifamurtide infusion) and will have standard clinical assessment. They will also have a formal visit every 3 months. Follow-up assessment of the disease will be performed (from last treatment administration) every 3 months for 3 years, then every 6 months for 2 years, then every year up to 10 years for all patients. Follow-up assessment of late treatment effects will be performed 1 and 3 years after the end of postoperative chemotherapy.

All medications for the treatment of symptoms are authorised, and their type, posology and duration of administration will be recorded. Patients should receive full supportive care during the study, including transfusions and analgesics, as appropriate.

As mifamurtide has a lipophilic formulation, it is recommended to separate the times of administration of mifamurtide and doxorubicin, or other lipophilic drugs.

Prohibited concomitant treatments are:
- Immuno depressive treatment.
- High-dose non-steroidal anti-inflammatory drugs.
- Chronic use of steroid treatment.
- Systematic use of steroids as antiemetic prophylaxis.

Premature treatment discontinuation may be due to the following reasons: disease progression, unacceptable toxicity, intercurrent conditions that preclude the continuation of treatment, patient choice or physician decision. Except in case of consent withdrawal for the participation in the study, patient follow-up will continue in compliance with the protocol and follow-up data will be collected until the end of the trial.

**Outcome measures**

**Study objectives**

The primary objective of the trial is to evaluate the survival benefit (EFS) of mifamurtide administered during 36 weeks combined with postoperative chemotherapy compared with postoperative chemotherapy alone. The secondary objectives are to evaluate: (1) the impact on OS of mifamurtide, (2) the feasibility and safety of mifamurtide administration during and after postoperative chemotherapy, (3) the mifamurtide effect on antitumour immunity in patients with sequential surgery of lung metastases, (4) biomarkers that could be surrogate of mifamurtide pharmacological effect or predictive factors of efficacy and/or toxicity of mifamurtide, (5) the tumour microenvironment in osteosarcoma and its correlation with clinical characteristics and outcome and (v6) the potential new therapeutic targets for future combinations (Whole Exon Sequencing, RNAseq, Immune-Histo-Chemistry, flow cytometry, ELISA).

**Statistical considerations**

**Sample size**

Due to the rare disease setting, we relaxed the alpha level of the one-sided log-rank test to 10%, and a pragmatic recruitment target has been set of accruing 126 patients over 3 years. If this target is met, the power is 80% if the true HR=0.55 (18% improvement of the 3-year EFS, 52% vs 70%), which requires 51 events if the analysis is performed at the end of study. The minimum follow-up will be 2 years from randomisation, leading to a total duration of the trial of 5 years. A long-term follow-up is also planned beyond the primary analysis, up to 10 years from randomisation. The sample size was computed using EAST (EAST V6.4, Cytel) assuming a piecewise exponential survival model with two knots at 0.3 and 1.6 years, consistent with the observed 1-year EFS=74%, 2-year EFS=56%, 3-year EFS=52% and 5-year EFS=45% in high-risk patients from Sarcome-09/OS2006 trial (hazard rates: λ1=0.1065, λ2=0.3723 and λ3=0.0878).

In the previous Sarcome-09/OS2006 trial, the proportion of patients lost to follow-up was <1% at 3 years. This has a very limited impact on the sample size calculation; it only requests a slightly longer follow-up duration from 2.048 to 2.091 for the last accrued patient. Based on Sarcome-09/OS2006 data, the estimated proportion of patients fulfilling eligibility criteria for the randomised trial is 43% among all patients enrolled in the study. As we expect an attrition rate of 25% (refusal of participation in the randomised trial), we estimate that 31.5% of the patients enrolled at diagnosis will be enrolled in the randomised trial. Consequently, 390 registered patients (126/ (0.43*0.75)=390) may be required for the study to reach the 126 patients and 51 events for the randomised trial.

**Statistical analysis**

EFS curves will be estimated using the Kaplan-Meier method. Difference between EFS curves will be tested using a log-rank test at a one-sided alpha=10%. As a first step of the analysis, the relative treatment effect of mifamurtide in terms of EFS and its CI will be estimated using a Cox model with adjustment or stratification on stratification of patients lost to follow-up was <1% at 3 years. This has a very limited impact on the sample size calculation; it only requests a slightly longer follow-up duration from 2.048 to 2.091 for the last accrued patient. Based on Sarcome-09/OS2006 data, the estimated proportion of patients fulfilling eligibility criteria for the randomised trial is 43% among all patients enrolled in the study. As we expect an attrition rate of 25% (refusal of participation in the randomised trial), we estimate that 31.5% of the patients enrolled at diagnosis will be enrolled in the randomised trial. Consequently, 390 registered patients (126/ (0.43*0.75)=390) may be required for the study to reach the 126 patients and 51 events for the randomised trial.

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The proportional hazards assumption will be evaluated using graphical methods and by extension of the Cox model including treatment-time interaction. As an exploratory analysis, the interaction between mifamurtide and the stratification factors (chemotherapy strata and risk group) will be investigated by adding an interaction term between treatment and each factor separately in the multivariable Cox model, in order to evaluate the heterogeneity of treatment effect across the different subgroups. Treatment effect will be estimated by subgroup (first (M)-API vs EI chemotherapy regimens; then the five risk groups) in the multivariable analysis, and reported in a forest plot. A similar approach will be used to evaluate the predictive value of some clinical and biological factors (exploratory analyses). The main analysis will be based on all patients included in the trial, regardless of protocol compliance (intention-to-treat analysis). Sensitivity analyses of EFS will be performed (1) on the per-protocol population, after exclusion of patients who have switched from one arm to the other as well as patients who could not undergo removal of all initially identified metastatic sites; (2) on the intention-to-treat population with adjustment or stratification on stratification variables and also other possible confounders. Absolute gain in EFS time will also be estimated using the restricted mean survival time difference which remains valid if the PH assumption appears violated or questionable.

Similar statistical analyses will be used for the OS analysis. No interim analysis of efficacy is planned.

As some historical information is available, the usual hypothesis-driven approach (frequentist approach) will be completed with a Bayesian analysis incorporating the aggregate treatment effect estimated from a fixed-effect meta-analysis from the previous INT-0133 trial (in patients with localised osteosarcoma and metastatic disease) leading to an overall HR=0.786 (SE=0.110). We also plan to incorporate individual historical data, from Sarcome-09/OS2006 subgroup of patients who fulfilled the planned Sarcome-13/OS2016 eligibility criteria, on the control arm of the current trial (See figure 2).

We will use the approach proposed by Brard et al (BMC Medical Research Methodology—BMRM, in press) combining a mixture prior to incorporate the aggregate treatment effect from the INT-0133 trial, and a power prior to incorporate the individual historical data from Sarcome-09/OS2006 subgroup of patients. The weights allocated to historical data (ω and α₀ for historical aggregate treatment effect and individual control data, respectively) have been calibrated based on a simulation study (see online supplementary materials). The trial will be considered successful if the posterior probability of (HR<1) is ≥ 90% (prespecified decision rule). Considering the chosen set of parameters resulting of the simulation study (ω = 0.1 and α₀ = 0.3, the power is increased from 80% (frequentist approach) to 98% (Bayesian approach) if the true treatment effect is HR=0.55. It increases from 34% to 65% if the true treatment effect is consistent with the published INT-0133 data (HR=0.786) (table 2).

The posterior distribution of the HR will be described including the probability of HR<1 or HR below different thresholds. Given the higher probability of positive conclusion of the planned Bayesian approach compared with the frequentist approach, it is possible the Bayesian success criteria will be met while the frequentist analysis is non-significant. In this case, we will claim a positive outcome for the trial based on the results of the Bayesian analysis. The ‘power’ of the proposed Bayesian approach is the probability of a positive conclusion considering the prespecified decision rule.

At the end of the trial, we may consider incorporating data from other relevant trials evaluating mifamurtide, which are currently in progress. A non-informative prior will also be considered to describe the distribution of the treatment effect when considering current trial data only. The Bayesian analysis will allow the description of the probability distribution of the treatment effect, such as the probability that HR is lower than various thresholds, not only HR<1. The statistical analytical plan will be amended before data base lock to prespecify all sources of historical data.

For each type of AE, the worst grade observed across the safety observational period (4 weeks after last treatment administration, and up to 24 weeks after last chemotherapy for control arm), will be tabulated by treatment arm, and the percentages of severe AE (grade ≥4 haematological AE and grade ≥3 extrahaematological AE) will be provided. A butterfly plot will be used to illustrate the difference in proportion of patients experiencing AE and severe AE between treatment groups. Relative risks of severe AE associated with mifamurtide will be estimated.

The Standard Protocol Items: Recommendations for Interventional Trials reporting guidelines have been used to report this trial protocol.
Ethics and Dissemination

The study is approved by the “Comité de Protection des Personnes Ile de France I” (CPP) and authorised by the National Agency for Medicines and Health Products Safety (ANSM) that reviewed the trial protocol, patient information sheets, informed consent forms, and other trial-related documents. If changes with substantial modifications occur, they will have to be submitted to the Agency for Medicines and Health Products Safety and to the Comité de Protection des Personnes Ile de France I by the sponsor. Data recorded during this trial are subject to a computerised treatment in compliance with the French law. The collection of biological samples has been notified to the French Ministry of Research. An independent data monitoring committee (IDMC), with expertise and experience in the pathology (two clinicians and one statistician), and without direct involvement in the conduct of the trial, will be set up. The IDMC will meet every 12 months and may recommend the early termination of the trial if an unacceptable toxicity occurred, or if the available data from the trial are sufficiently convincing to influence the therapeutic practices of the majority of clinicians. Investigators will make available to the authorised persons the documents and the patients’ individual data that are essential to monitor the trial on an ongoing basis, to perform quality control and audit of this research in accordance with national regulatory requirements.

Table 2

Comparison of power and alpha error of the Sarcome-13/OS2016 trial between the frequentist approach and the Bayesian approach with $\omega = 0.1$ and $\alpha_0 = 0.3$ for different ‘true treatment effect’

| Scenario regarding the true treatment effect | Frequentist approach, % | Bayesian approach, % |
|---------------------------------------------|-------------------------|----------------------|
| HR=0.55 (anticipated effect scenario)       | 80                      | 98                   |
| HR=0.786 (historical effect scenario)       | 34                      | 65                   |
| HR=0.886 (disappointing effect scenario)    | 20                      | 42                   |
| HR=1 (null effect scenario)                 | 10                      | 21                   |

Figure 2

Individual historical data, from Sarcome-09/OS2006 subgroup of patients who fulfilled the planned Sarcome-13/OS2016 eligibility criteria, on the control arm of the current trial.
Patient and public involvement

The protocol was designed and developed by paediatricians and medical oncologists who work in the InterSARC network that includes patients and parents, such as the ‘Info Sarcomes’ organisation. Patients were not directly involved in the design of the study; however, the protocol was discussed and approved by ‘Info Sarcomes’ organisation. The protocol has also been examined by the Patient Committee of the National League against Cancer who assessed the burden of the proposed intervention. Patients will be involved in recruitment to the study as the trial will be advertised on various websites (Info Sarcomes, Unicancer, hospitals’ websites) promoting clinical studies to patients. Study results will be made available to study participants by being posted on ClinicalTrials.gov website.

Dissemination

The trial results, even if they are inconclusive, as well as biological ancillary studies, will be presented at appropriate international congresses and published in international peer review journals.

Trial financing

This study is sponsored by UNICANCER. It is mainly funded by the French Ministry of Health through the Hospital Clinical Research Program (PHRC-K-2016-130 grant). The study drug will be provided by Takeda to the sponsor.

DISCUSSION

This trial is a randomised phase II trial designed to detect an EFS benefit due to mifamurtide combined with postoperative chemotherapy compared with postoperative chemotherapy alone in patients aged ≥2 and ≤50 with high-risk osteosarcoma. Considering the rare disease setting, a large phase III trial designed with the usual level of evidence would not have been feasible in a reasonable time frame at a national level, contrasting with the proposed design with a relaxed alpha error and a smaller sample size. The participation of other European centres has been discussed to increase the sample size. However, we had to cope with some major differences regarding the backbone chemotherapy, as most European countries use M-AP without ifosfamide whereas ifosfamide-based chemotherapy is used in all French centres. The possible interaction between mifamurtide and chemotherapy regimen (mainly ifosfamide based or not) could jeopardise the findings.

At the current stage of trial design, we cannot be sure that this trial will provide a definitive answer regarding the controversial role of mifamurtide in osteosarcoma, in particular, because of the relatively limited sample size (phase II trial). However, we think that it will contribute to address this issue in an appropriate way, as mifamurtide will be allocated by randomisation. The proposed Bayesian analysis will help in combining evidence from the various trials to get the best treatment estimate and will increase the level of evidence on mifamurtide efficacy.

If the efficacy of mifamurtide is confirmed, this trial may contribute to allow the use of this promising drug, including in an extended population (extension to metastatic disease at diagnosis and older patients >30 to 50 years old).

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Contributors

All authors designed the study; CB, GLT, M-CLD and NG contributed to the drafting of the manuscript. NG, SP-N, LB and JD contributed to the trial set-up. JD is responsible for data collection and for administrative support. M-CLD, CB, GLT and LVH have proposed the statistical design and have performed the preliminary simulation study. CB, GLT and LVH will contribute to statistical analyses. M-CLD is responsible for data management and statistical analyses. All authors will contribute to data interpretation. All authors contributed to the revision of the manuscript and approved it for submission.

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Competing interests

None declared.

Patient consent for publication

Not required.

Ethics approval

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Provenance and peer review

Not commissioned; externally peer reviewed.

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