

Oxford Brookes University – Research Archive and Digital Asset Repository (RADAR)

Peinemann, F, Smith, L, Kromp, M, Bartel, C, Kröger, N and Kulig, M Autologous hematopoietic stem cell transplantation following high-dose chemotherapy for non-rhabdomyosarcoma soft tissue sarcomas (Review).

Peinemann, F, Smith, L, Kromp, M, Bartel, C, Kröger, N and Kulig, M (2011) Autologous hematopoietic stem cell transplantation following high-dose chemotherapy for non-rhabdomyosarcoma soft tissue sarcomas (Review). *Cochrane Library*, (2) [doi]

This version is available: https://radar.brookes.ac.uk/radar/items/cd790492-7e37-658f-e18f-ba82ca2d7065/1/ Available on RADAR: July 2012

Copyright © and Moral Rights are retained by the author(s) and/ or other copyright owners. A copy can be downloaded for personal non-commercial research or study, without prior permission or charge. This item cannot be reproduced or quoted extensively from without first obtaining permission in writing from the copyright holder(s). The content must not be changed in any way or sold commercially in any format or medium without the formal permission of the copyright holders.

This document is the published version of the journal article. Some differences between the published version and this version may remain and you are advised to consult the published version if you wish to cite from it.



Autologous hematopoietic stem cell transplantation following high-dose chemotherapy for non-rhabdomyosarcoma soft tissue sarcomas (Review)

Peinemann F, Smith LA, Kromp M, Bartel C, Kröger N, Kulig M



This is a reprint of a Cochrane review, prepared and maintained by The Cochrane Collaboration and published in *The Cochrane Library* 2011, Issue 2

http://www.thecochranelibrary.com



TABLE OF CONTENTS

HEADER	1
ABSTRACT	1
PLAIN LANGUAGE SUMMARY	2
BACKGROUND	2
OBJECTIVES	4
METHODS	4
RESULTS	8
Figure 1	9
Figure 2	12
DISCUSSION	13
AUTHORS' CONCLUSIONS	14
ACKNOWLEDGEMENTS	14
REFERENCES	14
CHARACTERISTICS OF STUDIES	40
DATA AND ANALYSES	76
ADDITIONAL TABLES	76
APPENDICES	84
HISTORY	93
CONTRIBUTIONS OF AUTHORS	93
DECLARATIONS OF INTEREST	93
SOURCES OF SUPPORT	94
INDEX TERMS	94

[Intervention Review]

Autologous hematopoietic stem cell transplantation following high-dose chemotherapy for non-rhabdomyosarcoma soft tissue sarcomas

Frank Peinemann¹, Lesley A Smith², Mandy Kromp³, Carmen Bartel⁴, Nicolaus Kröger⁵, Michael Kulig⁶

¹Department of Non-Drug Interventions, Institute for Quality and Efficiency in Health Care (IQWiG), Cologne, Germany. ²School of Health and Social Care, Oxford Brookes University, Oxford, UK. ³Medical Biometry, Institute for Quality and Efficiency in Health Care, Cologne, Germany. ⁴Quality of Health Care, Institute for Quality and Efficiency in Health Care (IQWiG), Cologne, Germany. ⁵Interdisciplinary Clinic for Stem Cell Transplantation, University Hospital Hamburg-Eppendorf, Hamburg, Germany. ⁶Non-drug Interventions, Institute for Quality and Efficiency in Health Care, Cologne, Germany

Contact address: Frank Peinemann, Department of Non-Drug Interventions, Institute for Quality and Efficiency in Health Care (IQWiG), Dillenburger Str. 27, Cologne, 51105, Germany. frank.peinemann@iqwig.de.

Editorial group: Cochrane Gynaecological Cancer Group. Publication status and date: New, published in Issue 2, 2011. Review content assessed as up-to-date: 3 January 2011.

Citation: Peinemann F, Smith LA, Kromp M, Bartel C, Kröger N, Kulig M. Autologous hematopoietic stem cell transplantation following high-dose chemotherapy for non-rhabdomyosarcoma soft tissue sarcomas. *Cochrane Database of Systematic Reviews* 2011, Issue 2. Art. No.: CD008216. DOI: 10.1002/14651858.CD008216.pub3.

Copyright © 2011 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

ABSTRACT

Background

Soft tissue sarcomas (STS) are a highly heterogeneous group of rare malignant solid tumors. Non-rhabdomyosarcoma soft tissue sarcomas (NRSTS) comprise all STS except rhabdomyosarcoma. In patients with advanced local or metastatic disease, autologous hematopoietic stem cell transplantation (HSCT) applied after high-dose chemotherapy (HDCT) is a planned rescue therapy for HDCT-related severe hematologic toxicity.

Objectives

To assess the effectiveness and safety of HDCT followed by autologous HSCT for all stages of soft tissue sarcomas in children and adults.

Search methods

We searched the electronic databases CENTRAL (*The Cochrane Library* 2010, Issue 2), MEDLINE and EMBASE (February 2010). Online trial registers, congress abstracts and reference lists of reviews were searched and expert panels and authors were contacted.

Selection criteria

Terms representing STS and autologous HSCT were required in the title, abstract or keywords. In studies with aggregated data, participants with NRSTS and autologous HSCT had to constitute at least 80% of the data. Comparative non-randomized studies were included because randomized controlled trials (RCTs) were not expected. Case series and case reports were considered for an additional descriptive analysis.

Autologous hematopoietic stem cell transplantation following high-dose chemotherapy for non-rhabdomyosarcoma soft tissue sarcomas (Review)

Data collection and analysis

Study data were recorded by two review authors independently. For studies with no comparator group, we synthesised results for studies reporting aggregate data and conducted a pooled analysis of individual participant data using the Kaplan-Meyer method. The primary outcomes were overall survival (OS) and treatment-related mortality (TRM).

Main results

We included 54 studies, from 467 full texts articles screened (11.5%), reporting on 177 participants that received HSCT and 69 participants that received standard care. Only one study reported comparative data. In the one comparative study, OS at two years after HSCT was estimated as statistically significantly higher (62.3%) compared with participants that received standard care (23.2%). In a single-arm study, the OS two years after HSCT was reported as 20%. In a pooled analysis of the individual data of 54 participants, OS at two years was estimated as 49% (95% CI 34% to 64%). Data on TRM, secondary neoplasia and severe toxicity grade 3 to 4 after transplantation were sparse. All 54 studies had a high risk of bias.

Authors' conclusions

Due to a lack of comparative studies, it is unclear whether participants with NRSTS have improved survival from autologous HSCT following HDCT. Owing to this current gap in knowledge, at present HDCT and autologous HSCT for NRSTS should only be used within controlled trials.

PLAIN LANGUAGE SUMMARY

Hematopoietic stem cell transplantation following chemotherapy for non-rhabdomyosarcoma soft tissue sarcomas

Non-rhabdomyosarcoma soft tissue sarcomas are a group of rare cancers. Patients with inoperable or metastatic disease have a poor prognosis. It was believed higher doses of chemotherapy might improve patients' survival. However, high doses of chemotherapy stop the production of blood cells in the bone marrow and are not compatible with life. Stem cells collected from patients before high-dose chemotherapy can be transplanted back to the patient if the blood cell count gets too low. Due to a lack of research studies, it has not been proven that patients treated with this procedure lived longer than patients treated with standard chemotherapy.

We reviewed the published research on this treatment to investigate how effective and safe it is. Unfortunately we identified only one comparative study and the results of this study were not credible. Studies with aggregated data showed that two years after treatment between 20% to 60% of patients were still alive but the treatment had a high level of toxic side effects.

While the results of this systematic review may not be conclusive, they provide a summary of the current knowledge and highlight that more research is needed. Currently the research evidence says that patients with non-rhabdomyosarcoma soft tissue sarcomas should only be treated with high-dose chemotherapy and then autologous hematopoietic stem cell transplantation except within clinical trials.

BACKGROUND

Description of the condition

Soft tissue sarcomas (STS) are a highly heterogeneous group of rare malignant solid tumors of non-epithelial extraskeletal body tissue and are classified on a histogenetic basis (Enzinger 2001). STS have a significant risk of distant metastasis in addition to the potential for locally destructive growth and recurrence. Non-rhabdomyosarcoma soft tissue sarcomas (NRSTS) comprise all STS

except rhabdomyosarcoma, which primarily affects children and young adults. In this review we investigated NRSTS which are categorized as malignant according to the World Health Organization (WHO) 2002 classification (Fletcher 2002) as adopted by the European Society for Medical Oncology (ESMO) Guidelines Working Group (Casali 2009). This classification excludes the Ewing family of tumors (EFT).

NRSTS usually originate de novo and rarely from benign tumors. In most cases the pathogenesis is unknown; however, some factors

have been found to be associated with the development of NRSTS (Enzinger 2001). These include exposure to ionizing radiation, environmental carcinogenic substances, oncogenic viruses and immunologic factors. Genetic factors can also play a role since some inherited diseases such as neurofibromatosis type 1 are associated with a higher risk of NRSTS (Tsao 2000).

In Western countries about four new cases of NRSTS are estimated per 100,000 population every year (Casali 2009), with rhabdomyosarcoma and the Ewing family of tumors excluded from this statistic. STS constitute about 1% of malignancies in adults and 7% in children (NCI 2009a). Rhabdomyosarcoma represents about 50% of STS in children (Gurney 1997; Miller 1995). NRSTS are rare in both children and adults and the distribution of NRSTS differs significantly between children and adults (Table 1) according to (Spunt 2006).

Based on the Surveillance, Epidemiology and End Results (SEER) cancer statistics review (1975 to 2005) of the National Cancer Institute (NCI), in the US 10,390 new cases and 3680 deaths from STS were estimated for the year 2008 (NCI 2008a). Separate data were not available for rhabdomyosarcoma and NRSTS. The distribution of STS increased with age from 2001 to 2005, according to SEER data. Of all STS cases, 10.3% were in children and young adults less than 20 years of age (NCI 2008b). The median age at diagnosis of STS, including tumors of the heart, was 57 years (NCI 2008c).

Staging

Disease progression may be dichotomized into the two categories of limited and extensive disease. Limited disease is typically a localized, small-sized, low-grade and operable accessible tumor that has no regional lymph node involvement and no distant metastases. Extensive disease can also be denoted as advanced disease defined as localized, large-sized and high-grade tumor that may not be completely removed by surgery, may be invasive and may have regional lymph node involvement or distant metastases. Both categories differ significantly in terms of prognosis and treatment. Where many patients with limited disease may be cured by surgery, extensive disease is associated with a poor outcome and many patients receive chemotherapy as palliative therapy.

The American Joint Committee on Cancer (AJCC) tumor-node-metastasis (TNM) staging system combines grade, depth and size of the tumor as well as regional lymph node involvement and distant metastases and describes the extent of a cancer's spread from Stage 0 to IV (AJCC 2002). A review reported the 5-year overall survival (OS) estimates for stage I (low-grade, superficial and deep), II (high-grade, superficial and deep) and IV (any metastasis to lymph nodes or distant sites) as approximately 90%, 70%, 50% and 10% to 20%, respectively; information on treatment was not given (Clark 2005). In a multicentre study a total of 2185 participants with advanced STS revealed a median survival of 12 months (Van Glabbeke 1999). In

the same study, of the 1922 (26%) eligible participants who responded to chemotherapy, the 5-year OS was 10%; in univariate analyses response to chemotherapy was not predicted by the same factors as was OS.

Symptoms

The location of the primary tumor can involve any area of the body. The distribution is 40% lower limb and girdle, 20% upper limb and girdle, 20% abdominal sites, 10% trunk and 10% head and neck (Clark 2005). NRSTS can involve any type of tissue and typically affect muscles, tendons, adipose tissue, blood vessels, joints (Sondak 2001) and commonly present as a painless mass. The symptoms depend on the anatomical site of origin, the size of the mass and other aspects. Retroperitoneal sarcomas are most often asymptomatic, until the mass grows large enough to be clinically obvious or presses on vital organs and causes pain (Dileo 2005).

Patients who relapse or suffer progressive disease after therapy, or metastasis, are commonly called high-risk patients because these signs are associated with shorter survival time. Spontaneous recovery of NRSTS is unknown.

Description of the intervention

Standard therapy

Surgery is the standard treatment for localized NRSTS (Casali 2009) and can be curative if distant dissemination is not present (Kotilingam 2006). Chemotherapy is a standard treatment for patients with distant metastasis (Casali 2009) and is regarded mainly as a palliative treatment for high-risk patients who are characterized by inoperable, locally advanced and metastatic disease. Doxorubicin, ifosfamide, gemcitabine, dacarbacine, docetaxel and trabectedin are used in monotherapy or in combinations (Casali 2009).

High-dose chemotherapy (HDCT) followed by autologous hematopoietic stem cell transplantation (HSCT)

Autologous hematopoietic stem cell transplantation (HSCT) is defined as the transplantation of stem cells that have been collected previously from bone marrow or peripheral blood of the same person. High-dose chemotherapy (HDCT) uses higher doses of chemotherapeutic agents than is usually applied in standard-dose chemotherapy. HDCT may be tolerated by the patient or it may ablate the patient's bone marrow reserves and create an absolute requirement for stem cell rescue. Instead of HDCT, high-dose radiation therapy may be used to treat NRSTS patients. Autologous HSCT applied after HDCT or high-dose radiation is a

planned rescue therapy for HDCT-related severe hematologic toxicity (Banna 2007). Ideally, a mega-therapy regimen should be used consisting of several non-crossresistant agents that have a steep dose-response curve and little extramedullary toxicity (Ladenstein 1997).

HDCT and autologous HSCT are not a standard treatment option; they are an experimental approach. HDCT and autologous HSCT may be used in special cases after careful consideration, usually for patients who respond well to standard chemotherapy according to RECIST (Therasse 2000) criteria (Kasper 2005; Kasper 2007). Carboplatin, cisplatin, cyclophosphamide, etoposide, ifosfamide, melphalan, mitoxantrone and thiotepa, for example, have been used in HDCT regimens. HDCT and autologous HSCT are an experimental approach mainly used to treat high-risk patients with an unfavourable prognosis (stage IV with distant metastases). Independent of the disease status, HDCT and autologous HSCT are hazardous interventions that carry the risk of life-threatening organ failure.

Autologous HSCT and preceding HDCT were adopted to treat high-risk patients because it was believed that escalating doses in chemotherapy might increase survival by capturing putatively remnant malignant cells and might overcome resistance to standard-dose chemotherapy (Banna 2007).

Adverse events

Non-hematological adverse events, such as short-term and long-term organ toxicities, must be considered when using HDCT (Ladenstein 1997). Hematological adverse events as a result of autologous HSCT are usually manageable but life-threatening consequences of pancytopenia. They generally affect all patients and include, for example, graft failure, severe infections and bleeding.

Frequency

Of a total of 15,278 autologous HSCTs that were registered in 2005 by the European Group for Blood and Marrow Transplantation (EBMT), 69 were indicated for STS (Gratwohl 2007).

How the intervention might work

Escalating doses of chemotherapy may increase survival by capturing putatively remnant malignant cells and thus overcome cell resistance to standard chemotherapy (Banna 2007). High-dose chemotherapy may also cause severe hematologic and non-hematologic toxicity. Autologous HSCT is a planned rescue therapy for the HDCT-related demise of hematopoietic stem cells.

Why it is important to do this review

The potential benefit of this treatment option has not been investigated sufficiently in comparative studies (Pedrazzoli 2006). Some authors have warned against the use of HDCT with autologous HSCT, indicating the possibility of repositioning of malignant cells (Woods 1999). Others have questioned the rationale of HDCT with reference to the potential existence of refractory cancer stem cells (Banna 2007; Bonnet 1997; Sanchez-Garcia 2007). The question has not been answered whether autologous HSCT preceded by HDCT is able to increase OS in patients with NRSTS when compared to standard-dose chemotherapy. Randomized controlled trials (RCTs) have not been published. The rationale for this intervention, as described above, was based on non-comparative studies. We summarized and described the present available evidence to provide an evidence base to inform the design of future comparative studies.

OBJECTIVES

To assess the effectiveness and safety of autologous high-dose chemotherapy (HDCT) followed by autologous hematopoietic stem cell transplantation (HDCT) for all stages of non-rhabdomyosarcoma soft tissue sarcomas (NRSTS) in children and adults.

METHODS

Criteria for considering studies for this review

Types of studies

Inclusion criteria

• Randomized controlled trials (RCTs).

Since we expected to find few, if any, RCTs non-RCTs were also included as follows.

• Quasi-RCTs, non-RCTs, phase I and II prospective studies, prospective and retrospective cohort studies, case-control studies, case series and case reports.

Results from RCTs and controlled clinical trials may provide data for estimation of effects on overall survival (OS) and answer the question: "Has the intervention a significantly better survival than the control and does the quality of the studies fit with the assumption that the intervention is better than the control?"

Data from non-comparative studies (phase I and II prospective studies, case series and case reports) were collected to estimate treatment-related mortality (TRM) within a cohort of participants, as

a descriptive analysis. Due to the lack of a control group the studies do not provide data for estimation of treatment effect. review (Table 2) and we present the terms for tumors that were not considered (Table 3).

Exclusion criteria

None

Rationale for including non-RCTs

Authors of studies on HDCT with autologous HSCT have stated that RCTs are both necessary and feasible. However, NRSTS is a rare disease and, according to the results of a preview literature search, currently there are no published RCTs available. In addition, controlled clinical trials or studies with any comparative data may be unlikely or rare. If they do exist they may be of low methodological quality. Based on the assumption that it is unlikely that the intervention has been or will be studied in RCTs in the near future, this systematic report of the findings and limitations of all available published studies will be useful, for example, for informing the design of appropriate RCTs and providing a summary of all of the evidence on the topic to date.

Types of participants

Inclusion criteria

We have adopted the WHO classification of soft tissue tumors to define the population of patients with NRSTS (Fletcher 2002) with the exception of the Ewing family of tumors (see 'Exclusion criteria'). Studies were included as long as at least 80% of patients had NRSTS. Children as well as adults were investigated and age limits did not apply. Participants were included regardless of the severity of the disease and of clinical staging information, as long as they received autologous (from either a peripheral or bone marrow source, or both) HSCT.

Exclusion criteria

Whilst the WHO classification of NRSTS includes the Ewing family of tumors, that is extraosseous tumor types, we excluded these because they are primarily bone sarcomas. Because extraosseous types are rarely diagnosed and share common features, they were regarded with osseous types as one entity and were excluded.

The clear delineation of soft tissue sarcomas to be included in the present report and the grounds for exclusion of some tumor types was hindered by the presence of more than 30 heterogenous tumor entities, the distinction between malignant tumors and two categories of intermediate malignancies as described in the WHO classification (Fletcher 2002), and a complicated histology and terminology. Therefore, we present the designation of tumors that were regarded as (malignant) soft tissue sarcomas in the present

Types of interventions

Intervention: autologous hematopoietic stem cell transplantation (HSCT), stem cells from peripheral source or the bone marrow, serving as a rescue therapy usually applied after high-dose chemotherapy (HDCT).

Comparison: standard-dose chemotherapy, which is defined as chemotherapy at a lower dose than HDCT without the need for stem cell rescue.

Allogenic HSCT was excluded.

Types of outcome measures

Primary outcomes

- Overall survival (OS): survival until death, from all causes. Survival was assessed starting from the time when participants received autologous HSCT.
- Treatment-related mortality (TRM): deaths that were classified as treatment related or the participants died of complications after autologous HSCT.

Secondary outcomes

- Disease-free survival (DFS): time free of disease after receiving autologous HSCT; the events were death due to all causes or any sign of the disease. The extent of disease was evaluated by clinical, histologic and imaging studies.
- Progression-free survival (PFS): time staying free of disease progression after receiving autologous HSCT. Participants may still have the disease but their disease is stable or showed a partial response to treatment; the events are death from all causes or any progression of the disease.
- Event-free survival (EFS): time staying free of any of a particular group of defined events after receiving autologous HSCT. Participants may still have the disease; the events are death from all causes, any sign of the disease in participants who had a complete response to treatment, any relapse or progression of the disease, or events that were defined by the individual study protocol.
- Failure-free survival (FFS): time staying free of treatment failure after receiving autologous HSCT; the events are disease-or treatment-related death, any sign of the disease in participants who had a complete response to treatment, refractory disease with no response to treatment, stable disease, or progression of the disease after treatment.
- Toxicity: adverse events classified according to the common toxicity criteria (NCI 2009b) within 90 days of autologous HSCT; grades 3 and 4 of toxicity were extracted and grouped as

hematological (leukopenia, neutropenia, thrombocytopenia) and non-hematological (nausea, kidney, liver, nervous system, heart) toxicities.

- Secondary neoplasia: as classified by the study authors.
- Health-related quality of life (HRQoL): measured using a questionnaire that has been validated through reporting of norms in a peer-reviewed publication.

Search methods for identification of studies

The search methods suggested in the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2009) and by the Cochrane Gynaecological Cancer Review Group were used. Articles in any language were included. Translations were carried out as necessary.

The literature sources and search steps are shown in Table 4. In the first step, three different bibliographic databases were searched electronically to find topic-related articles. In the second step, online registers were searched to find additional information on completed or ongoing comparative studies that have not been published. References cited in 98 identified reviews (Appendix 4), including three systematic reviews, were evaluated.

Electronic searches

The following electronic databases were searched: the Cochrane Central Register of Controlled Trials (CENTRAL) (*The Cochrane Library* 2010, Issue 2), Ovid MEDLINE (from 1950 to February 2010), Ovid EMBASE (from 1980 to February 2010). See Appendix 1, Appendix 2, and Appendix 3 for the appropriate medical subject headings (MeSH) and text words for the search strategies. An updated search was run in PubMed (6 June 2010) using the following terms: ("Transplantation, Autologous" [Mesh] OR "Peripheral Blood Stem Cell Transplantation" [Mesh]) AND "Sarcoma" [Mesh].

Searching other resources

Information about trials not listed in CENTRAL, MEDLINE or EMBASE, either published or unpublished, were located by searching the reference lists of relevant articles and review articles. We also electronically searched the abstracts of the conference proceedings of the American Society of Clinical Oncology (ASCO) annual meetings (from 2004 to 2009). We searched for ongoing trials by scanning online registers listed in Table 4. We also searched for ongoing trials by contacting researchers involved in the area.

Data collection and analysis

Selection of studies

All titles and abstracts retrieved by electronic searching were down-loaded to the reference management database (Reference Manager Version 11) (Thomson Reuters Corp 2009); duplicates were removed and the remaining references were examined by two review authors (FP and TBH) independently. Those studies which clearly did not meet the inclusion criteria were excluded and copies of the full text of potentially relevant references were obtained. The eligibility of retrieved papers was assessed independently by two review authors (FP and TBH). Disagreements were resolved by discussion between the two review authors and consultation with a third review author (MK), if necessary. Reasons for exclusion were documented.

Data extraction and management

For included studies, data on characteristics of studies, participants and interventions; risk of bias; duration of follow up; outcomes and deviations from protocol were abstracted independently by two review authors (FP and MaKr). Differences between review authors were resolved by discussion or by appeal to a third review author (CB).

Characteristics of studies

- Study type (RCTs, non-RCTs, non-randomised trials with no control group (phase I or II study), cohort studies, case-control studies, case series, case reports)
- Design (randomization, sequence generation and concealment of allocation, blinding, prospective, retrospective, consecutive enrolment sample selection)
 - Observation period (calendar years)
- Inclusion and exclusion criteria (Ewing family of tumors, other excluded sarcomas, other solid tumors)
 - Number and location of participating centres

Characteristics of participants

- Age
- Gender
- Type of histological category
- Status of metastasis
- Number of recruited and analyzed participants

Characteristics of interventions

- High-dose chemotherapy
- Autologous peripheral blood stem cells
- Autologous bone marrow stem cells

Survival measures

- Time to event from treatment with HSCT
- Number of events and participants at risk
- Kaplan-Meier survival estimate
- Hazard ratio (HR)
- 95% confidence interval (CI)
- Log rank P value
- Duration of follow up (median; range)

Treatment-related mortality

- Number of events and participants at risk
- Number of recruited and analyzed participants
- Cause of death

Secondary neoplasia

- Number of events and participants at risk
- Number of recruited and analyzed participants
- Type of secondary neoplasia

Toxicity

- Number of WHO grade 3 or 4 adverse events and participants at risk
 - Number of recruited and analyzed participants
 - Organ system affected

Quality of life

- Scale
- Number analysed
- Mean or median
- Standard deviation or range

Where relevant and if reported, both unadjusted and adjusted summary statistics were extracted. Where possible, all data extracted were those relevant to an intention-to-treat (ITT) analysis in which participants were analyzed in the groups to which they were assigned. The time points at which outcomes were collected and reported were noted.

Assessment of risk of bias in included studies

The assessment of risks of bias in included controlled studies was independently applied at the study level by two review authors (FP and MaKr) according to Table 5 and to the guidelines of the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2009). Differences were resolved by discussion or by appeal to a third review author (CB). Results of data syntheses were interpreted in light of the findings with respect to risk of bias. Assessment of blinding of the care provider and blinding of the

participants was not applicable as blinding is not ethically accepted for studies on stem cell transplantation.

Measures of treatment effect

We used the following measures of the effect of treatment.

- For time to event data we used the HR, if possible.
- For dichotomous outcomes we used the risk ratio (RR) or odds ratio (OR).
- For continuous outcomes we used the mean difference between treatment arms on the condition that the distribution characteristics had been evaluated.

Studies reporting aggregate data that combined the results of several participants (including results from separately reported subpopulations that fulfilled the inclusion criteria) were distinguished from studies with individual data of single participants. Data from these studies were described as narrative summaries.

In some studies diagnoses of NRSTS were mixed with non-NRSTS solid tumors and rhabdomyosarcomas to such an extent that the proportion of NRSTS participants was less than 80% of the study population. In this case, if data on single participants were identified that fulfilled the inclusion criteria of the present review we included the study and data for the individual participant in data analysis.

Estimates of OS were considered for the evaluation if the use of the Kaplan-Meier method was reported in the study. A survival analysis was conducted of individual participant level data based on the Kaplan-Meier method. Data were not used for survival analysis if the follow-up data were only available for selected participants and if the beginning of the follow-up period was not reported clearly, or reported as starting from the time of diagnosis. Statistical analyses of time to event data were performed using SAS Version 9.2 (SAS Institute Corp 2009).

Unit of analysis issues

None

Dealing with missing data

Information on the outcome status and on the follow-up period had to be complete for all participants in each study. We did not impute missing outcome data for the primary outcome. If data were missing or only imputed data were reported we contacted trial authors to request data on the outcomes among participants who were assessed.

Assessment of heterogeneity

The data were entered in Review Manager Version 5 (Review Manager 2008) and analyzed according to the guidelines of the Cochrane Handbook for Systematic Reviews of Interventions (

Higgins 2009). We looked for sources of clinical heterogeneity due to differences in:

- risk factors of participants studied, i.e. tumor subdiagnosis or histology, presence or absence of metastasis;
 - study design; and
 - likelihood of bias.

Heterogeneity between studies was assessed by inspection of the study methods and participants' characteristics. Forest plots and formal statistical tests could not be conducted because the body of studies did not contain sufficient numbers of comparative studies.

Data synthesis

Aggregate data reported in controlled studies or case series were synthesized narratively. In contrast, individual data were pooled and available time-to-event data were analyzed in a Kaplan Meier survival analysis.

Subgroup analysis and investigation of heterogeneity

No subgroup analyses were carried out.

Sensitivity analysis

No sensitivity analyses were carried out.

RESULTS

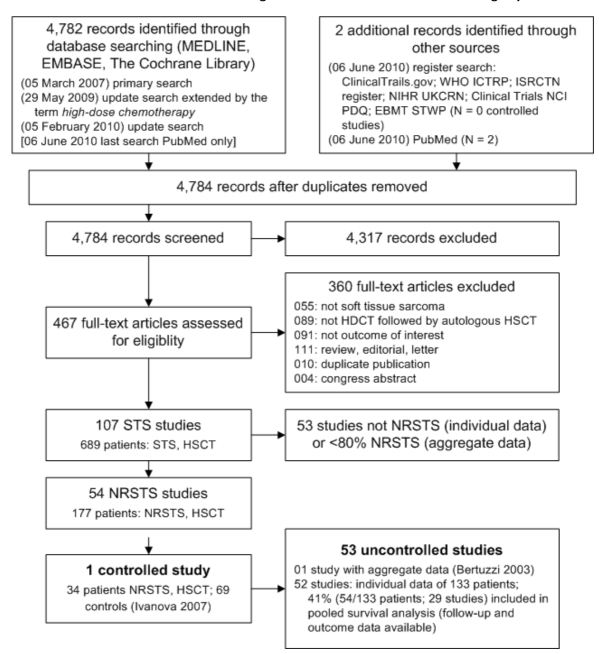
Description of studies

See: Characteristics of included studies; Characteristics of excluded studies.

Results of the search

Considering all sources as shown in Table 4, 4784 different articles (duplicates removed) were identified (Figure 1). The titles and abstracts of 4317 articles did not fulfil the inclusion criteria and 10% (467 of 4784) of the retrieved articles were evaluated in detail using the full text. Of these a total of 11% (54 of 467 references) of the full text articles were included in the present review and the other 413 studies were excluded.

Figure I. Literature search and study flow. Abbreviations: EBMT STWP: European Group for Blood and Marrow Transplantation Soft Tissue Working Party; HDCT: high-dose chemotherapy; HSCT: hematopoietic stem cell transplantation; N: number; NCI PDQ: National Cancer Institute Physician Data Query Clinical Trials; NIHR UKCRN: National Institute for Health Research (NIHR) UK Clinical Research Network's Portfolio Database; NRSTS: non-rhabdomyosarcoma soft tissue sarcoma; RMS: rhabdomyosarcoma; STS: soft tissue sarcoma. WHO ICTRP: World Health Organization International Clinical Trials Registry Platform.



We were unable to identify any additional studies from screening the reference lists of included studies and reviews or from institutions and authors. Relevant studies were not identified from online trial registries or congress proceedings, described in Table 4, either.

Included studies

We identified 54 studies including a total of 246 participants: 177 that received HDCT followed by autologous HSCT transplant and 69 that received standard therapy. The characteristics of all 54 included studies are described in the section Characteristics of included studies.

We included one retrospective study in which 34 participants who received HDCT followed by autologous HSCT were compared with a historical control group consisting of 69 participants with STS matched for age, gender, histological diagnosis and stage of disease and who received standard care (Ivanova 2007). This study did not provide sufficient detailed information about the control group to allow a meaningful evaluation. One study had a prospective design and included 10 participants with NRSTS who received HDCT and peripheral HSCT (Bertuzzi 2003). The remaining 52 studies comprised 16 phase I or II prospective trials with no control groups and 36 retrospective case series or individual case reports. Whilst not all participants had NRSTS in the studies that were phase I or II prospective trials, data were reported for individual participants so that we were able to include these studies. RCTs and clinical controlled studies with a concurrent control group were not identified.

Excluded studies

A total of 88% (413 of 467 references) of the potentially relevant articles were excluded (Figure 1) based on:

- diagnosis not STS according to Table 2 (n = 55);
- not NRSTS (individual data) or less than 80% NRSTS (aggregate data) (n = 53);
 - intervention not autologous HSCT (n = 89);
 - primary outcome survival not reported (n = 91);
 - study design was a review, editorial, abstract (n = 111);
 - duplicate publication (n = 10);
 - congress abstract (n = 4).

Excluded studies are described in the Characteristics of excluded studies table.

Risk of bias in included studies

Comparative studies

The design of the study by Ivanova 2007 was not clearly reported; we categorised it as a retrospective comparative study that used historical data to create a control group. Results of 34 participants that received HSCT were compared to 69 participants that did not receive HSCT. The author reported that the groups were matched for age, gender, histology and stage, though we were unable to verify this as insufficient data were reported in the article. Assignment to treatment groups was not described, the participant flow was unclear and loss to follow up was not addressed. Selective outcome reporting was unclear in this retrospective study, study results were reported amidst reports of results from other studies and information about statistical methods was sparse. Therefore, the risk of bias was very large. Data for individual participants were not reported.

Non-comparative studies

The risk of bias was high in all 53 studies due to the study design being either phase I and II non-comparative trials, case reports or case series.

Aggregate data

In one study (Bertuzzi 2003) the characteristics of the participants and the intervention were described in detail and the participants were described as a consecutive sample. The primary outcome, OS, was clearly reported in the text and shown in an appropriate graph; the Kaplan-Meier method was used to conduct a survival analysis. All 10 included participants had the histologic subtype of desmoplastic small round-cell tumor. The prognosis of this subtype may be regarded as specific and not comparable to other subtypes. Data for individual participants were not reported.

Individual participant data

In 52 studies, the characteristics of the participants and the intervention were described in detail. The primary outcome of OS was clearly reported in the text or tables, however the start of the follow-up period varied between the studies and was not reported in every study. We defined the start of the follow up as the time of treatment, either transplantation or high-dose chemotherapy. The start of follow up as the time of diagnosis was not accepted because the time lag between diagnosis and intervention can be considerable, as demonstrated in a study on 22 participants with STS including 11 NRSTS and 11 RMS: "Median delay between diagnosis and intensification was between 4 to 39 months" (Dumontet 1992). Therefore, OS with treatment was estimated using only data with complete outcome and follow-up information for the pooled analysis. Considerable data were incomplete or inappropriate, resulting in 59% (79 of 133 participants) of the pooled

individual data being excluded from the analysis, therefore only 41% (54 of 133 participants in 29 studies) of the data were used in the pooled survival analysis.

Allocation

The allocation of participants to the two alternative treatment groups was not described in the one comparative study.

Blinding

Blinding was not assessed as it is unlikely that blinding of participants or investigators would be adopted in studies evaluating HSCT.

Incomplete outcome data

There was insufficient reporting of attrition and exclusions to permit judgement whether incomplete outcome data were adequately addressed. Participant flow, prospective comparative study design, characteristics of both treatment groups and outcomes were not described in detail. Consecutive recruitment of participants was described in the one case series with aggregate data (Bertuzzi 2003).

Selective reporting

There was insufficient information to permit judgement of whether the reports were free of selective outcome reporting. It is likely that all studies fall into this category. The start of follow up was not stated in one (Ivanova 2007) of two studies that presented aggregate outcome data. Follow-up data were not reported

adequately for 59% (79 of 133) of participants of studies that presented individual data, therefore we could not use the data. TRM was not addressed in every study. Secondary neoplasia is a long-term outcome that was not addressed in every study. Data on toxicity outcomes could not be extracted from most studies because they were not reported individually for the population of interest.

Other potential sources of bias

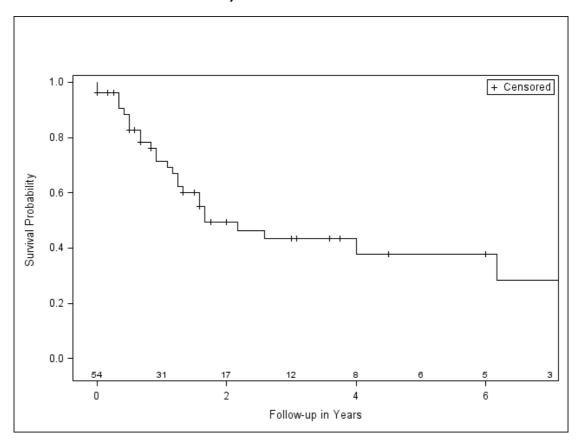
All 54 studies had a potential source of bias related to the specific study design used, such as lack of a control group or no description of the characteristics of a control group, no reporting of how the participants were selected and no reporting of reasons for loss to follow up.

Effects of interventions

Overall survival (OS)

In one comparative study (Ivanova 2007), OS at two years after HSCT was estimated as 62.3% when compared with 23.2% for participants that received standard care. The difference was reported to be statistically significant (Table 6) although the statistical test was not described and a P value was not reported. The Kaplan-Meier estimator for OS at two years after transplantation was reported as 20% in another study (Bertuzzi 2003) with aggregate data of 10 participants with desmoplastic small round-cell tumors (Table 6). In the pooled survival analysis of 54 individuals who received HSCT, OS at two years was estimated as 49% (95% CI 34% to 64%) (Table 6; Figure 2).

Figure 2. Meta-analysis of overall survival of 54 individual patients with complete follow-up information pooled from 29 case series and case reports. Number of patients at risk at 1 year intervals. Abbreviations: HDCT: high-dose chemotherapy; HSCT: hematopoietic stem cell transplantation; NRSTS: non-rhabdomyosarcoma soft tissue sarcoma.



Treatment-related mortality (TRM)

TRM was addressed in eight non-comparative studies (67 transplanted participants) and a procedure-related death was described for 11 participants (Table 7). Severe infection was the main cause of death (4 cases). TRM was not addressed in the study by Ivanova 2007.

Disease-free survival

Not reported

Progression-free survival (PFS)

PFS at two years after transplantation was reported as 0% in one study (Bertuzzi 2003) with aggregate data of participants with desmoplastic small round-cell tumors (Table 6).

Event-free survival

Not reported

Failure-free survival

Not reported

Secondary neoplasia

Secondary neoplasia was addressed in one case report (Table 8) and was not addressed in the comparative study (Ivanova 2007).

Toxicity

Data on hematological and non-hematological severe toxicity grade 3 to 4 (NCI CTEP 2006) after transplantation was sparse and extracted from five non-comparative studies only (Table 9).

Subgroup analysis

Subgroup analyses were not conducted.

Health-related quality of life

Studies on health-related quality of life were not identified.

DISCUSSION

Summary of main results

We identified only one comparative study, with a high likelihood of bias. OS at two years for participants in the HSCT group was significantly higher than for participants in the control group. In one case series of 10 participants with desmoplastic small roundcell tumors, OS at two years was low at 20%, which may reflect that participants with this special tumor type have a lower risk of survival than participants with other NRSTS. Our estimated OS of 49% at two years for the remaining studies with participant level data falls within the range reported above for the two studies with aggregate data. For TRM even the more conservative estimate, 6.2%, is considerably higher than the 2.0% TRM within the first days following HSCT reported by the EBMT registry (EBMT 2009) for the year 1998 (Rosti 2002). Secondary neoplasia was reported for 0.6% of participants and was probably an extreme underestimation of the true frequency because of relatively short follow up in the included studies and the fact that the included studies were not designed to specifically detect secondary neoplasia. The detection of secondary neoplasia depends on a long follow up, which might be provided by cancer registers. This reported incidence compares with a frequency of 4.0% for secondary neoplasia based on register data (Neglia 2001) and 6.9% (Baker 2003) after a long observation period of 20 years. Severe toxicity grade 3 to 4 was sparsely reported.

Overall completeness and applicability of evidence

Many of the studies we identified had to be excluded because they included participants with a mixture of heterogenous tumors and the proportion of participants with NRSTS was fewer than 80%. Furthermore, most treatments were performed 10 to 20 years ago. Thus, the results may not be applicable to patients who are treated today. It is also a possibility that the results reflect the course of the disease and an effect of the prior therapy rather than the effect of the test intervention.

Quality of the evidence

All 54 studies have a high risk of bias. The one comparative study lacks information on the characteristics of participants, the intervention and applied statistical methods. Thus, these data do not provide any evidence that HDCT followed by autologous HSCT has additional benefit over standard therapy. An additional descriptive analysis of 52 phase I and II studies, case series or case reports showed that there are many different tumor types of NRSTS treated by transplantation, though each individual NRSTS entity was scarce. We found that a pooled survival analysis of individual participant data from phase I and II studies and case reports was considerably hampered. Specifically, the majority of required follow-up information was incomplete or missing and, therefore, could not be included in the survival analysis. One requirement for survival analysis is a unique definition of the start of the follow-up period for all included participants. This was missing from many studies. As suggested by the low OS for desmoplastic small round-cell tumor, each entity may carry an individual risk profile and, therefore, ideally should be evaluated separately. The body of evidence does not allow robust conclusions in relation to the objectives of the review.

Potential biases in the review process

Strengths

The search strategy was broad and it is likely that all relevant studies were identified. The WHO classification of NRSTS was adopted and modified to define a clear terminology for the study selection process. Studies were excluded if the proportion of non-eligible participants were greater or equal to 20% of the total population. The follow up of participants with individual data had to begin at the same time point to be considered in the pooled analysis of survival. Authors were contacted to ask for additional data.

Limitations

RCTs and non-controlled trials with low risk of bias were not identified and only one comparative study with high risk of bias was identified. No database is available for a conclusive comparison. Many studies were excluded because participants with NRSTS were included with participants with other malignant diseases. The heterogeneity of NRSTS and the possible different terminology used in publications may have led us to overlook studies with eligible participants. This may be more an issue for case series but it is highly unlikely for controlled trials. The pooled survival analysis of individual data was based on less than half of all the individual data available and the exclusion of these data whilst intending to reduce bias may also have introduced bias.

Agreements and disagreements with other studies or reviews

In the last two decades, from 1986 to 2007, the lack of evidence and need to conduct randomized controlled trials was stated in at least 20 publications (Blay 2000; Carvajal 2005; Dumontet 1992; Ek 2006; Elias 1998; Hale 2005; Kasper 2004; Kasper 2007; Kavan 1997; Ladenstein 1997; Mackall 2001; Meyers 2004; Michon 1999; Pinkerton 1986; Reichardt 2002; Rosti 2002; Schlemmer 2006; Seeger 1991; Weigel 2001; Woods 1999) seeking to clarify the relevance of HDCT followed by autologous HSCT in high-risk patients with STS. We identified reviews on OS at five years after chemotherapy without transplantation. The estimates were 6% (Ramanathan 1999), less than 10% (Banna 2007; Van Glabbeke 1999), 14% (Tumorregister München 2007) and 10% to 20% (Clark 2005). In the control group of Ivanova 2007, OS at two years was 23%. A systematic review (Verma 2008) was performed to determine whether first-line dose-intensive chemotherapy supported by growth factor or autologous bone marrow or stem cell transplantation improves outcomes compared with standard-dose chemotherapy in patients with inoperable, locally advanced or metastatic soft tissue sarcoma. In this review, only one case series (Schlemmer 2006) with HDCT followed by autologous HSCT was reported, which was excluded from the present review. In a narrative review (Banna 2007) of HDCT followed by autologous HSCT in patients with solid tumors, outcomes were reported for three sarcomas (rhabdomyosarcoma, Ewing sarcoma and osteosarcoma) not included in the present review. Predictive values were not reported in a review (Clark 2005) on soft tissue sarcomas in adults. Kasper 2005 concluded that the use of HDCT for locally advanced or metastatic adult (soft tissue and bone) sarcomas still remains highly investigational and should not be performed outside clinical trials.

AUTHORS' CONCLUSIONS

Implications for practice

The evidence base does not support the use of HDCT followed by autologous HSCT in high-risk patients with NRSTS except in prospective concurrent, preferably randomized, controlled trials.

Implications for research

Randomized controlled trials are needed to clarify the relevance of HDCT followed by autologous HSCT in patients with NRSTS. If non-randomized controlled studies are conducted, a low risk of bias should be achieved. Case series and case reports are not helpful. Criteria of included tumor types should adhere to the WHO classification.

ACKNOWLEDGEMENTS

We thank the Cochrane Gynaecological Cancer Review Group for their assistance during the preparation of the protocol and we express our gratitude especially to Clare Jess for coordinating the protocol and review and Jane Hayes for comments on the search strategy. We thank Irit Ben-Aharon, Liat Vidal and Kathie Godfrey for their comments on the protocol and the review. We thank Tatjana Burkhardt-Hammer for screening retrieved papers against eligibility criteria.

All the work on this analysis was funded by the Institute for Quality and Efficiency in Health Care (IQWiG), an independent non-profit and non-government organization that evaluates the quality and efficiency of healthcare services in Germany. All work on this study was supported by IQWiG within the framework of a systematic review on autologous stem cell transplantation for STS.

REFERENCES

References to studies included in this review

Al Balushi 2009 {published data only}

Al Balushi Z, Bulduc S, Mulleur C, Lallier M. Desmoplastic small round cell tumor in children: a new therapeutic approach. *Journal of pediatric surgery* 2009;44(5):949–52.

Andres 2006 {published data only}

Andres R, Mayordomo JI, Isla D, Ramon y cajal S, Tejero E, Navarro A, et al.Desmoplastic small round cell tumor of the abdomen. *Oncologia* 2006;**29**(2):73–5.

Bernbeck 2007 {published data only}

Bernbeck B, Bahci S, Meisel R, Troeger A, Schönberger S, Laws H-J, et al. Serial intense chemotherapy combining topotecan, etoposide, carboplatin and cyclophosphamide (TECC) followed by autologous hematopoietic stem cell

support in patients with high risk soft tissue sarcoma (STS). Klinische Pädiatrie 2007;**219**(6):318–22.

Bertuzzi 2003 {published data only}

Bertuzzi A, Castagna L, Quagliuolo V, Ginanni V, Compasso S, Magagnoli M, et al.Prospective study of high-dose chemotherapy and autologous peripheral stem cell transplantation in adult patients with advanced desmoplastic small round-cell tumour. *British Journal of Cancer* 2003;**89**(7):1159–61.

Bley 2004 {published data only}

Bley TA, Zeiser R, Juttner E, Windfuhr-Blum M, Ghanem N, Kotter E, et al. Thrombectomy discloses intravascular growth of chondroid liposarcoma mimicking a long distance vena cava thrombosis. *In Vivo* 2004;**18**(4):463–4.

Bölke 2005 {published data only}

Bölke E, Ruf L, Budach W, Reinecke P, Röhrborn A, Pape H, et al. Tandem high-dose chemotherapy supported by autologous peripheral blood stem-cell transplantation and radiotherapy for recurrent malignant fibrous histiocytoma. Wiener Klinische Wochenschrift 2005;117(23-24):833–6.

Cole 1999 {published data only}

Cole P, Ladanyi M, Gerald WL, Cheung NK, Kramer K, LaQuaglia MP, et al. Synovial sarcoma mimicking desmoplastic small round-cell tumor: critical role for molecular diagnosis. *Medical and Pediatric Oncology* 1999; **32**(2):97–101.

Doros 2008 {published data only}

Doros L, Kaste SC, Rodriguez-Galindo C. Sister Mary Joseph's nodule as presenting sign of a desmoplastic small round cell tumor. *Pediatric Blood and Cancer* 2008;**50**(2): 388–90.

Endo 1996 {published data only}

Endo M, Yokoyama J, Ikawa H, Watanabe K, Ueda M, Kitajima M, et al.Treatment of high-risk solid tumors of childhood with myeloablative chemotherapy and autologous stem cell transplantation: A single institution experience. *Oncology Reports* 1996;**3**:519–25.

Engelhardt 2007 {published data only}

Engelhardt M, Zeiser R, Ihorst G, Finke J, Müller CI. High-dose chemotherapy and autologous peripheral blood stem cell transplantation in adult patients with high-risk or advanced Ewing and soft tissue sarcoma. *Journal of Cancer Research and Clinical Oncology* 2007;**133**(1):1–11.

Fang 2008 {published data only}

Fang X, Rodabaugh K, Penetrante R, Wong M, Wagner T, Sait S, et al.Desmoplastic small round cell tumor (DSRCT) with ovarian involvement in 2 young women. *Applied Immunohistochemistry and Molecular Morphology* 2008;**16** (1):94–9.

Farruggia 2008 {published data only}

Farruggia P, D'Angelo P, Lo Cascio M, Solazzo L, Montemaggi P, Novara V, et al. Synovial sarcoma of the neck in a child: a multidisciplinary approach. *Pediatric Hematology and Oncology* 2008;**25**(5):431–7.

Fetscher 1997 {published data only}

Fetscher S, Brugger W, Bertz H, Krieger G, Kanz L, Mertelsmann R, et al. High-dose chemotherapy with autologous peripheral blood stem cell transplantation for metastatic gastric leiomyosarcoma. *Bone Marrow Transplantation* 1997;**20**(9):787–8.

Frapier 1998 {published data only}

Frapier J-M, Marty-Ane CH, Marguerite G, Serre I, Taylor N. Primary sarcoma of the lung with left atrial involvement. A case of combined resection in a pediatric patient. *International Journal of Pediatric Hematology and Oncology* 1998;**5**(5):367–72.

Fraser 2006 {published data only}

Fraser CJ, Weigel BJ, Perentesis JP, Dusenbery KE, DeFor TE, Baker KS, et al. Autologous stem cell transplantation for

high-risk Ewing's sarcoma and other pediatric solid tumors. Bone Marrow Transplantation 2006;**37**(2):175–81.

Garrido 1998 {published data only}

Garrido SM, Chauncey TR. Neuroleptic malignant syndrome following autologous peripheral blood stem cell transplantation. *Bone Marrow Transplantation* 1998;**21**(4): 427–8.

Graham 1997 {published data only}

Graham ML, Herndon JE 2nd, Casey JR, Chaffee S, Ciocci GH, Krischer JP, et al. High-dose chemotherapy with autologous stem-cell rescue in patients with recurrent and high-risk pediatric brain tumors. *Journal of Clinical Oncology* 1997;**15**(5):1814–23.

Hawkins 2002 {published data only}

Hawkins DS, Felgenhauer J, Park J, Kreissman S, Thomson B, Douglas J, et al.Peripheral blood stem cell support reduces the toxicity of intensive chemotherapy for children and adolescents with metastatic sarcomas. *Cancer* 2002;**95** (6):1354–65.

Hoogerbrugge 1997 {published data only}

Hoogerbrugge PM, Egeler RM. Autologous transplantation of G-CSF mobilized bone marrow cells in a child with disseminated fibrosarcoma. *Bone Marrow Transplantation* 1997;**20**(7):613–4.

Ivanova 2007 {published data only}

Ivanova NM, Aliev MD, Shvarova AV, Dzampaev AZ, Mentkevich GL. Innovation approaches to treatment of children with malignant tumors of the musculoskeletal system [Article in Russian]. *Vestnik Rossiiskoi Akademii Meditsinskikh Nauk* 2007;**10**:26–32.

Kaminski 2000 {published data only}

Kaminski JM, Yang CC, Yagmai F, Movsas B, Lee M, Barrett JT. Intracranial fibrosarcoma arising 5 years after chemotherapy alone for glioblastoma multiforme in a child. *Pediatric Neurosurgery* 2000;**33**(5):257–60.

Kasper 2007 {published data only}

Kasper B, Dietrich S, Mechtersheimer G, Ho AD, Egerer G. Large institutional experience with dose-intensive chemotherapy and stem cell support in the management of sarcoma patients. *Oncology* 2007;73(1-2):58–64.

Kasper 2010 {published data only}

Kasper B, Scharrenbroich I, Schmitt T, Wuchter P, Dietrich S, Ho AD, et al. Consolidation with high-dose chemotherapy and stem cell support for responding patients with metastatic soft tissue sarcomas: prospective, single-institutional phase II study. *Bone Marrow Transplantation* 2010;45(7):1234–8.

Kozuka 2002 {published data only}

Kozuka T, Kiura K, Katayama H, Fujii N, Ishimaru F, Ikeda K, et al. Tandem high-dose chemotherapy supported by autologous peripheral blood stem cell transplantation for recurrent soft tissue sarcoma. *Anticancer Research* 2002;**22** (5):2939–44.

Kretschmar 1996 {published data only}

Kretschmar CS, Colbach C, Bhan I, Crombleholme TM. Desmoplastic small cell tumor: a report of three cases and a review of the literature. *Journal of Pediatric Hematology and Oncology* 1996;**18**(3):293–8.

Krskova 2007 {published data only}

Krskova L, Sumerauer D, Stejskalova E, Kodet R. A novel variant of SYT-SSX1 fusion gene in a case of spindle cell synovial sarcoma. *Diagnostic Molecular Pathology* 2007;**16** (3):179–83.

Kurre 2000 {published data only}

Kurre P, Felgenhauer JL, Miser JS, Patterson K, Hawkins DS. Successful dose-intensive treatment of desmoplastic small round cell tumor in three children. *Journal of Pediatric Hematology and Oncology* 2000;**22**(5):446–50.

Kushner 1996 {published data only}

Kushner BH, LaQuaglia MP, Wollner N, Meyers PA, Lindsley KL, Ghavimi F, et al.Desmoplastic small round-cell tumor: prolonged progression-free survival with aggressive multimodality therapy. *Journal of Clinical Oncology* 1996; **14**(5):1526–31.

Kushner 2001 {published data only}

Kushner BH, Cheung NK, Kramer K, Dunkel IJ, Calleja E, Boulad F. Topotecan combined with myeloablative doses of thiotepa and carboplatin for neuroblastoma, brain tumors, and other poor-risk solid tumors in children and young adults. *Bone Marrow Transplantation* 2001;**28**(6):551–6.

Kushner 2008 {published data only}

Kushner BH, LaQuaglia MP, Gerald WL, Kramer K, Modak S, Cheung NKV. Solitary relapse of desmoplastic small round cell tumor detected by positron emission tomography/computed tomography. *Journal of Clinical Oncology* 2008;**26**(30):4995–6.

Kühne 2000 {published data only}

Kühne T, Staehelin F, Avoledo P, Tichelli A, Passweg J, Imbach P, et al. Single and double high-dose chemotherapy with autologous stem cell transplantation in children with advanced solid tumors: first experiences [Article in German: Einfache und doppelte Hochdosis—Chemotherapie mit autologer Stammzelltransplantation bei Kindern mit fortgeschrittenen soliden Tumoren: erste Erfahrungen]. Schweizerische Medizinische Wochenschrift 2000;130(12): 419–25.

Lafay-Cousin 2000 {published data only}

Lafay-Cousin L, Hartmann, Plouvier P, Mechinaud F, Boutard P, Oberlin O. High-dose thiotepa and hematopoietic stem cell transplantation in pediatric malignant mesenchymal tumors: a phase II study. *Bone Marrow Transplantation* 2000;**26**(6):627–32.

Lashkari 2009 {published data only}

Lashkari A, Chow WA, Valdes F, Leong L, Phan V, Twardowski P, et al. Tandem high-dose chemotherapy followed by autologous transplantation in patients with locally advanced or metastatic sarcoma. *Anticancer Research* 2009;**29**(8):3281–8.

Lippe 2003 {published data only}

Lippe P, Berardi R, Cappelletti C, Massacesi C, Mattioli R, Latini L, et al.Desmoplastic small round cell tumour:

a description of two cases and review of the literature. *Oncology* 2003;**64**(1):14–7.

Livaditi 2006 {published data only}

Livaditi E, Mavridis G, Soutis M, Papandreou E, Moschovi M, Papadakis V, et al.Diffuse intraabdominal desmoplastic small round cell tumor: a ten-year experience. *European Journal of Pediatric Surgery* 2006;**16**(6):423–7.

Madigan 2007 {published data only}

Madigan CE, Armenian SH, Malogolowkin MH, Mascarenhas L. Extracranial malignant rhabdoid tumors in childhood: The Childrens Hospital Los Angeles experience. *Cancer* 2007;**110**(9):2061–6.

Matsuzaki 2002 {published data only}

Matsuzaki A, Suminoe A, Hattori H, Hoshina T, Hara T. Immunotherapy with autologous dendritic cells and tumor-specific synthetic peptides for synovial sarcoma. *Journal of Pediatric Hematology and Oncology* 2002;**24**(3):220–3.

Mazuryk 1998 {published data only}

Mazuryk M, Paterson AH, Temple W, Arthur K, Crabtree T, Stewart DA. Benefit of aggressive multimodality therapy with autologous stem cell support for intra-abdominal desmoplastic small round cell tumor. *Bone Marrow Transplantation* 1998;**21**(9):961–3.

Mesia 1994 {published data only}

Mesia R, Sola C, Lopez Pousa A, Mendoza L, Bellet M, Andres L, et al.High-dose chemotherapy and autologous bone marrow transplantation in high-grade metastatic sarcomas [Article in Spanish: Quimioterapia con altas dosis y trasplante autologo de medula osea en sarcomas de alto grado metastasicos]. *Revista Clinica Espaniola* 1994;**194** (11):960–5.

Mitchell 1994 {published data only}

Mitchell PL, Shepherd VB, Proctor HM, Dainton M, Cabral SD, Pinkerton CR. Peripheral blood stem cells used to augment autologous bone marrow transplantation. *Archives of Disease in Childhood* 1994;**70**(3):237–40.

Nakamura 2008 {published data only}

Nakamura K, Kaga H, Ogita K, Shiga K, Hikita T, Wakita S, et al. Autologous peripheral blood stem cell transplantation in five advanced pediatric cancer patients [Article in Japanese]. *Teikyo Medical Journal* 2008;**31**(6): 319–30.

Navid 2006 {published data only}

Navid F, Santana VM, Billups CA, Merchant TE, Furman WL, Spunt SL, et al.Concomitant administration of vincristine, doxorubicin, cyclophosphamide, ifosfamide, and etoposide for high-risk sarcomas. *Cancer* 2006;**106**(8): 1846–56.

Patel 2004 {published data only}

Patel SR, Papadopolous N, Raymond AK, Donato M, Seong CM, Yasko AW, et al.A phase II study of cisplatin, doxorubicin, and ifosfamide with peripheral blood stem cell support in patients with skeletal osteosarcoma and variant bone tumors with a poor prognosis. *Cancer* 2004;**101**(1): 156–63.

Peters 1986 {published data only}

Peters WP, Eder JP, Henner WD, Schryber S, Wilmore D, Finberg R, et al. High-dose combination alkylating agents with autologous bone marrow support: a Phase 1 trial. *Journal of Clinical Oncology* 1986;4(5):646–54.

Peters 1989 {published data only}

Peters WP, Stuart A, Klotman M, Gilbert C, Jones RB, Shpall EJ, et al. High-dose combination cyclophosphamide, cisplatin, and melphalan with autologous bone marrow support. *Cancer Chemotherapy and Pharmacology* 1989;**23** (6):377–83.

Recchia 2006 {published data only}

Recchia F, Saggio G, Amiconi G, Di Blasio A, Cesta A, Candeloro G, et al.Cardiac metastases in malignant fibrous histiocytoma. *Tumori* 2006;**92**(1):76–8.

Ronghe 2004 {published data only}

Ronghe MD, Moss TH, Lowis SP. Treatment of CNS malignant rhabdoid tumors. *Pediatric Blood and Cancer* 2004;**42**(3):254–60.

Saab 2007 {published data only}

Saab R, Khoury JD, Krasin M, Davidoff AM, Navid F. Desmoplastic small round cell tumor in childhood: The St. Jude Children's Research Hospital experience. *Pediatric Blood and Cancer* 2007;**49**(3):274–9.

Shaw 1996 {published data only}

Shaw PJ, Pinkerton CR, Yaniv I. Melphalan combined with a carboplatin dose based on glomerular filtration rate followed by autologous stem cell rescue for children with solid tumours. *Bone Marrow Transplantation* 1996;**18**(6): 1043–7.

Slease 1988 {published data only}

Slease RB, Benear JB, Selby GB, Reitz CL, Hughes WL, Watkins CL, et al. High-dose combination alkylating agent therapy with autologous bone marrow rescue for refractory solid tumors. *Journal of Clinical Oncology* 1988;**6**(8): 1314–20.

Sung 2003 {published data only}

Sung KW, Yoo KH, Chung EH, Jung HL, Koo HH, Shin HJ, et al. Successive double high-dose chemotherapy with peripheral blood stem cell rescue collected during a single leukapheresis round in patients with high-risk pediatric solid tumors: a pilot study in a single center. *Bone Marrow Transplantation* 2003;**31**(6):447–52.

Watanabe 2006 {published data only}

Watanabe H, Watanabe T, Kaneko M, Suzuya H, Onishi T, Okamoto Y, et al. Treatment of unresectable malignant rhabdoid tumor of the orbit with tandem high-dose chemotherapy and gamma-knife radiosurgery. *Pediatric Blood and Cancer* 2006;47(6):846–50.

Yamamura 2003 {published data only}

Yamamura R, Yamane T, Aoyama Y, Nakamae H, Makita K, Shima E, et al. Development of chronic myelocytic leukemia after chemotherapy for malignant fibrous histiocytoma. *Acta Haematologica* 2003;**109**(3):141–4.

Yonemoto 1999 {published data only}

Yonemoto T, Tatezaki S, Ishii T, Satoh T. High-dose chemotherapy with autologous peripheral blood stem cell transplantation (PBSCT) for refractory bone and soft tissue sarcomas [Article in Japanese]. *Gan To Kagaku Ryoho* 1999; **26**(10):1431–5.

References to studies excluded from this review

Abdel-Dayem 1999 {published data only}

Abdel-Dayem HM, Rosen G, El-Zeftawy H, Naddaf S, Kumar M, Atay S, et al.Fluorine-18 fluorodeoxyglucose splenic uptake from extramedullary hematopoiesis after granulocyte colony-stimulating factor stimulation. *Clinical Nuclear Medicine* 1999;**24**(5):319–22.

Abidi 2007 {published data only}

Abidi MH, Tove I, Ibrahim RB, Maria D, Peres E. Thalidomide for the treatment of histiocytic sarcoma after hematopoietic stem cell transplant. *American Journal of Hematology* 2007;**82**(10):932–3.

ABMTR 1986 {published data only}

Advisory Committee of the International ABMTR Autologous Bone Marrow Transplant Registry. Bonemarrow autotransplantation in man: report of an international cooperative study. *Lancet* 1986;**2**(8513): 960–2.

ABMTR 1989 {published data only}

Advisory Committee of the International ABMTR Autologous Bone Marrow Transplant Registry. Autologous bone marrow transplants: different indications in Europe and North America. *Lancet* 1989;**2**(8658):317–8.

Abrahamsen 2000 {published data only}

Abrahamsen JF, Kristoffersen EK, Hervig T, Ekanger R, Nesthus I, Ulvestad E. High dose chemotherapy with autologous stem cell support in cancer patients [Article in Norwegian: Hoydosebehandling med autolog stamcellestotte hos kreftpasienter]. *Tidsskrift for den Norske Laegeforening* 2000;**120**(13):1523–8.

Admiraal 2007 {published data only}

Admiraal R, Van der Paardt M, Kobes J, Kremer LCM, Bisogno G, Merks JHM. High dose chemotherapy for children with stage IV rhabdomyosarcoma [Protocol]. *Cochrane Database of Systematic Reviews* 2007, Issue 3. [Art. No.: CD006301. DOI: 10.1002/14651858.CD006301.pub2]

Aleinikova 2002 {published data only}

Aleinikova OV, Strongin IuS, Pochetukhin KV. High-dose chemotherapy with autologous bone marrow transplantation in children with high-risk malignant neoformations [Article in Russian: Vysokodoznaia khimioterapiia s autologichnoi transplantatsiei kostnogo mozga u detei gruppy vysokogo riska so zlokachestvennymi novoobrazovaniiami]. *Voprosy Onkologii* 2002;48(3):327–30.

Alpers 1982 {published data only}

Alpers CE, Biava CG, Salvatierra O Jr. Angiosarcoma following renal transplantation. *Transplantion Proceedings* 1982;**14**(2):448–51.

Anderson 2005 {published data only}

Anderson BD, Schoenfeldt M. Clinical trials referral resource: current phase III clinical trials investigating pediatric cancers. *Oncology* 2005;**19**(1):69–78.

Antman 1987 {published data only}

Antman K, Eder JP, Frei E 3rd. High-dose chemotherapy with bone marrow support for solid tumors. *Important Advances in Oncology* 1987;1:221–35.

Antman 1990 {published data only}

Antman KH, Elias A. Dana-Farber Cancer Institute studies in advanced sarcoma. *Seminars in Oncology* 1990;**17 Suppl 2**(1):7–15.

Antman 2001 {published data only}

Antman KH. New biology and therapies in soft tissue sarcomas. *Biomedicine & Pharmacotherapy* 2001;**55**(9-10): 553–7

Ashihara 2002 {published data only}

Ashihara E, Shimazaki C, Okano A, Hatsuse M, Okamoto A, Shimura K, et al.Infusion of a high number of CD34+ cells provides a rapid hematopoietic recovery and cost savings in autologous peripheral blood stem cell transplantation. *Japanese Journal of Clinical Oncology* 2002; **32**(4):135–9.

Atra 1996 {published data only}

Atra A, Pinkerton R. Autologous stem cell transplantation in solid tumours of childhood. *Annals of Medicine* 1996;**28** (2):159–64.

Atra 2002 {published data only}

Atra A, Pinkerton R. High-dose chemotherapy in soft tissue sarcoma in children. *Critical Reviews in Oncology/Hematology* 2002;**41**(2):191–6.

Avramova 2006 {published data only}

Avramova B, Jordanova M, Michailov G, Konstantinov D, Christosova I, Bobev Dr. Myeloablative chemotherapy with autologous peripheral blood stem cell transplantation in patients with poor-prognosis solid tumors - Bulgarian experience. *Journal of Balkan Union of Oncology* 2006;11 (4):433–8.

Bader 1989 {published data only}

Bader JL, Horowitz ME, Dewan R, Watkins E, Triche TJ, Tsokos M, et al.Intensive combined modality therapy of small round cell and undifferentiated sarcomas in children and young adults: local control and patterns of failure. *Radiotherapy and Oncology* 1989;16(3):189–201.

Bagnulo 1985 {published data only}

Bagnulo S, Perez DJ, Barrett A. High dose melphalan and autologous bone marrow transplantation for solid tumours of childhood. *European Paediatric Haematology/Oncology* 1985;**2**(3):129–33.

Bambakidis 2002 {published data only}

Bambakidis NC, Robinson S, Cohen M, Cohen AR. Atypical teratoid/rhabdoid tumors of the central nervous system: clinical, radiographic and pathologic features. *Pediatric Neurosurgery* 2002;**37**(2):64–70.

Banna 2007 {published data only}

Banna GL, Simonelli M, Santoro A. High-dose chemotherapy followed by autologous hematopoietic stemcell transplantation for the treatment of solid tumors in adults: a critical review. *Current Stem Cell Research and Therapy* 2007;**2**(1):65–82.

Barfield 2008 {published data only}

Barfield RC, Kasow KA, Hale GA. Advances in pediatric hematopoietic stem cell transplantation. *Cancer Biology and Therapy* 2008;7(10):1533–9.

BCBS MAP 1999 {published data only}

Blue Cross and Blue Shield Association. Medical Advisory Panel. Salvage high-dose chemotherapy with allogeneic stem-cell support for relapse following high dose chemotherapy with autologous stem-cell support for solid tumors. *Technology Evaluation Center Assessment Program* 1999;14(1):1–3.

Beaujean 1989 {published data only}

Beaujean F, Hartmann O, Benhamou E, Lemerle J, Duedari N. Hemopoietic reconstitution after repeated autologous transplantation with mafosfamide-purged marrow. *Bone Marrow Transplantation* 1989;4(5):537–41.

Bellmunt 1997 {published data only}

Bellmunt J, Eres N, Ribas A, Casado S, Albanell J, Baselga J. Feasibility trial of high-dose 7-day continuous-infusion ifosfamide given on an outpatient basis. *Cancer Chemotherapy and Pharmacology* 1997;**40**(3):273–6.

Bertuzzi 2002 {published data only}

Bertuzzi A, Castagna L, Nozza A, Quagliuolo V, Siracusano L, Balzarotti M, et al.High-dose chemotherapy in poorprognosis adult small round-cell tumors: clinical and molecular results from a prospective study. *Journal of Clinical Oncology* 2002;**20**(8):2181–8.

Beschorner 2006 {published data only}

Beschorner R, Mittelbronn M, Koerbel A, Ernemann U, Thal DR, Scheel-Walter HG, et al. Atypical teratoid-rhabdoid tumor spreading along the trigeminal nerve. *Pediatric Neurosurgery* 2006;**42**(4):258–63.

Bezwoda 1994 {published data only}

Bezwoda WR, Dansey R, Seymour L, Glencross D. Non-cryopreserved, limited number (1 or 2) peripheral blood progenitor cell (PBPC) collections following GCSF administration provide adequate hematologic support for high dose chemotherapy. *Hematological Oncology* 1994;**12** (3):101–10.

Bickert 2002 {published data only}

Bickert BM. Treatment of common childhood malignancies. *Journal of Pharmacy Practice* 2002;**15**(1):42–51.

Bien 2007 {published data only}

Bien E, Stachowicz-Stencel T, Sierota D, Polczynska K, Szolkiewicz A, Stefanowicz J, et al.Sarcomas in children with neurofibromatosis type 1 - Poor prognosis despite aggressive combined therapy in four patients treated in a single oncological institution. *Child's Nervous System* 2007; 23(10):1147–53.

Bini-Antunes 2006 {published data only}

Bini-Antunes M, Roncon S, Campilho F, Barbosa I, Leal H, Avila A, et al. Peripheral blood progenitor cells collection in paediatric patients [Article in Portuguese: Colheita de celulas progenitoras hematopoieticas perifericas em doentes pediatricos]. *Arquivos de Medicina* 2006;**20**(1-2):25–9.

Bisogno 2009 {published data only}

Bisogno G, Ferrari A, Prete A, Messina C, Basso E, Cecchetto G, et al. Sequential high-dose chemotherapy for children with metastatic rhabdomyosarcoma. *European Journal of Cancer* 2009;45(17):3035–41.

Blay 1994 {published data only}

Blay JY, Bouhour D, Brunat-Mentigny M, Rivoire M, Philip I, Philip T, et al.High-dose chemotherapy (VIC) and bone marrow support in advanced sarcomas. *Bone Marrow Transplantation* 1994;**14 Suppl** 1:55.

Blay 2000 {published data only}

Blay JY, Bouhour D, Ray-Coquard I, Dumontet C, Philip T, Biron P. High-dose chemotherapy with autologous hematopoietic stem-cell transplantation for advanced soft tissue sarcoma in adults. *Journal of Clinical Oncology* 2000; **18**(21):3643–50.

Bode-Lesniewska 2005 {published data only}

Bode-Lesniewska B, Hodler J, Von Hochstetter A, Guillou L, Exner U, Caduff R. Late solitary bone metastasis of a primary pulmonary synovial sarcoma with SYT-SSX1 translocation type: case report with a long follow-up. *Virchows Archiv* 2005;446(3):310–5.

Bodey 1981 {published data only}

Bodey GP, Rodriguez V, Murphy WK, Burgess A, Benjamin RS. Protected environment: prophylactic antibiotic program for malignant sarcomas; randomized trial during remission induction chemotherapy. *Cancer* 1981;47(10): 2422–9

Bojko 2002 {published data only}

Bojko P, Pawloski D, Stellberg W, Schröder JK, Seeber S. Flt3 ligand and thrombopoietin serum levels during peripheral blood stem cell mobilization with chemotherapy and recombinant human glycoslated granulocyte colonystimulating factor (rhu-G-CSF, lenograstim) and after highdose chemotherapy. *Annals of Hematology* 2002;**81**(9): 522–8.

Bokemeyer 1997 {published data only}

Bokemeyer C, Franzke A, Hartmann JT, Schober C, Arseniev L, Metzner B, et al. A phase I/II study of sequential, dose-escalated, high dose ifosfamide plus doxorubicin with peripheral blood stem cell support for the treatment of patients with advanced soft tissue sarcomas. *Cancer* 1997; **80**(7):1221–7.

Borden 1987 {published data only}

Borden EC, Amato DA, Rosenbaum C, Enterline HT, Shiraki MJ, Creech RH, et al.Randomized comparison of three adriamycin regimens for metastatic soft tissue sarcomas. *Journal of Clinical Oncology* 1987;**5**(6):840–50.

Boulad 1998 {published data only}

Boulad F, Kernan NA, LaQuaglia MP, Heller G, Lindsley KL, Rosenfield NS, et al. High-dose induction chemoradiotherapy followed by autologous bone marrow transplantation as consolidation therapy in rhabdomyosarcoma, extraosseous Ewing's sarcoma, and undifferentiated sarcoma. *Journal of Clinical Oncology* 1998; **16**(5):1697–706.

Bramwell 1986 {published data only}

Bramwell VH, Mouridsen HT, Santoro A, Blackledge G, Somers R, Thomas D, et al. Cyclophosphamide versus ifosfamide: preliminary report of a randomized phase II trial in adult soft tissue sarcomas. *Cancer Chemotherapy and Pharmacology* 1986;**18 Suppl 2**:13–6.

Bramwell 1987 {published data only}

Bramwell VH, Mouridsen HT, Santoro A, Blackledge G, Somers R, Verwey J, et al. Cyclophosphamide versus ifosfamide: final report of a randomized phase II trial in adult soft tissue sarcomas. *European Journal of Cancer and Clinical Oncology* 1987;**23**(3):311–21.

Bramwell 2001 {published data only}

Bramwell V, Anderson D, Charette M. Doxorubicin-based chemotherapy for the palliative treatment of adult patients with locally advanced or metastatic soft tissue sarcoma. *Cochrane Database of Systematic Reviews* 2001, Issue 4. [Art. No.: CD003293. DOI: 10.1002/14651858.CD003293]

Breitfeld 2001 {published data only}

Breitfeld PP, Lyden E, Raney RB, Teot LA, Wharam M, Lobe T, et al.Ifosfamide and etoposide are superior to vincristine and melphalan for pediatric metastatic rhabdomyosarcoma when administered with irradiation and combination chemotherapy: a report from the Intergroup Rhabdomyosarcoma Study Group. *Journal of Pediatric Hematology/Oncology* 2001;23(4):225–33.

Brugger 1995 {published data only}

Brugger W, Heimfeld S, Berenson RJ, Mertelsmann R, Kanz L. Reconstitution of hematopoiesis after high-dose chemotherapy by autologous progenitor cells generated ex vivo. *New England Journal of Medicine* 1995;**333**(5):283–7.

Brugieres 1988 {published data only}

Brugieres L, Hartmann O, Benhamou E, Patte C, Kalifa C, Lemerle J. Hepatic complications after high-dose chemotherapy and bone marrow transplantation for solid tumors in children. *Presse Medicale* 1988;17(25):1305–8.

Brun 1984 {published data only}

Brun del Re G, Baumgartner C, Bleher EA, Bucher U, Greiner R, Hirt A, et al.Autologe Knochenmarktransplantation in der Behandlung von fortgeschrittenen bosartigen Tumoren bei Kindern und Jugendlichen. *Folia Haematologica* 1984;**111**(2):243–8.

Bylund 2008 {published data only}

Bylund KC, Giampoli E, Singh D, Doerr T, Sahasrabudhe D, Liesveld J, et al.Soft tissue sarcoma in the setting of chronic cutaneous graft versus host disease after allogenic bone marrow transplantation. *Cancer Investigation* 2008;**26** (6):638–41.

Cacchione 2008 {published data only}

Cacchione A, LeMaitre A, Valteau Couanet D, Benhamou E, Amoroso L, Simonnard N, et al.Risk factors for hepatic veno-occlusive disease: a retrospective unicentric study in 116 children autografted after a high-dose BU-thiotepa regimen. *Bone Marrow Transplantation* 2008;42(7):449–54.

Caglar 2002 {published data only}

Caglar K, Kinalp C, Arpaci F, Turan M, Saglam K, Ozturk B, et al. Cumulative prior dose of cisplatin as a cause of the nephrotoxicity of high-dose chemotherapy followed by autologous stem-cell transplantation. *Nephrology Dialysis Transplantation* 2002;**17**(11):1931–5.

Carli 1988 {published data only}

Carli M, Pastore G, Perilongo G, Grotto P, De Bernardi B, Ceci A, et al. Tumor response and toxicity after single highdose versus standard five-day divided-dose dactinomycin in childhood rhabdomyosarcoma. *Journal of Clinical Oncology* 1988;**6**(4):654–8.

Carli 1999 {published data only}

Carli M, Colombatti R, Oberlin O, Bisogno G, Treuner J, Koscielniak E, et al.European intergroup studies (MMT4-89 and MMT4-91) on childhood metastatic rhabdomyosarcoma: final results and analysis of prognostic factors. *Journal of Clinical Oncology* 2004;22(23): 4787–4794.

* Carli M, Colombatti R, Oberlin O, Stevens M, Masiero L, Frascella E, et al. High-dose melphalan with autologous stem-cell rescue in metastatic rhabdomyosarcoma. *Journal of Clinical Oncology* 1999;17(9):2796–803.

Casado 2004 {published data only}

Casado Herraez A, Moreno Anton F. Chemotherapy of soft tissue sarcomas. *Revisiones en Cancer* 2004;**18**(6):316–29.

Casanova 2009 {published data only}

Casanova M, Meazza C, Favini F, Fiore M, Morosi C, Ferrari A. Rhabdomyosarcoma of the extremities: A focus on tumors arising in the hand and foot. *Pediatric Hematology and Oncology* 2009;**26**(5):321–331.

Casper 1991 {published data only}

Casper ES, Gaynor JJ, Hajdu SI, Magill GB, Tan C, Friedrich C, et al.A prospective randomized trial of adjuvant chemotherapy with bolus versus continuous infusion of doxorubicin in patients with high-grade extremity soft tissue sarcoma and an analysis of prognostic factors. *Cancer* 1991; **68**(6):1221–9.

Ceschel 2006 {published data only}

Ceschel S, Casotto V, Valsecchi MG, Tamaro P, Jankovic M, Hanau G, et al. Survival after relapse in children with solid tumors: a follow-up study from the Italian off-therapy registry. *Pediatric Blood and Cancer* 2006;47:560–6.

Chan 1991 {published data only}

Chan KW, Rogers PC, Fryer CJ. Breast metastases after bone marrow transplantation for rhabdomyosarcoma. *Bone Marrow Transplantation* 1991;7(2):171–2.

Chan 1999 {published data only}

Chan LL, Lin HP, Ariffin WA, Ariffin H, Saw MH. Treating high risk childhood solid tumours with autologous peripheral blood stem cell transplantation: early experience in University Hospital, Kuala Lumpur. *Medical Journal of Malaysia* 1999;**54**(2):175–9.

Chang 1979 {published data only}

Chang AE, Shiling DJ, Stillman RC. A prospective randomized trial of delta-9-tetrahydrocannabinol (THC) as an antiemetic in patients receiving high dose methotrexate (MTX). Proceeding of the American Association of Cancer Research 1979;20:abstract No. C-357.

Chang 1979a {published data only}

Chang AE, Shiling DJ, Stillman RC. Delta-9-tetrahydrocannabinol as an antiemetic in cancer patients receiving high-dose methotrexate: a prospective, randomized evaluation. *Annals of Internal Medicine* 1979; **91**(6):819–24.

Chang 1981 {published data only}

Chang AE, Shiling DJ, Stillman RC, Goldberg NH, Seipp CA, Barofsky I, et al.A prospective evaluation of delta-9-tetrahydrocannabinol as an antiemetic in patients receiving adriamycin and cytoxan chemotherapy. *Cancer* 1981;47(7): 1746–51.

Chang 1988 {published data only}

Chang AE, Kinsella T, Glatstein E, Baker AR, Sindelar WF, Lotze MT, et al. Adjuvant chemotherapy for patients with high-grade soft-tissue sarcomas of the extremity. *Journal of Clinical Oncology* 1988;6(9):1491–500.

Chauvin 1991 {published data only}

Chauvin F, Ladenstein R, Lasset C, Pinzani V, Abdelbost Z, Bartolomucci A, et al. European Bone Marrow Registry in solid tumors: 7 years of experience. *Bone Marrow Transplantation* 1991;**7 Suppl** 2:157.

Chen 1999 {published data only}

Chen AR. High-dose therapy with stem cell rescue for pediatric solid tumors: rationale and results. *Pediatric Transplantation* 1999;**3 Suppl** 1:78–86.

Chen 2008 {published data only}

Chen IL, Yang SN, Hsiao CC, Wu KS, Sheen JM. Treatment with high-dose methylprednisolone for hepatic veno-occlusive disease in a child with rhabdomyosarcoma. *Pediatrics and Neonatology* 2008;**49**(4):141–4.

Childs 2004 {published data only}

Childs RW. Evolving trends in hematopoietic cell transplantation for solid tumors: tempering enthusiasm with clinical reality. *Annals of Oncology* 2004;**15**(4):543–4.

Cho 2005 {published data only}

Cho HJ, Jung HK, Sung KW, Ku HH, Lee SH, Kim DW. Autologous peripheral blood stem cell collections in children weighing less than 10 Kg with solid tumors: experience of a single center. *Journal of Clinical Apheresis* 2005;**20**(2):65–71.

Chuman 2000 {published data only}

Chuman H. Evidence-based chemotherapy for patients with bone and soft part sarcoma [Japanese]. *Gan to Kagaku Ryoho [Japanese Journal of Cancer and Chemotherapy*] 2000; **27**(2):192–202.

Clausen 1993 {published data only}

Clausen N, Schroder H. Autologous bone marrow transplantation in children [Article in Danish: Autolog knoglemarvstransplantation til born]. *Ugeskrift for Laeger* 1993;**155**(20):1531–5.

Corbett 2009 {published data only}

Corbett RP. Childhood solid tumours occurring in adolescents and young adults. *Cancer Forum* 2009;**33**(1): 13–7.

Coulibalya 2008 {published data only}

Coulibalya B, Liprandia A, Le Hemon A, Fernandez C, Hardwigsen J, Berthet B, et al.Desmoplastic small round-cell tumor: two cases of diffuse abdominopelvic infiltration [Article in French: Tumeur desmoplastique a petites cellules rondes: deux cas d'atteinte abdominopelvienne diffuse]. Gastroenterologie Clinique et Biologique 2008;32(3):278–81.

Couzin 2007 {published data only}

Couzin J. In their prime, and dying of cancer. *Science* 2007; **317**(5842):1160–2.

Czyzewski 1999 {published data only}

Czyzewski EA, Goldman S, Mundt AJ, Nachman J, Rubin C, Hallahan DE. Radiation therapy for consolidation of metastatic or recurrent sarcomas in children treated with intensive chemotherapy and stem cell rescue. *International Journal of Radiation Oncology, Biology, Physics* 1999;44(3): 569–77.

Dagher 1997 {published data only}

Dagher R, Robertson KA, Lucas KG, Emanuel D, Smith FO. Outpatient total body irradiation for pediatric patients undergoing stem cell transplantation. *Bone Marrow Transplantation* 1997;**19**(11):1065–7.

Dallorso 1996 {published data only}

Dallorso S, Dini G, Miano M, Rivabella L, Scarso L, Martinengo M, et al.G-CSF primed peripheral blood progenitor cells (PBPC) autotransplantation in stage IV Neuroblastoma and poor risk solid tumors. *Bone Marrow Transplantation* 1996;**18 Suppl 2**:182–4.

Dallorso 2000 {published data only}

Dallorso S, Manzitti C, Morreale G, Faraci M. High dose therapy and autologous hematopoietic stem cell transplantation in poor risk solid tumors of childhood. *Haematologica* 2000;**85 Suppl**(11):66–70.

Dantonello 2008 {published data only}

Dantonello TM, Int-Veen C, Winkler P, Leuschner I, Schuck A, Schmidt BF, et al.Initial patient characteristics can predict pattern and risk of relapse in localized rhabdomyosarcoma. *Journal of Clinical Oncology* 2008;**26** (3):406–13.

De Kraker 1984 {published data only}

De Kraker J, Voute PA, Behrendt H. Intensive chemotherapy followed by autologous bone marrow transplantation in solid tumors in childhood [Article in Dutch: Intensieve chemotherapie gevolgd door autologe beenmergtransplantatie bij solide tumoren op de kinderleeftijd]. Nederlands Tijdschrift voor Geneeskunde 1984;128(49):2302–5.

De Pasquale 2003 {published data only}

De Pasquale MD, Cacchione A, Foco M, Gozzer M, Libera F, Soscia F, et al. Peripheral blood stem cell collection in pediatric malignancies: comparison between three mobilizing regimens [Article in Italian: La raccolta di cellule staminali periferiche nei tumori pediatrici: confronto fra tre sistemi di mobilizzazione]. *La Clinica Terapeutica* 2003;154 (5):305–9.

De Sio 2006 {published data only}

De Sio L, Milano GM, Castellano A, Jenkner A, Fidani P, Dominici C, et al. Temozolomide in resistant or relapsed pediatric solid tumors. *Pediatric Blood & Cancer* 2006;47 (1):30–6.

De Terlizzi 1988 {published data only}

De Terlizzi M, Philip T, Toma MG, Colella R, Ceci A. Massive therapy and transplant of autologous bone marrow in childhood lymphomas and solid tumors: state of art and future perspectives [Article in Italian: Terapia massiva e trapianto di midollo autologo nei linfomi e tumori solidi pediatrici. Stato dell'arte e prospettive]. *La Pediatria Medica e Chirurgica* 1988;**10**(4):359–64.

De Vries 1995 {published data only}

De Vries EG, de Graaf H, Boonstra A, van der Graaf WT, Mulder NH. High-dose chemotherapy with stem cell reinfusion and growth factor support for solid tumors. *Stem Cells* 1995;**13**(6):597–606.

Demirci 2003 {published data only}

Demirci H, Shields JA, Shields CL, Maguire AM. Atypical presentation of bone marrow transplant retinopathy. *Journal of Pediatric Ophthalmology and Strabismus* 2003;**40** (6):361–3.

Demirer 2008 {published data only}

Demirer T, Barkholt L, Blaise D, Pedrazzoli P, Aglietta M, Carella AM, et al.Transplantation of allogeneic hematopoietic stem cells: an emerging treatment modality for solid tumors. *Nature Clinical Practice. Oncology* 2008;**5** (5):256–67.

Devalck 1992 {published data only}

Devalck C, Ferster A, De Laet MH, Nafa S, Bujan W, Azzi N, et al. Autologous bone marrow graft in solid tumors in childhood [Article in French: Greffe de moelle autologue dans les tumeurs solides des enfants]. *Revue Medicale de Bruxelles* 1992;**13**(6):201–6.

Diaz 1999 {published data only}

Diaz MA, Vicent MG, Madero L. High-dose busulfan/melphalan as conditioning for autologous PBPC transplantation in pediatric patients with solid tumors. *Bone Marrow Transplantation* 1999;**24**(11):1157–9.

Dicke 1984 {published data only}

Dicke KA, Jagannath S, Spitzer G, Poynton C, Zander A, Vellekoop L, et al. The role of autologous bone marrow transplantation in various malignancies. *Seminars in Hematology* 1984;**21**(2):109–22.

Dicke 1986 {published data only}

Dicke KA, Spitzer G. Evaluation of the use of high-dose cytoreduction with autologous marrow rescue in various malignancies. *Transplantation* 1986;**41**(1):4–20.

Dileo 2005 {published data only}

Dileo P, Demetri GD. Update on new diagnostic and therapeutic approaches for sarcomas. *Clinical Advances in Hematology and Oncology* 2005;**3**(10):781–91.

Dillman 1995 {published data only}

Dillman RO, Barth NM, Mahdavi K, VanderMolen LA, Nayak SK, O'Connor A. The integration of high-dose chemotherapy and biotherapy: initial 5-year experience with autologous bone marrow transplantation in a comprehensive community cancer center. *Cancer Biotherapy* 1995;**10**(1):25–36.

Dincol 2000 {published data only}

Dincol D, Samur M, Pamir A, Sencan O, Akbulut H, Yalcin B, et al. Prospective randomized comparison of morning versus night daily single subcutaneous administration of granulocyte-macrophage-colony stimulating factor in patients with soft tissue or bone sarcoma. *Cancer* 2000;**88** (9):2033–6.

Donaldson 2001 {published data only}

Donaldson SS, Meza J, Breneman JC, Crist WM, Laurie F, Qualman SJ, et al.Results from the IRS-IV randomized trial of hyperfractionated radiotherapy in children with rhabdomyosarcoma: a report from the IRSG. *International Journal of Radiation Oncology, Biology, Physics* 2001;**51**(3): 718–28.

Donker 2009 {published data only}

Donker AE, Hoogerbrugge PM, Mavinkurve-Groothuis AMC, van de Kar NCAJ, Boetes C, Hulsbergen-van de Kaa CA, et al.Metastatic rhabdomyosarcoma cured after chemotherapy and allogeneic SCT. *Bone Marrow Transplantation* 2009;**43**(2):179–80.

Drabko 2006 {published data only}

Drabko K, Choma M, Zaucha-Prazmo A, Wojcik B, Gorczynska E, Kalwak K, et al.Megachemotherapy and autologous hematopoietic stem cell transplantation in children with solid tumours excluding neuroblastoma-experience of Polish paediatric centres [Article in Polish: Megachemotherapia i przeszczepianie autologicznych hematopoetycznych komorek macierzystych u dzieci z guzami litymi innymi niz neuroblastoma—doswiadczenia polskich osrodkow pediatrycznych]. *Medycyna Wieku Rozwojowego* 2006;**10**(3 Pt 1):785–92.

Dumontet 1992 {published data only}

Dumontet C, Biron P, Bouffet E, Blay JY, Meckenstock R, Chauvin F, et al. High dose chemotherapy with ABMT in soft tissue sarcomas: a report of 22 cases. *Bone Marrow Transplantation* 1992;**10**(5):405–8.

Ederhy 2007 {published data only}

Ederhy S. Prevention of high-dose chemotherapy-induced cardiotoxicity in high-risk patients by angiotensin-converting enzyne inhibition [Prevention de la cardiotoxicite de chimiotherapies intensives par les inhibiteurs de l'enzyme

de conversion chez des patients a haute risque]. *Medecine Therapeutique Cardio* 2007;**3**(1):45–7.

Eggermont 1997 {published data only}

Eggermont AM, Schraffordt Koops H, Klausner JM, Lienard D, Kroon BB, Schlag PM, et al.Isolation limb perfusion with tumor necrosis factor alpha and chemotherapy for advanced extremity soft tissue sarcomas. Seminars in Oncology 1997;24(5):547–55.

Ek 2006 {published data only}

Ek ETH, Choong PFM. The role of high-dose therapy and autologous stem cell transplantation for pediatric bone and soft tissue sarcomas. *Expert Review of Anticancer Therapy* 2006;**6**(2):225–37.

Ekert 1982 {published data only}

Ekert H, Ellis WM, Waters KD, Tauro GP. Autologous bone marrow rescue in the treatment of advanced tumors of childhood. *Cancer* 1982;49(3):603–9.

Ekert 1984 {published data only}

Ekert H, Tiedemann K, Waters KD, Ellis WM. Experience with high dose multiagent chemotherapy and autologous bone marrow rescue in the treatment of twenty-two children with advanced tumours. *Australian Paediatric Journal* 1984; **20**(3):195–201.

Elias 1998 {published data only}

Elias AD. High-dose therapy for adult soft tissue sarcoma: dose response and survival. *Seminars in Oncology* 1998;**25 Suppl 4**(2):19–23.

Emminger 1991 {published data only}

Emminger W, Emminger-Schmidmeier W, Peters C, Hawliczek R, Hocker P, Gadner H. Is treatment intensification by adding etoposide and carboplatin to fractionated total body irradiation and melphalan acceptable in children with solid tumors with respect to toxicity? *Bone Marrow Transplantation* 1991;8(2):119–23.

Endo 1995 {published data only}

Endo M, Tanosaki R. Myeloablative chemotherapy with autologous bone marrow and/or peripheral blood stem cell transplantation in children with high-risk solid tumor [Article in Japanese]. *Gan to Kagaku Ryoho* 1995;**22**(12): 1762–70.

Erkisi 1996 {published data only}

Erkisi M, Erkurt E, Ozbarlas S, Burgut R, Doran F, Seyrek E. The use of recombinant human granulocyte colony-stimulating factor in combination with single or fractionated doses of ifosfamide and doxorubicin in patients with advanced soft tissue sarcoma. *Journal of Chemotherapy (Florence, Italy)* 1996;**8**(3):224–8.

Espinosa 2001 {published data only}

Espinosa AE, Gonzalez BM. High-dose chemotherapy in nonhematologic tumors [Article in Spanish: Quimioterapia a altas dosis en el tratamiento de tumores no hematologicos]. *Revista Clinica Espanola* 2001;**201**(2):93–4.

Fazekas 2008 {published data only}

Fazekas T, Wiesbauer P, Kronberger M, Wank H, Gadner H, Dworzak M. Nodular pulmonary lesions in children

after autologous stem cell transplantation: a source of misinterpretation. *British Journal of Haematology* 2008;**140** (4):429–32.

Fekrat 1993 {published data only}

Fekrat S, Miller NR, Loury MC. Alveolar rhabdomyosarcoma that metastasized to the orbit. *Archives of Ophthalmology* 1993;**111**(12):1662–4.

Ferrari 2005 {published data only}

Ferrari A, Casanova M, Collini P, Meazza C, Luksch R, Massimino M, et al. Adult-type soft tissue sarcomas in pediatric-age patients: Experience at the Istituto Nazionale Tumori in Milan. *Journal of Clinical Oncology* 2005;**23**(18): 4021–30.

Fetscher 1996 {published data only}

Fetscher S, Kiani A, Kanz L, Brugger W, Lange W, Mertelsmann R. Neo-adjuvant high-dose chemotherapy with autologous peripheral blood stem cell transplantation for inoperable relapse of nuchal liposarcoma resistant to standard-dose chemotherapy. *Annals of Oncology* 1996;7(8): 871.

Figuerres 2000 {published data only}

Figuerres E, Haut PR, Olzewski M, Kletzel M. Analysis of parameters affecting engraftment in children undergoing autologous peripheral blood stem cell transplants. *Bone Marrow Transplantation* 2000;**25**(6):583–8.

Fizazi 1994 {published data only}

Fizazi K, Cojean I, Le Cesne A, Kayitalire L, Le Chevalier T, Tursz T, et al. Soft tissue sarcomas: general review [Article in French: Sarcomes des tissus mous: revue generale]. *Bulletin du Cancer* 1994;**81**(10):835–52.

Flamant 1998 {published data only}

Flamant F, Rodary C, Rey A, Praquin MT, Sommelet D, Quintana E, et al.Treatment of non-metastatic rhabdomyosarcomas in childhood and adolescence: results of the second study of the International Society of Paediatric Oncology; MMT84. *European Journal of Cancer* 1998;**34** (7):1050–62.

Foncillas 2004 {published data only}

Foncillas MA, Diaz MA, Sevilla J, Gonzalez VM, Fernandez-Plaza S, Perez A, et al.Engraftment syndrome emerges as the main cause of transplant-related mortality in pediatric patients receiving autologous peripheral blood progenitor cell transplantation. *Journal of Pediatric Hematology/ Oncology* 2004;**26**(8):492–6.

Frustaci 2001 {published data only}

Frustaci S, Gherlinzoni F, De Paoli A, Bonetti M, Azzarelli A, Comandone A, et al. Adjuvant chemotherapy for adult soft tissue sarcomas of the extremities and girdles: results of the Italian randomized cooperative trial. *Journal of Clinical Oncology* 2001;**19**(5):1238–47.

Fujita 2005 {published data only}

Fujita M, Sato M, Nakamura M, Kudo K, Nagasaka T, Mizuno M, et al. Multicentric atypical teratoid/rhabdoid tumors occurring in the eye and fourth ventricle of an infant: case report. *Journal of Neurosurgery* 2005;**102 Suppl** (3):299–302.

Gadner 2002 {published data only}

Gadner H. Is there evidence-based benefit of autologous stem cell transplantation in children with solid tumors?. *Onkologie* 2002;**25**(3):278–81.

Garaventa 1986 {published data only}

Garaventa A, Lanino E, Dallorso S, Dini G, Buffa P, Baldelli I, et al. Total parenteral nutrition in children with cancer under aggressive chemotherapy with or without autologous bone marrow transplantation [Article in Italian: Nutrizione parenterale in bambini con neoplasia maligna sottoposti a chemioterapia aggressiva]. *Rivista Italiana di Nutrizione Parenterale ed Enterale* 1986;4(3):159–64.

Garaventa 1987 {published data only}

Garaventa A, Lanino E, Dini G, Dallorso S, Viscoli C, Loy A, et al. Autologous bone marrow transplantation in children. Use of parenteral nutrition [Trapianto di midollo autologo in bambini affetti da neoplasie maligne. Impiego della nutrizione parenterale]. *La Pediatria Medica e Chirurgica* 1987;9(3):259–62.

Gardner 2008 {published data only}

Gardner SL, Asgharzadeh S, Green A, Horn B, McCowage G, Finlay J. Intensive induction chemotherapy followed by high dose chemotherapy with autologous hematopoietic progenitor cell rescue in young children newly diagnosed with central nervous system atypical teratoid rhabdoid tumors. *Pediatric Blood and Cancer* 2008;**51**(2):235–40.

Gebhardt 1999 {published data only}

Gebhardt MC, Parekh SG, Rosenberg AE, Rosenthal DI. Extraskeletal myxoid chondrosarcoma of the knee. *Skeletal Radiology* 1999;**28**(6):354–8.

Geisler 2003 {published data only}

Geisler CH, Daugaard KG, Dickmeiss E, Ifversen M, Knudsen LM. Treatment of cancer with high-dose chemotherapy and autologous stem cell transplantation [Article in Danish: Behandling af kraeftsygdomme med hojdosis kemoterapi og autolog stamcelletransplatation]. *Ugeskrift for Laeger* 2003;**165**(50):4846–9.

Geissler 1984 {published data only}

Geissler K, Lenzhofer R, Schneeweiss B, Moser K. Clinical experiences with high-dose methotrexate [Article in German: Klinische Erfahrungen mit hochdosiertem Methotrexat]. Wiener Klinische Wochenschrift 1984;96(10): 381–9.

Gentet 1993 {published data only}

Gentet JC, Plouvier E, Coze C. Bone marrow autograft and cancer in children [Article in French: Autogreffes de moelle osseuse et cancers pediatriques]. *La Revue du Praticien* 1993; 43(17):2213–7.

Ghalie 1994 {published data only}

Ghalie R, Reynolds J, Valentino LA, Manson S, Korenblit AD, Feingold JM, et al. Busulfan-containing pre-transplant regimens for the treatment of solid tumors. *Bone Marrow Transplantation* 1994;14(3):437–42.

Ghavamzadeh 2009 {published data only}

Ghavamzadeh A, Alimogaddam K, Jahani M, Mousavi SA, Iravani M, Bahar B, et al. Stem cell transplantation: Iranian experience. *Archives of Iranian Medicine* 2009;**12**(1):69–72.

Glenn 1985 {published data only}

Glenn J, Sindelar WF, Kinsella T, Glatstein E, Tepper J, Costa J, et al.Results of multimodality therapy of resectable soft-tissue sarcomas of the retroperitoneum. *Surgery* 1985; **97**(3):316–25.

Gonzalez 1989 {published data only}

Gonzalez Rivero MA, Mulet JF, Martin JC, Illa J, Melo M, Pardo N, et al.Genitourinary rhabdomyosarcoma in childhood [Article in Spanish: Rabdomiosarcomas genitourinarios en la infancia]. *Cirugia Pediatrica* 1989;**2** (4):186–90.

Gortzak 2001 {published data only}

Gortzak E, Azzarelli A, Buesa J, Bramwell VH, van Coevorden F, van Geel AN, et al. A randomised phase II study on neo-adjuvant chemotherapy for 'high-risk' adult soft-tissue sarcoma. *European Journal of Cancer* 2001;**37**(9): 1096–103.

Goto 2004 {published data only}

Goto T, Kosaku H, Kobayashi H, Hozumi T, Kondo T. Soft tissue sarcoma: postoperative chemotherapy [Article in Japanese]. *Gan to Kagaku Ryoho* 2004;**31**(9):1324–30.

Graham 1992 {published data only}

Graham ML, Yeager AM, Leventhal BG, Wiley JM, Civin CI, Strauss LC, et al.Treatment of recurrent and refractory pediatric solid tumors with high-dose busulfan and cyclophosphamide followed by autologous bone marrow rescue. *Journal of Clinical Oncology* 1992;**10**(12):1857–64.

Graham-Pole 1995 {published data only}

Graham-Pole J. Pediatric malignancies workshop: section on sarcomas and PNETs. *Bone Marrow Transplantation* 1995;**15 Suppl 1**:247–52.

Gratwohl 2002 {published data only}

Gratwohl A, Baldomero H, Passweg J, Urbano-Ispizua A. Increasing use of reduced intensity conditioning transplants: report of the 2001 EBMT activity survey. *Bone Marrow Transplantation* 2002;**30**(12):813–31.

Gratwohl 2004 {published data only}

Gratwohl A, Schmid O, Baldomero H, Horisberger B, Urbano-Ispizua A. Haematopoietic stem cell transplantation (HSCT) in Europe 2002. *Bone Marrow Transplantation* 2004;**34**(10):855–75.

Gratwohl 2004a {published data only}

Gratwohl A, Baldomero H, Demirer T, Rosti G, Dini G, Ladenstein R, et al. Hematopoetic stem cell transplantation for solid tumors in Europe. *Annals of Oncology* 2004;**15**(4): 653–60.

Gratwohl 2004b {published data only}

Gratwohl A. Overview of transplant activity in Europe. *The Hematology Journal* 2004;**5 Suppl** 3:29–33.

Gratwohl 2006 {published data only}

Gratwohl A, Baldomero H, Frauendorfer K, Urbano-Ispizua A. EBMT activity survey 2004 and changes in disease indication over the past 15 years. *Bone Marrow Transplantation* 2006;**37**(12):1069–85.

Gratwohl 2007 {published data only}

Gratwohl A. Activity survey and historical perspective of autologous stem cell transplantation in Europe. *Seminars in Hematology* 2007;44(4):220–6.

Gratwohl 2007a {published data only}

Gratwohl A, Baldomero H, Frauendorfer K, Urbano-Ispizua A, Niederwieser D. Results of the EBMT activity survey 2005 on haematopoietic stem cell transplantation: focus on increasing use of unrelated donors. *Bone Marrow Transplantation* 2007;**39**(2):71–87.

Grundy 2001 {published data only}

Grundy R, Anderson J, Gaze M, Gerrard M, Glaser A, Gordon A, et al. Congenital alveolar rhabdomyosarcoma: clinical and molecular distinction from alveolar rhabdomyosarcoma in older children. *Cancer* 2001;**91**(3): 606–12.

Haas 1990 {published data only}

Haas R, Ho AD, Bredthauer U, Cayeux S, Egerer G, Knauf W, et al. Successful autologous transplantation of blood stem cells mobilized with recombinant human granulocytemacrophage colony-stimulating factor. *Experimental Hematology* 1990;**18**(2):94–8.

Hale 2005 {published data only}

Hale GA. Autologous hematopoietic stem cell transplantation for pediatric solid tumors. *Expert Review of Anticancer Therapy* 2005;5(5):835–46.

Hara 1998 {published data only}

Hara J, Osugi Y, Ohta H, Matsuda Y, Nakanishi K, Takai K, et al.Double-conditioning regimens consisting of thiotepa, melphalan and busulfan with stem cell rescue for the treatment of pediatric solid tumors. *Bone Marrow Transplantation* 1998;**22**(1):7–12.

Harmon 2001 {published data only}

Harmon DC. Advances in chemotherapeutic treatments for bone and soft tissue sarcomas. *Current Opinion in Orthopaedics* 2001;**12**(6):499–504.

Hartmann 1986 {published data only}

Hartmann O, Benhamou E, Beaujean F, Pico JL, Kalifa C, Patte C, et al. High-dose busulfan and cyclophosphamide with autologous bone marrow transplantation support in advanced malignancies in children: a phase II study. *Journal of Clinical Oncology* 1986;4(12):1804–10.

Hartmann 1997 {published data only}

Hartmann O, Le Carroller AG, Blaise D, Michon J, Philip I, Norol F, et al. Peripheral blood stem cell and bone marrow transplantation for solid tumors and lymphomas: Hematologic recovery and costs: A randomized, controlled trial. *Annals of Internal Medicine* 1997;**126**(8):600–7.

Hartmann 2001 {published data only}

Hartmann JT, Von Vangerow A, Fels LM, Knop S, Stolte H, Kanz L, et al.A randomized trial of amifostine in patients with high-dose VIC chemotherapy plus autologous blood stem cell transplanation. *British Journal of Cancer* 2001;**84** (3):313–20.

Hartmann 2005 {published data only}

Hartmann O. High dose chemotherapy for childhood solid tumours: lessons to learn and future developments. European Journal of Cancer Supplements 2005;3(3):438–40.

He 1999 {published data only}

He FX, Fan SH, Ge LZ. Combination therapy in the prevention of postoperative recurrence of soft tissue sarcomas: field-in-field radiotherapy and high dose cisplatin-hydration chemotherapy. *Chinese Journal of Radiation Oncology* 1999;**8**(4):219–21.

Hensel 2002 {published data only}

Hensel M, Egerer G, Schneeweiss A, Goldschmidt H, Ho AD. Quality of life and rehabilitation in social and professional life after autologous stem cell transplantation. *Annals of Oncology* 2002;**13**(2):209–17.

Herzog 2005 {published data only}

Herzog CE. Sarcomas in adolescents and young adults: a summary of a recent symposium. *Journal of Pediatric Hematology/Oncology* 2005;**27**(4):177–8.

Hilden 1998 {published data only}

Hilden JM, Watterson J, Longee DC, Moertel CL, Dunn ME, Kurtzberg J, et al. Central nervous system atypical teratoid tumor/rhabdoid tumor: response to intensive therapy and review of the literature. *Journal of Neuro-oncology* 1998;**40**(3):265–75.

Hilden 2004 {published data only}

Hilden JM, Meerbaum S, Burger P, Finlay J, Janss A, Scheithauer BW, et al. Central nervous system atypical teratoid/rhabdoid tumor: results of therapy in children enrolled in a registry. *Journal of Clinical Oncology* 2004;**22** (14):2877–84.

Hiraiwa 1983 {published data only}

Hiraiwa A, Akao Y, Sao H. Autologous bone marrow transplantation after high-dose combination chemotherapy in the treatment of selective malignancies. *Nippon Gan Chiryo Gakkai Shi* 1983;18(5):1167–74.

Hoekstra 1994 {published data only}

Hoekstra HJ, Schraffordt Koops H, Oldhoff J. Soft tissue sarcoma of the extremity. *European Journal of Surgical Oncology* 1994;**20**(1):3–6.

Holtta 2005 {published data only}

Holtta P, Alaluusua S, Saarinen-Pihkala UM, Peltola J, Hovi L. Agenesis and microdontia of permanent teeth as late adverse effects after stem cell transplantation in young children. *Cancer* 2005;**103**(1):181–90.

Holtta 2005a {published data only}

Holtta P, Hovi L, Saarinen-Pihkala UM, Peltola J, Alaluusua S. Disturbed root development of permanent teeth after

pediatric stem cell transplantation: dental root development after SCT. *Cancer* 2005;103(7):1484–93.

Horn 2002 {published data only}

Horn B, Reiss U, Matthay K, McMillan A, Cowan M. Veno-occlusive disease of the liver in children with solid tumors undergoing autologous hematopoietic progenitor cell transplantation: a high incidence in patients with neuroblastoma. *Bone Marrow Transplantation* 2002;**29**(5): 409–15.

Horowitz 1993 {published data only}

Horowitz ME, Kinsella TJ, Wexler LH, Belasco J, Triche T, Tsokos M, et al. Total-body irradiation and autologous bone marrow transplant in the treatment of high-risk Ewing's sarcoma and rhabdomyosarcoma. *Journal of Clinical Oncology* 1993;**11**(10):1911–8.

Hosoi 2007 {published data only}

Hosoi H, Teramukai S, Matsumoto Y, Tsuchiya K, Iehara T, Hara J, et al.A review of 331 rhabdomyosarcoma cases in patients treated between 1991 and 2002 in Japan. *International Journal of Clinical Oncology* 2007;**12**(2): 137–45.

Hotte 2004 {published data only}

Hotte SJ, Smith AM, Bramwell VHC, Howson-Jan K. High-dose chemotherapy followed by peripheral and/or bone marrow stem cell transplant in patients with advanced sarcoma: Experience of a Canadian Centre. *Sarcoma* 2004; **8**(2-3):63–9.

Hoy 2007 {published data only}

Hoy SM, Lyseng-Williamson KA. Intravenous busulfan: in the conditioning treatment of pediatric patients prior to hematopoietic stem cell transplantation. *Pediatric Drugs* 2007;**9**(4):271–8.

Huang 2006 {published data only}

Huang HQ, Cai QC, Shi YX, Lin XB, Wei J, Guo Y, et al. Preliminary assessment of immune reconstitution after autologous peripheral hematopoietic stem cell transplantation (AHSCT). *Ai Zheng* 2006;**25**(8):1023–8.

Huttmann 2005 {published data only}

Huttmann A, Schirsafi K, Seeber S, Bojko P. Comparison of lenograstim and filgrastim: effects on blood cell recovery after high-dose chemotherapy and autologous peripheral blood stem cell transplantation. *Journal of Cancer Research and Clinical Oncology* 2005;**131**(3):152–6.

Höffken 1997 {published data only}

Höffken K, Kath R, Fricke HJ, Blumenstengel K, Vogel W, Sayer HG. High-dose chemotherapy of solid tumors [Article in German: Hochdosischemotherapie bei soliden Tumoren]. *Medizinische Klinik* 1997;**92**(7):410–4.

Iankelevich 2000 {published data only}

Iankelevich MIa, Dolgopolov IS, Andreeva LIu, Ravshanova RS, Izhogin DG, Mkheidze DM, et al. Use of subgrafting doses of peripheral stem cells is a new approach to overcoming hematological toxicity of multiple intensive courses of chemotherapy in children [Article in Russian: Ispol'zovanie subtransplantatsionnykh doz perifericheskikh stvolovykh kletok – novyi podkhod k preodoleniiu

gematologicheskoi toksichnosti mnogokratnykh intensivnykh kursov khimioterapii u detei]. Vestnik Rossiiskoi Akademii Meditsinskikh Nauk 2000;**6**:21–4.

ICR 1994 {published data only}

ICR Institute of Cancer Research. Small-round-cell tumours of childhood. *Lancet* 1994;**344**(8924):725–9.

Imbach 1979 {published data only}

Imbach P, Odavic R, Bleher EA, Bucher U, Deubelbeiss KA, Wagner HP. Autologous bone marrow reimplantation in children with advanced tumor [Article in German: Autologe Knochenmark–Reimplantation bei Kindern mit fortgeschrittenem Tumor]. Schweizerische Medizinische Wochenschrift Journal Suisse de Medecine 1979;109(8):

Irle 1989 {published data only}

Irle C. Massive chemotherapy of solid tumors with bone marrow transplantation [Article in French: Chimiotherapie lourde avec greffe de moelle dans les tumouers solides]. *Medecine et Hygiene* 1989;47(1816):3377–82.

Issels 1995 {published data only}

Issels RD. Soft tissue sarcomas: what is currently being done. *European Journal of Surgical Oncology* 1995;**21**(5): 471–4.

Issels 2002 {published data only}

Issels RD, Schlemmer M. Current trials and new aspects in soft tissue sarcoma of adults. *Cancer Chemotherapy and Pharmacology* 2002;**49 Suppl** 1:4–8.

Issels 2004 {published data only}

Issels R. Knochentumoren und Weichteilsarkome: Empfehlungen zur Diagnostik, Therapie und Nachsorge. München: Zuckschwerdt, 2004.

Jamil 2004 {published data only}

Jamil A, Bayoumy M, Termuhlen AM, Wrona S. Pediatric Autologous Stem Cell Transplantation: A Comparison between Peripheral Blood Stem Cell and Bone Marrow. *International Pediatrics* 2004;**19**(1):28–33.

Jelic 1994 {published data only}

Jelic S, Vuletic L, Kovcin V, Kreacic M, Milanovic N, Radosavlevic D, et al. High-dose epirubicin vs. high-dose epirubicin-cisplatin chemotherapy for advanced soft tissue sarcoma: an interim report. *Annals of Oncology* 1994;**5**Suppl 8:176: abstract no. p887.

Jelic 1997 {published data only}

Jelic S, Kovcin V, Milanovic N, Babovic N, Kreacic M, Ristovic Z, et al.Randomised study of high-dose epirubicin versus high-dose epirubicin-cisplatin chemotherapy for advanced soft tissue sarcoma. *European Journal of Cancer* 1997;**33**(2):220–5.

Kaatsch 2009 {published data only}

Kaatsch P, Debling D, Blettner M, Spix C. Second malignant neoplasms after childhood cancer in Germany - Results from the long-term follow-up of the German Childhood Cancer Registry. *Strahlentherapie und Onkologie* 2009;**185 Suppl 2**:8–10.

Kabickova 2003 {published data only}

Kabickova E. High-dose chemotherapy with autologous hematopoietic stem cell transplantation for pediatric solid tumors [Article in Czech]. *Klinicka Onkologie* 2003;**16** Suppl 1:119–21.

Kadan-Lottick 2008 {published data only}

Kadan-Lottick NS, Dinu I, Wasilewski-Masker K, Kaste S, Meacham LR, Mahajan A, et al.Osteonecrosis in adult survivors of childhood cancer: a report from the childhood cancer survivor study. *Journal of Clinical Oncology* 2008;**26** (18):3038–45.

Kaizer 1979 {published data only}

Kaizer H, Wharam MD, Munoz LL, Johnson RJ, Elfenbein GJ, Tutschka PJ, et al. Autologous bone marrow transplantation in the treatment of selected human malignancies: The Johns Hopkins Oncology Center Program. Experimental Hematology 1979;7 Suppl 5: 309–20.

Kaizer 1980 {published data only}

Kaizer H, Wharam MD, Johnson RJ, Economou JG, Shin HS, Santos GW, et al.Requirements for the successful application of autologous bone marrow transplantation in the treatment of selected malignancies. *Haematology and Blood Transfusion* 1980;**25**:285–96.

Kaizer 1984 {published data only}

Kaizer H, Chow HS. Autologous bone marrow transplantation (ABMT) in the treatment of cancer. *Cancer Investigation* 1984;**2**(3):203–13.

Kalwak 2002 {published data only}

Kalwak K, Gorczynska E, Toporski J, Turkiewicz D, Slociak M, Ussowicz M, et al.Immune reconstitution after haematopoietic cell transplantation in children: Immunophenotype analysis with regard to factors affecting the speed of recovery. *British Journal of Haematology* 2002; **118**(1):74–89.

Kanabar 1995 {published data only}

Kanabar DJ, Attard-Montalto S, Saha V, Kingston JE, Malpas JE, Eden OB. Quality of life in survivors of childhood cancer after megatherapy with autologous bone marrow rescue. *Pediatric Hematology and Oncology* 1995;**12** (1):29–36.

Kasper 2004 {published data only}

Kasper B, Lehnert T, Bernd L, Mechtersheimer G, Goldschmidt H, Ho AD, et al. High-dose chemotherapy with autologous peripheral blood stem cell transplantation for bone and soft-tissue sarcomas. *Bone Marrow Transplantation* 2004;**34**(1):37–41.

Kasper 2005 {published data only}

Kasper B, Ho AD, Egerer G. Is there an indication for high-dose chemotherapy in the treatment of bone and soft-tissue sarcoma?. *Oncology* 2005;**68**(2-3):115–21.

Kasper 2006 {published data only}

Kasper B, Ho AD, Egerer G. Dose-intensive chemotherapy with stem cell support as a treatment strategy for bone and soft-tissue sarcomas. *Current Stem Cell Research and Therapy* 2006;**1**(1):29–35.

Katzenstein 2003 {published data only}

Katzenstein HM, Kletzel M, Reynolds M, Superina R, Gonzalez-Crussi F. Metastatic malignant rhabdoid tumor of the liver treated with tandem high-dose therapy and autologous peripheral blood stem cell rescue. *Medical and Pediatric Oncology* 2003;**40**(3):199–201.

Kavan 1997 {published data only}

Kavan P, Koutecky J. Current results with myeloablative therapy followed by hematopoietic stem cell rescue in pediatric solid tumors [Article in Czech: Soucasny pohled na myeloablativni lecbu s naslednou transplantaci hematopoetickych kmenovych bunek u detskych solidnich nadoru]. Klinicka Onkologie 1997;10(4):106–9.

Kavan 1997a {published data only}

Kavan P, Stankova J, Koutecky J, Gajdos P. High-dose chemotherapy with subsequent autologous stem cell transplantation in children and adolescents for high-risk rhabdomyosarcoma. A 3-year survival outcome [Article in Czech: Vysokodavkovana terapies naslednou autologni transplantaci hematopoetickych kmenovych bunek u deti a mladistvychs rabdomyosarkomem vysokeho rizika. Vysledky trileteho prezivani]. *Klincka Onkologie* 1997;10 (5):148–51.

Kinsella 1988 {published data only}

Kinsella TJ, Miser JS, Triche TJ, Horvath K, Glatstein E. Treatment of high-risk sarcomas in children and young adults: analysis of local control using intensive combined modality therapy. *NCI Monographs* 1988;**6**:291–6.

Kinsella 1988a {published data only}

Kinsella TJ, Sindelar WF, Lack E, Glatstein E, Rosenberg SA. Preliminary results of a randomized study of adjuvant radiation therapy in resectable adult retroperitoneal soft tissue sarcomas. *Journal of Clinical Oncology* 1988;**6**(1): 18–25.

Klaritsch 2006 {published data only}

Klaritsch P, Reich O, Regauer S, Bauernhofer T. Recurrent endometrial stromal sarcoma after treatment with high-dose chemotherapy and autologous stem-cell support: a case report. *European Journal of Gynaecological Oncology* 2006; **27**(3):297–8.

Kletzel 1997 {published data only}

Kletzel M, Kim AR. Autologous bone marrow transplantation in pediatric solid tumors. *Cancer Treatment and Research* 1997;77:333–56.

Kletzel 1998 {published data only}

Kletzel M, Longino R, Rademaker AW, nner-Koptik KE, Olszewski M, Morgan ER. Peripheral blood stem cell transplantation in young children: experience with harvesting, mobilization and engraftment. *Pediatric Transplantation* 1998;**2**(3):191–6.

Kletzel 2005 {published data only}

Kletzel M, Hewlett B. Pediatric transplantation: results in solid tumors. *Current Hematology Reports* 2005;4(4):260–9.

Klingebiel 1989 {published data only}

Klingebiel T, Dopfer R, Handgretinger R, Niethammer D. Indications for autologous bone marrow transplantation in

pediatric oncology. Results of the 5th meeting of experts of the Kind-Philipp Foundation, Riesensburg, November 1988 [Article in German: Indikation zur autologen Knochenmarktransplantation in der Padiatrischen Onkologie: Ergebnisse der 5. Expertentagung der Kind-Philipp-Stiftung, Reisenburg, November 1988]. Klinische Padiatrie 1989;201(4):304–10.

Klingebiel 1994 {published data only}

Klingebiel T, Handgretinger R, Niethammer D. Autologous bone marrow transplantation [Article in German: Autologe Knochenmarktransplantation]. *Infusionstherapie und Transfusionsmedizin* 1994;**21 Suppl** 3:42–5.

Klingebiel 2008 {published data only}

Klingebiel T, Boos J, Beske F, Hallmen E, Int-Veen C, Dantonello T, et al. Treatment of children with metastatic soft tissue sarcoma with oral maintenance compared to high dose chemotherapy: report of the HD CWS-96 trial. *Pediatric Blood and Cancer* 2008;**50**(4):739–45.

Kook 1998 {published data only}

Kook H, Kim KM, Choi SH, Choi BS, Kim HJ, Chung SY, et al.Life-threatening carboplatin hypersensitivity during conditioning for autologous PBSC transplantation: successful rechallenge after desensitization. *Bone Marrow Transplantation* 1998;**21**(7):727–9.

Korfel 2001 {published data only}

Korfel A, Fischer L, Foss HD, Koch HC, Thiel E. Testicular germ cell tumor with rhabdomyosarcoma successfully treated by disease-adapted chemotherapy including high-dose chemotherapy: case report and review of the literature. *Bone Marrow Transplantation* 2001;**28**(8):787–9.

Koscielniak 1992 {published data only}

Koscielniak E, Rodary C, Flamant F, Carli M, Treuner J, Pinkerton CR, et al. Metastatic rhabdomyosarcoma and histologically similar tumors in childhood: A retrospective European multi-center analysis. *Medical and Pediatric Oncology* 1992;**20**(3):209–14.

Koscielniak 1997 {published data only}

Koscielniak E, Klingebiel TH, Peters C, Hermann J, Burdach ST, Bender-Götze C, et al.Do patients with metastatic and recurrent rhabdomyosarcoma benefit from high-dose therapy with hematopoietic rescue? Report of the German/Austrian Pediatric Bone Marrow Transplantation Group. *Bone Marrow Transplantation* 1997;19(3):227–31.

Koscielniak 1999 {published data only}

Koscielniak E. The role of high dose therapy (HDC) with stem cell rescue in the treatment of high-risk rhabdomyosarcoma. *Rivista Italiana di Pediatria* 1999;**25** Suppl(3):106–8.

Koscielniak 2001 {published data only}

Koscielniak E. Soft tissue sarcoma in children. Diagnosis and therapeutic modalities. *Advances in Clinical and Experimental Medicine* 2001;**10**(1):3–8.

Koscielniak 2002 {published data only}

Koscielniak E, Morgan M, Treuner J. Soft tissue sarcoma in children: prognosis and management. *Paediatric Drugs* 2002;4(1):21–8.

Koscielniak 2005 {published data only}

Koscielniak E. Therapy for soft tissue sacrcoma: more questions than answers? [Article in German: Therapie der Weichteilsarkome: mehr Fragen offen als beantwortet?]. Wiener Klinische Wochenschrifte 2005;117(5-6):176–9.

Kuroiwa 2009 {published data only}

Kuroiwa M, Sakamoto J, Shimada A, Suzuki N, Hirato J, Park MJ, et al.Manifestation of alveolar rhabdomyosarcoma as primary cutaneous lesions in a neonate with Beckwith-Wiedemann syndrome. *Journal of Pediatric Surgery* 2009;44 (3):e31–5.

Kushner 2000 {published data only}

Kushner BH, Kramer K, Meyers PA, Wollner N, Cheung NK. Pilot study of topotecan and high-dose cyclophosphamide for resistant pediatric solid tumors. *Medical and Pediatric Oncology* 2000;**35**:468–74.

Kwan 1996 {published data only}

Kwan WH, Choi PH, Li CK, Shing MK, Chik KW, Yuen P, et al.Breast metastasis in adolescents with alveolar rhabdomyosarcoma of the extremities: report of two cases. *Pediatric Hematology and Oncology* 1996;**13**(3):277–85.

Kwon 2010 {published data only}

Kwon SY, Won SC, Han JW, Shin YJ, Lyu CJ. Feasibility of sequential high-dose chemotherapy in advanced pediatric solid tumors. *Pediatric Hematology and Oncology* 2010;**27** (1):1–12.

Ladenstein 1993 {published data only}

Ladenstein R, Hartmann O, Pinkerton CR. The role of megatherapy with autologous bone marrow rescue in solid tumours of childhood. *Annals of Oncology* 1993;**4**(Suppl 1): S45–58.

Ladenstein 1997 {published data only}

Ladenstein R, Philip T, Gardner H. Autologous stem cell transplantation for solid tumors in children. *Current Opinion in Pediatrics* 1997;**9**(1):55–69.

Lal 2005 {published data only}

Lal DR, Su WT, Wolden SL, Loh KC, Modak S, La Quaglia MP. Results of multimodal treatment for desmoplastic small round cell tumors. *Journal of Pediatric Surgery* 2005;**40**(1): 251–5.

Lang 2006 {published data only}

Lang P, Pfeiffer M, Müller I, Schumm M, Ebinger M, Koscielniak E, et al. Haploidentical stem cell transplantation in patients with pediatric solid tumors: preliminary results of a pilot study and analysis of graft versus tumor effects. *Klinische Pädiatrie* 2006;**218**(6):321–6.

Lange 2004 {published data only}

Lange T, Niederwieser D. Stem cell transplantation with reduced conditioning [Stammzelltransplantation mit reduzierter Konditionierung]. *Onkologisch* 2004;**3**:6–7.

Larsen 2000 {published data only}

Larsen E, Bramson RT, De Leval L, Ebb DH. Case Records of the Massachusetts General Hospital. Case 33-2000. Presentation of case. *New England Journal of Medicine* 2000; **343**(17):1249–57.

Le Cesne 2000 {published data only}

Le Cesne A, Judson I, Crowther D, Rodenhuis S, Keizer HJ, Van Hoesel Q, et al.Randomized phase III study comparing conventional-dose doxorubicin plus ifosfamide versus high-dose doxorubicin plus ifosfamide plus recombinant human granulocyte-macrophage colony-stimulating factor in advanced soft tissue sarcomas: a trial of the European Organization for Research and Treatment of Cancer/Soft Tissue and Bone Sarcoma Group. *Journal of Clinical Oncology* 2000;18(14):2676–84.

Le Corroller 1997 {published data only}

Le Corroller AG, Faucher C, Auperin A, Blaise D, Fortanier C, Benhamou E, et al. Autologous peripheral blood progenitor-cell transplantation versus autologous bone marrow transplantation for adults and children with non-leukaemic malignant disease. *Pharmacoeconomics* 1997;11 (5):454–63.

Lehrnbecher 2006 {published data only}

Lehrnbecher T, Becker M, Schwabe D, Kohl U, Kriener S, Hunfeld K-P, et al. Primary intestinal aspergillosis after high-dose chemotherapy and autologous stem cell rescue. *Pediatric Infectious Disease Journal* 2006;**25**(5):465–6.

Lessnick 2009 {published data only}

Lessnick SL, Dei Tos AP, Sorensen PHB, Dileo P, Baker LH, Ferrari S, et al.Small round cell sarcomas. *Seminars in Oncology* 2009;**36**(4):338–46.

Liseth 2004 {published data only}

Liseth K, Abrahamsen JF, Ekanger R, Nesthus I, Sjo MS. Survival after high-dose therapy with autologous stem cell support [Overlevelse etter hoydosebehandling med autolog stamcellestotte]. *Tidsskrift for den Norske Laegeforening* 2004;**124**(10):1374–5.

Locatelli 2008 {published data only}

Locatelli F, Giorgiani G, Di-Cesare-Merlone A, Merli P, Sparta V, Moretta F. The changing role of stem cell transplantation in childhood. *Bone Marrow Transplantation* 2008;**41 Suppl 2**:3–7.

Lorenz 1999 {published data only}

Lorenz F, Skotnicki AB. Autotransplantation for solid tumors [Article in Polish: Autotransplantacja w gutzach litych]. *Przeglad Lekarski* 1999;**56 Suppl 1**:101–7.

Lorigan 2007 {published data only}

Lorigan P, Verweij J, Papai Z, Rodenhuis S, Le Cesne A, Leahy MG, et al. Phase III trial of two investigational schedules of ifosfamide compared with standard-dose doxorubicin in advanced or metastatic soft tissue sarcoma: a European Organisation for Research and Treatment of Cancer Soft Tissue and Bone Sarcoma Group Study. *Journal of Clinical Oncology* 2007;25(21):3144–50.

Lucidarme 1998 {published data only}

Lucidarme N, Valteau-Couanet D, Oberlin O, Couanet D, Kalifa C, Beaujean F, et al. Phase II study of high-dose thiotepa and hematopoietic stem cell transplantation in children with solid tumors. *Bone Marrow Transplantation* 1998;**22**(6):535–40.

Mace 2003 {published data only}

Mace JR, Keohan ML, Bernardy H, Junge K, Niebch G, Romeis P, et al. Crossover randomized comparison of intravenous versus intravenous/oral mesna in soft tissue sarcoma treated with high-dose ifosfamide. *Clinical Cancer Research* 2003;**9**(16 Pt 1):5829–34.

Machado 2007 {published data only}

Machado M, Moreb JS, Khan SA. Six cases of permanent alopecia after various conditioning regimens commonly used in hematopoietic stem cell transplantation. *Bone Marrow Transplantation* 2007;**40**(10):979–82.

Mack 1995 {published data only}

Mack TM. Sarcomas and other malignancies of soft tissue, retroperitoneum, peritoneum, pleura, heart, mediastinum, and spleen. *Cancer* 1995;75 **Suppl**(1):211–44.

Mackall 2001 {published data only}

Mackall CL, Helman LJ. High-dose chemotherapy for rhabdomyosarcoma: where do we go from here. *Journal of Pediatric Hematology/Oncology* 2001;**23**(5):266–7.

Madero 1995 {published data only}

Madero L, Muonz A, az de Heredia A, Martinez A, Badell I, Esquembre C, et al.G-CSF after autologous bone marrow transplantation for malignant diseases in children. *Bone Marrow Transplantation* 1995;**15**(3):349–51.

Maeda 2008 {published data only}

Maeda M. Late effects of childhood cancer: life-threatening issues. *Journal of Nippon Medical School* 2008;**75**(6):320–4.

Mankin 2004 {published data only}

Mankin HJ, Hornicek FJ, Temple HT, Gebhardt MC. Malignant tumors of the pelvis: an outcome study. *Clinical Orthopaedics and Related Research* 2004;**425**:212–7.

Marina 1997 {published data only}

Marina NM. Biology and treatment of pediatric malignant solid tumors. *Cancer Chemotherapy and Biological Response Modifiers* 1997;**17**:642–71.

Matsubara 2003 {published data only}

Matsubara H, Makimoto A, Higa T, Kawamoto H, Takayama J, Ohira M, et al. Possible benefits of high-dose chemotherapy as intensive consolidation in patients with high-risk rhabdomyosarcoma who achieve complete remission with conventional chemotherapy. *Pediatric Hematology and Oncology* 2003;**20**(3):201–10.

Matsuyama 2000 {published data only}

Matsuyama T. Autologous bone marrow transplantation for pediatric malignancies. *Biotherapy* 2000;**14**(3):207–42.

Matthews 2007 {published data only}

Matthews RH, Emami M, Connaghan DG, Holland HK, Morris LE. Home administration of high-dose oral busulfan in patients undergoing hematopoietic stem cell transplantation. *Bone Marrow Transplantation* 2007;**39**(7): 397–400.

Medioni 2003 {published data only}

Medioni J, Fernandez-Bruno P, Meignin V, Bourrier P, Laurence V, Ramdani M, et al. Testicular metastasis of alveolar rhabdomyosarcoma: clinical case of a 19-year-

old man [Article in French: Metastase testiculaire d'un rhabdomyosarcome alveolaire: cas clinique d'un homme age de 19 ans]. *Progres en Urologie* 2003;**13**(4):700–2.

Mesia 1995 {published data only}

Mesia R, Sola C, Mendoza L, Tabernero JM, Brunet J, Maroto P, et al. High dose chemotherapy in autologous bone marrow transplantation in lymphomas and solid metastatic tumors [Article in Spanish: Quimioterapia a altas dosis con trasplante autologa de medula osea en linfomas y tumores solidos metastasicos]. *Neoplasia* 1995;12(2):43–50.

Meyers 2004 {published data only}

Meyers Paul A. High-dose therapy with autologous stem cell rescue for pediatric sarcomas. *Current Opinion in Oncology* 2004;**16**(2):120–5.

Michailov 2001 {published data only}

Michailov G, Avramova B. Autologous transplantation of hemopoietic cells by solid tumors: clinical experience of transplantation unit, hospital for treatment of children's oncohaematological diseases, Sofia 1997-2001. *Pediatriya* 2001;41(3):30–4.

Michon 1999 {published data only}

Michon J, Schleiermacher G. Autologous haematopoietic stem cell transplantation for paediatric solid tumours. *Bailliere's best practice and research. Clinical Haematology* 1999;**12**(1-2):247–59.

Mikhailova 1998 {published data only}

Mikhailova NB, Darskaia EI, Morozova EV, Shavva SA, Soldatenkov VE, Katyshev AV, et al. The transplantation of hemopoietic cells in patients with solid tumors [Article in Russian: Transplantatsiia gemopoeticheskikh kletok u bol'nykh s solidnymi opukholiami]. *Terapevticheskii Arkhiv* 1998;**70**(7):63–7.

Miliauskas 1993 {published data only}

Miliauskas JR, Abbott RL, Sarre R. Intra-abdominal desmoplastic small round cell tumour. *Australian and New Zealand Journal of Surgery* 1993;**63**(2):157–9.

Mimeault 2008 {published data only}

Mimeault M, Batra SK. Targeting of cancer stem/progenitor cells plus stem cell-based therapies: the ultimate hope for treating and curing aggressive and recurrent cancers. *Panminerva Medica* 2008;**50**:3–18.

Minard-Colin 2004 {published data only}

Minard-Colin V, Kalifa C, Guinebretiere JM, Brugieres L, Dubousset J, Habrand JL, et al.Outcome of flat bone sarcomas (other than Ewing's) in children and adolescents: a study of 25 cases. *British Journal of Cancer* 2004;**90**(3): 613–9.

Mingo 2005 {published data only}

Mingo L, Seguel F, Rollan V. Intraabdominal desmoplastic small round cell tumour. *Pediatric Surgery International* 2005;**21**(4):279–81.

Miyagi 2003 {published data only}

Miyagi T, Nagasaki A, Nakazato T, Hyakuna N, Takasu N, Masuda M. Emergence of secondary acute lymphoblastic leukemia of Burkitt type after treatment of rhabdomyosarcoma. *Medical and Pediatric Oncology* 2003; **41**(5):471–2.

Moore 2009 {published data only}

Moore AS, Shaw PJ, Hallahan AR, Carter TL, Kilo T, Nivison-Smith I, et al. Haemopoietic stem cell transplantation for children in Australia and New Zealand, 1998-2006: a report on behalf of the Australasian Bone Marrow Transplant Recipient Registry and the Australian and New Zealand Children's Haematology Oncology Group. *Medical Journal of Australia* 2009;190(3):121–5.

Morikawa 2005 {published data only}

Morikawa Y. Childhood rhabdomyosarcoma [Article in Japanese]. Nippon Geka Gakkai Zasshi Journal of Japan Surgical Society 2005;106(7):431–6.

Munoz 1983 {published data only}

Munoz LL, Wharam M, Kaiser H, Leventhal BG, Ruymann F. Magna-field irradiation and autologous marrow rescue in the tratment of pediatric solid tumors. *International Journal of Radiation Oncology, Biology, Physics* 1983;**9**(12):1951–4.

Müller 2002 {published data only}

Müller HL, Marx A, Trusen M, Schneider P, Kühl J. Disseminated malignant ectomesenchymoma (MEM): case report and review of the literature. *Pediatric Hematology and Oncology* 2002;**19**(1):9–17.

Nachbaur 1994 {published data only}

Nachbaur D, Schwaighofer H, Thaler J, Weyrer W, Fink M, Nussbaumer W, et al.Results of bone marrow transplantation for hematologic malignancies and solid tumours in Innsbruck [Article in German: Innsbrucker Ergebnisse mit der Knochenmarktransplantation in der Behandlung hämatologischer Neoplasien und solider Tumoren]. Wiener Klinische Wochenschrift 1994;106(7): 201–7.

Nag 1995 {published data only}

Nag S, Olson T, Ruymann F, Teich S, Pieters R. High-doserate brachytherapy in childhood sarcomas: a local control strategy preserving bone growth and function. *Medical and Pediatric Oncology* 1995;**25**:463–9.

Nath 2005 {published data only}

Nath SV, Prince HM, Choong PFM, Toner GC. Durable remissions are rare following high dose therapy with autologous stem cell transplantation for adults with 'paediatric' bone and soft tissue sarcomas. *International Seminars in Surgical Oncology* 2005;**2**(1):12.

Nenadov 1995 {published data only}

Nenadov Beck M, Meresse V, Hartmann O, Gaultier C. Long-term pulmonary sequelae after autologous bone marrow transplantation in children without total body irradiation. *Bone Marrow Transplantation* 1995;**16**(6): 771–5.

Nieboer 2005 {published data only}

Nieboer P, De Vries EGE, Mulder NH, Van der Graaf WTA. Relevance of high-dose chemotherapy in solid tumours. *Cancer Treatment Reviews* 2005;**31**(3):210–25.

Nieto 1999 {published data only}

Nieto Y, Shpall EJ. Autologous stem-cell transplantation for solid tumors in adults. *Hematology/Oncology Clinics of North America* 1999;**13**(5):939–68.

Nieto 2004 {published data only}

Nieto Y, Jones RB, Shpall EJ. Stem-cell transplantation for the treatment of advanced solid tumors. *Springer Seminars in Immunopathology* 2004;**26**(1-2):31–56.

Nieto 2007 {published data only}

Nieto Y, Aldaz A, Rifon J, Perez-Calvo J, Zafra A, Zufia L, et al. Phase I and Pharmacokinetic Study of Gemcitabine Administered at Fixed-Dose Rate, Combined with Docetaxel/Melphalan/Carboplatin, with Autologous Hematopoietic Progenitor-Cell Support, in Patients with Advanced Refractory Tumors. *Biology of Blood and Marrow Transplantation* 2007;13(11):1324–37.

Nivison-Smith 2005 {published data only}

Nivison-Smith I, Bradstock KF, Dodds AJ, Hawkins PA, Szer J. Haemopoietic stem cell transplantation in Australia and New Zealand, 1992-2001: Progress report from the Australasian Bone Marrow Transplant Recipient Registry. *Internal Medicine Journal* 2005;**35**(1):18–27.

Nivison-Smith 2007 {published data only}

Nivison-Smith I, Bradstock KF, Dodds AJ, Hawkins PA, Ma DDF, Moore JJ, et al.Hematopoietic stem cell transplantation in Australia and New Zealand, 1992-2004. Biology of Blood and Marrow Transplantation 2007;13(8): 905–12.

Nobile 1984 {published data only}

Nobile MT, Rosso R, Brema F, Cinquegrana A, Santi L. Phase II study of ifosfamide combined with Mesna uroprotection in advanced non-small-cell lung carcinoma and other solid tumors. *Tumori* 1984;**70**(5):433–7.

Notteghem 2003 {published data only}

Notteghem P, Soler C, Dellatolas G, Kieffer-Renaux V, Valteau-Couanet D, Raimondo G, et al. Neuropsychological outcome in long-term survivors of a childhood extracranial solid tumor who have undergone autologous bone marrow transplantation. *Bone Marrow Transplantation* 2003;**31**(7): 599–606.

Oeffinger 2008 {published data only}

Oeffinger KC, Nathan PC, Kremer LCM. Challenges after curative treatment for childhood cancer and long-term follow up of survivors. *Pediatric Clinics of North America* 2008;**55**(1):251–73.

Ohira 1983 {published data only}

Ohira M, Shibata T, Ise T. Autologous bone marrow transplantation in solid tumors in children [Article in Japanese]. *Gan to Kagaku Ryoho* 1983;**10**(6):1428–37.

Ohira 1990 {published data only}

Ohira M. Autologous bone marrow transplantation in pediatric cancer [Article in Japanese]. *Gan to Kagaku Ryoho* 1990;**17**(12):2299–306.

Ohta 2001 {published data only}

Ohta S, Suzuki A, Yagi K, Taga T, Higashino K, Narita T, et al. Three-year-old boy with primary rhabdomyosarcoma of

the diaphragm presenting with a hemothorax. *International Journal of Pediatric Hematology/Oncology* 2001;7(5-6): 305–9.

Oppenheim 2002 {published data only}

Oppenheim D, Valteau-Couanet D, Vasselon S, Hartmann O. How do parents perceive high-dose chemotherapy and autologous stem cell transplantation for their children. *Bone Marrow Transplantation* 2002;**30**(1):35–9.

Ortega 1991 {published data only}

Ortega JA, Wharam M, Gehan EA, Ragab AH, Crist W, Webber B, et al. Clinical features and results of therapy for children with paraspinal soft tissue sarcoma: a report of the Intergroup Rhabdomyosarcoma Study. *Journal of Clinical Oncology* 1991;**9**(5):796–801.

Osugi 2000 {published data only}

Osugi Y, Tokimasa S, Fujisaki H, Takai K, Nakanishi K, Matsuda Y, et al.Intra-arterial high-dose chemotherapy consisting of melphalan and thio-TEPA with stem cell rescue for advanced maxillary tumors. *International Journal of Pediatric Hematology/Oncology* 2000;7(2):109–16.

Oue 2003 {published data only}

Oue T, Kubota A, Okuyama H, Kawahara H, Inoue M, Yagi K, et al. Megatherapy with hematopoietic stem cell rescue as a preoperative treatment in unresectable pediatric malignancies. *Journal of Pediatric Surgery* 2003;**38**(1): 130–3.

Ozkaynak 1990 {published data only}

Ozkaynak MF, Nolta J, Parkman R. In vitro purging of human rhabdomyosarcoma cells using 4-hydroperoxycyclophosphamide. *Cancer Research* 1990;**50** (5):1455–8.

Ozkaynak 1998 {published data only}

Ozkaynak MF, Matthay K, Cairo M, Harris RE, Feig S, Reynolds CP, et al.Double-alkylator non-total-body irradiation regimen with autologous hematopoietic stemcell transplantation in pediatric solid tumors. *Journal of Clinical Oncology* 1998;**16**(3):937–44.

Ozkaynak 2008 {published data only}

Ozkaynak MF, Sahdev I, Gross TG, Levine JE, Cheerva AC, Richards MK, et al.A pilot study of addition of amifostine to melphalan, carboplatin, etoposide, and cyclophosphamide with autologous hematopoietic stem cell transplantation in pediatric solid tumors-A pediatric blood and marrow transplant consortium study. *Journal of Pediatric Hematology/Oncology* 2008;**30**(3):204–9.

Pasetto 2003 {published data only}

Pasetto LM, Basso U, Brandes AA. Improved tolerability of chemotherapy in soft tissue sarcomas: old and new strategies. *Expert Reviews of Anticancer Therapy* 2003;**3**(2): 167–78.

Patel 1992 {published data only}

Patel S, Benjamin RS. Standard and high dose chemotherapy for advanced soft tissue sarcomas. *Annals of Oncology* 1992; **3 Suppl 2**:81–3.

Patel 1994 {published data only}

Patel SR, Benjamin RS. The role of chemotherapy in soft tissue sarcomas. *Cancer Control* 1994;**1**(6):599–605.

Patzer 1999 {published data only}

Patzer L, Ringelmann F, Kentouche K, Zintl F, Misselwitz J. Unusual course of renal Fanconi syndrome after ifosfamide therapy [Ungewöhnlicher Verlauf eines de Toni–Debre–Fanconi–Syndroms nach Ifosfamidtherapie]. *Pädiatrie und Grenzgebiete* 1999; **38**(1-2):173–83.

Paulides 2006 {published data only}

Paulides M, Stohr W, Langer T, Kremer LCM, van Dalen EC. Cyclophosphamide versus ifosfamide for paediatric and young adult sarcoma patients. *Cochrane Database of Systematic Reviews* 2006, Issue 4. [DOI: 10.1002/14651858.CD006300.pub2.]

Pedrazzoli 2006 {published data only}

Pedrazzoli P, Ledermann JA, Lotz JP, Leyvraz S, Aglietta M, Rosti G, et al. High dose chemotherapy with autologous hematopoietic stem cell support for solid tumors other than breast cancer in adults. *Annals of Oncology* 2006;17(10): 1479–88.

Perentesis 1999 {published data only}

Perentesis J, Katsanis E, DeFor T, Neglia J, Ramsay N. Autologous stem cell transplantation for high-risk pediatric solid tumors. *Bone Marrow Transplantation* 1999;**24**(6): 609–15.

Pession 1999 {published data only}

Pession A, Prete A, Locatelli F, Bella S, Melchionda F, Garaventa A, et al. Phase I study of high-dose thiotepa with busulfan, etoposide, and autologous stem cell support in children with disseminated solid tumors. *Medical and Pediatric Oncology* 1999;**33**(5):450–4.

Philip 1984 {published data only}

Philip T, Bouffet E, Biron P. High-dose chemotherapy and bone marrow autograft in solid tumors and non-leukemic lymphomas in pediatrics [Article in French: Chimiotherapie massive et autogreffe de moelle dans les tumeurs solides et les lymphomes non leucemiques en pediatrie]. *Annales de Pediatrie* 1984;31(9):745–52.

Pick 1988 {published data only}

Pick TE. Autologous bone marrow transplantation in children. *Critical Reviews in Oncology/Hematology* 1988;**8** (4):311–37.

Pico 1993 {published data only}

Pico JL, Ibrahim A, Castagna L, Bourhis JH, Chazard M, Maraninchi D, et al. Escalating high-dose carboplatin and autologous bone marrow transplantation in solid tumors. *Oncology* 1993;**50 Suppl 2**:47–52.

Pinedo 1987 {published data only}

Pinedo HM, Mouridsen HT, Somers R, Santor A, Mulder YH, Blanwell V, et al.A randomized trial comparing the effect of epirubicin and doxorubicin in soft tissue sarcoma. *Clinical Trials Journal* 1987;24 Suppl 1:231–41.

Pinkerton 1986 {published data only}

Pinkerton R, Philip T, Bouffet E, Lashford L, Kemshead J. Autologous bone marrow transplantation in paediatric solid tumours. *Clinical Haematology* 1986;**15**(1):187–203.

Pinkerton 1987 {published data only}

Pinkerton R, Philip T. Autologous bone marrow transplantation in paediatric solid tumours. *Haematology and Blood Transfusion* 1987;**31**:92–6.

Pinkerton 1991 {published data only}

Pinkerton CR. Megatherapy for soft tissue sarcomas. *Bone Marrow Transplantation* 1991;**7 Suppl** 3:120–2.

Pinkerton 1991a {published data only}

Pinkerton CR, Groot-Loonen J, Barrett A, Meller ST, Tait D, Ashley S, et al. Rapid VAC high dose melphalan regimen, a novel chemotherapy approach in childhood soft tissue sarcomas. *British Journal of Cancer* 1991;**64**(2):381–5.

Pinkerton 1995 {published data only}

Pinkerton CR. Intensive chemotherapy with stem cell support-experience in pediatric solid tumours. *Bulletin du Cancer* 1995;**82 Suppl** 1:61–5.

Pohar-Marinsek 2001 {published data only}

Pohar-Marinsek Z, Anzic J, Jereb B. Evolving strategies in the treatment of childhood rhabdomyosarcoma: Slovenian experience. *Radiology and Oncology* 2001;**35**(4):259–66.

Pohar-Marinsek 2003 {published data only}

Pohar-Marinsek Z, Anzic J, Jereb B. Twenty-three years of experience in the management of childhood rhabdomyosarcoma in Slovenia. *Medical and Pediatric Oncology* 2003;**40**(2):118–9.

Raben 1994 {published data only}

Raben D, Williams J, Abrams RA. The clinical use of multimodality therapy in the management of cancer. *In Vivo* 1994;**8**(5):635–42.

Radeva 2005 {published data only}

Radeva JI, VanScoyoc E, Smith FO, Curtis LH, Breitfeld PP. National estimates of the use of hematopoietic stem-cell transplantation in children with cancer in the United States. *Bone Marrow Transplantation* 2005;**36**(5):397–404.

Raja 2003 {published data only}

Raja V, Lin JT, Xiao SY. Intra-abdominal desmoplastic small round cell tumor. *Journal of Clinical Oncology* 2003;**21**(5): 951–3.

Raney 1997 {published data only}

Raney RB, Asmar L, Newton WAJ, Bagwell C, Breneman JC, Crist W, et al. Ewing's sarcoma of soft tissues in childhood: a report from the Intergroup Rhabdomyosarcoma Study, 1972 to 1991. *Journal of Clinical Oncology* 1997;**15**(2):574–82.

Raney 2001 {published data only}

Raney RB, Anderson JR, Barr FG, Donaldson SS, Pappo AS, Qualman SJ, et al.Rhabdomyosarcoma and undifferentiated sarcoma in the first two decades of life: a selective review of intergroup rhabdomyosarcoma study group experience and rationale for Intergroup Rhabdomyosarcoma Study

V. Journal of Pediatric Hematology/Oncology 2001;**23**(4): 215–20.

Rapidis 2008 {published data only}

Rapidis AD. Sarcomas of the head and neck in adult patients: current concepts and future perspectives. *Expert Review of Anticancer Therapy* 2008;**8**(8):1271–97.

Ray-Coquard 2001 {published data only}

Ray-Coquard I, Biron P, Blay JY. High-dose chemotherapy in soft tissue sarcomas of adults. *Bulletin du Cancer* 2001; **88**(9):858–62.

Recchia 1996 {published data only}

Recchia F, Ginaldi L, Discepoli S, De Martinis M, Coloni F, Quaglino D. Autologous bone marrow transplantation for uterine sarcoma. Case report. *European Journal of Cancer* 1996;**32A**(3):553–4.

Recchia 2003 {published data only}

Recchia F, De Fillipis S, Piccinini M, Rea S. Highdose carboplatin, cyclophosphamide, etoposide with hematological growth factors, without stem cell support in patients with advanced cancer. *Anticancer Research* 2003;**23** (5 B):4141–7.

Reich 2001 {published data only}

Reich G, Mapara MY, Reichardt P, Dorken B, Maschmeyer G. Infectious complications after high-dose chemotherapy and autologous stem cell transplantation: Comparison between patients with lymphoma or multiple myeloma and patients with solid tumors. *Bone Marrow Transplantation* 2001;**27**(5):525–9.

Reichardt 1997 {published data only}

Reichardt P, Verweij J, Crowther D. Should high-dose chemotherapy be used in the treatment of soft tissue sarcoma?. *European Journal of Cancer* 1997;**33**(9):1351–60.

Reichardt 2002 {published data only}

Reichardt P. High-dose chemotherapy in adult soft tissue sarcoma. *Critical Reviews in Oncology/Hematology* 2002;**41** (2):157–67.

Rill 1994 {published data only}

Rill DR, Santana VM, Roberts WM, Nilson T, Bowman LC, Krance RA, et al.Direct demonstration that autologous bone marrow transplantation for solid tumors can return a multiplicity of tumorigenic cells. *Blood* 1994;**84**(2):380–3.

Ritchie 2004 {published data only}

Ritchie DS, Grigg AP, Roberts AW, Rosenthal MA, Fox RM, Szer J. Staged autologous peripheral blood progenitor cell transplantation for Ewing sarcoma and rhabdomyosarcoma. *Internal Medicine Journal* 2004;**34**(7):431–4.

Rivera-Luna 2001 {published data only}

Rivera-Luna R, Olaya-Vargas A, Meza-Coria C, Cardenas-Cardos R, Leal-Leal C, Amador-Zarco J. Is autologous bone marrow transplant (ABMT) and high-dose chemotherapy an approach that can rescue some children with advanced cancer disease?. *Pediatric Hematology and Oncology* 2001;**18** (7):443–51.

Rodenhuis 1999 {published data only}

Rodenhuis S, De Vries EG. High-dose chemotherapy with stem cell support for solid tumors in adults [Article in Dutch: Hooggedoseerde chemotherapie met stamcelondersteuning bij solide tumoren van volwassenen]. *Nederlands Tijdschrift voor Geneeskunde* 1999;**143**(14):731–8.

Roh 2001 {published data only}

Roh MS, Huh GY, Jeong JS, Lee GD, Hong SH. Left atrial myxosarcoma with systemic metastasis: a case report. *Journal of Korean Medical Science* 2001;**16**(1):111–4.

Roman-Unfer 1996 {published data only}

Roman-Unfer S, Bitran JD. High-dose chemotherapy for malignancies: a review. *Comprehensive Therapy* 1996;**22**: 107–20.

Rosenberg 1981 {published data only}

Rosenberg SA. Treatment of soft tissue and bone sarcomas: review of studies at the National Cancer Institute. *National Cancer Institute Monograph* 1981;**56**:241–4.

Rosenberg 1982 {published data only}

Rosenberg SA, Tepper J, Glatstein E, Costa J, Baker A, Brennan M, et al. The treatment of soft-tissue sarcomas of the extremities: prospective randomized evaluations of (1) limb-sparing surgery plus radiation therapy compared with amputation and (2) the role of adjuvant chemotherapy. *Annals of Surgery* 1982;**196**(3):305–15.

Rosenberg 1983 {published data only}

Rosenberg SA, Tepper J, Glatstein E, Costa J, Young R, Baker A, et al. Prospective randomized evaluation of adjuvant chemotherapy in adults with soft tissue sarcomas of the extremities. *Cancer* 1983;**52**(3):424–34.

Rosman 2008 {published data only}

Rosman IS, Lloyd BM, Hayashi RJ, Bayliss SJ. Cutaneous effects of thiotepa in pediatric patients receiving high-dose chemotherapy with autologous stem cell transplantation. Journal of the American Academy of Dermatology 2008;58(4): 575–8.

Rossbach 1999 {published data only}

Rossbach HC, Lacson A, Grana NH, Barbosa JL. Duchenne muscular dystrophy and concomitant metastatic alveolar rhabdomyosarcoma. *Journal of Pediatric Hematology and Oncology* 1999;**21**(6):528–30.

Rosti 2002 {published data only}

Rosti G, Ferrante P, Ledermann J, Leyvraz S, Ladenstein R, Koscielniak E, et al.High-dose chemotherapy for solid tumors: results of the EBMT. *Critical Reviews in Oncology/Hematology* 2002;**41**(2):129–40.

Rousselet 1994 {published data only}

Rousselet MC, Francois S, Croue A, Maigre M, Saint-Andre JP, Ifrah N. A lymph node interdigitating reticulum cell sarcoma. *Archives of Pathology and Laboratory Medicine* 1994;**118**(2):183–8.

Rubie 2003 {published data only}

Rubie H, Doz F, Vassal G, Chastagner P, Gentet JC, Urien S, et al.Individual dosing of carboplatin based on drug monitoring in children receiving high-dose chemotherapy. *European Journal of Cancer* 2003;**39**(10):1433–8.

Rzepecki 2006 {published data only}

Rzepecki P, Sarosiek T, Deptala A, Szczylik C. Autologous hematopoietic cell transplantation in adult patients with certain solid tumors. *Acta Haematologica Polonica* 2006;**37** (2):159–66.

Rzepecki 2006a {published data only}

Rzepecki P, Sarosiek T, Szczylik C. Autologous hematopoietic cell transplantation in adult patients with germ cell tumors and soft tissue sarcomas. *Wspolczesna Onkologia* 2006;**10**(1):7–12.

Saikawa 2006 {published data only}

Saikawa Y, Tone Y, Ikawa Y, Maeba H, Koizumi S, Minato H. Hemophagocytic alveolar rhabdomyosarcoma. *Journal of Clinical Oncology* 2006;**24**(36):5783–4.

Sajedi 2002 {published data only}

Sajedi M, Wolff JEA, Egeler RM, Pinto A, Hughes R, Anderson RA, et al. Congenital extrarenal non-central nervous system malignant rhabdoid tumor. *Journal of Pediatric Hematology/Oncology* 2002;**24**(4):316–20.

Sakayama 2008 {published data only}

Sakayama K, Tauchi H, Sugawara Y, Kidani T, Tokuda K, Miyazaki T, et al.A complete remission of sclerosing rhabdomyosarcoma with multiple lung and bone metastases treated with multi-agent chemotherapy and peripheral blood stem cell transplantation (PBSCT): a case report. *Anticancer Research* 2008;**28**(4C):2361–7.

Salutari 1998 {published data only}

Salutari P, Sica S, Laurenti L, Leone F, Chiusolo P, Piccirillo N, et al.Incidence of sepsis after peripheral blood progenitor cells transplantation: analysis of 86 consecutive hemato oncological patients. *Leukemia and Lymphoma* 1998;**30**(1-2):193–7.

Sanchez 1986 {published data only}

Sanchez de Toledo Codina J, Gallego Melcon S, Perez Payarols J, Prats Vinas J, Javier G, Massuet L, et al. Highdosage melphalan followed by bone marrow autotransplant in solid tumors of childhood: our experience [Article in Spanish: Melfalan a altas dosis seguido de autotrasplante de medula osea en tumores solidos de la infancia: nuestra experiencia]. *Medicina Clinica* 1986;86(15):621–3.

Sanchez-Garcia 2007 {published data only}

Sanchez-Garcia I, Vicente-Duenas C, Cobaleda C. The theoretical basis of cancer-stem-cell-based therapeutics of cancer: can it be put into practice?. *BioEssays* 2007;**29**(12): 1269–80.

Santana 1992 {published data only}

Santana VM, Schell MJ, Williams R, Bowman LC, Thompson EI, Brenner MK, et al. Escalating sequential high-dose carboplatin and etoposide with autologous marrow support in children with relapsed solid tumors. *Bone Marrow Transplantation* 1992;**10**(5):457–62.

Sanz 1997 {published data only}

Sanz N, de Mingo L, Florez F, Rollan V. Rhabdomyosarcoma of the biliary tree. *Pediatric Surgery International* 1997;**12**: 200–1.

Sato 1998 {published data only}

Sato A, Imaizumi M, Saisho T, Saito T, Yoshinari M, Cui Y, et al.Improved survival of children with advanced tumors by myeloablative chemotherapy and autologous peripheral blood stem cell transplantation in complete remission.

Tohoku Journal of Experimental Medicine 1998;186(4): 255–65

Sauer 1998 {published data only}

Sauer H. Adjuvant chemotherapy in early soft tissue sarcoma and palliative chemotherapy in advanced soft tissue sarcoma in adults [Article in German: Adjuvante Chemotherapie bei lokoregional begrenzten Weichteilsarkomen und palliative Chemotherapie bei fortgeschrittenen Weichteilsarkomen im Erwachsenenalter]. Schweizerische Rundschau fur Medizinische Praxis 1998;87(34):1066–71.

Sauer 1998a {published data only}

Sauer M, Gruhn B, Fuchs D, Altermann W, Zintl F. Heparin-associated thrombocytopenia (HAT II) in a patient undergoing high dose chemotherapy with consecutive stem cell rescue [Article in German: Heparin-induzierte Thrombozytopenie Typ II im Rahmen einer Hochdosis-Chemotherapie mit anschliessender Stammzellrescue]. Klinische Padiatrie 1998;210(3):102–5.

Savasan 2005 {published data only}

Savasan Sureyya, Abella Esteban M. Current issues in pediatric stem cell transplantation. *Clinics in Laboratory Medicine* 2005;**25**(3):519–40.

Savolainen 2005 {published data only}

Savolainen H, Lautenschlager I, Piiparinen H, Saarinen-Pihkala U, Hovi L, Vettenranta K. Human herpesvirus-6 and -7 in pediatric stem cell transplantation. *Pediatric Blood* and Cancer 2005;45(6):820–5.

Sawyer 1999 {published data only}

Sawyer M, Bramwell V. The treatment of distant metastases in soft tissue sarcoma. *Seminars in Radiation Oncology* 1999; **9**(4):389–400.

Schimmer 2002 {published data only}

Schimmer AD, Dranitsaris G, Ali V, Falconer M, Keating A. The autologous blood and marrow transplant long-term follow-up clinic: A model of care for following and treating survivors of autotransplant. *Supportive Care in Cancer* 2002; **10**(3):247–52.

Schlemmer 2006 {published data only}

Schlemmer M, Wendtner CM, Falk M, Abdel-Rahman S, Licht T, Baumert J, et al. Efficacy of consolidation high-dose chemotherapy with ifosfamide, carboplatin and etoposide (HD-ICE) followed by autologous peripheral blood stem cell rescue in chemosensitive patients with metastatic soft tissue sarcomas. *Oncology* 2006;71(1-2):32–9.

Schmidt 1994 {published data only}

Schmidt D, Köster E, Harms D. Intraabdominal desmoplastic small-cell tumor with divergent differentiation: clinicopathological findings and DNA ploidy. *Medical and Pediatric Oncology* 1994;**22**(2):97–102.

Schulz 1991 {published data only}

Schulz G, Frisch J, Greifenberg B, Nicolay U, Oster W. New therapeutic modalities for the clinical use of rhGM-CSF in patients with malignancies. *American Journal of Clinical Oncology* 1991;**14 Suppl** 1:19–26.

Schuster 2008 {published data only}

Schuster MW, Shore TB, Harpel JG, Greenberg J, Jalilizeinali B, Possley S, et al.Safety and tolerability of velafermin (CG53135-05) in patients receiving high-dose chemotherapy and autologous peripheral blood stem cell transplant. *Supportive Care in Cancer* 2008;**16**(5):477–83.

Schwella 1998 {published data only}

Schwella N, Rick O, Meyer O, Loffel J, Schleicher J, Serke S, et al. Mobilization of peripheral blood progenitor cells by disease-specific chemotherapy in patients with soft tissue sarcoma. *Bone Marrow Transplantation* 1998;**21**(9):863–8.

Secondino 2007 {published data only}

Secondino S, Carrabba M G, Pedrazzoli P, Castagna L, Spina F, Grosso F, et al. Reduced intensity stem cell transplantation for advanced soft tissue sarcomas in adults: a retrospective analysis of the European Group for Blood and Marrow Transplantation. *Haematologica* 2007;**92**(3):418–20.

Seeger 1991 {published data only}

Seeger RC, Reynolds CP. Treatment of high-risk solid tumors of childhood with intensive therapy and autologous bone marrow transplantation. *Pediatric Clinics of North America* 1991;**38**(2):393–424.

Segura 2001 {published data only}

Segura Huerta A, Lopez Tendero P, Yuste Izquierdo A, Girones Sarrio R, Perez Fidalgo J, Aparicio Urtasun J. Embryonal rhabdomyosarcoma of the prostate [Article in Spanish: Rabdomiosarcoma embrionario de prostata. Presentacion de un caso y revision de la bibliografia]. *Actas Urologica Espaniolas* 2001;25(2):144–9.

Seregard 2002 {published data only}

Seregard S. Management of alveolar rhabdomyosarcoma of the orbit. *Acta Ophthalmologica Scandinavica* 2002;**80**(6): 660–4.

Seynaeve 1999 {published data only}

Seynaeve C, Verweij J. High-dose chemotherapy in adult sarcomas: no standard yet. *Seminars in Oncology* 1999;**26** (1):119–33.

Shea 1995 {published data only}

Shea T, Graham M, Bernard S, Steagall A, Wiley J, Serody J, et al. A clinical and pharmacokinetic study of high-dose carboplatin, paclitaxel, granulocyte colony-stimulating factor, and peripheral blood stem cells in patients with unresectable or metastatic cancer. *Seminars in Oncology* 1995;**22 Suppl 12**(5):80–5.

Shen 1993 {published data only}

Shen BJ. Human umbilical cord blood transplantation in 4 cases with advanced solid tumors [Article in Chinese]. *Chung Hua Chung Liu Tsa Chih* 1993;**15**(3):152–4.

Shen 1994 {published data only}

Shen BJ, Hou HS, Zhang HQ, Sui XW. Unrelated, HLA-mismatched multiple human umbilical cord blood transfusion in four cases with advanced solid tumors: initial studies. *Blood Cells* 1994;**20**(2-3):285–92.

Shimizu 2008 {published data only}

Shimizu S, Yasui C, Minauchi K, Tsuchiya K. Metastatic rhabdomyosarcoma of the skin. *Journal of the American Academy of Dermatology* 2008;**58 Suppl 1**(5):118–20.

Shinkoda 2009 {published data only}

Shinkoda Y, Nagatoshi Y, Fukano R, Nishiyama K, Okamura J. Rhabdomyosarcoma masquerading as acute leukemia. *Pediatric Blood and Cancer* 2009;**52**(2):286–7.

Simon 2007 {published data only}

Simon Arne, Besuden Mette, Vezmar Sandra, Hasan Carola, Lampe Dagmar, Kreutzberg Sigrid, et al.Itraconazole prophylaxis in pediatric cancer patients receiving conventional chemotherapy or autologous stem cell transplants. *Supportive Care in Cancer* 2007;15(2):213–20.

Skinner 1974 {published data only}

Skinner DG. Management of extensive, localized neoplasms of lower abdominal wall. *Urology* 1974;**3**(1):34–7.

Sola 1999 {published data only}

Sola C, Maroto P, Salazar R, Mesia R, Mendoza L, Brunet J, et al. Prognostic factors of peripheral blood stem cell mobilization with cyclophosphamide and Filgrastim (rmetHuG-CSF): the CD34+ cell dose positively affects the time to hematopoietic recovery and supportive requirements after high-dose chemotherapy. *Hematology* 1999;4(3): 195–209.

Somlo 1995 {published data only}

Somlo G, Doroshow JH, Lev Ran A, Ahn DC, Hwang L, Raschko JW, et al. Effect of low-dose prophylactic dopamine on high-dose cisplatin-induced electrolyte wasting, ototoxicity, and epidermal growth factor excretion: a randomized, placebo-controlled, double-blind trial. *Journal of Clinical Oncology* 1995;**13**(5):1231–7.

Spitzer 1980 {published data only}

Spitzer G, Dicke KA, Litam J. High-dose combination chemotherapy with autologous bone marrow transplantation in adult solid tumors. *Cancer* 1980;45(12):3075–85.

Spitzer 1984 {published data only}

Spitzer G, Dicke K, Zander AR, Jagannath S, Vellekoop L, Freireich EJ. High-dose chemotherapy with autologous bone marrow transplantation. *Cancer* 1984;**54**(6):1216–25.

Spitzer 1994 {published data only}

Spitzer G, Dunphy FR, Bowers CE, Adkins DR. High-dose therapy with stem cell support in solid tumors. *Medical Oncology* 1994;**11**(2):53–62.

Spitzer 1995 {published data only}

Spitzer TR, Cirenza E, McAfee S, Foelber R, Zarzin J, Cahill R, et al. Phase I-II trial of high-dose cyclophosphamide, carboplatin and autologous bone marrow or peripheral blood stem cell rescue. *Bone Marrow Transplantation* 1995; **15**(4):537–42.

Spruce 1983 {published data only}

Spruce WE. Bone marrow transplantation. *American Journal of Pediatric Hematology and Oncology* 1983;**5**(3):

Stea 1987 {published data only}

Stea B, Kinsella TJ, Triche TJ, Horvath K, Glatstein E, Miser JS. Treatment of pelvic sarcomas in adolescents and young adults with intensive combined modality therapy. *International Journal of Radiation Oncology, Biology, Physics* 1987;**13**(12):1797–805.

Steinbrenner 2005 {published data only}

Steinbrenner M, Hafer R, Gruhn B, Müller A, Fuchs D, Hermann J, et al.T-cell independent production of salivary secretory IgA after hematopoietic stem cell transplantation in children. *Oral Microbiology and Immunology* 2005;**20**(5): 282–8.

Stöhr 2006 {published data only}

Stöhr W, Paulides M, Brecht I, Kremers A, Treuner J, Langer T, et al. Comparison of epirubicin and doxorubicin cardiotoxicity in children and adolescents treated within the German Cooperative Soft Tissue Sarcoma Study (CWS). *Journal of Cancer Research and Clinical Oncology* 2006;132 (1):35–40.

Suita 2005 {published data only}

Suita S, Noguchi S, Takamatsu H, Mizote H, Nagasaki A, Inomata Y, et al. Clinical characteristics and the prognosis of rhabdomyosarcoma - A report from the Study Group for Pediatric Solid Malignant Tumors in the Kyushu Area, Japan. *European Journal of Pediatric Surgery* 2005;**15**(6): 409–13.

Sussman 2008 {published data only}

Sussman N. Maintaining quality of life for seriously III children and young adults. *Primary Psychiatry* 2008;**15**(7): 21–2.

Takata 1997 {published data only}

Takata M, Hatta N, Takehara K, Fujiwara H. Absence of human herpesvirus-8 DNA in angiosarcomas and other skin tumours in immunocompetent patients, and in graft-versus-host disease in the immunosuppressed recipients of bone marrow transplants. *British Journal of Dermatology* 1997;137(1):156–7.

Takaue 2002 {published data only}

Takaue Y. Mini-transplantation strategy for solid tumors. International Journal of Hematology 2002;76 Suppl 2:13–4.

Takenaka 2007 {published data only}

Takenaka M, Okamoto Y, Ikeda K, Hashimoto R, Ueda T, Kurokawa N, et al. Comparison of antiemetic efficacy of 5-HT3 receptor antagonists in orthopedics cancer patients receiving high-dose chemotherapy [Article in Japanese]. *Gan to Kagaku Ryoho* 2007;**34**(3):403–7.

Tang 2009 {published data only}

Tang JY, Pan C, Xu M, Xue HL, Chen J, Dong L, et al. Effect of protocol RS-99 for childhood rhabdomyosarcoma [Article in Chinese]. *Chung Hua I Hsueh Tsa Chih* 2009;**89** (2):121–3.

Thomson 1999 {published data only}

Thomson B, Hawkins D, Felgenhauer J, Radich J. RT-PCR evaluation of peripheral blood, bone marrow and peripheral blood stem cells in children and adolescents undergoing VACIME chemotherapy for Ewing's sarcoma and alveolar rhabdomyosarcoma. *Bone Marrow Transplantation* 1999;24 (5):527–33.

Toma 1992 {published data only}

Toma S, Palumbo R, Sogno G, Venturino A, Santi L. Doxorubicin (or epidoxorubicin) combined with ifosfamide in the treatment of adult advanced soft tissue sarcomas. *Annals of Oncology* 1992;**3 Suppl** 2:119–23.

Trigg 2002 {published data only}

Trigg ME. Milestones in the development of pediatric hematopoietic stem cell transplantation: 50 years of progress. *Pediatric Transplantion* 2002;**6**(6):465–74.

Unal 2006 {published data only}

Unal E, Yen C, Saiman L, George D, Della-Latta P, Van de Ven C, et al. A low incidence of nontuberculous mycobacterial infections in pediatric hematopoietic stem cell transplantation recipients. *Biology of Blood and Marrow Transplantation* 2006;**12**(11):1188–97.

Urban 1997 {published data only}

Urban C, Schwinger W, Benesch M, Lackner H, Kerbl R, Gilli R, et al.Feasibility of peripheral blood stem cell (PBSC) and peripheral blood mononuclear cell (PBMNC) separation in children with a body weight below 20 KG. *Medical and Pediatric Oncology* 1997;**29**(2):115–20.

Urbano-Ispizua 2002 {published data only}

Urbano-Ispizua A, Schmitz N, de WT, Frassoni F, Rosti G, Schrezenmeier H, et al. Allogeneic and autologous transplantation for haematological diseases, solid tumours and immune disorders: definitions and current practice in Europe. *Bone Marrow Transplantation* 2002;**29**(8):639–46.

Vadhan 1996 {published data only}

Vadhan-Raj S, Patel S, Broxmeyer HF, Bueso-Ramos C, Reddy SP, Papadopolous N, et al. Phase I-II investigation of recombinant human thrombopoietin (rhTPO) in patients with sarcoma receiving high dose chemotherapy (CT) with adriamycin (A) and ifosfamide (I). *Blood* 1996;**88 Suppl** (10, Pt 1):448a.

Vadhan-Raj {published data only}

Vadhan-Raj S. GM-CSF after high-dose chemotherapy yields improved bone marrow protection. *Hospital Formulary* 1991;**26**(8):628.

Valkova 2003 {published data only}

Valkova J, Sramkova L, Smelhaus V, Cerny M, Kodet R, Svabova V, et al. Neonatal neoplasms. *Ceskoslovenska Pediatrie* 2003;**58**(8):478–83.

Valteau-Couanet 2007 {published data only}

Valteau-Couanet D, Dufour C, Hartmann O. High-dose chemotherapy and autologous stem cell transplantation in treating paediatric malignancies [Article in French]. *Oncologie* 2007;**9**(12):827–31.

Valteau-Couanet 2007a {published data only}

Valteau-Couanet D, Minard V. Poor prognosis childhood cancers [Article in French: Les cancers de l'enfant de mauvais pronostic]. *Revue Praticien* 2007;57(10):1087–91.

Van Dalen 2009 {published data only}

Van Dalen EC, Raphael MF, Caron HN, Kremer LCM. Treatment including anthracyclines versus treatment not including anthracyclines for childhood cancer. *Cochrane Database of Systematic Reviews* 2009, Issue 1. [Art. No.: CD006647. DOI: 10.1002/14651858.CD006647.pub2]

Van den Berg 2006 {published data only}

Van den Berg H. Biology and therapy of solid tumors in childhood. *Update on Cancer Therapeutics* 2006;**1**(3): 367–83

Van den Berg 2007 {published data only}

Van den Berg H. Biology and treatment of malignant solid tumors in childhood. *Update on Cancer Therapeutics* 2007; **2**(4):177–91.

Van den Berg 2008 {published data only}

Van den Berg H, Van Rijn RR, Merks JH. Management of tumors of the chest wall in childhood: a review. *Journal of Pediatric Hematology and Oncology* 2008;**30**(3):214–21.

Van Glabbeke 1993 {published data only}

Van Glabbeke M, van Oosterom AT, Steward W, Verweij J, Mouridsen H. Selection of large and objectively measurable target lesions in EORTC phase II trials: impact on recruitment and response rate. *European Journal of Cancer* 1993;**29A**(14):1943–7.

Varterasian 1997 {published data only}

Varterasian M, Zalupski M, Karanes C. The heterogeneity of leukemia occurring after treatment for sarcoma. *American Journal of Clinical Oncology* 1997;**20**(6):585–6.

Vassal 2005 {published data only}

Vassal G. Has chemotherapy reached its limits in pediatric cancers?. *European Journal of Cancer* 2005;**41**(4):564–75.

Vaughan 2001 {published data only}

Vaughan WP. Applications of high-dose chemotherapy with bone marrow/stem cell support in solid tumors. *Cancer Control* 2001;**8 Suppl** 2(6):50–2.

Verma 2002 {published data only}

Verma S, Bramwell V. Dose-intensive chemotherapy in advanced adult soft tissue sarcoma. Expert Review of Anticancer Therapy 2002;2(2):201–15.

Verma 2008 {published data only}

Verma S, Younus J, Stys-Norman D, Haynes AE, Blackstein M. Dose-intensive chemotherapy with growth factor or autologous bone marrow/stem cell transplant support in first-line treatment of advanced or metastatic adult soft tissue sarcoma: a systematic review. *Cancer* 2008;**112**(6): 1197–205.

Verma 2008a {published data only}

Verma S, Younus J, Haynes AE, Stys-Norman D, Blackstein M. Dose-intensive chemotherapy with growth factor or autologous bone marrow or stem-cell transplant support in first-line treatment of advanced or metastatic adult

soft tissue sarcoma: a clinical practice guideline. *Current Oncology* 2008;**15**(2):31–5.

Wachowiak 2008 {published data only}

Wachowiak J, Labopin M, Miano M, Chybicka A, Stary J, Sterba J, et al.Haematopoietic stem cell transplantation in children in eastern European countries 1985-2004: development, recent activity and role of the EBMT/ESH Outreach Programme. *Bone Marrow Transplantation* 2008; **41 Suppl** 2:112–7.

Walterhouse 1999 {published data only}

Walterhouse DO, Hoover ML, Marymont MA, Kletzel M. High-dose chemotherapy followed by peripheral blood stem cell rescue for metastatic rhabdomyosarcoma: the experience at Chicago Children's Memorial Hospital. *Medical and Pediatric Oncology* 1999;**32**(2):88–92.

Wasserman 1997 {published data only}

Wasserman E, Hidalgo M, Hornedo J, Cortes-Funes H. Octreotide (SMS 201-995) for hematopoietic support-dependent high-dose chemotherapy (HSD-HDC)-related diarrhoea: dose finding study and evaluation of efficacy. *Bone Marrow Transplantation* 1997;**20**(9):711–4.

Watanabe 2006a {published data only}

Watanabe H, Watanabe T, Suzuya H, Wakata Y, Kaneko M, Onishi T, et al. Peripheral blood stem cell mobilization by granulocyte colony-stimulating factor alone and engraftment kinetics following autologous transplantation in children and adolescents with solid tumor. *Bone Marrow Transplantation* 2006;37(7):661–8.

Weaver 1997 {published data only}

Weaver CH, Schwartzberg LS, Hainsworth J, Greco FA, Li W, Buckner CD, et al. Treatment-related mortality in 1000 consecutive patients receiving high-dose chemotherapy and peripheral blood progenitor cell transplantation in community cancer centers. *Bone Marrow Transplantation* 1997;**19**(7):671–8.

Weh 1995 {published data only}

Weh HJ, Hossfeld DK. Systemic therapy of disseminated soft tissue sarcomas. *Recent Results in Cancer Research* 1995; **138**:147–59.

Weh 1996 {published data only}

Weh HJ, De Wit M, Zornig C, Hossfeld DK. Treatment of adult metastatic soft-tissue sarcoma with doxorubicin/ifosfamide: better hematologic tolerance by G-CSF?. *Onkologie* 1996;**19**(2):159–62.

Weigel 2001 {published data only}

Weigel BJ, Breitfeld PP, Hawkins D, Crist WM, Baker KS. Role of high-dose chemotherapy with hematopoietic stem cell rescue in the treatment of metastatic or recurrent rhabdomyosarcoma. *Journal of Pediatric Hematology/ Oncology* 2001;**23**(5):272–6.

Werchniak 2005 {published data only}

Werchniak AE, Chaffee S, Dinulos JGH. Methotrexate-induced bullous acral erythema in a child. *Journal of the American Academy of Dermatology* 2005;**52 Suppl 1**(5): 93–5.

Wexler 1996 {published data only}

Wexler LH, Andrich MP, Venzon D, Berg SL, Weaver McClure L, Chen CC, et al.Randomized trial of the cardioprotective agent ICRF-187 in pediatric sarcoma patients treated with doxorubicin. *Journal of Clinical Oncology* 1996;**14**(2):362–72.

Willenbacher 1998 {published data only}

Willenbacher W, Mumm A, Ruther A, Weis J, Bartsch HH. Somatic risk factors for intermediate and long-term sequelae after hematological stem cell therapy predictive for feasibility of a rehabilitation program. *Onkologie* 1998;**21** (3):217–23.

Williams 2004 {published data only}

Williams BA, Williams KM, Doyle J, Stephens D, Greenberg M, Malkin D, et al. Metastatic rhabdomyosarcoma: a retrospective review of patients treated at the hospital for sick children between 1989 and 1999. *Journal of Pediatric Hematology and Oncology* 2004;**26**(4):243–7.

Womer 1996 {published data only}

Womer RB. Problems and controversies in the management of childhood sarcomas. *British Medical Bulletin* 1996;**52**(4): 826–43.

Womer 2000 {published data only}

Womer RB, Pressey JG. Rhabdomyosarcoma and soft tissue sarcoma in childhood. *Current Opinion in Oncology* 2000; **12**(4):337–44.

Woods 1999 {published data only}

Woods WG. Myeloablative therapy followed by stem cell rescue for pediatric solid tumors: A non-transplanter's perspective. *Cancer Research Therapy and Control* 1999;**9**(1-2):95–9.

Worden 2005 {published data only}

Worden FP, Taylor JM, Biermann JS, Sondak VK, Leu KM, Chugh R, et al.Randomized phase II evaluation of 6 g/m2 of ifosfamide plus doxorubicin and granulocyte colonystimulating factor (G-CSF) compared with 12 g/m2 of ifosfamide plus doxorubicin and G-CSF in the treatment of poor-prognosis soft tissue sarcoma. *Journal of Clinical Oncology* 2005;23(1):105–12.

Yamada 2007 {published data only}

Yamada K, Takahashi M, Ogura M, Kagami Y, Taji H, Kamiya Y, et al. High-dose chemotherapy and autologous peripheral blood stem cell transfusion for adult and adolescent patients with small round cell sarcomas. *Bone Marrow Transplantation* 2007;**39**(8):471–6.

Yaniv 1990 {published data only}

Yaniv I, Bouffet E, Irle C, Negrier S, Biron P, Favrot M, et al. Autologous bone marrow transplantation in pediatric solid tumors. *Pediatric Hematology and Oncology* 1990;7(1): 35–46.

Yaniv 2000 {published data only}

Yaniv I. Lymphokines post autologous peripheral blood stem cell transplantation in children. *Pediatric Hematology and Oncology* 2000;**17**(1):9–13.

Yaqoob 2006 {published data only}

Yaqoob N, Hasan SH. Desmoplastic small round cell tumor. Journal of the College Physicians and Surgeons - Pakistan 2006;16(9):614–6.

Yin 2009 {published data only}

Yin X, Zhang H, Wu T, Yan Y, Bu H. Treatment for leiomyosarcoma and leiomyoma in children with HIV infection. *Cochrane Database of Systematic Reviews* 2009, Issue 1. [DOI: 10.1002/14651858.CD007665.pub2.]

Young 1989 {published data only}

Young MM, Kinsella TJ, Miser JS, Triche TJ, Glaubiger DL, Steinberg SM, et al. Treatment of sarcomas of the chest wall using intensive combined modality therapy. *International Journal of Radiation Oncology, Biology, Physics* 1989;**16**(1):49–57.

Zoubek 1994 {published data only}

Zoubek A, Holzinger B, Mann G, Peters C, Emminger W, Perneczky-Hintringer E, et al. High-dose cyclophosphamide, adriamycin, and vincristine (HD-CAV) in children with recurrent solid tumor. *Pediatric Hematology and Oncology* 1994;**11**(6):613–23.

Additional references

AICC 2002

American Joint Committee on Cancer. AJCC cancer staging handbook. 6. Berlin: Springer, 2002.

Baker 2003

Baker KS, DeFor TE, Burns LJ, Ramsay NK, Neglia JP, Robison LL. New malignancies after blood or marrow stem-cell transplantation in children and adults: incidence and risk factors. *Journal of Clinical Oncology* 2003;**21**(7): 1352–8.

Banna 2007

Banna GL, Simonelli M, Santoro A. High-dose chemotherapy followed by autologous hematopoietic stemcell transplantation for the treatment of solid tumors in adults: A critical review. *Current Stem Cell Research and Therapy* 2007;**2**(1):65–82.

Bonnet 1997

Bonnet D, Dick JE. Human acute myeloid leukemia is organized as a hierarchy that originates from a primitive hematopoietic cell. *Nature Medicine* 1997;3(7):730–7.

Carvajal 2005

Carvajal R, Meyers P. Ewing's sarcoma and primitive neuroectodermal family of tumors. *Hematology/Oncology Clinics of North America* 2005;**19**(3):501–525.

Casali 2009

Casali PG, Jost L, Sleijfer S, Verweij J, Blay JY, ESMO Guidelines Working Group. Soft tissue sarcomas: ESMO clinical recommendations for diagnosis, treatment and follow-up. *Annals of Oncology* 2009;**20 Suppl 4**:32–6.

Clark 2005

Clark MA, Fisher C, Judson I, Thomas JM. Soft-tissue sarcomas in adults. *New England Journal of Medicine* 2005; **353**(7):701–11.

ClinicalTrials.gov 2010

ClinicalTrials.gov. National Institutes of Health, Bethesda, MD, USA 2010, issue http://clinicaltrials.gov/.

Dileo 2005

Dileo P, Demetri GD. Update on new diagnostic and therapeutic approaches for sarcomas. *Clinical Advances in Hematology & Oncology* 2005;**3**(10):781–91.

EBMT 2009

EBMT. European Group for Blood and Marrow Transplantation. EBMT Central Registry Office, London, UK 2009, issue http://www.ebmt.org.

EBMT STWP 2010

EBMT European Group for Blood and Marrow Transplantation, STWP Solid Tumor Working Party. Ongoing studies. EBMT Central Registry Office, London, UK 2010, issue http://www.ebmt.org/5WorkingParties/ STWP/wparties-st5.html.

Enzinger 2001

Weiss SW, Goldblum JR. General considerations. In: Weiss SW, Goldblum JR editor(s). *Enzinger and Weiss's soft tissue tumors*. 4. St Louis: Mosby, 2001:1–19.

Fletcher 2002

Fletcher CDM, Rydholm A, Singer S, Sundaram M, Coindre JM. Soft tissue tumours: WHO classification, epidemiology, clinical features, histopathological typing and grading. In: Fletcher CDM, Unni KK, Mertens F editor (s). World Health Organization classification of tumours. Pathology and genetics of tumours of soft tissue and bone. Lyon: International Agency for Research on Cancer, 2002:12–8.

Gurney 1997

Gurney JG, Young JL, Roffers SD, Smith MA, Bunin GR. Soft tissue sarcomas. In: Ries LAG, Smith MA, Gurney JG, Linet M, Tamra T, Young JL, Bunin GR editor(s). Cancer incidence and survival among children and adolescents: United States SEER program 1975-1995 (SEER Pediatric Monograph) NIH Pub. No. 99-4649. Bethesda: National Cancer Institute, 1997:111–24.

Higgins 2009

Higgins JPT, Green S, editors. Cochrane Handbook for Systematic Reviews of Interventions 5.0.0 [updated September 2009]. The Cochrane Collaboration, 2008. Available from www.cochrane-handbook.org.

ICTRP 2010

International Clinical Trials Registry Platform. World Health Organization, Geneva, Switzerland 2010, issue http://www.who.int/ictrp/en/.

ISRCTN 2010

International Standard Randomised Controlled Trial Number Register. Current Controlled Trials Ltd, London, UK 2010, issue http://www.controlled-trials.com/.

Kasper 2007

Kasper B. Standards and novel therapeutic options in the treatment of patients with soft tissue sarcoma. *Reviews on Recent Clinical Trials* 2007;**2**(3):206–11.

Kotilingam 2006

Kotilingam D, Lev DC, Lazar AJ, Pollock RE. Staging soft tissue sarcoma: evolution and change. *CA: A Cancer Journal for Clinicians* 2006;**56**(5):282–91.

Miller 1995

Miller RW, Young JL Jr, Novakovic B. Childhood cancer. Cancer 1995;75 Suppl(1):395–405.

NCI 2008a

SEER Cancer Statistics Review, 1975-2005. Table I-1. Estimated new cancer cases and deaths for 2008, National Cancer Institute; 2008 [2008]. National Cancer Institute (NCI), Bethesda, MD, USA issue http://seer.cancer.gov/csr/1975`2005/results`merged/sect`01`overview.pdf.

NCI 2008b

SEER Cancer Statistics Review, 1975-2005. Table I-10. Age distribution (%) of incidence cases by site, 2001-2005. National Cancer Institute (NCI), Bethesda, MD, USA 2008, issue http://seer.cancer.gov/csr/1975 2005/results merged/topic age dist.pdf.

NCI 2008c

SEER Cancer Statistics Review, 1975-2005. Table I-11. Median age of cancer patients at diagnosis, 2001-2005, N. National Cancer Institute (NCI), Bethesda, MD, USA 2008, issue http://seer.cancer.gov/csr/1975*2005/results*merged/topic*med*age.pdf.

NCI 2009a

A Snapshot of Sarcoma, National Cancer Institute; 2009. National Cancer Institute (NCI), Bethesda, MD, USA 2009, issue http://www.cancer.gov/aboutnci/servingpeople/snapshots/sarcoma.pdf.

NCI 2009b

Common Terminology Criteria for Adverse Events (CTCAE) and Common Toxicity Criteria (CTC). National Cancer Institute (NCI), Bethesda, MD, USA 2009, issue http://ctep.cancer.gov/protocolDevelopment/electronic applications/ctc.htm.

NCI CTEP 2006

NCI National Cancer Institute. CTCAE Common terminology criteria for adverse events. CTEP Cancer Therapy Evaluation Program 2006, issue http://ctep.cancer.gov/protocoldevelopment/electronic applications/docs/ctcaev3.pdf.

NCI PDQ 2010

Clinical Trials, Physician Data Query (PDQ). National Cancer Institute (NCI), Bethesda, MD, USA 2010, issue http://www.cancer.gov/clinicaltrials/search.

Neglia 2001

Neglia JP, Friedman DL, Yasui Y, Mertens AC, Hammond S, Stovall M, et al. Second malignant neoplasms in five-year survivors of childhood cancer: Childhood Cancer Survivor Study. *Journal of the National Cancer Institute* 2001;**93**(8): 618–29

NIHR UKCRN 2010

United Kingdom Clinical Research Network's (UKCRN) Portfolio Database. National Institute for Health

Research (NIHR), London, UK 2010, issue http://public.ukcrn.org.uk/search/.

Pedrazzoli 2006

Pedrazzoli P, Ledermann JA, Lotz JP, Leyvraz S, Aglietta M, Rosti G, et al. High dose chemotherapy with autologous hematopoietic stem cell support for solid tumors other than breast cancer in adults. *Annals of Oncology* 2006;17(10): 1479–88.

Pinkerton 1986

Pinkerton R, Philip T, Bouffet E, Lashford L, Kemshead J. Autologous bone marrow transplantation in paediatric solid tumours. *Clinics in Haematology* 1986;**15**(1):187–203.

Ramanathan 1999

Ramanathan RC, A'Hern R, Fisher C, Thomas JM. Modified staging system for extremity soft tissue sarcomas. Annals of Surgical Oncology 1999;6(1):57–69.

Reichardt 2002

Reichardt P. High-dose chemotherapy in adult soft tissue sarcoma. *Critical Reviews in Oncology/Hematology* 2002;**41** (2):157–67.

Review Manager 2008

Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration. Review Manager (RevMan). 5. Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2008.

Rosti 2002

Rosti G, Ferrante P, Ledermann J, Leyvraz S, Ladenstein R, Koscielniak E, et al. High-dose chemotherapy for solid tumors: results of the EBMT. *Critical Reviews in Oncology/Hematology* 2002;**41**(2):129–40.

Sanchez-Garcia 2007

Sanchez-Garcia I, Vicente-Duenas C, Cobaleda C. The theoretical basis of cancer-stem-cell-based therapeutics of cancer: Can it be put into practice?. *BioEssays* 2007;**29**(12): 1269–80.

SAS Institute Corp 2009

SAS Institute Inc.,100 SAS Campus Drive, Cary, NC 27513-2414, USA. English Title:. SAS Institute Inc.,100 SAS Campus Drive, Cary, NC 27513–2414, USA, 2009.

Seeger 1991

Seeger RC, Reynolds CP. Treatment of high-risk solid tumors of childhood with intensive therapy and autologous bone marrow transplantation. *Pediatric Clinics of North America* 1991;**38**(2):393–424.

Sondak 2001

Sondak VK, Chang AE. Clinical evaluation and treatment of soft tissue tumors. In: Weiss SW, Goldblum JR editor(s). *Enzinger and Weiss's soft tissue tumors*. 4. St Louis: Mosby, 2001:21–44.

Spunt 2006

Spunt SL, Pappo AS. Childhood nonrhabdomyosarcoma soft tissue sarcomas are not adult-type tumors. *Journal of Clinical Oncology* 2006;**24**(12):1958–9.

Therasse 2000

Therasse P, Arbuck SG, Eisenhauer EA, Wanders J, Kaplan RS, Rubinstein L, et al.New guidelines to evaluate the response to treatment in solid tumors. *Journal of the National Cancer Institute* 2000;**92**(3):205–16.

Thomson Reuters Corp 2009

Thomson Reuters, 3 Times Square, New York City, NY 10036, USA. Reference Manager Version 11. Thomson Reuters, 3 Times Square, New York City, NY 10036, USA, 2009.

Tsao 2000

Tsao H. Update on familial cancer syndromes and the skin. *Journal of the American Academy of Dermatology* 2000;**42**(6): 939–69.

Tumorregister München 2007

Tumorregister München. Überlebenszeitanalysen für Weichteilsarkome klassifiziert nach der Histologie. Tumorzentrum München. München: Tumorregister, 2007, issue http://www.tumorregister-muenchen.de/facts/surv/surv hST · · G.pdf.

Van Glabbeke 1999

Van Glabbeke M, van Oosterom AT, Oosterhuis JW, Mouridsen H, Crowther D, Somers R, et al. Prognostic factors for the outcome of chemotherapy in advanced soft tissue sarcoma: an analysis of 2,185 patients treated with anthracycline-containing first-line regimens--a European Organization for Research and Treatment of Cancer Soft Tissue and Bone Sarcoma Group Study. *Journal of Clinical Oncology* 1999;17(1):150–7.

Woods 1999

Woods WG. Myeloablative therapy followed by stem cell rescue for pediatric solid tumors: A non-transplanter's perspective. *Cancer Research Therapy & Control* 1999;**9**(1-2):95–9.

* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Al Balushi 2009

Methods	Retrospective report of cases without control in a single-centre, observed in Canada from 2000 to 2007
Participants	5 children with desmoplastic small round-cell tumor with metasteses; 4 male, 1 female; mean age 11 years
Interventions	HDCT followed by autologous HSCT in 3 of 5 children
Outcomes	overall survival; toxicity
Notes	individual data for each child presented

Andres 2006

Methods	Retrospective report of a single case without control in a single-centre, observed in Spain; no information on observation period available
Participants	1 female 21 years of age with desmoplastic small round-cell tumor
Interventions	HDCT followed by autologous HSCT
Outcomes	overall survival; toxicity
Notes	

Bernbeck 2007

Methods	Retrospective report of cases without control in a single-centre study, observed in Germany from 2001 to 2005
Participants	9 participants with high-risk soft tissue sarcomas, 8 of whom had rhabdomyosarcoma or synovial sarcoma at various clinical stages. Children and young adults (3 male, 5 female) ranging from 1 to 21 years of age
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival; toxicity
Notes	individual data for each participant presented

Bertuzzi 2003

Methods	Prospective study of consecutive cases without control in a single-centre, observed in Italy from 1997 to 2002
Participants	10 adults (10 male) aged 15 to 60 years, median 29 years of age with advanced desmoplastic small round-cell tumor at various clinical stages, n=4 with metastases
Interventions	HDCT (melphalan, mitoxantrone, thiotepa) followed by autologous peripheral HSCT
Outcomes	overall survival; progression-free survival; toxicity (NCI-CTC)
Notes	aggregate data

Bley 2004

Methods	Retrospective report of a single case without control in a single-centre, observed in Germany; no information on observation period available
Participants	1 female 22 years of age with liposarcoma
Interventions	HDCT followed by autologous HSCT
Outcomes	overall survival; toxicity
Notes	

Bölke 2005

Methods	Retrospective report of a single case without control in a single-centre, observed in Germany in 1993
Participants	1 female 33 years of age with recurrent malignant fibrous histiocytoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	overall survival; adverse events
Notes	

Cole 1999

Methods	Retrospective report of a single case without control in a single-centre, observed in the United States; information on observation period not available
Participants	1 male 26 years of age with synovial sarcoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	overall survival; adverse events

Cole 1999 (Continued)

Notes	
Doros 2008	
Methods	Retrospective report of a single case without control in a single-centre, observed in the United States; information on observation period not available
Participants	1 male 14 years of age with desmoplastic small round-cell tumor
Interventions	HDCT followed by autologous HSCT
Outcomes	overall survival; adverse events
Notes	
Endo 1996	
Methods	Retrospective report of a case series without controls in a single-centre, observed in Japan from 1987 to 1995
Participants	16 participants with high-risk solid tumors; of which 1 male 19 years of age with undifferentiated sarcoma
Interventions	HDCT followed by autologous peripheral and bone marrow HSCT
Outcomes	overall survival; adverse events
Notes	individual data available for each case
Engelhardt 20	07
Methods	Retrospective report of a consecutive series of cases without control, observed from 1992 to 2003; the number of centres and the countries were not specified (probably in Germany and the United States of America)
Participants	35 participants with Ewing sarcoma and soft tissue sarcomas, of which 23 had NRSTS (anaplastic soft tissue sarcoma, angiosarcoma, fibrosarcoma, leiomyosarcoma, liposarcoma, malignant fibrous histiocytoma, malignant haemangioperiocytoma, synovial sarcoma); 12 male, 11 female, 21 to 56 years of age
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival; toxicity
Notes	individual data for participants with rhabdomyosarcoma and other soft tissue sarcoma. The reported aggregate data were not considered because the proportion of participants with NRSTS plus 3 rhabdomyosarcomas was less then 80% of all participants therefore did not meet our eligibility criteria

Fang 2008

Methods	Retrospective report of cases without control in a single-centre, observed in the United States in 2006
Participants	2 cases, 1 of whom had a desmoplastic small round-cell tumor; female 23 years of age
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival
Notes	individual data presented for each participant

Farruggia 2008

Methods	Retrospective report of a single case in a single-centre, observed in Italy; information on observation period not available
Participants	1 male 10 years of age with synovial sarcoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	overall survival; toxicity
Notes	

Fetscher 1997

Methods	Retrospective report of a single case in a single-centre, observed in Germany in 1994
Participants	1 female 23 years of age with metastatic leiomyosarcoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	overall survival
Notes	

Frapier 1998

Methods	Retrospective report of a single case in a single-centre, observed in France; information on observation period not available
Participants	1 male 11 years of age with high grade undifferentiated sarcoma
Interventions	HDCT followed by autologous bone marrow HSCT
Outcomes	survival; adverse events

Frapier 1998 (Continued)

Notes	
Fraser 2006	
Methods	Report of two prospective phase I/II studies of cases without control in a single-centre, observed in the United States from 1995 to 2004
Participants	36 participants with solid tumors that were metastatic or relapsed following therapy, treated on two consecutive trial protocols n=11 and n=25; 5 had desmoplastic small round-cell tumor and rhabdoid tumor; aged from 2 to 20 years
Interventions	HDCT followed by autologous peripheral or bone marrow HSCT
Outcomes	survival; toxicity
Notes	individual data for each participant reported on

Garrido 1998

Methods	Retrospective report of cases in a single centre; observed in the United States; 1991 to 1995	
Participants	2 participants, n=1 45 year old man with metastatic liposarcoma	
Interventions	HDCT followed by autologous HSCT	
Outcomes	adverse events	
Notes	individual data available for each case	

Graham 1997

Methods	Prospective phase I/II trial in a single centre, observed in the United States from 1991 to 1995	
Participants	49 participants with recurrent and high-risk pediatric brain tumors 1 of whom had fibrosarcoma; aged 1 to 32 years, median 12 years	
Interventions	HDCT followed by autologous bone marrow HSCT	
Outcomes	survival; toxicity	
Notes	individual data available for participants	

Hawkins 2002

Methods	Prospective phase I/II trial without controls in a 3-centre study, observed in United States from 1996 to 1998
Participants	23 children and adolescents with metastatic sarcomas of whom 6 had NRSTS (desmoplastic small round-cell tumor, fibromyxoid sarcoma, leiomyosarcoma, undifferentiated sarcoma) 5 male; 1 female, ranging from 5 to 19 years of age
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival, toxicity
Notes	individual data available for each participant

Hoogerbrugge 1997

Methods	Retrospective report of a single case in a single centre, observed in the Netherlands; information on observation period not available
Participants	1 child 1 year of age with fibrosarcoma; information on gender not available
Interventions	HDCT followed by autologous bone marrow HSCT
Outcomes	overall survival; toxicity
Notes	

Ivanova 2007

Methods	Retrospective report of cases with results from a control group in a single-centre study, observed in Russia from 1990 to 2006
Participants	103 patients with various soft tissue sarcomas were investigated in two treatment groups; individual information on diagnoses, age and gender not available
Interventions	34 children were treated with HDCT (carboplatin, cyclophosphamide, etoposide) followed by autologous HSCT; 69 children as a control; information on the control therapy not available
Outcomes	overall survival, adverse events
Notes	aggregate data

Risk of bias

Bias	Authors' judgement	Support for judgement
Assignment of patients to treatment groups	High risk	no information about assignment; study report results about transplanted patients

Ivanova 2007 (Continued)

		such as a case series; then adds OS of an unspecified control group; control group was matched to the test group, however, the description of this procedure was not comprehensible
Concurrent control	High risk	control group was probably selected from historical data
Comparable baseline characteristics	Unclear risk	no report about patients' characteristics, except the average tumor volume 180 cm ³ which was comparable in both groups; no other baseline characteristics; information insufficient to assess comparability between groups
Loss to follow-up	Unclear risk	loss to follow up was not addressed; no patient flow described
Selective outcome reporting	Unclear risk	the study has a retrospective design and a study protocol is not available
Other causes for high risk of bias	Unclear risk	study results were reported amidst reports of results from other studies; the article resembles a narrative review; information about statistical methods was sparse

Kaminski 2000

Methods	Retrospective report of a single case in a single centre, observed in the United States in 1993
Participants	1 child (female) 6 years of age with fibrosarcoma
Interventions	HDCT followed by autologous bone marrow HSCT
Outcomes	overall survival; adverse events
Notes	

Kasper 2007

Methods	Retrospective report of cases without control in a single-centre study, observed in Germany from 1998 to 2007
Participants	38 participants with soft tissue and bone sarcomas, of whom 14 had NRSTS (leiomyosarcoma; liposarcoma; malignant fibrous histiocytoma; not otherwise specified soft tissue sarcomas; synovial sarcoma) aged 23 to 65 years; no individivual information available on gender

Kasper 2007 (Continued)

Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	toxicity (WHO)
Notes	mainly aggregate data reported

Kasper 2010

Methods	Prospective controlled clinical trial; single-institutional phase II study, observed in Germany from 2003 to 2008
Participants	34 patients with various solid tumors were investigated in two treatment groups
Interventions	9 participants with solid tumors were treated with HDCT (ifosfamide, carboplatin, etoposide) followed by autologous HSCT (peripheral blood stem cell rescue), of whom 7 had NRSTS (leiomyosarcoma; malignant fibrous histiocytoma; not otherwise specified soft tissue sarcomas; synovial sarcoma); aged 29 to 65 years of age 25 participants with solid tumors were treated in a control group with standard-dose chemotherapy (ifosfamide, doxorubicin), of whom 20 had NRSTS (leiomyosarcoma; liposarcoma; malignant fibrous histiocytoma; not otherwise specified soft tissue sarcomas; synovial sarcoma); age was not reported
Outcomes	overall survival; progression-free survival; adverse events
Notes	aggregate data included less than 80% NRSTS patient data; individual data for transplanted data were available but not included in the meta-analysis because follow up did not start at intervention

Risk of bias

Bias	Authors' judgement	Support for judgement
Assignment of patients to treatment groups	High risk	no information about assignment; all par- ticipants had metastatic disease; study re- ports results about transplanted and con- trol patients such as a case series
Concurrent control	Low risk	all participants were treated bewteen 2003 and 2008
Comparable baseline characteristics	Unclear risk	tumor classification and metastatic site re- ported for all participants; age reported for 9 transplanted participants; no other baseline characteristics; information insuf- ficient to assess comparability between groups

Kasper 2010 (Continued)

Loss to follow-up	Unclear risk	loss to follow up was not addressed; no patient flow described
Selective outcome reporting	Unclear risk	overall survival was assessed from time of study inclusion and not from time of transplantation; the gap between study inclusion and transplantation may be up to 4 months; reporting makes comparision to other studies difficult; treatment-related mortality not reported; toxicity reported scarcely
Other causes for high risk of bias	Unclear risk	duplicate data because some patients were reported in Kasper 2007 and Kasper 2010

Kozuka 2002

Methods	Retrospective report of cases without control in a single-centre study, observed in Japan from 1999 to 2000
Participants	2 adults (2 male) 21 and 37 years of age with recurrent NRSTS (malignant fibrous histiocytoma or malignant hemangiopericytoma)
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival; adverse events
Notes	individual data for each participant reported

Kretschmar 1996

Methods	Retrospective report of cases in a single-centre study, observed in the United States; information on observation period not available
Participants	3 adolescent males with desmoplastic small round-cell tumor, 1 of whom received HSCT (13 years of age)
Interventions	HDCT followed by autologous bone marrow HSCT
Outcomes	survival; toxicity
Notes	

Krskova 2007

Methods	Retrospective report of a single case in a single centre, observed in the Czech Republic in 1998
Participants	1 child (male) 9 years of age with synovial sarcoma
Interventions	HDCT followed by autologous HSCT
Outcomes	overall survival; adverse events
Notes	

Kurre 2000

Methods	Retrospective report of 3 cases in a single-centre study, observed in the United States from 1994 to 1998
Participants	3 cases with desmoplastic small round-cell tumor, 2 of whom received HSCT: 1 male aged 5 years and 1 female aged 2.5 years
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival; adverse events
Notes	individual data reported for each participant

Kushner 1996

Methods	Prospective phase I/II study without controls in a single centre, observed in the United States; information on observation period not available
Participants	12 patients with desmoplastic small round-cell tumor of whom 4 boys from 10 to 14 years of age received HSCT
Interventions	HDCT (P6 protocol) followed by autologous bone marrow HSCT
Outcomes	Survival; toxicity (NCI CTC criteria)
Notes	individual data reported for each participant

Kushner 2001

Methods	Preliminary results from a prospective phase II/III study in a single-centre study, observed in the United States; information on observation period not available
Participants	21 participants with neuroblastoma, brain tumors and other poor-risk solid tumors, of whom 1 male 29 years of age with desmoplastic small round-cell tumor
Interventions	HDCT followed by autologous peripheral HSCT

Kushner 2001 (Continued)

Outcomes	overall survival; toxicity according to NCI CTC criteria
Notes	individual data for each participant available

Kushner 2008

Methods	Retrospective report of a case in a single-centre study, observed in the United States; information on observation period not available
Participants	1 adult (male) 18 years of age with desmoplastic small round-cell tumor
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival; adverse events
Notes	individual data

Kühne 2000

Methods	Prospective phase I/II study without control group in a single centre, observed in Switzerland from 1997 to 1999
Participants	11 children with brain tumors, soft tissue sarcomas, germ-cell tumors and neuroblastomas, of whom 1 child (male) 3 years of age with rhabdoid sarcoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	overall survival; adverse events
Notes	individual data available for each participant

Lafay-Cousin 2000

Methods	Prospective phase II study without controls in a 4-centre study, observed in France from 1986 to 1998
Participants	18 children with recurrent mesenchymal tumors, of whom 4 (2 male; 2 female) from 10 to 16 years of age with NRSTS (desmoplastic small round-cell tumor; undifferentiated sarcoma)
Interventions	HDCT followed by autologous peripheral or bone marrow HSCT
Outcomes	overall survival; adverse events
Notes	individual data reported for each participant

Lashkari 2009

Methods	Prospective phase II study without controls in a single centre, observed in the USA from 1995 to 1999
Participants	13 children with locally advanced or metastatic sarcoma; of whom 2 (1 male; 1 female) 40 and 30 years of age with metastatic NRSTS (malignant fibrous histiocytoma; soft tissue sarcoma without histologic subtype information)
Interventions	HDCT followed by autologous HSCT
Outcomes	survival
Notes	individual data reported for each participant

Lippe 2003

Methods	Retrospective report of 2 cases without controls in a single-centre study; information on observation period not available
Participants	Two participants with desmoplastic small round-cell tumor; of whom 1 adult (male) 27 years of age received HSCT
Interventions	HDCT followed by autologous HSCT
Outcomes	survival; toxicity
Notes	individual data reported for each participant

Livaditi 2006

Methods	Retrospective report of 5 cases without controls in a single centre, observed in Greece; information on observation period not available
Participants	5 children with desmoplastic small round-cell tumor, of whom 2 (1 male aged 7 years and 1 female aged 13 years) received HSCT
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival; adverse events
Notes	individual data reported for each participant

Madigan 2007

Methods	Retrospective report of 14 cases without controls in a single centre, observed in the United States from 1983 to 2003
Participants	14 children with extracranial rhabdoid tumors, of whom 2 (1 male aged 6 months and 1 female aged 30 months) received HSCT for rhabdoid sarcoma
Interventions	HDCT followed by autologous HSCT

Madigan 2007 (Continued)

Outcomes	survival; adverse events
Notes	some individual data reported for each participant and aggregate data on survival time in Kaplan-Meier curve

Matsuzaki 2002

Methods	Retrospective report of a single case in a single centre, observed in Japan in 1999
Participants	1 child (female) 11 years of age with synovial sarcoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	overall survival; adverse events
Notes	individual data

Mazuryk 1998

Methods	Retrospective report of a single case in a single centre, observed in Canada in 1996
Participants	1 female 19 years of age with desmoplastic small round-cell tumor
Interventions	HDCT followed by autologous HSCT
Outcomes	overall survival; adverse events
Notes	

Mesia 1994

Methods	Retrospective report of a series of cases without controls in a single centre, observed in Spain from 1989 to 1992
Participants	9 patients with metastatic sarcomas, of whom 1 male 21 years of age had undifferentiated sarcoma
Interventions	HDCT followed by autologous bone marrow HSCT
Outcomes	survival; adverse events
Notes	individual data reported for individual patients

Mitchell 1994

Methods	Unclear if retrospective or prospective study without controls in a single centre, observed in the United Kingdom; information on observation period not available
Participants	11 patients with malignant disease, of whom 1 male 16 years of age with angiosarcoma
Interventions	HDCT followed by autologous peripheral and bone marrow HSCT
Outcomes	survival; toxicity
Notes	individual data reported for each participant

Nakamura 2008

Methods	Retrospective report of a case without control in a single-centre study, observed in Japan; information on observation period not available
Participants	1 child (male) 11 years of age with undifferentiated soft tissue sarcoma
Interventions	HDCT followed by autologous HSCT
Outcomes	overall survival; adverse events
Notes	

Navid 2006

Methods	Prospective phase II study without controls in a single centre, observed in the United States from 1996 to 2000
Participants	24 patients with high-risk sarcomas, of whom 4 had desmoplastic small round-cell tumor and 2 received HSCT(2 males aged 14 and 21 years)
Interventions	HDCT followed by autologous peripheral and bone marrow HSCT
Outcomes	survival; adverse events
Notes	individual data reported for each participant

Patel 2004

Methods	Prospective phase II study without controls in a single-centre, observed in the United States from 1994 to 2001
Participants	37 patients with skeletal osteosarcoma and variant bone tumors with poor prognosis, of whom 6 adults had malignant fibrous histiocytoma; individual information on age and gender not available, median age 38 years, range 18-63 years
Interventions	HDCT followed by autologous peripheral HSCT

Patel 2004 (Continued)

Outcomes	toxicity according to NCI CTC
Notes	individual data available for toxicity

Peters 1986

Methods	Prospective phase I study without controls in a single centre, observed in the United States; information on observation period not available
Participants	29 patients with metastatic cancer and sarcoma, of whom 2 females aged 24 and 38 years of age with fibrosarcoma and leiomyosarcoma
Interventions	HDCT followed by autologous bone marrow HSCT
Outcomes	overall survival; toxicity
Notes	individual data reported for each participant

Peters 1989

Methods	Prospective phase I study without controls in a single centre, observed in the United States; information on observation period not available
Participants	23 patients with metastatic cancer and sarcoma, of whom 2 patients aged 15 and 26 years with synovia sarcoma; individual information on gender not available
Interventions	HDCT followed by autologous bone marrow HSCT
Outcomes	adverse events
Notes	individual data

Recchia 2006

Methods	Retrospective report of a single case in a single centre, observed in Italy; information on observation period not available
Participants	1 adult (male) 40 years of age with malignant fibrous histiocytoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	overall survival; toxicity
Notes	individual data

Ronghe 2004

Methods	Retrospective report of 2 cases in a single centre, observed in the United Kingdom; information on observation period not available
Participants	2 children with malignant rhabdoid tumors, of whom 1 female 14 months of age received HSCT
Interventions	HDCT followed by autologous bone marrow HSCT
Outcomes	overall survival; adverse events
Notes	individual data reported for each patient

Saab 2007

Methods	Retrospective report of cases in a single centre, observed in the United States; information on observation period not available
Participants	11 participants with desmoplastic small round-cell tumor, of whom 4 males from 5 to 21 years of age received HSCT
Interventions	HDCT followed by autologous HSCT
Outcomes	survival; toxicity
Notes	individual data reported for each participant

Shaw 1996

Methods	Prospective phase I study without controls in Australia, UK and Israel; information on the number of centres not available
Participants	30 patients with malignant solid tumors, of whom 2 children aged 1 and 2 years with NRSTS (rhabdoid sarcoma; not otherwise specified); individual information, gender not available
Interventions	HDCT followed by autologous peripheral or bone marrow HSCT
Outcomes	adverse events
Notes	individual data for each participant reported

Slease 1988

Methods	Phase I trial in a single-centre study, observed in the United States; information on observation period not available
Participants	26 adults with refractory malignant solid tumors, of whom 3 adult males from 41 to 47 years of age with NRSTS (leiomyosarcoma; malignant fibrous histiocytoma)
Interventions	HDCT followed by autologous bone marrow HSCT

Slease 1988 (Continued)

Outcomes	survival; adverse events
Notes	individual data for each participant reported

Sung 2003

Methods	Prospective phase I study in a single centre in Korea from 1998 to 2001
Participants	26 participants with high-risk malignant solid tumors, of whom 2 children (1 male aged 20 months and 1 female aged 47 months) with NRSTS (malignant fibrous histiocytoma; rhabdoid sarcoma)
Interventions	successive double HDCT followed by autologous peripheral HSCT
Outcomes	overall survival; adverse events
Notes	individual data

Watanabe 2006

Methods	Retrospective report of a case without control in a single-centre study, observed in Japan; information on observation period not available
Participants	1 child (male) 1 year of age with rhabdoid sarcoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	survival; adverse events
Notes	

Yamamura 2003

Methods	Retrospective report of a single case in a single centre, observed in Japan in 1996
Participants	1 adult (male) 33 years of age with malignant fibrous histiocytoma
Interventions	HDCT followed by autologous peripheral HSCT
Outcomes	overall survival; adverse events
Notes	reports on the development of chronic myelocytic leukemia following chemotherapy

Yonemoto 1999

Methods	Retrospective report of cases without control in a single-centre study, observed in Japan from 1995
Participants	3 young adults (3 male; 0 female) from 17 to 40 years of age with synovial sarcoma, of a total of 10 participants with sarcomas
Interventions	HDCT followed by autologous HSCT
Outcomes	adverse events
Notes	individual data

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Abdel-Dayem 1999	not intervention of interest
Abidi 2007	not diagnosis of interest
ABMTR 1986	not study design of interest (review)
ABMTR 1989	not diagnosis of interest
Abrahamsen 2000	data of interest not described separately
Admiraal 2007	not diagnosis of interest (rhabdomyosarcoma)
Aleinikova 2002	not diagnosis of interest (rhabdomyosarcoma)
Alpers 1982	not intervention of interest
Anderson 2005	not study design of interest
Antman 1987	not study design of interest (review)
Antman 1990	not intervention of interest
Antman 2001	not study design of interest (review)
Ashihara 2002	not diagnosis of interest (rhabdomyosarcoma)
Atra 1996	not study design of interest (review)
Atra 2002	not study design of interest (review)

Avramova 2006	follow-up of study Michalov 2001
Bader 1989	not diagnosis of interest (rhabdomyosarcoma)
Bagnulo 1985	not diagnosis of interest (rhabdomyosarcoma)
Bambakidis 2002	not diagnosis of interest
Banna 2007	not study design of interest (review)
Barfield 2008	not study design of interest (review)
BCBS MAP 1999	not diagnosis of interest
Beaujean 1989	not diagnosis of interest (rhabdomyosarcoma)
Bellmunt 1997	data of interest not described separately
Bertuzzi 2002	data of interest not described separately
Beschorner 2006	not diagnosis of interest
Bezwoda 1994	not diagnosis of interest
Bickert 2002	not study design of interest (review)
Bien 2007	not diagnosis of interest (rhabdomyosarcoma)
Bini-Antunes 2006	not diagnosis of interest
Bisogno 2009	not disease of interest (rhabdomyosarcoma)
Blay 1994	not study design of interest (abstract)
Blay 2000	not disease of interest for 11 of 30 (36%) patients: 5 patients with rhabdomyosarcoma, 3 patients with unclassified sarcoma, 1 paraganglioma, 1 Schwannosarcoma, and 1 peripheral neuroectodarmal tumorwere included in a total of 30 analyzed patients
Bode-Lesniewska 2005	not intervention of interest
Bodey 1981	not diagnosis of interest
Bojko 2002	not diagnosis of interest
Bokemeyer 1997	not diagnosis of interest
Borden 1987	not intervention of interest

Boulad 1998	data of interest not described separately
Bramwell 1986	not intervention of interest
Bramwell 1987	not intervention of interest
Bramwell 2001	not intervention of interest
Breitfeld 2001	not intervention of interest
Brugger 1995	not outcome of interest; paper retracted
Brugieres 1988	not diagnosis of interest (rhabdomyosarcoma)
Brun 1984	not intervention of interest
Bylund 2008	not intervention of interest
Cacchione 2008	data of interest not described separately
Caglar 2002	not diagnosis of interest
Carli 1988	not intervention of interest
Carli 1999	not diagnosis of interest (rhabdomyosarcoma)
Casado 2004	not study design of interest (review)
Casanova 2009	not disease of interest
Casper 1991	not intervention of interest
Ceschel 2006	data of interest not described separately
Chan 1991	not diagnosis of interest (rhabdomyosarcoma)
Chan 1999	not intervention of interest
Chang 1979	not intervention of interest
Chang 1979a	not intervention of interest
Chang 1981	not intervention of interest
Chang 1988	not intervention of interest
Chauvin 1991	not study design of interest (abstract)

Chen 1999 not study design of interest (review) Chen 2008 not intervention of interest Childs 2004 not study design of interest (review) Cho 2005 data of interest not described separately Chuman 2000 not study design of interest (review) Clausen 1993 data of interest not described separately Corbett 2009 not study design of interest (review) Coulibalya 2008 not intervention of interest Couzin 2007 not study design of interest (review)	
Childs 2004 not study design of interest (review) Cho 2005 data of interest not described separately Chuman 2000 not study design of interest (review) Clausen 1993 data of interest not described separately Corbett 2009 not study design of interest (review) Coulibalya 2008 not intervention of interest	
Cho 2005 data of interest not described separately Chuman 2000 not study design of interest (review) Clausen 1993 data of interest not described separately Corbett 2009 not study design of interest (review) Coulibalya 2008 not intervention of interest	
Chuman 2000 not study design of interest (review) Clausen 1993 data of interest not described separately Corbett 2009 not study design of interest (review) Coulibalya 2008 not intervention of interest	
Clausen 1993 data of interest not described separately Corbett 2009 not study design of interest (review) Coulibalya 2008 not intervention of interest	
Corbett 2009 not study design of interest (review) Coulibalya 2008 not intervention of interest	
Coulibalya 2008 not intervention of interest	
· · · · · · · · · · · · · · · · · · ·	
Couzin 2007 not study design of interest (review)	
Czyzewski 1999 data of interest not described separately	
Dagher 1997 not intervention of interest	
Dallorso 1996 not diagnosis of interest (rhabdomyosarcoma)	
Dallorso 2000 not study design of interest (review)	
Dantonello 2008 not intervention of interest	
De Kraker 1984 not diagnosis of interest	
De Pasquale 2003 not outcome of interest	
De Sio 2006 not diagnosis of interest (rhabdomyosarcoma)	
De Terlizzi 1988 not study design of interest (review)	
De Vries 1995 not diagnosis of interest	
Demirci 2003 not diagnosis of interest	
Demirer 2008 not intervention of interest	
Devalck 1992 not study design of interest (review)	
Diaz 1999 not diagnosis of interest (rhabdomyosarcoma)	
Dicke 1984 not study design of interest (review)	

-	
Dicke 1986	not study design of interest (review)
Dileo 2005	not study design of interest (review)
Dillman 1995	data of interest not described separately
Dincol 2000	not intervention of interest
Donaldson 2001	not intervention of interest
Donker 2009	not intervention of interest (allogeneic)
Drabko 2006	not diagnosis of interest (rhabdomyosarcoma)
Dumontet 1992	not diagnosis of interest (rhabdomyosarcoma)
Ederhy 2007	not diagnosis of interest
Eggermont 1997	not intervention of interest
Ek 2006	not study design of interest (review)
Ekert 1982	duplicate publication of study Ekert 1984
Ekert 1984	not diagnosis of interest (rhabdomyosarcoma)
Elias 1998	not study design of interest (review)
Emminger 1991	not diagnosis of interest (rhabdomyosarcoma)
Endo 1995	not study design of interest (review)
Erkisi 1996	not intervention of interest
Espinosa 2001	not diagnosis of interest
Fazekas 2008	not diagnosis of interest
Fekrat 1993	not diagnosis of interest (rhabdomyosarcoma)
Ferrari 2005	not outcome of interest
Fetscher 1996	not study design of interest (abstract)
Figuerres 2000	data of interest not described separately
Fizazi 1994	not intervention of interest

Flamant 1998	data of interest not described separately
Foncillas 2004	not diagnosis of interest (rhabdomyosarcoma)
Frustaci 2001	not intervention of interest
Fujita 2005	not diagnosis of interest
Gadner 2002	not study design of interest (review)
Garaventa 1986	not diagnosis of interest (rhabdomyosarcoma)
Garaventa 1987	not diagnosis of interest (rhabdomyosarcoma)
Gardner 2008	not diagnosis of interest
Gebhardt 1999	not intervention of interest
Geisler 2003	not diagnosis of interest
Geissler 1984	not intervention of interest
Gentet 1993	not study design of interest (review)
Ghalie 1994	not diagnosis of interest
Ghavamzadeh 2009	not diagnosis of interest (rhabdomyosarcoma)
Glenn 1985	not intervention of interest
Gonzalez 1989	not diagnosis of interest (rhabdomyosarcoma)
Gortzak 2001	not intervention of interest
Goto 2004	not study design of interest (review)
Graham 1992	not diagnosis of interest (rhabdomyosarcoma)
Graham-Pole 1995	not study design of interest (abstract)
Gratwohl 2002	not outcome of interest
Gratwohl 2004	not study design of interest (review)
Gratwohl 2004a	not study design of interest (review)
Gratwohl 2004b	not outcome of interest

Gratwohl 2006	not outcome of interest
Gratwohl 2007	not study design of interest (review)
Gratwohl 2007a	not study design of interest (review)
Grundy 2001	not diagnosis of interest (rhabdomyosarcoma)
Haas 1990	not diagnosis of interest
Hale 2005	not study design of interest (review)
Hara 1998	not diagnosis of interest (rhabdomyosarcoma)
Harmon 2001	not intervention of interest
Hartmann 1986	not diagnosis of interest (rhabdomyosarcoma)
Hartmann 1997	data of interest not described separately
Hartmann 2001	not diagnosis of interest
Hartmann 2005	not diagnosis of interest
He 1999	not intervention of interest
Hensel 2002	not diagnosis of interest
Herzog 2005	not study design of interest (review)
Hilden 1998	not diagnosis of interest
Hilden 2004	not diagnosis of interest
Hiraiwa 1983	not diagnosis of interest
Hoekstra 1994	not study design of interest (review)
Holtta 2005	not diagnosis of interest (rhabdomyosarcoma)
Holtta 2005a	not diagnosis of interest (rhabdomyosarcoma)
Horn 2002	data of interest not described separately
Horowitz 1993	not diagnosis of interest (rhabdomyosarcoma)
Hosoi 2007	not diagnosis of interest (rhabdomyosarcoma)

Hotte 2004	data of interest not described separately
Hoy 2007	not diagnosis of interest
Huang 2006	not diagnosis of interest (rhabdomyosarcoma)
Huttmann 2005	not diagnosis of interest
Höffken 1997	not study design of interest (review)
Iankelevich 2000	not outcome of interest
ICR 1994	not diagnosis of interest (rhabdomyosarcoma)
Imbach 1979	not diagnosis of interest
Irle 1989	not study design of interest (review)
Issels 1995	not study design of interest (review)
Issels 2002	not intervention of interest
Issels 2004	not study design of interest (review)
Jamil 2004	data of interest not described separately
Jelic 1994	not intervention of interest
Jelic 1997	not intervention of interest
Kaatsch 2009	not test intervention of interest
Kabickova 2003	data of interest not described separately
Kadan-Lottick 2008	not outcome of interest
Kaizer 1979	not diagnosis of interest (rhabdomyosarcoma)
Kaizer 1980	duplicate paper of Kaizer 1979
Kaizer 1984	not study design of interest (review)
Kalwak 2002	not diagnosis of interest (rhabdomyosarcoma)
Kanabar 1995	not diagnosis of interest (rhabdomyosarcoma)
Kasper 2004	data included in follow-up paper Kasper 2008

Kasper 2005	not study design of interest (review)
Kasper 2006	data included in follow-up paper Kasper 2008
Katzenstein 2003	not study design of interest
Kavan 1997	not study design of interest (review)
Kavan 1997a	not diagnosis of interest (rhabdomyosarcoma)
Kinsella 1988	not diagnosis of interest (rhabdomyosarcoma)
Kinsella 1988a	data of interest not described separately
Klaritsch 2006	not diagnosis of interest
Kletzel 1997	not study design of interest (review)
Kletzel 1998	not diagnosis of interest (rhabdomyosarcoma)
Kletzel 2005	not study design of interest (review)
Klingebiel 1989	not study design of interest (review)
Klingebiel 1994	not study design of interest (review)
Klingebiel 2008	not diagnosis of interest (rhabdomyosarcoma)
Kook 1998	not diagnosis of interest
Korfel 2001	not diagnosis of interest (rhabdomyosarcoma)
Koscielniak 1992	data of interest not described separately
Koscielniak 1997	not diagnosis of interest (rhabdomyosarcoma)
Koscielniak 1999	not study design of interest (review)
Koscielniak 2001	not study design of interest (review)
Koscielniak 2002	not study design of interest (review)
Koscielniak 2005	not study design of interest (review)
Kuroiwa 2009	not diagnosis of interest (rhabdomyosarcoma)

Kushner 2000	not study design of interest (review)
Kwan 1996	not diagnosis of interest (rhabdomyosarcoma)
Kwon 2010	not disease of interest (rhabdomyosarcoma, neuroblastoma)
Ladenstein 1993	not study design of interest (review)
Ladenstein 1997	not study design of interest (review)
Lal 2005	data of interest not described separately
Lang 2006	not diagnosis of interest (rhabdomyosarcoma)
Lange 2004	not intervention of interest (allogeneic HSCT)
Larsen 2000	not diagnosis of interest (rhabdomyosarcoma)
Le Cesne 2000	not intervention of interest
Le Corroller 1997	data of interest not described separately
Lehrnbecher 2006	not diagnosis of interest
Lessnick 2009	not study design of interest (review)
Liseth 2004	data of interest not described separately
Locatelli 2008	not intervention of interest (allogeneic HSCT)
Lorenz 1999	not study design of interest (review)
Lorigan 2007	not intervention of interest
Lucidarme 1998	not diagnosis of interest (rhabdomyosarcoma)
Mace 2003	not intervention of interest
Machado 2007	not diagnosis of interest
Mack 1995	not study design of interest (review)
Mackall 2001	not study design of interest (review)
Madero 1995	not diagnosis of interest (rhabdomyosarcoma)
Maeda 2008	not study design of interest (review)

Mankin 2004	not intervention of interest
Marina 1997	not study design of interest (review)
Matsubara 2003	not diagnosis of interest (rhabdomyosarcoma)
Matsuyama 2000	not study design of interest (review)
Matthews 2007	not diagnosis of interest
Medioni 2003	not intervention of interest
Mesia 1995	not diagnosis of interest (rhabdomyosarcoma)
Meyers 2004	not study design of interest (review)
Michailov 2001	not diagnosis of interest (rhabdomyosarcoma)
Michon 1999	not study design of interest (review)
Mikhailova 1998	not outcome of interest
Miliauskas 1993	not intervention of interest
Mimeault 2008	not study design of interest (review)
Minard-Colin 2004	not intervention of interest
Mingo 2005	not intervention of interest
Miyagi 2003	not diagnosis of interest (rhabdomyosarcoma)
Moore 2009	not diagnosis of interest (rhabdomyosarcoma)
Morikawa 2005	not intervention of interest
Munoz 1983	not diagnosis of interest (rhabdomyosarcoma)
Müller 2002	not intervention of interest
Nachbaur 1994	not diagnosis of interest
Nag 1995	not diagnosis of interest (rhabdomyosarcoma)
Nath 2005	not diagnosis of interest (rhabdomyosarcoma)
Nenadov 1995	not diagnosis of interest (rhabdomyosarcoma)

not study design of interest (review)
not study design of interest (review)
not study design of interest (review)
data of interest not described separately
not outcome of interest
not outcome of interest
not intervention of interest
not diagnosis of interest (rhabdomyosarcoma)
not study design of interest (review)
not diagnosis of interest (rhabdomyosarcoma)
not diagnosis of interest (rhabdomyosarcoma)
not diagnosis of interest (rhabdomyosarcoma)
not diagnosis of interest
not diagnosis of interest (rhabdomyosarcoma)
not diagnosis of interest (rhabdomyosarcoma)
not diagnosis of interest (rhabdomyosarcoma)
not intervention of interest
not diagnosis of interest (rhabdomyosarcoma)
data of interest not described separately
not study design of interest (review)
not study design of interest (review)
not study design of interest (review)
not diagnosis of interest (rhabdomyosarcoma)
not intervention of interest

Pedrazzoli 2006	not study design of interest (review)
Perentesis 1999	not diagnosis of interest (rhabdomyosarcoma)
Pession 1999	not diagnosis of interest (rhabdomyosarcoma)
Philip 1984	not study design of interest (review)
Pick 1988	not study design of interest (review)
Pico 1993	not diagnosis of interest (rhabdomyosarcoma)
Pinedo 1987	not intervention of interest
Pinkerton 1986	not study design of interest (review)
Pinkerton 1987	not study design of interest (review)
Pinkerton 1991	data of interest not described separately
Pinkerton 1991a	not diagnosis of interest (rhabdomyosarcoma)
Pinkerton 1995	not study design of interest (review)
Pohar-Marinsek 2001	not diagnosis of interest (rhabdomyosarcoma)
Pohar-Marinsek 2003	not diagnosis of interest (rhabdomyosarcoma)
Raben 1994	not intervention of interest
Radeva 2005	not study design of interest
Raja 2003	not intervention of interest
Raney 1997	not diagnosis of interest (rhabdomyosarcoma)
Raney 2001	not study design of interest (review)
Rapidis 2008	not diagnosis of interest
Ray-Coquard 2001	not study design of interest (review)
Recchia 1996	not diagnosis of interest
Recchia 2003	not diagnosis of interest
Reich 2001	data of interest not described separately

Reichardt 1997	not study design of interest (review)
Reichardt 2002	not study design of interest (review)
Rill 1994	not diagnosis of interest
Ritchie 2004	not diagnosis of interest (rhabdomyosarcoma)
Rivera-Luna 2001	not diagnosis of interest
Rodenhuis 1999	not study design of interest (review)
Roh 2001	not diagnosis of interest
Roman-Unfer 1996	not study design of interest (review)
Rosenberg 1981	not intervention of interest
Rosenberg 1982	not intervention of interest
Rosenberg 1983	not intervention of interest
Rosman 2008	data of interest not described separately
Rossbach 1999	not diagnosis of interest (rhabdomyosarcoma)
Rosti 2002	not study design of interest (review)
Rousselet 1994	not diagnosis of interest
Rubie 2003	not diagnosis of interest
Rzepecki 2006	not study design of interest (review)
Rzepecki 2006a	not study design of interest (review)
Saikawa 2006	not diagnosis of interest (rhabdomyosarcoma)
Sajedi 2002	not intervention of interest (allogeneic HSCT)
Sakayama 2008	not diagnosis of interest (rhabdomyosarcoma)
Salutari 1998	not diagnosis of interest (rhabdomyosarcoma)
Sanchez 1986	not study design of interest (review)

-	
Sanchez-Garcia 2007	not diagnosis of interest
Santana 1992	not diagnosis of interest (rhabdomyosarcoma)
Sanz 1997	not diagnosis of interest (rhabdomyosarcoma)
Sato 1998	not diagnosis of interest (rhabdomyosarcoma)
Sauer 1998	not study design of interest (review)
Sauer 1998a	not study design of interest (review)
Savasan 2005	not study design of interest (review)
Savolainen 2005	not diagnosis of interest (rhabdomyosarcoma)
Sawyer 1999	not study design of interest (review)
Schimmer 2002	data of interest not described separately
Schlemmer 2006	not disease of interest for 22 of 55 (40%) patients: 3 patients with rhabdomyosarcoma, 3 patients with peripheral neuroectodarmal tumor, and 16 patients with not identified tumors were included in a total of 55 analyzed patients
Schmidt 1994	not intervention of interest (allogeneic HSCT)
Schulz 1991	data of interest not described separately
Schuster 2008	data of interest not described separately
Schwella 1998	data of interest not described separately
Secondino 2007	not intervention of interest (allogeneic HSCT)
Seeger 1991	not study design of interest (review)
Segura 2001	not diagnosis of interest (rhabdomyosarcoma)
Seregard 2002	not diagnosis of interest (rhabdomyosarcoma)
Seynaeve 1999	not study design of interest
Shea 1995	not diagnosis of interest
Shen 1993	duplicate paper of Shen 1994

Shen 1994	not intervention of interest (allogeneic HSCT)
Shimizu 2008	not diagnosis of interest (rhabdomyosarcoma)
Shinkoda 2009	not diagnosis of interest (rhabdomyosarcoma)
Simon 2007	data of interest not described separately
Skinner 1974	not intervention of interest
Sola 1999	not diagnosis of interest
Somlo 1995	not intervention of interest
Spitzer 1980	not diagnosis of interest
Spitzer 1984	not diagnosis of interest
Spitzer 1994	not study design of interest (review)
Spitzer 1995	not diagnosis of interest
Spruce 1983	not study design of interest (review)
Stea 1987	not diagnosis of interest (rhabdomyosarcoma)
Steinbrenner 2005	data of interest not described separately
Stöhr 2006	not intervention of interest
Suita 2005	not diagnosis of interest (rhabdomyosarcoma)
Sussman 2008	not study design of interest (review)
Takata 1997	not intervention of interest
Takaue 2002	not diagnosis of interest (rhabdomyosarcoma)
Takenaka 2007	not intervention of interest
Tang 2009	not disease of interest (rhabdomyosarcoma)
Thomson 1999	not diagnosis of interest (rhabdomyosarcoma)
Toma 1992	not intervention of interest
Trigg 2002	not study design of interest (review)

Unal 2006	data of interest not described separately
Urban 1997	not outcome of interest
Urbano-Ispizua 2002	not outcome of interest
Vadhan 1996	not study design of interest (abstract)
Vadhan-Raj	not intervention of interest
Valkova 2003	not intervention of interest
Valteau-Couanet 2007	not study design of interest (review)
Valteau-Couanet 2007a	not diagnosis of interest
Van Dalen 2009	not intervention of interest
Van den Berg 2006	not study design of interest (review)
Van den Berg 2007	not study design of interest (review)
Van den Berg 2008	not intervention of interest
Van Glabbeke 1993	not intervention of interest
Varterasian 1997	not intervention of interest
Vassal 2005	not study design of interest (review)
Vaughan 2001	not study design of interest (review)
Verma 2002	not study design of interest (review)
Verma 2008	not study design of interest (review)
Verma 2008a	not study design of interest (review)
Wachowiak 2008	not outcome of interest
Walterhouse 1999	not diagnosis of interest (rhabdomyosarcoma)
Wasserman 1997	not diagnosis of interest
Watanabe 2006a	not study design of interest
Weaver 1997	data of interest not described separately

Weh 1995	not study design of interest (review)
Weh 1996	not intervention of interest
Weigel 2001	not study design of interest (review)
Werchniak 2005	not diagnosis of interest
Wexler 1996	not intervention of interest
Willenbacher 1998	data of interest not described separately
Williams 2004	not diagnosis of interest (rhabdomyosarcoma)
Womer 1996	not study design of interest (review)
Womer 2000	not study design of interest (review)
Woods 1999	not study design of interest (review)
Worden 2005	not intervention of interest
Yamada 2007	not diagnosis of interest (rhabdomyosarcoma)
Yaniv 1990	not study design of interest (review)
Yaniv 2000	not study design of interest (review)
Yaqoob 2006	not study design of interest (review)
Yin 2009	not intervention of interest
Young 1989	data of interest not described separately
Zoubek 1994	not intervention of interest

DATA AND ANALYSES

This review has no analyses.

ADDITIONAL TABLES

Table 1. Frequency of subtypes of included NRSTS in patients of young versus advanced age

Young age (< 20 years)		Advanced (≥ 20 years)	
Subtype	%	Subtype	%
Synovial sarcoma	7.7	Leiomyosarcoma	13.7
Malignant fibrous histiocytoma	4.9	Malignant fibrous histiocytoma	10.1
Fibrosarcoma	4.5	Liposarcoma	8.0
Liposarcoma	2.8	Hemangiosarcoma	2.5
Epitheloid sarcoma	2.0	Spindle cell sarcoma	2.3

^{*}according to Spunt 2006

Table 2. Included non-rhabdomyosarcoma soft tissue sarcomas (NRSTS)

Diagnosis(*)		
Alveolar soft part sarcoma		
Anaplastic soft tissue sarcoma		
Angiosarcoma 1. Angiosarcoma of soft tissue 2. Hemangiosarcoma 3. Hemangiopericytoma 4. Lymphangiosarcoma		
Clear cell myomelanocytic tumor		
Clear cell sarcoma of soft tissue		
Desmoplastic small round cell tumor		
Epithelioid sarcoma		

Table 2. Included non-rhabdomyosarcoma soft tissue sarcomas (NRSTS) (Continued)

Fibrosarcoma

- 1. Adult fibrosarcoma
- 2. Myxofibrosarcoma
- 3. Low grade fibromyxoid sarcoma; hyalinizing spindle cell tumor
- 4. Sclerosing epithelioid fibrosarcoma

Fibromyxoid sarcoma

Epithelioid hemangioendothelioma

Intimal sarcoma

Leiomyosarcoma

1. Leiomyosarcoma (excluding skin)

Liposarcoma

- 1. Dedifferentiated liposarcoma
- 2. Myxoid liposarcoma
- 3. Round cell liposarcoma
- 4. Pleomorphic liposarcoma
- 5. Mixed-type liposarcoma
- 6. Liposarcoma, not otherwise specified

Mesenchymal sarcoma

Malignant glomus tumor

Malignant fibrous histiocytoma

- 1. Pleomorphic malignant fibrous histiocytoma; undifferentiated pleomorphic sarcoma
- 2. Giant cell malignant fibrous histiocytoma; undifferentiated pleomorphic sarcoma with giant cells
- 3. Inflammatory malignant fibrous histiocytoma; undifferentiated pleomorphic sarcoma with prominent inflammation
- 4. Undifferentiated pleomorphic sarcoma
- 5. Spindle cell sarcoma

Malignant haemangiopericytoma

Malignant mesenchymoma

Neoplasms with perivascular epithelioid cell differentiation (PEComa)

Rhabdoid sarcoma

1. Extra-renal rhabdoid tumor

Synovial sarcoma

Unclassified sarcoma

Undifferentiated sarcoma

Table 3. Excluded tumor types

Diagnosis	Reason for exclusion(*)
Atypical teratoid/rhabdoid tumors	WHO classification of tumors of the central nervous system
Chondrosarcoma 1. Mesenchymal chondrosarcoma 2. Extraskeletal myxoid chondrosarcoma ('chordoid type')	Extraskeletal types are difficult to separate
Dermatofibrosarcoma protuberance	WHO classification of tumors of the skin
Endometrial stroma sarcoma	WHO classification of tumors: pathology and genetics of tumors of the breast and female genital organs
Ewing family of tumors 1. Ewing sarcoma 2. Skeletal Ewing's sarcoma 3. Extraskeletal Ewingsarcoma 4. Peripheral primitive neuroectodermal tumour (pPNET) 5. Extraskeletal peripheral primitive neuroectodermal tumor (pPNET) 6. Askin tumor	Extraskeletal types are difficult to separate; the Ewing family of tumors is one entity
Extragonadal germ cell sarcoma	WHO classification of tumors: pathology and genetics of tumors of the urinary system and male genital organs WHO classification of tumors: pathology and genetics of tumors of the breast and female genital organs
Follicular dendritic cell sarcoma	WHO classification of tumors of hematopoietic and lymphoid tissues
Ganglioneuroblastoma	WHO classification of nervous system tumors
Gastrointestinal stromal tumor	WHO classification of tumors: pathology and genetics of tumors of the digestive system
Giant cell fibroblastoma	WHO classification of tumors of the skin
Giant cell tumour of bone	WHO classification for tumors of bone tissue
Histiocytic sarcoma	WHO classification of tumors of hematopoietic and lymphoid tissues
Interdigitating dendritic cell sarcoma	WHO classification of tumors of hematopoietic and lymphoid tissues

^{*} category of malignant tumors according to the World Health Organization Classification of Tumours: Pathology and Genetics of Tumours of Soft Tissue and Bone (Fletcher 2002)

Table 3. Excluded tumor types (Continued)

Interdigitating reticulum cell sarcoma	WHO classification of tumors of hematopoietic and lymphoid tissues
Kaposi sarcoma	Intermediate malignancy (rarely metastasizing)
Lymphoblastic lymphosarcoma	WHO classification of tumors of hematopoietic and lymphoid tissues
Medulloblastoma	WHO classification of tumors of the central nervous system
Myeloid sarcoma	WHO classification of tumors of hematopoietic and lymphoid tissues
Myxosarcoma (cardiac tumor)	WHO classification of tumors: pathology and genetics of tumors of the lung, pleura, thymus and heart
Nephroblastoma	WHO classification of tumors: pathology and genetics of tumors of the urinary system and male genital organs
Neuroblastoma (Wilms tumor)	WHO classification of tumors: pathology and genetics of tumors of the urinary system and male genital organs
Osteosarcoma 1. Extraskeletal osteosarcoma	Extraskeletal types are difficult to separate
Peripheral nerve sheath tumor, malignant (neurofibrosarcoma)	WHO classification of nervous system tumors
Rhabdoid tumour, renal	WHO classification of tumors: pathology and genetics of tumors of the urinary system and male genital organs
Rhabdoid tumour, cerebral	WHO classification of tumors of the central nervous system
Rhabdomyosarcoma 1. Embryonal rhabdomyosarcoma (including spindle cell, botryoid, anaplastic) 2. Alveolar rhabdomyosarcoma (including solid, anaplastic) 3. Pleomorphic rhabdomyosarcoma 4. Undifferentiated rhabdomyosarcoma	A soft tissue sarcoma that is excluded to separate rhabdomyosarcomas from non-rhabdomyosarcoma soft tissue sarcomas
Schwannoma, malignant	WHO classification of nervous system tumors
Uterine endometrial stromal sarcoma	WHO classification of tumors: pathology and genetics of tumors of the breast and female genital organs

^{*} WHO: World Health Organization

Table 4. Literature sources and search steps

Category	Sources
Step 1	
Bibliographic databases	 MEDLINE via Ovid; via PubMed, includes Clinical Queries EMBASE via Ovid The Cochrane Library via Wiley InterScience Cochrane central register of controlled trials Cochrane database of systematic reviews (CDSR; Cochrane reviews) database of abstracts of reviews of effects (DARE; other reviews) health technology assessment database (HTA; technology assessments) National Health Services economic evaluation database (NHSEED; economic evaluations)
Step 2	
Online trial registers	 ClinicalTrials.gov (ClinicalTrials.gov 2010) International Standard Randomised Controlled Trial Number (ISRCTN 2010) Register National Institute for Health Research UK Clinical Research Network's (NIHR UKCRN 2010) Portfolio Database National Cancer Institute Physician Data Query (NCI PDQ 2010) Clinical Trials European Group for Blood and Marrow Transplantation Solid Tumor Working Party (EBMT STWP 2010) World Health Organization International Clinical Trials Registry Platform (ICTRP 2010)
Step 3	
Reviews	systematic reviews (rhabdomyosarcoma included): Admiraal 2007 (Cochrane Protocol); Verma 2008; Weigel 2001 narrative reviews (rhabdomyosarcoma included): 95 articles from 1983 to 2008
Step 4	
Congress proceedings	Blood (American Society of Hematology Annual Meeting Abstracts) 2004 to 2007
Step 5	
Institutions(*)	 Scientific Institute San Raffaele, Milan, Italy Istituto Nazionale dei Tumori, Milan, Italy Deutsche Gesellschaft für Hämatologie und Onkologie (DGHO), Berlin, Germany St. Jude Children's Research Hospital, Memphis, Tennessee, USA Universitätsklinikum Hamburg-Eppendorf, Hamburg, Germany National CancerCenter Hospital, Tokyo, Japan Memorial Sloan-Kettering Cancer Center, New York City, New York, USA European Group for Blood and Marrow Transplantation (EBMT), Leipzig, Germany Ospedale Niguarda Ca'Granda, Milano, Italy Universitätsklinikum Charite, Berlin, Germany Medizinische Universitätsklinik, Ulm, Germany Fred Hutchinson Cancer Research Center, Seattle, Washington, USA

Table 4. Literature sources and search steps (Continued)

	13. Italian sarcoma Group (ISG), Bologna, Italy
Step 6	
Authors(*)	Blay 2000; Ivanova 2007; Kasper 2007; Kasper 2009; Schlemmer 2006; Simon 2007; Suita 2005

^{*} direct enquiries by e-mail and post

Table 5. Assessment of risks of bias

Type of comparative study	Randomized controlled intervention trial	Non-randomized comparative intervention studies • non-randomized controlled clinical intervention trial • prospective cohort study • retrospective cohort study • case-control study		
Assignment of patients to treatment groups	Was the allocation sequence adequately generated? • yes • e.g. participants assigned to treatments on basis of a computergenerated random sequence or a table of random numbers • unclear • e.g. not reported, information not available An answer no would mark a non-randomized study and, therefore, this option is not provided	Were relevant details of criteria for assignment of patients to treatment groups provided? • yes • e.g. participants assigned alternating to treatments on basis of date of birth, clinic id-number or surname, at the discretion of the responsible physician, or no attempt to randomise participants • no • unclear		
Concurrent control	not applicable	Were data of the control group collected during the same time period as data of the test group? • yes • no • historical control data collected earlier than for the test group • unclear		
Concealment of allocation	Was allocation adequately concealed? • yes • e.g. where the allocation sequence could not be foretold • no • e.g. allocation sequence could be foretold by patients, investigators or	not applicable		

Table 5. Assessment of risks of bias (Continued)

	treatment providers • unclear • e.g. not reported
Comparable baseline characteristics	Were the two treatment groups comparable? Were the groups balanced in respect to confounders? Were there no differences of baseline characteristics between the two treatment groups or were differences controlled for, in particular with reference to prognostic factors, such as, age, gender, histological diagnosis, year of transplantation? • yes • groups were comparable or differences between groups were considered (e.g. adjusted for), or factors were matched and groups were balanced in respect to confounders • no • if the two groups differed and differences were not controlled for • unclear
Loss to follow-up	Was loss to follow-up less than 20% and were the reasons for loss to follow-up similar in both arms? • yes • no • unclear
Selective outcome reporting	Are reports of the study free of suggestion of selective outcome reporting? • yes • no • e.g if protocol reports all outcomes specified in the protocol • unclear
Other causes for high risk of bias	Was the study apparently free of other problems that could put it at a high risk of bias? • yes • no • unclear

Table 6. Overall survival (OS) and progression-free survival (PFS)

Study	Follow up started at	os	PFS			
		2 years	2 years			
Controlled trials (HSC	Γ after HDCT versus s	tandard-dose chemotherapy)				
Ivanova 2007	unclear	62.3% vs. 23.2%*	-			
No control group (only patients with HSCT after HDCT)						
Bertuzzi 2003 "therapy" $20\%^{\dagger}$ $0\%^{\dagger}$						
Pooled meta-analysis of individual data (only patients with HSCT after HDCT)						

Table 6. Overall survival (OS) and progression-free survival (PFS) (Continued)

|--|

^{* &}quot;statistically significant", no detailed description of statistical method

Table 7. Treatment-related mortality (TRM) in transplanted patients

Study	TRM, n	Cause of death				
No control group	No control group, individual data (only patients with HSCT)					
Doros 2008	1	not specified				
Engelhardt 2007	3	(1) sepsis; (2) sepsis; (3) pulmonary metastases, pneumonia, respiratory failure				
Kasper 2007	1	cardiac arrest of unknown origin				
Navid 2006	1	hepatic and renal failure				
Saab 2007	2	(1) acute myocardial infarction; (2) veno-occlusive disease				
Shaw 1996	1	veno-occlusive disease and necrotising interstitial pneumonitis				
Slease 1988	2	(1) progressive encephalopathy; (2) sepsis				

Table 8. Secondary neoplasia in transplanted patients

Study	Secondary neoplasia, n Diagnosis						
No control group, individual data (only patients with HSCT)							
Yamamura 2003 1 chronic myelogenous leukemia							

Table 9. Toxicity, National Cancer Institute (NCI) common terminology criteria for adverse events (CTCAE) grade 3-4, in transplanted patients

Study	· ·	Hematological toxicity (Number of affected / total patients)		Non-hematological toxicity (Number of affected / total patients)				
	Leukopenia	Neutrope- nia	Throm- bopenia	Nausea	Kidney	Liver	Nervous system	Heart
No control group, individual data (only patients with HSCT)								

 $^{^{\}dagger}$ reading from Kaplan-Meier curve

Table 9. Toxicity, National Cancer Institute (NCI) common terminology criteria for adverse events (CTCAE) grade 3-4, in transplanted patients (Continued)

Kasper 2007	14 / 14	14 / 14	14 / 14	-	-	-	-	-
Kozuka 2002	-	1 / 1	1 / 1	1 / 1	-	-	-	-
Kushner 2001	-	-	-	-	-	-	1/1	-
Patel 2004	-	-	-	-	1 / 1	1 / 1	-	-
Yonemoto 1999	-	-	-	-	-	1 / 4	-	-

APPENDICES

Appendix I. MEDLINE/Ovid search strategy

- 1. exp SARCOMA/
- 2. (sarcom\$ or sarkom\$).mp.
- 3. exp LIPOSARCOMA/
- 4. liposar#om\$.mp.
- 5. exp FIBROSARCOMA/
- 6. fibrosar#om\$.mp.
- 7. exp HISTIOCYTOMA, MALIGNANT FIBROUS/
- 8. malign\$ fibrous histio#ytom\$.mp.
- 9. exp LEIOMYOSARCOMA/
- 10. leiomyosar#om\$.mp.
- 11. malign\$ glom\$ tumo\$.mp.
- 12. exp RHABDOMYOSARCOMA/
- 13. rhabdomyosar#om\$.mp.
- 14. exp HEMANGIOENDOTHELIOMA/
- 15. (hemangioendotheliom\$).mp.
- 16. exp HEMANGIOSARCOMA/
- 17. (angiosar#om\$ or hemangiosar#om\$ or haemangiosar#om\$).mp.
- 18. exp SARCOMA, SYNOVIAL/
- 19. synovia\$ sar#om\$.mp.
- 20. epithelioid sar#om\$.mp.
- 21. exp SARCOMA, ALVEOLAR SOFT PART/
- 22. (alveolar soft part sar#om\$ or alveolar soft tissue sar#om\$).mp.
- 23. exp SARCOMA, CLEAR CELL/
- 24. clear cell sar#om\$.mp.
- 25. exp SARCOMA, SMALL CELL/
- 26. (desmoplastic and small round cell tumo\$ or small cell tumo\$)).mp.
- 27. exp RHABDOID TUMOR/
- 28. ((extrarenal or extra-renal) and rhabdoid tumo\$).mp.

- 29. (malignan\$ and mesenchymom\$).mp.
- 30. clear cell myomelano#ytic tumo\$.mp.
- 31. intima\$ sar#om\$.mp.
- 32. exp STEM CELL TRANSPLANTATION/
- 33. exp BONE MARROW TRANSPLANTATION/
- 34. exp TRANSPLANTATION, AUTOLOGOUS/
- 35. exp TRANSPLANTATION, HOMOLOGOUS/
- 36. exp TRANSPLANTATION, CONDITIONING/
- 37. (autolog\$ hemato\$ or autolog\$ stem cell or autolog\$ bone marrow or autolog\$ periph\$ or autolog\$ transplant\$ or autolog\$ graft\$ or autotransplant\$ or autotransplant\$ or autograft\$ or auto-graft\$).mp.
- 38. (homolog\$ hemato\$ or homolog\$ haemato\$ or homolog\$ stem cell or homolog\$ bone marrow or homolog\$ cord or homolog\$ umbilical or homolog\$ peripheral or homolog\$ transplant\$ or homolog\$ graft\$ or homolog\$ transplant\$).mp.
- 39. (stem cell transplant\$ or bone marrow transplant\$ or periph\$ blood stem cell or periph\$ stem cell or cord blood transplant\$).mp.
- 40. (reduced intens\$ or myeloablat\$ or nonmyeloablat\$ or non-myeloablat\$).mp.
- 41. high dose chemotherapy.mp.
- 42. or/1-31
- 43. or/32-41
- 44. and/42-43
- 45. (ANIMALS not (ANIMALS and HUMANS)).sh.
- 46. 44 not 45

Appendix 2. EMBASE/Ovid search strategy

- 1. exp SARCOMA/
- 2. (sarcom\$ or sarkom\$).mp.
- 3. exp LIPOSARCOMA/
- 4. liposar#om\$.mp.
- 5. exp FIBROSARCOMA/
- 6. fibrosar#om\$.mp.
- 7. exp MALIGNANT FIBROUS HISTIOCYTOMA/
- 8. malign\$ fibrous histio#ytom\$.mp.
- 9. exp LEIOMYOSARCOMA/
- 10. leiomyosar#om\$.mp.
- 11. malign\$ glom\$ tumo\$.mp.
- 12. exp RHABDOMYOSARCOMA/
- 13. rhabdomyosar#om\$.mp.
- 14. exp HEMANGIOENDOTHELIOMA/
- 15. (hemangioendotheliom\$).mp.
- 16. exp HEMANGIOENDOTHELIOSARCOMA/
- 17. (hemangioendotheliosar#om\$ or haemangioendotheliosar#om\$).mp.
- 18. exp ANGIOSARCOMA/
- 19. angiosar#om\$.mp.
- 20. exp SYNOVIAL SARCOMA/
- 21. synovia\$ sar#om\$.mp.
- 22. exp EPITHELIOID SARCOMA/
- 23. (epithelioid\$ sar#om\$ or epitheloid\$ sar#om\$).mp.
- 24. exp ALVEOLAR SOFT PART SARCOMA/
- 25. (alveolar soft part sar#om\$ or alveolar soft tissue sar#om\$).mp.
- 26. exp CLEAR CELL SARCOMA/
- 27. clear cell sar#om\$.mp.
- 28. exp DESMOPLASTIC SMALL ROUND CELL TUMOR/
- 29. exp SMALL CELL SARCOMA/

- 30. (desmoplastic and (small round cell tumo\$ or small cell tumo\$)).mp.
- 31. ((extrarenal\$ or extra-renal\$) and rhabdoid\$ tumo\$).mp.
- 32. (malign\$ and mesenchymom\$).mp.
- 33. clear cell myomelano#yt\$ tumo\$.mp.
- 34. intima\$ sar#om\$.mp.
- 35. exp STEM CELL TRANSPLANTATION/
- 36. exp BONE MARROW TRANSPLANTATION/
- 37. exp NONMYELOABLATIVE STEM CELL TRANSPLANTATION/
- 38. exp NONMYELOBLATIVE CONDITIONING/
- 39. exp REDUCED INTENSITY CONDITIONING/
- 40. exp MYELOABLATIVE CONDITIONING/
- 41. (autolog\$ hemato\$ or autolog\$ stem cell or autolog\$ bone marrow or autolog\$ periph\$ or autolog\$ transplant\$ or autolog\$ graft\$ autotransplant\$ or auto-transplant\$ or autograft\$ or auto-graft\$).mp.
- 42. (homolog\$ hemato\$ or homolog\$ haemato\$ or homolog\$ stem cell or homolog\$ bone marrow or homolog\$ cord or homolog\$ umbilical or homolog\$ periph\$ or homolog\$ transplant\$ or homolog\$ graft\$).mp.
- 43. (stem cell transplant\$ or bone marrow transplant\$ or periph\$ blood stem cell or periph\$ stem cell or cord blood transplant\$).mp.
- 44. (reduced intens\$ or myeloablat\$ or non-myeloablat\$).mp.
- 45. high dose chemotherapy.mp.
- 46. or/1-34
- 47. or/35-45
- 48. and/46-47
- 49. (ANIMALS not (ANIMALS and HUMANS)).sh.
- 50. 48 not 49

Appendix 3. Cochrane/Wiley search strategy

- 1. exp SARCOMA/
- 2. (sarcom\$ or sarkom\$).mp.
- 3. exp LIPOSARCOMA/
- 4. liposar#om\$.mp.
- 5. exp FIBROSARCOMA/
- 6. fibrosar#om\$.mp.
- 7. exp HISTIOCYTOMA, MALIGNANT FIBROUS/
- 8. malign\$ fibrous histio#ytom\$.mp.
- 9. exp LEIOMYOSARCOMA/
- 10. leiomyosar#om\$.mp.
- 11. malign\$ glom\$ tumo\$.mp.
- 12. exp RHABDOMYOSARCOMA/
- 13. rhabdomyosar#om\$.mp.
- 14. exp HEMANGIOENDOTHELIOMA/
- 15. (hemangioendotheliom\$).mp.
- 16. exp HEMANGIOSARCOMA/
- 17. (angiosar#om\$ or hemangiosar#om\$).mp.
- 18. exp SARCOMA, SYNOVIAL/
- 19. synovia\$ sar#om\$.mp.
- 20. (epithelioid sar#om\$).mp.
- 21. exp SARCOMA, ALVEOLAR SOFT PART/
- 22. (alveolar soft part sar#om\$ or alveolar soft tissue sar#om\$).mp.
- 23. exp SARCOMA, CLEAR CELL/
- 24. clear cell sar#om\$.mp.
- 25. exp SARCOMA, SMALL CELL/
- 26. (desmoplastic and (small round cell tumo\$ or small cell tumo\$)).mp.

- 27. exp RHABDOID TUMOR/
- 28. ((extrarenal or extra-renal) and rhabdoid tumo\$).mp.
- 29. (malignan\$ and mesenchymom\$).mp.
- 30. clear cell myomelano#ytic tumo\$.mp.
- 31. intima\$ sar#om\$.mp.
- 32. exp STEM CELL TRANSPLANTATION/
- 33. exp BONE MARROW TRANSPLANTATION/
- 34. exp TRANSPLANTATION, AUTOLOGOUS/
- 35. exp TRANSPLANTATION, HOMOLOGOUS/
- 36. exp TRANSPLANTATION, CONDITIONING/
- 37. (autolog\$ hemato\$ or autolog\$ stem cell or autolog\$ bone marrow or autolog\$ periph\$ or autolog\$ transplant\$ or autolog\$ graft\$ or autotransplant\$ or auto-transplant\$ or autograft\$ or auto-graft\$).mp.
- 38. (homolog\$ hemato\$ or homolog\$ stem cell or homolog\$ bone marrow or homolog\$ cord or homolog\$ umbilical or homolog\$ periph\$ or homolog\$ transplant\$ or homolog\$ graft\$).mp.
- 39. (stem cell transplant\$ or bone marrow transplant\$ or periph\$ blood stem cell or periph\$ stem cell or cord blood transplant\$).mp.
- 40. (reduced intens\$ or myeloablat\$ or non-myeloablat\$).mp.
- 41. high dose chemotherapy.mp.
- 42. or/1-31
- 43. or/32-41
- 44. and/42-43
- 45. (ANIMALS not (ANIMALS and HUMANS)).sh.
- 46. 44 not 45

Appendix 4. Reviews (n = 98) checked for additional studies

Bone-marrow autotransplantation in man. Lancet 1986;2:960-2.

Admiraal R, Van der Paardt M, Kobes J, Kremer LCM, Bisogno G, Merks JHM. High dose chemotherapy for children with stage IV rhabdomyosarcoma. [Protocol]. Cochrane Database of Systematic Reviews 2007, Issue 3

Antman K, Eder JP, Frei E. High-dose chemotherapy with bone marrow support for solid tumors. Important Advances in Oncology 1987;221-35

Antman KH. New biology and therapies in soft tissue sarcomas. Biomedicine & Pharmacotherapy 2001;55(9-10):553-7

Atra A, Pinkerton R. Autologous stem cell transplantation in solid tumours of childhood. Annals of Medicine 1996;28(2):159-64

Atra A, Pinkerton R. High-dose chemotherapy in soft tissue sarcoma in children. Critical Reviews in Oncology/Hematology 2002;41 (2):191-6

Banna GL, Simonelli M, Santoro A. High-dose chemotherapy followed by autologous hematopoietic stem-cell transplantation for the treatment of solid tumors in adults: A critical review. Current Stem Cell Research and Therapy 2007;2:65-82

Bickert BM. Treatment of common childhood malignancies. Journal of Pharmacy Practice 2002;15(1):42-51

Casado HA, Moreno AF. Chemotherapy of soft tissue sarcomas. Revisiones en Cancer 2004;18(6):316-29

Chen AR. High-dose therapy with stem cell rescue for pediatric solid tumors: rationale and results. Pediatric Transplantation 1999;3 (Suppl 1):78-86

Childs RW. Evolving trends in hematopoietic cell transplantation for solid tumors: Tempering enthusiasm with clinical reality. Annals of Oncology 2004;15(4):543-4

Chuman H. Evidence-based chemotherapy for patients with bone and soft part sarcoma [Japanese]. Gan to Kagaku Ryoho [Japanese Journal of Cancer and Chemotherapy] 2000;27(2):192-202

Dallorso S, Manzitti C, Morreale G, Faraci M. High dose therapy and autologous hematopoietic stem cell transplantation in poor risk solid tumors of childhood. Haematologica 2000;85:66-70

De Terlizzi M, Philip T, Toma MG, Colella R, Ceci A. Massive therapy and transplant of autologous bone marrow in childhood lymphomas and solid tumors: State of art and future perspectives [Terapia massiva e trapianto di midollo autologo nei linfomi e tumori solidi pediatrici. Stato dell'arte e prospettive]. La Pediatria Medica e Chirurgica 1988;10:359-364

Devalck C, Ferster A, De Laet MH, Nafa S, Bujan W, Azzi N, et al. Autologous bone marrow graft in solid tumors in childhood [Greffe de moelle autologue dans les tumeurs solides des enfants]. Revue Medicale de Bruxelles 1992;13:201-6

Dicke KA, Spitzer G. Evaluation of the use of high-dose cytoreduction with autologous marrow rescue in various malignancies. Transplantation 1986;41(1):4-20

Dicke KA, Jagannath S, Spitzer G, Poynton C, Zander A, Vellekoop L, et al. The role of autologous bone marrow transplantation in various malignancies. Seminars in Hematology 1984;21(2):109-22

Dileo P, Demetri GD. Update on new diagnostic and therapeutic approaches for sarcomas. Clinical Advances in Hematology and Oncology 2005;3(10):781-91

Ek ETH, Choong PFM. The role of high-dose therapy and autologous stem cell transplantation for pediatric bone and soft tissue sarcomas. Expert Review of Anticancer Therapy 2006;6(2):225-37

Elias AD. High-dose therapy for adult soft tissue sarcoma: dose response and survival. Seminars in Oncology 1998;25(2 Suppl 4): 19-23

Endo M, Tanosaki R. Myeloablative chemotherapy with autologous bone marrow and/or peripheral blood stem cell transplantation in children with high-risk solid tumor. Gan To Kagaku Ryoho 1995;22:1762-70

Gadner H. Is there evidence-based benefit of autologous stem cell transplantation in children with solid tumors? Onkologie 2002;25: 278-81

Gentet JC, Plouvier E, Coze C. Bone marrow autograft and cancer in children [Autogreffes de moelle osseuse et cancers pediatriques]. La Revue du Praticien 1993;43(17):2213-7

Goto T, Kosaku H, Kobayashi H, Hozumi T, Kondo T. Soft tissue sarcoma: postoperative chemotherapy. Gan To Kagaku Ryoho 2004;31:1324-30

Gratwohl A. Activity Survey and Historical Perspective of Autologous Stem Cell Transplantation in Europe. Seminars in Hematology 2007;44:220-6

Gratwohl A, Schmid O, Baldomero H, Horisberger B, Urbano-Ispizua A, Accreditation Committee of the European Group for Blood and Marrow Transplantation. Haematopoietic stem cell transplantation (HSCT) in Europe 2002. Bone Marrow Transplantation 2004;34:855-75

Gratwohl A, Baldomero H, Demirer T, Rosti G, Dini G, Ladenstein R, et al. Hematopoetic stem cell transplantation for solid tumors in Europe. Annals of Oncology 2004;15:653-60

Gratwohl A. Overview of transplant activity in Europe. The Hematology Journal 2004;5(Suppl 3):S29-S33

Gratwohl A, Baldomero H, Frauendorfer K, Urbano-Ispizua A, Niederwieser D. Results of the EBMT activity survey 2005 on haematopoietic stem cell transplantation: Focus on increasing use of unrelated donors. Bone Marrow Transplantation 2007;39:71-87

Hale GA. Autologous hematopoietic stem cell transplantation for pediatric solid tumors. Expert Review of Anticancer Therapy 2005;5 (5):835-46

Herzog CE. Sarcomas in adolescents and young adults: A summary of a recent symposium. Journal of Pediatric Hematology/Oncology 2005;27(4):177-8

Hoekstra HJ, Schraffordt KH, Oldhoff J. Soft tissue sarcoma of the extremity. European Journal of Surgical Oncology 1994;20:3-6

Höffken K, Kath R, Fricke HJ, Blumenstengel K, Vogel W, Sayer HG. High dose chemotherapy of solid tumors [Hochdosis-chemotherapie bei soliden Tumoren]. Medizinische Klinik 1997;92(7):410-4

Imanguli MM, Childs RW. Hematopoietic stem cell transplantation for solid tumors. Update on Cancer Therapeutics 2006;1:343-52

Irle C. Massive chemotherapy of solid tumors with bone marrow transplantation [Article in French: Chimiotherapie lourde avec greffe de moelle dans les tumouers solides]. Medecine et Hygiene 1989;47(1816):3377-82

Issels R. Knochentumoren und Weichteilsarkome: Empfehlungen zur Diagnostik, Therapie und Nachsorge. München: Zuckschwerdt, 2004

Issels RD. Soft tissue sarcomas--what is currently being done. European Journal of Surgical Oncology 1995;21:471-4

Kaizer H, Chow HS. Autologous bone marrow transplantation (ABMT) in the treatment of cancer. Cancer Investigation 1984;2: 203-13

Kasper B, Ho AD, Egerer G. Is there an indication for high-dose chemotherapy in the treatment of bone and soft-tissue sarcoma? Oncology 2005;68:115-21

Kavan P, Koutecky J. Current results with myeloablative therapy followed by hematopoietic stem cell rescue in pediatric solid tumors [Soucasny pohled na myeloablativni lecbu s naslednou transplantaci hematopoetickych kmenovych bunek u detskych solidnich nadoru]. Klinicka Onkologie 1997;10(4):106-9

Kletzel M, Kim AR. Autologous bone marrow transplantation in pediatric solid tumors. Cancer Treatment and Research 1997;77: 333-56

Kletzel M, Hewlett B. Pediatric transplantation: results in solid tumors. Current Hematology Reports 2005;4:260-9

Klingebiel T, Handgretinger R, Niethammer D. Autologous bone marrow transplantation [Autologe Knochenmarktransplantation]. Infusionstherapie und Transfusionsmedizin 1994;21(Suppl 3):42-5

Klingebiel T, Dopfer R, Handgretinger R, Niethammer D. Indications for autologous bone marrow transplantation in pediatric oncology. Results of the 5th meeting of experts of the Kind-Philipp Foundation, Riesensburg, November 1988 [Indikation zur autologen Knochenmarktransplantation in der Padiatrischen Onkologie. Ergebnisse der 5. Expertentagung der Kind-Philipp-Stiftung, Reisensburg, November 1988]. Klinische Pädiatrie 1989;201:304-10

Koscielniak E. Therapy for soft tissue sacrcoma: more questions than answers? [Article in German: Therapie der Weichteilsarkome: mehr Fragen offen als beantwortet?]. Wiener Klinische Wochenschrift 2005;117(5-6):176-79

Koscielniak E. Soft tissue sarcoma in children. Diagnosis and therapeutic modalities. Advances in Clinical and Experimental Medicine 2001;10:3-8

Koscielniak E. The role of high dose therapy (HDC) with stem cell rescue in the treatment of high-risk rhabdomyosarcoma. Rivista Italiana di Pediatria 1999;25:106-8

Ladenstein R, Philip T, Gardner H. Autologous stem cell transplantation for solid tumors in children. Current Opinion in Pediatrics 1997;9:55-69

Ladenstein R, Hartmann O, Pinkerton CR. The role of megatherapy with autologous bone marrow rescue in solid tumours of childhood. Annals of Oncology 1993;4(Suppl 1):45-58

Lorenz F, Skotnicki AB. Autotransplantation for solid tumors [Article in Polish: Autotransplantacja w guzach litych]. Przeglad Lekarski 1999;56(Suppl 1):101-7

Mack TM. Sarcomas and other malignancies of soft tissue, retroperitoneum, peritoneum, pleura, heart, mediastinum and spleen. Cancer 1995;75:211-44

Mackall CL, Helman LJ. High-dose chemotherapy for rhabdomyosarcoma: where do we go from here. Journal of Pediatric Hematology/Oncology 2001;23:266-7

Marina NM. Biology and treatment of pediatric malignant solid tumors. Cancer Chemotherapy and Biological Response Modifiers 1997;17:642-71

Matsuyama T. Autologous bone marrow transplantation for pediatric malignancies. Biotherapy (Tokyo) 2000;14:207-42

Meyers PA. High-dose therapy with autologous stem cell rescue for pediatric sarcomas. Current Opinion in Oncology 2004;16:120-5

Michon J, Schleiermacher G. Autologous haematopoietic stem cell transplantation for paediatric solid tumours. Bailliere's Best Practice & Research. Clinical Haematology 1999;12:247-59

Mimeault M, Batra SK. Targeting of cancer stem/progenitor cells plus stem cell-based therapies: the ultimate hope for treating and curing aggressive and recurrent cancers. Panminerva Medica 2008;50:3-18

Nieboer P, de Vries EGE, Mulder NH, van der Graaf WTA. Relevance of high-dose chemotherapy in solid tumours. Cancer Treatment Reviews 2005;31:210-25

Nieto Y, Shpall EJ. Autologous stem-cell transplantation for solid tumors in adults. Hematology/Oncology Clinics of North America 1999;13:939-68

Nieto Y, Jones RB, Shpall EJ. Stem-cell transplantation for the treatment of advanced solid tumors. Springer Seminars in Immunopathology 2004;26:31-56

Oeffinger KC, Nathan PC, Kremer LCM. Challenges after curative treatment for childhood cancer and long-term follow up of survivors. Pediatric Clinics of North America 2008;55(1):251-73

Pasetto LM, Basso U, Brandes AA. Improved tolerability of chemotherapy in soft tissue sarcomas: old and new strategies. Expert Review of Anticancer Therapy 2003;3:167-78

Patel S, Benjamin RS. Standard and high dose chemotherapy for advanced soft tissue sarcomas. Annals of Oncology 1992;3(Suppl 2):S81-3

Pedrazzoli P, Ledermann JA, Lotz JP, Leyvraz S, Aglietta M, Rosti G, et al. High dose chemotherapy with autologous hematopoietic stem cell support for solid tumors other than breast cancer in adults. Annals of Oncology 2006;17:1479-88

Pick TE. Autologous bone marrow transplantation in children. Critical Reviews in Oncology/Hematology 1988;8:311-37

Pinkerton CR. Intensive chemotherapy with stem cell support-experience in pediatric solid tumours. Bulletin du Cancer 1995;82 (Suppl 1):61s-5s

Pinkerton R, Philip T, Bouffet E, Lashford L, Kemshead J. Autologous bone marrow transplantation in paediatric solid tumours. Clinics in Haematology 1986;15:187-203

Pinkerton R, Philip T. Autologous bone marrow transplantation in paediatric solid tumours. Haematology and Blood Transfusion 1987;31:92-6

Rajic L. Hematopoietic stem cell transplantation in children with solid tumors. Paediatria Croatica 2003;(Suppl):103-6

Raney RB anderson JR, Barr FG, Donaldson SS, Pappo AS, Qualman SJ, et al. Rhabdomyosarcoma and undifferentiated sarcoma in the first two decades of life: a selective review of intergroup rhabdomyosarcoma study group experience and rationale for Intergroup Rhabdomyosarcoma Study V. Journal of Pediatric Hematology/Oncology 2001;23:215-20

Ray-Coquard I, Biron P, Blay JY. High-dose chemotherapy in soft tissue sarcomas of adults [Article in French: Chimiotherapie a hautes doses dans les sarcomes des tissus mous de l'adulte]. Bulletin du Cancer 2001;88:858-62

Reichardt P. High-dose chemotherapy in adult soft tissue sarcoma. Critical Reviews in Oncology/Hematology 2002;41:157-67

Rodenhuis S, de Vries EG. High-dose chemotherapy with stem cell support for solid tumors in adults [Hooggedoseerde chemotherapie met stamcelondersteuning bij solide tumoren van volwassenen]. Nederlands Tijdschrift voor Geneeskunde 1999;143:731-8

Rosti G, Ferrante P, Ledermann J, Leyvraz S, Ladenstein R, Koscileniak E, et al. High-dose chemotherapy for solid tumors: results of the EBMT. Critical Reviews in Oncology/Hematology 2002;41:129-40

Rzepecki P, Sarosiek T, Deptala A, Szczylik C. Autologous hematopoietic cell transplantation in adult patients with certain solid tumors [Przeszczepianie autologicznych krwiotworczych komorek macierzystych w leczeniu wybranych guzow litych u dorosłych]. Acta Haematologica Polonica 2006;37:159-66

Rzepecki P, Sarosiek T, Szczylik C. Autologous hematopoietic cell transplantation in adult patients with germ cell tumors and soft tissue sarcomas [Article in Polish: Rola przeszczepienia autologicznych krwiotworczych komorek macierzystych w leczeniu guzow zarodkowych oraz miesakow tkanek miekkich]. Wspolczesna Onkologia 2006;10:7-12

Sanchez-Garcia I, Vicente-Duenas C, Cobaleda C. The theoretical basis of cancer-stem-cell-based therapeutics of cancer: Can it be put into practice? BioEssays 2007;29:1269-80

Sauer H. Adjuvant chemotherapy in early soft tissue sarcoma and palliative chemotherapy in advanced soft tissue sarcoma in adults [Article in German: Adjuvante Chemotherapie bei lokoregional begrenzten Weichteilsarkomen und palliative Chemotherapie bei fortgeschrittenen Weichteilsarkomen im Erwachsenenalter]. Schweizerische Rundschau für Medizin Praxis 1998;87:1066-71

Sauer M, Gruhn B, Fuchs D, Altermann W, Zintl F. Heparin-induced type II thrombocytopenia within the scope of high dose chemotherapy with subsequent stem cell rescue [Heparin-induzierte Thrombozytopenie Typ II im Rahmen einer Hochdosis-Chemotherapie mit anschliessender Stammzellrescue]. Klinische Pädiatrie 1998;210:102-5

Savasan S, Abella EM. Current issues in pediatric stem cell transplantation. Clinics in Laboratory Medicine 2005;25:519-40

Seeger RC, Reynolds CP. Treatment of high-risk solid tumors of childhood with intensive therapy and autologous bone marrow transplantation. Pediatric Clinics in North America 1991;38:393-424

Spitzer G, Dunphy FR, Bowers CE, Adkins DR. High-dose therapy with stem cell support in solid tumors. Medical Oncology 1994;11:53-62

Spruce WE. Bone marrow transplantation. The American Journal of Pediatric Hematology/Oncology 1983;5:287-94

Trigg ME. Milestones in the development of pediatric hematopoietic stem cell transplantation--50 years of progress. Pediatric Transplantation 2002;6:465-74

Valteau-Couanet D, Dufour C, Hartmann O. High-dose chemotherapy and autologous stem cell transplantation in treating paediatric malignancies [Article in French: Chimiotherapie a hautes doses et autogreffe en oncologie pediatrique]. Oncologie 2007;9:827-31

van den Berg H. Biology and treatment of malignant solid tumors in childhood. Update on Cancer Therapy 2007;177-91

van den Berg H. Biology and therapy of solid tumors in childhood. Update on Cancer Therapy 2006;1:367-83

Vassal G. Has chemotherapy reached its limits in pediatric cancers? European Journal of Cancer 2005;41:564-75

Verma S, Bramwell V. Dose-intensive chemotherapy in advanced adult soft tissue sarcoma. Expert Review on Anticancer Therapy 2002;2:201-15

Verma S, Younus J, Stys-Norman D, Haynes AE, Blackstein M and the members of the Sarcoma Disease Site Group of Cancer Care Ontario's Program in Evidence-based Care. Dose-intensive chemotherapy with growth factor or autologous bone marrow/stem cell transplant support in first-line treatment of advanced or metastatic adult soft tissue sarcoma: a systematic review. Cancer 2008;112: 1197-205

Weh HJ, Hossfeld DK. Systemic therapy of disseminated soft tissue sarcomas. Recent Results in Cancer Research 1995;138:147-59

Weigel BJ, Breitfeld PP, Hawkins D, Crist WM, Baker KS. Role of high-dose chemotherapy with hematopoietic stem cell rescue in the treatment of metastatic or recurrent rhabdomyosarcoma. Journal of Pediatric Hematology/Oncology 2001; 23: 272-276

Womer RB. Problems and controversies in the management of childhood sarcomas. British Medical Bulletin 1996;52:826-43

Womer RB, Pressey JG. Rhabdomyosarcoma and soft tissue sarcoma in childhood. Current Opinion in Oncology 2000;12:337-44

Woods WG. Myeloablative therapy followed by stem cell rescue for pediatric solid tumors: A non-transplanter's perspective. Cancer Research Therapy and Control 1999;9:95-9

Yaniv I, Bouffet E, Irle C, Negrier S, Biron P, Favrot M, et al. Autologous bone marrow transplantation in pediatric solid tumors. Pediatric Hematology and Oncology 1990;7:35-46

Yaniv I. Lymphokines post autologous peripheral blood stem cell transplantation in children. Pediatric Hematology and Oncology 2000;17:9-13

Yaqoob N, Hasan SH. Desmoplastic small round cell tumor. Journal of the College of Physicians and Surgeons - Pakistan 2006;16: 614-6

HISTORY

Protocol first published: Issue 1, 2010 Review first published: Issue 2, 2011

CONTRIBUTIONS OF AUTHORS

FP: designing and coordinating the review, data collection for the review, designing search strategies, undertaking searches, screening search results, organizing retrieval of papers, screening retrieved papers against eligibility criteria, appraising quality of papers, extracting data from papers, writing to authors of papers for additional information, data management for the review, entering data into RevMan, analysis of data, interpretation of data, writing the review and the protocol.

LAS: providing methodological advice, screening included papers to verify data, interpretation of data, writing the review and the protocol

MaKr: analysis of data, interpretation of data, appraising quality of papers

CB: screening retrieved papers against eligibility criteria, extracting data from papers

NK: providing a clinical perspective

MiKu: appraising quality of papers, interpretation of data, providing a methodological perspective

DECLARATIONS OF INTEREST

The authors declare that they have no competing interests.

SOURCES OF SUPPORT

Internal sources

• IQWiG Institute of Quality and Efficiency in Health Care, Germany. Computer and programs, fulltext of articles

External sources

• No sources of support supplied

INDEX TERMS

Medical Subject Headings (MeSH)

Antineoplastic Combined Chemotherapy Protocols [*administration & dosage; adverse effects]; Hematopoietic Stem Cell Transplantation [*methods; mortality]; Salvage Therapy [*methods; mortality]; Sarcoma [*drug therapy; mortality]; Transplantation, Autologous

MeSH check words

Adult; Humans