

De-implementation of low-value care in hip and knee arthroplasty Voorn, V.M.A.

Citation

Voorn, V. M. A. (2019, February 26). *De-implementation of low-value care in hip and knee arthroplasty*. Retrieved from https://hdl.handle.net/1887/68705

Version: Not Applicable (or Unknown)

License: License agreement concerning inclusion of doctoral thesis in the

Institutional Repository of the University of Leiden

Downloaded from: https://hdl.handle.net/1887/68705

Note: To cite this publication please use the final published version (if applicable).

Cover Page



Universiteit Leiden



The following handle holds various files of this Leiden University dissertation: http://hdl.handle.net/1887/68705

Author: Voorn, V.M.A.

Title: De-implementation of low-value care in hip and knee arthroplasty

Issue Date: 2019-02-26

Chapter 2

Cell Salvage in Hip and Knee Arthroplasty: A Meta-Analysis of Randomized Controlled Trials

Leti van Bodegom-Vos, Veronique MA Voorn, Cynthia So-Osman, Thea PM Vliet Vlieland,
Albert Dahan, Ankie WMM Koopman-van Gemert, Stephan B Vehmeijer, Rob GHH
Nelissen, Perla J Marang-van de Mheen

J Bone Joint Surg Am. 2015 Jun 17;97(12):1012-21

Abstract

Background: Cell salvage is used to reduce allogeneic red blood-cell (RBC) transfusions in total hip arthroplasty (THA) and total knee arthroplasty (TKA). We performed a meta-analysis to assess the effectiveness of cell salvage to reduce transfusions in THA and TKA separately, and to examine whether recent trials change the conclusions from previous meta-analyses.

Methods: We searched MEDLINE through January 2013 for randomized clinical trials evaluating the effects of cell salvage in THA and TKA. Trial results were extracted using standardized forms and pooled using a random-effects model. Methodological quality of the trials was evaluated using the Cochrane Collaboration's tool for risk-of-bias assessment.

Results: Forty-three trials (5631 patients) were included. Overall, cell salvage reduced the exposure to allogeneic RBC transfusion in THA (risk ratio [RR], 0.66; 95% confidence interval [CI], 0.51 to 0.85) and TKA (RR, 0.51; 95% CI, 0.39 to 0.68). However, trials published in 2010 to 2012, with a lower risk of bias, showed no significant effect of cell salvage in THA (RR, 0.82; 95% CI, 0.66 to 1.02) and TKA (RR, 0.91; 95% CI, 0.63 to 1.31), suggesting that the treatment policy regarding transfusion may have changed over time.

Conclusions: Looking at all trials, cell salvage still significantly reduced the RBC exposure rate and the volume of RBCs transfused in both THA and TKA. However, in trials published more recently (2010 to 2012), cell salvage reduced neither the exposure rate nor the volume of RBCs transfused in THA and TKA, most likely explained by changes in blood transfusion management.

Introduction

Blood loss in total hip arthroplasty (THA) and total knee arthroplasty (TKA) may necessitate allogeneic red blood cell (RBC) transfusion. Concerns regarding the safety of allogeneic RBC transfusions have led to the use of perioperative cell salvage, intended to reduce allogeneic blood use.¹

Previous meta-analyses of randomized controlled trials concluded that cell salvage is effective at reducing the need for allogeneic RBC transfusion, without adverse impact on clinical outcomes in orthopaedic surgery. 1-3 None of those meta-analyses compared the effectiveness of cell salvage in THA with those in TKA. However, it can be hypothesized that the effects in THA and TKA might be different, given differences in anatomy, size of the wound, and surgical technique. Furthermore, as there is less surrounding tissue that can absorb blood lost in TKA, reinfusion drains are likely to collect blood more effectively in TKA than in THA, leading to a larger reduction in the risk for allogeneic RBC transfusion in TKA

Furthermore, several large randomized controlled trials that have been published more recently indicated that cell salvage did not reduce the need for allogeneic RBC transfusion. 4-6 Various developments in orthopaedic surgery may have resulted in these different outcomes of recent trials. First, there has been a trend toward using more restrictive transfusion thresholds. In the last decade there has been an increased awareness that the traditional transfusion trigger, a haemoglobin concentration of <10 g/dL (\sim 6.2 mmol/L),⁷ is no longer tenable because of transfusion risks and escalating costs. Therefore, transfusion in many centres is now based on clinical symptoms, overall patient health, and a more restrictive haemoglobin level of 8 g/dL (~5.0 mmol/L) in uncomplicated cases.⁸ Second, the treatment policy in control groups may be different in recent trials, particularly with respect to the routine use of closed suction drainage since Parker et al.⁹ showed in 2007 that this was associated with higher transfusion rates in THA and TKA without any effect on the rates of wound infections or hematomas compared with using no drain. Third, changes in the timing of cell salvage potentially affected the outcomes of recent trials. Currently, cell salvage devices can reinfuse blood collected both intraoperatively and postoperatively (i.e., perioperatively), whereas the first cell salvage devices could only reinfuse blood collected during surgery. Finally, surgical techniques might have changed. For example, concerns have been raised about the use of tourniquet control in TKA as complications due to its use can delay recovery. 10 Because of these concerns, more recent studies may not have had routine tourniquet use, leading to lower effectiveness of cell salvage in TKA. All of these developments underline the need to update the available evidence.

The aims of the present study were 1) to assess the effectiveness of cell salvage in reducing allogeneic RBC transfusion in THA and TKA separately, and 2) to examine whether the addition of recent trials changes the conclusions regarding the effectiveness of cell salvage as described by Carless et al.¹ To our knowledge, the meta-analysis by Carless et al. was not only the largest meta-analysis but also the most complete one, as the other meta-analyses only reviewed specific types of cell salvage or patient groups.^{2,3}

Materials and Methods

Study selection

All articles involving orthopaedic procedures identified by Carless et al. were retrieved. Next, we searched MEDLINE from January 2009 through January 2013 using the same search strategy as Carless et al. (see Appendix 1). Furthermore, the references of included articles were checked and experts in the field were contacted for additional studies.

Articles were eligible for inclusion if they reported results of randomized controlled trials using cell salvage in THA and/or TKA in adult patients (at least eighteen years old). Studies with a combination of active comparisons were only included if both the intervention and control groups were equally exposed to the active treatment (active treatment plus cell salvage compared with active treatment only), as was done by Carless et al.¹ There were no language restrictions.

Data extraction and outcome measures

Study characteristics and outcomes were extracted for all thirty-five studies involving orthopaedic procedures from Carless et al.¹, using standardized forms, to show the results for THA and TKA separately. If data could not be extracted separately for THA and TKA, the authors of the study were contacted. If they did not respond, the article was placed in the category "not able to split or other orthopaedic procedures."

Next, the titles of newly identified trials from our search strategy were screened by two reviewers, and full-text articles were retrieved. The reviewers independently selected trials that met the inclusion criteria, with disagreements resolved by consensus. For each selected trial, the reviewers independently extracted the following study characteristics: type of surgery (THA or TKA), transfusion threshold used (none, \leq 8.0 g/dL [\sim 5.0 mmol/L] [restrictive], or >8.0 g/dL [\sim 5.0 mmol/L] [traditional]), treatment policy in the control group (no drain, use of closed suction wound drainage after surgery, or another active intervention), timing of cell salvage (intraoperative, perioperative, or postoperative), use

of tourniquet control (in TKA), and primary outcomes (the number of patients exposed to allogeneic RBC transfusion, and the volume of RBCs transfused per patient [with transfusion data expressed in millilitres converted to RBC units by dividing by 300]).

Risk of bias assessment

Included studies were assessed for methodological quality, using the Cochrane Collaboration's tool for assessing risk of bias, by two independent reviewers. The domains assessed were sequence generation, allocation concealment, and blinding 11. Disagreement was resolved by consensus.

Statistical Analysis

Data were extracted and entered into Review Manager (RevMan) (version 5.2.13; The Nordic Cochrane Center, The Cochrane Collaboration, Copenhagen, Denmark). Dichotomous and continuous data were pooled across trials using a random-effects model. Differences in outcome between the experimental group (receiving cell salvage) and the control group were expressed as a risk ratio (RR) for dichotomous outcomes and as a weighted mean difference (WMD) for continuous outcomes, along with a 95% confidence interval (CI). Thus, an RR of <1 indicates that cell salvage reduces the risk for allogeneic blood transfusion, and a negative WMD value indicates a reduction in the volume of RBCs transfused. If neither the standard deviation nor the standard error of the mean was reported for continuous data, the trial was not included. Differences were considered significant if p < 0.05. In addition, data in RevMan were arranged into three groups according to the decade of publication to assess changes in the effectiveness of cell salvage over time.

Subgroup Analysis and Investigation of Heterogeneity

Statistical heterogeneity was examined with the I² test. The I² test describes the percentage of the total variation across studies that is due to heterogeneity rather than chance (with 0% indicating no observed heterogeneity, and >50% indicating substantial heterogeneity)¹¹. Four exploratory analyses of subgroups (defined prior to the study) were performed; these involved the transfusion threshold used, treatment policy in the control group, timing of cell salvage, and use of tourniquet control (in TKA).

Source of funding

This study was funded by the Netherlands Organisation for Health Research and Development (ZonMw 837003001).

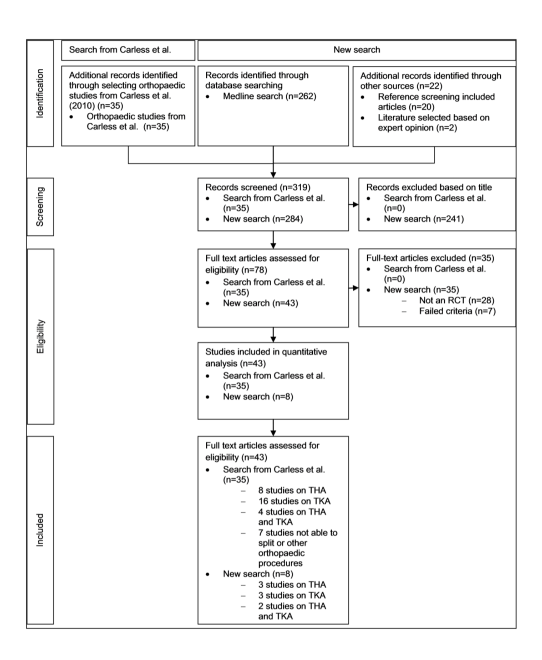


Figure 1: Literature search results

Results

We identified 284 titles in our search: 262 from MEDLINE and twenty-two after checking references and consulting experts (Figure 1). Review of these titles identified forty-three potentially eligible studies. Based on the full articles, eight studies fulfilled the inclusion criteria and were included in addition to the thirty-five studies identified by Carless et al.¹ Of these forty-three included studies (5631 patients), eleven included only THA,^{4,12-21} nineteen included only TKA,^{5,22-39} six included both THA and TKA,^{6,40-44} and seven studies could not be split up or included other orthopaedic procedures.⁴⁵⁻⁵¹ Appendix 2 summarizes the characteristics and the risk-of-bias assessment of all included studies.

Risk-of-Bias Assessment

The risk of bias due to inadequate sequence generation was judged to be low in fifteen studies (Table 1). Five studies had adequate allocation concealment (that is, low risk of bias). Three studies were judged to be double-blinded. Recent studies more often seemed to have lower risk of bias (that is, higher quality) compared with studies published before 2010, particularly with respect to sequence generation and allocation concealment.

Table 1: Risk of bias of included studies

		Studies from Carl	ess et al.	New search
	Total, N = 43	Published 1990- 1999, n=22	Published 2000- 2009, n=13	Published 2010- 2012, n=8
Adequate sequence generation				
- Yes, i.e. low risk of bias	15	6	4	5
- No, i.e. high risk of bias	5	2	2	1
- Unclear, i.e. uncertain risk of bias	23	14	7	2
Adequate allocation concealment				
- Yes, i.e. low risk of bias	5	-	-	5
- No, i.e. high risk of bias	11	5	5	1
- Unclear, i.e. uncertain risk of bias	27	17	8	2
Adequate blinding				
- Yes, i.e. low risk of bias	3	1	-	2
- No, i.e. high risk of bias	39	21	13	5
- Unclear, i.e. uncertain risk of bias	1	-	-	1

Effects of Cell Salvage in Orthopaedic Surgery

Figure 2 shows the effect of cell salvage on the RBC exposure rate in orthopaedic surgery from Carless et al.¹. In THA, cell salvage reduced the RBC exposure rate by 44% (RR, 0.56; 95% CI, 0.38 to 0.82; n = 11) and in TKA by 56% (RR, 0.44; 95% CI, 0.32 to 0.60; n = 18). Cell salvage did not significantly reduce the volume of RBCs transfused in either THA (WMD, -0.97; 95% CI, -1.94 to 0.00; n = 5) or TKA (WMD, -0.42; 95% CI, -0.92 to 0.09; n = 6).

Effects of Cell Salvage in THA

Overall, cell salvage still reduced the RBC exposure rate by 34% (RR, 0.66; 95% CI, 0.51 to 0.85) in THA when recent trials were included, without substantial heterogeneity among studies ($I^2 = 50\%$). However, as shown in figure 3, the date of the study appeared to have an effect, with more recent studies (2010 to 2012) showing no significant effect of cell salvage (RR, 0.82; 95% CI, 0.66 to 1.02), without any heterogeneity ($I^2 = 0\%$).

Overall, cell salvage in THA reduced the volume of RBCs transfused (WMD, -0.67; 95% CI, -1.08 to -0.27; $I^2 = 91\%$). Again, an effect of the study date was observed, with recently published studies (2010 to 2012) showing a nonsignificant reduction in the volume of RBCs transfused (WMD, -0.13; 95% CI, -0.30 to 0.04; $I^2 = 39\%$).

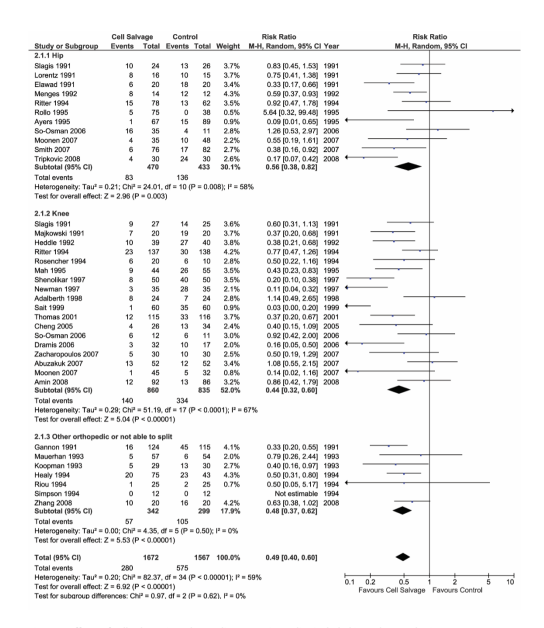


Figure 2: Effects of cell salvage in orthopaedic surgery in studies included in Carless et al: Hip versus Knee Arthroplasty

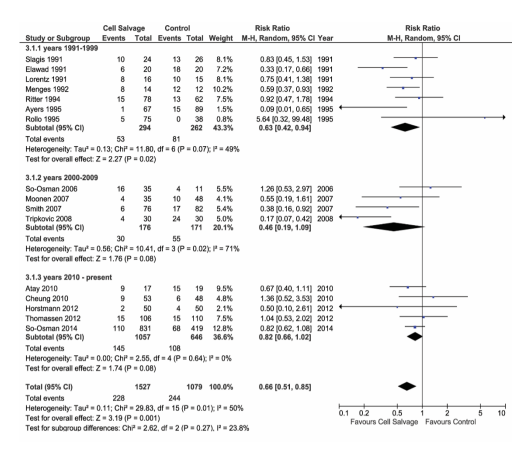


Figure 3: Effects of cell salvage in Hip Arthroplasty over time

Subgroup Analyses

To explain the time period effect described above, exploratory subgroup analyses were performed. Given the number of studies per time period, no further stratification was possible. Therefore, we included all studies from all time periods in the subgroup analyses and assessed 1) whether the effectiveness between subgroups was different, and 2) whether a possible explanatory variable (for example, a more strict transfusion threshold) was more frequently present in recent than in older studies. The variable was considered a possible explanation for a part of the observed change in effectiveness over time only if both criteria were true.

- In studies using a traditional transfusion threshold, cell salvage significantly reduced the RBC exposure rate (RR, 0.57; 95% CI, 0.36 to 0.89; $I^2 = 67\%$; n = 6 [1 recent]) and

the volume of RBCs transfused (WMD, -1.56; 95% CI, -2.16 to -0.95; $I^2 = 61\%$; n = 3 [none recent]). In studies with a more restrictive threshold, cell salvage resulted in a smaller reduction of the RBC exposure rate (RR, 0.72; 95% CI, 0.58 to 0.91; $I^2 = 0\%$; n = 5 [3 recent]) and did not significantly reduce the volume of RBCs transfused (WMD, -0.13; 95% CI, -0.30 to 0.04; $I^2 = 39\%$; n = 3 [all recent]).

- In studies using closed suction wound drainage in the control group, cell salvage significantly reduced the RBC exposure rate (RR, 0.78; 95% CI, 0.62 to 0.98, I² = 6%; n = 6 [3 recent]), but not the volume of RBCs transfused (WMD, −0.16; 95% CI, −0.45 to 0.13, I² = 61%; n = 4 [2 recent]). In studies using no drain in the control group, cell salvage did not significantly reduce the RBC exposure rate (RR, 0.69; 95% CI, 0.43 to 1.13, I² = 45%; n = 5 [2 recent]) or the volume of RBCs transfused (WMD, −1.04; 95% CI, −2.96 to 0.88; I² = 98%; n = 2 [1 recent]).
- Intraoperative cell salvage (only applied in one trial) reduced the RBC exposure rate (RR, 0.33; 95% CI, 0.17 to 0.66) and the volume of RBCs transfused (WMD, −2.04; 95% CI, −2.58 to −1.50). Postoperative cell salvage significantly reduced the RBC exposure rate (RR, 0.68; 95% CI, 0.49 to 0.93, I² = 55%; n = 13 [4 recent]) and the volume of RBCs transfused (WMD, −0.38; 95% CI, −0.72 to −0.04; I² = 86%; n = 6 [3 recent]). Perioperative cell salvage significantly reduced neither the RBC exposure rate (RR, 0.76; 95% CI, 0.58 to 1.00; I² = 0%; n = 4 [2 recent]) nor the volume of RBCs transfused (WMD, −0.28; 95% CI, −0.76 to 0.18; I² = 34%; n = 2 [1 recent]).

Effects of Cell Salvage in TKA

Overall, cell salvage still reduced the RBC exposure rate by 49% (RR, 0.51; 95% CI, 0.39 to 0.68) in TKA when recent trials were added (Figure 4), with substantial heterogeneity among studies ($I^2 = 75\%$). Again, a time period effect was observed, with more recent studies (2010 to 2012) showing no significant effect of cell salvage (RR, 0.91; 95% CI, 0.63 to 1.31; $I^2 = 54\%$).

Overall, cell salvage in total knee arthroplasty also reduced the volume of RBCs transfused (WMD, -0.33; 95% CI, -0.59 to -0.08; I² = 91%). Again, a time period effect was observed, with recently published studies (2010 to 2012) showing a nonsignificant reduction in the volume of RBCs transfused (WMD, -0.32; 95% CI, -0.63 to 0.00; I² = 95%).

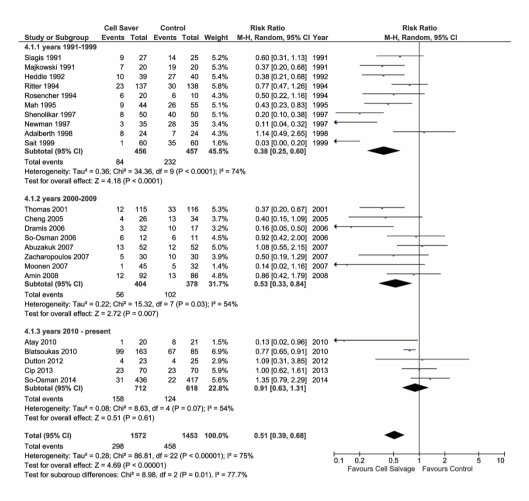


Figure 4: Effects of cell salvage in Knee Arthroplasty over time

Subgroup Analyses

To explain the time period effect described above, exploratory subgroup analyses similar to those for THA were performed.

- In studies using a traditional threshold, cell salvage significantly reduced the RBC exposure rate (RR, 0.54; 95% CI, 0.40 to 0.73; I² = 72%; n = 13 [2 recent]) and the volume of RBCs transfused (WMD, −0.60; 95% CI, −1.08 to −0.12; I² = 80%; n = 4 [1 recent]). In studies with a more restrictive threshold, cell salvage reduced neither the RBC exposure rate (RR, 0.54; 95% CI, 0.25 to 1.18, I² = 74%; n = 5 [2 recent]) nor the

- volume of RBCs transfused (WMD, -0.45; 95% CI, -1.07 to 0.18; $I^2 = 92\%$; n = 3 [2 recent]).
- In studies using closed suction wound drainage in the control group, cell salvage significantly reduced the RBC exposure rate (RR, 0.44; 95% CI, 0.27 to 0.72; $I^2 = 78\%$; n = 13 [3 recent]), but not the volume of RBCs transfused (WMD, -0.38; 95% CI, -0.82 to 0.05; $I^2 = 90\%$; n = 5 [1 recent]). In studies using no drain in the control group, cell salvage resulted in a smaller reduction of the RBC exposure rate (RR, 0.56; 95% CI, 0.37 to 0.85, $I^2 = 75\%$; n = 8 [2 recent]), and did not significantly reduce the volume of RBCs transfused (WMD, -0.24; 95% CI, -0.92 to 0.45; $I^2 = 96\%$; n = 3 [2 recent]).
- Postoperative cell salvage significantly reduced the RBC exposure rate (RR, 0.49; 95% CI, 0.37 to 0.66; I² = 73%; n = 22 [4 recent]) and the volume of RBCs transfused (WMD, −0.32; 95% CI, −0.55 to −0.08; I² = 92%; n = 10 [4 recent]). Perioperative cell salvage resulted in a smaller reduction of the RBC exposure rate (RR, 0.81; 95% CI, 0.68 to 0.97, I² = 0%; n = 2 [both recent]) and a reduction of the volume of RBCs transfused (WMD, −0.93; 95% CI, −1.21 to −0.65; n = 1 [recent]).
- In studies performing TKA under tourniquet control, cell salvage significantly reduced the RBC exposure rate (RR, 0.46; 95% CI, 0.33 to 0.65, I² = 71%; n = 20 [3 recent]), but did not reduce the volume of RBCs transfused (WMD, −0.22; 95% CI, −0.45 to 0.01; I² = 87%; n = 8 [3 recent]). In studies performing TKA without tourniquet control, cell salvage resulted in a smaller reduction of the RBC exposure (RR, 0.78; 95% CI, 0.67 to 0.91; I² = 0%; n = 3 [2 recent]) and a reduction in the volume of RBCs transfused (WMD, −0.85; 95% CI, −1.09 to −0.61; I² = 0%; n = 2 [1 recent]).

Discussion

Our meta-analysis showed that cell salvage significantly reduces the RBC exposure rate and the volume of RBCs transfused in both THA and TKA, with a larger effect in TKA than in THA based on group averages. However, in trials published more recently (2010 to 2012), cell salvage reduced neither the exposure rate nor the volume of RBCs transfused in both THA and TKA. We therefore conclude that, given changes in blood transfusion management, the effect of cell salvage may have changed over time and it may not be as effective as shown in previous meta-analyses.¹⁻³ This conclusion seems even stronger if the methodological quality of the studies is considered. Recent studies more often had a lower risk of bias and therefore higher quality of evidence.

Subgroup analyses showed that a more restrictive transfusion trigger (haemoglobin [Hb] ≤8.0 g/dL) was associated with a smaller effect of cell salvage. Cell salvage reduced the exposure rate only in THA and was not effective in TKA. Given that recent trials more

often used this restrictive transfusion threshold, this may partly explain the observed time period effect in effectiveness of cell salvage.

Similarly, using no drain as the standard treatment in the control group was also associated with smaller effects of cell salvage. Cell salvage was no longer effective in THA, and it reduced only the RBC exposure rate in TKA. These results are in line with the meta-analysis of Parker et al.,⁹ who showed that routine use of closed suction drainage in THA and TKA was associated with higher transfusion rates and did not have any effect on the rate of wound infections or hematomas compared with no drain use. However, as recent studies did not use 'no drain' as the control treatment more frequently than studies published before 2010, it does not explain the observed time period effect.

Subgroup analyses regarding the timing of cell salvage and use of tourniquet control established no clear reasons for the observed time period effect. Only a few studies, although proportionally more recent studies, performed TKA without tourniquet control. This is in line with the results of the 2009 review by Smith and Hing¹⁰ showing that the use of a tourniquet decreases intraoperative blood loss but could not influence postoperative blood loss in drains or affect transfusion rates.

Some relevant variables were not reported in a sufficient number of trials and could thus not be used in the meta-analysis: preoperative and postoperative haemoglobin levels, the exact timing of haemoglobin measurements resulting in the decision to transfuse or not, and the exact amount of blood given back to the patient, which differs among devices. Therefore, additional research is needed to be able to assess whether cell salvage may have benefit in raising haemoglobin levels for subgroups of patients and to interpret the effect of the timing of haemoglobin measurement and the volume of blood transfused on the effectiveness of cell salvage. Furthermore, we recommend that future studies report the utilized surgical techniques in more detail, enabling future meta-analyses to perform subgroup analyses to determine whether primary outcomes of cell salvage differ by surgical technique.

There are some important limitations of this meta-analysis. First, it included an insufficient number of high-quality studies to permit limiting our analyses to high-quality studies only. However, our risk-of-bias assessment showed that more recent studies seemed to have lower risk of bias compared with studies published before 2010, which strengthens our conclusion that cell salvage may no longer be effective in reducing the RBC exposure rate and the volume of RBCs transfused. Second, only three of the included studies were judged to be double-blinded. Although this is problematic for the quality of the studies, it is probably not possible to further improve blinding procedures given the nature of the intervention. However, as sequence generation and allocation concealment clearly

2

improved in recent studies, there is lower risk of bias and thus higher quality of evidence in recent studies. Third, the results of this meta-analysis only apply to cell salvage in THA and TKA. Cell salvage may still be effective for other surgical procedures (for example, during cardiac surgery), which could be a topic for further research. In addition, the results only allow us to draw conclusions about the effectiveness of perioperative collection and reinfusion of autologous blood (cell salvage) and not about preoperative autologous blood donation and reinfusion.

Given the results of this meta-analysis, the benefit of cell salvage in clinical practice in uncomplicated patients undergoing THA and TKA is questioned. Further research is needed to be able to definitely answer this question, as current trials have insufficient data on parameters such as haemoglobin levels. The current meta-analysis contributes to this debate by creating awareness among professionals that the effectiveness of cell salvage to reduce transfusion rates is minimized in recent studies, which have lower risk of bias and more often have used more restrictive transfusion triggers.

Acknowledgments

The authors thank Ms. C. Kooy-Verhoef for her help with the literature search and data collection.

References

- Carless PA, Henry DA, Moxey AJ, O'Connell D, Brown T, Fergusson DA. Cell salvage for minimising perioperative allogeneic blood transfusion. Cochrane Database Syst Rev. 2010;4(4):CD001888. Epub 2010 Apr 14.
- Haien Z, Yong J, Baoan M, Mingjun G, Qingyu F. Post-operative auto-transfusion in total hip or knee arthroplasty: a meta-analysis of randomized controlled trials. PLoS One. 2013;8(1):e55073. Epub 2013 Jan 25.
- Markar SR, Jones GG, Karthikesalingam A, Segaren N, Patel RV. Transfusion drains versus suction drains in total knee replacement: meta-analysis. Knee Surg Sports Traumatol Arthrosc. 2012 Sep;20(9):1766-72. Epub 2011 Nov 10.
- Cheung G, Carmont MR, Bing AJ, Kuiper JH, Alcock RJ, Graham NM. No drain, autologous transfusion drain
 or suction drain? A randomised prospective study in total hip replacement surgery of 168 patients. Acta
 Orthop Belg. 2010 Oct;76(5):619-27.
- Cip J, Widemschek M, Benesch T, Waibel R, Martin A. Does single use of an autologous transfusion system in TKA reduce the need for allogenic blood?: a prospective randomized trial. Clin Orthop Relat Res. 2013 Apr;471(4):1319-25. Epub 2012 Dec 11.
- 5. So-Osman C, Nelissen RG, Koopman-van Gemert AW, Kluyver E, Pöll RG, Onstenk R, Van Hilten JA, Jansen-Werkhoven TM, van den Hout WB, Brand R, Brand A. Patient blood management in elective total hip- and knee-replacement surgery (part 1): a randomized controlled trial on erythropoietin and blood salvage as transfusion alternatives using a restrictive transfusion policy in erythropoietin-eligible patients. Anesthesiology. 2014 Apr;120(4):839-51.
- Adams RC, Lundy JS. Anesthesia in cases of poor surgical risk: some suggestions for decreasing the risk. Surg Gynecol Obstet. 1942;74:1011-9.
- 7. Young SW, Marsh DJ, Akhavani MA, Walker CG, Skinner JA. Attitudes to blood transfusion post arthroplasty surgery in the United Kingdom: a national survey. Int Orthop. 2008 Jun;32(3):325-9. Epub 2007 Mar 30.
- 8. Parker MJ, Livingstone V, Clifton R, McKee A. Closed suction surgical wound drainage after orthopaedic surgery. Cochrane Database Syst Rev. 2007;(3):CD001825. Epub 2007 Jul 18.
- Smith TO, Hing CB. A meta-analysis of tourniquet assisted arthroscopic knee surgery. Knee. 2009 Oct;16(5):317-21. Epub 2009 Feb 23.
- Higgins JPT, Green S. Cochrane handbook for systematic reviews of interventions. 2011.http://handbook.cochrane.org/. Accessed 2015 Jan 14.
- 11. Ayers DC, Murray DG, Duerr DM. Blood salvage after total hip arthroplasty. J Bone Joint Surg Am. 1995 Sep;77(9):1347-51.
- 12. Ekbäck G, Schött U, Axelsson K, Carlberg M. Perioperative autotransfusion and functional coagulation analysis in total hip replacement. Acta Anaesthesiol Scand. 1995 Apr;39(3):390-5.
- Elawad AA, Ohlin AK, Berntorp E, Nilsson IM, Fredin H. Intraoperative autotransfusion in primary hip arthroplasty. A randomized comparison with homologous blood. Acta Orthop Scand. 1991 Dec;62(6):557-62
- Horstmann WG, Kuipers BM, Slappendel R, Castelein RM, Kollen BJ, Verheyen CC. Postoperative autologous blood transfusion drain or no drain in primary total hip arthroplasty? A randomised controlled trial. Int Orthop. 2012 Oct;36(10):2033-9. Epub 2012 Jul 13.
- 15. Lorentz A, Osswald PM, Schilling M, Jani L. [A comparison of autologous transfusion procedures in hip surgery]. Anaesthesist. 1991 Apr;40(4):205-13. German.
- Menges T, Rupp D, van Lessen A, Hempelmann G. [Measures for reducing the use of homologous blood. Effects on blood coagulation during total endoprosthesis]. Anaesthesist. 1992 Jan;41(1):27-33. German.
- 17. Rollo VJ, Hozack WJ, Rothman RH, Chao W, Eng KO. Prospective randomized evaluation of blood salvage techniques for primary total hip arthroplasty. J Arthroplasty. 1995 Aug;10(4):532-9.
- 18. Smith LK, Williams DH, Langkamer VG. Post-operative blood salvage with autologous retransfusion in primary total hip replacement. J Bone Joint Surg Br. 2007 Aug;89(8):1092-7.
- Thomassen BJ, Pilot P, Scholtes VA, Grohs JG, Holen K, Bisbe E, Poolman RW. Limit allogeneic blood use with routine re-use of patient's own blood: a prospective, randomized, controlled trial in total hip surgery. PLoS One. 2012;7(9):e44503. Epub 2012 Sep 13.
- Tripković B, Buković D, Sakić K, Sakić S, Buković N, Radaković B. Quality of the blood sampled from surgical drainage after total hip arthroplasty. Coll Antropol. 2008 Mar;32(1):153-60.

- Abuzakuk T, Senthil Kumar V, Shenava Y, Bulstrode C, Skinner JA, Cannon SR, Briggs TW. Autotransfusion drains in total knee replacement. Are they alternatives to homologous transfusion? Int Orthop. 2007 Apr;31(2):235-9. Epub 2006 Jun 8.
- 22. Adalberth G, Byström S, Kolstad K, Mallmin H, Milbrink J. Postoperative drainage of knee arthroplasty is not necessary: a randomized study of 90 patients. Acta Orthop Scand. 1998 Oct;69(5):475-8.
- 23. Altinel L, Kaya E, Kose KC, Fidan F, Ergan V, Fidan H. Effect of shed blood retransfusion on pulmonary perfusion after total knee arthroplasty: a prospective controlled study. Int Orthop. 2007 Dec;31(6):837-44. Epub 2006 Nov 4.
- 24. Amin A, Watson A, Mangwani J, Nawabi D, Ahluwalia R, Loeffler M. A prospective randomised controlled trial of autologous retransfusion in total knee replacement. J Bone Joint Surg Br. 2008 Apr;90(4):451-4.
- 25. Blatsoukas KS, Drosos Gl, Kazakos K, Papaioakim M, Gioka T, Chloropoulou P, Verettas DA. Prospective comparative study of two different autotransfusion methods versus control group in total knee replacement. Arch Orthop Trauma Surg. 2010 Jun;130(6):733-7. Epub 2010 Feb 18.
- Cheng SC, Hung TS, Tse PY. Investigation of the use of drained blood reinfusion after total knee arthroplasty: a prospective randomised controlled study. J Orthop Surg (Hong Kong). 2005 Aug;13(2):120-4.
- 27. Dramis A, Plewes J. Autologous blood transfusion after primary unilateral total knee replacement surgery. Acta Orthop Belg. 2006 Jan;72(1):15-7.
- Dutton T, De-Souza R, Parsons N, Costa ML. The timing of tourniquet release and 'retransfusion' drains in total knee arthroplasty: a stratified randomised pilot investigation. Knee. 2012 Jun;19(3):190-2. Epub 2011 Mar 25.
- 29. Heddle NM, Brox WT, Klama LN, Dickson LL, Levine MN. A randomized trial on the efficacy of an autologous blood drainage and transfusion device in patients undergoing elective knee arthroplasty. Transfusion. 1992 Oct;32(8):742-6.
- Kirkos JM, Krystallis CT, Konstantinidis PA, Papavasiliou KA, Kyrkos MJ, Ikonomidis LG. Postoperative reperfusion of drained blood in patients undergoing total knee arthroplasty: is it effective and cost-efficient? Acta Orthop Belg. 2006 Jan;72(1):18-23.
- 31. Mah ET, Davis R, Seshadri P, Nyman TL, Seshadri R. The role of autologous blood transfusion in joint replacement surgery. Anaesth Intensive Care. 1995 Aug;23(4):472-7.
- 32. Majkowski RS, Currie IC, Newman JH. Postoperative collection and reinfusion of autologous blood in total knee arthroplasty. Ann R Coll Surg Engl. 1991 Nov;73(6):381-4.
- 33. Newman JH, Bowers M, Murphy J. The clinical advantages of autologous transfusion. A randomized, controlled study after knee replacement. J Bone Joint Surg Br. 1997 Jul;79(4):630-2.
- 34. Rosencher N, Vassilieff V, Tallet F, Toulon P, Leoni J, Tomeno B, Conseiller C. [Comparison of Orth-Evac and Solcotrans Plus devices for the autotransfusion of blood drained after total knee joint arthroplasty]. Ann Fr Anesth Reanim. 1994;13(3):318-25. French.
- 35. Sait MS, Earnshaw P. Autotransfusion in total knee arthroplasty is it worth the effect? J Bone Joint Surg Br. 1999;81(Suppl 2):244.
- 36. Shenolikar A, Wareham K, Newington D, Thomas D, Hughes J, Downes M. Cell salvage auto transfusion in total knee replacement surgery. Transfus Med. 1997 Dec;7(4):277-80.
- 37. Thomas D, Wareham K, Cohen D, Hutchings H. Autologous blood transfusion in total knee replacement surgery. Br J Anaesth. 2001 May;86(5):669-73.
- 38. Zacharopoulos A, Apostolopoulos A, Kyriakidis A. The effectiveness of reinfusion after total knee replacement. A prospective randomised controlled study. Int Orthop. 2007 Jun;31(3):303-8. Epub 2006 Jun 30.
- Atay EF, Güven M, Altıntaş F, Kadıoğlu B, Ceviz E, Ipek S. Allogeneic blood transfusion decreases with postoperative autotransfusion in hip and knee arthroplasty. Acta Orthop Traumatol Turc. 2010;44(4):306-12.
- 40. Moonen AF, Knoors NT, van Os JJ, Verburg AD, Pilot P. Retransfusion of filtered shed blood in primary total hip and knee arthroplasty: a prospective randomized clinical trial. Transfusion. 2007 Mar;47(3):379-84.
- 41. Ritter MA, Keating EM, Faris PM. Closed wound drainage in total hip or total knee replacement. A prospective, randomized study. J Bone Joint Surg Am. 1994 Jan;76(1):35-8.
- So-Osman C, Nelissen RG, Eikenboom HC, Brand A. Efficacy, safety and user-friendliness of two devices for postoperative autologous shed red blood cell re-infusion in elective orthopaedic surgery patients: a randomized pilot study. Transfus Med. 2006 Oct;16(5):321-8.
- Slagis SV, Benjamin JB, Volz RG, Giordano GF. Postoperative blood salvage in total hip and knee arthroplasty. A randomised controlled trial. J Bone Joint Surg Br. 1991 Jul;73(4):591-4.

- 44. Gannon DM, Lombardi AV Jr, Mallory TH, Vaughn BK, Finney CR, Niemcryk S. An evaluation of the efficacy of postoperative blood salvage after total joint arthroplasty. A prospective randomized trial. J Arthroplasty. 1991 Jun;6(2):109-14.
- 45. Healy WL, Pfeifer BA, Kurtz SR, Johnson C, Johnson W, Johnston R, Sanders D, Karpman R, Hallack GN, Valeri CR. Evaluation of autologous shed blood for autotransfusion after orthopaedic surgery. Clin Orthop Relat Res. 1994 Feb;299:53-9.
- 46. Koopman-van Gemert AWMM. Processed autotransfusion and homologous red cell requirement in elective cardiac and orthopaedic surgery: a randomised prospective study. In: Perioperative autotransfusion by means of a blood cell separator. Den Haag: Cip-Data Koninklijke Bibliotheek; 1993. p 105-26.
- 47. Mauerhan DR, Nussman D, Mokris JG, Beaver WB. Effect of postoperative reinfusion systems on hemoglobin levels in primary total hip and total knee arthroplasties. A prospective randomized study. J Arthroplasty. 1993 Oct;8(5):523-7.
- Riou B, Arock M, Guerrero M, Ramos M, Thoreux P, Guillosson JJ, Roy-Camille R, Viars P. Haematological effects of postoperative autotransfusion in spinal surgery. Acta Anaesthesiol Scand. 1994 May;38(4):336-41
- 49. Simpson MB, Murphy KP, Chambers HG, Bucknell AL. The effect of postoperative wound drainage reinfusion in reducing the need for blood transfusions in elective total joint arthroplasty: a prospective, randomized study. Orthopedics. 1994 Feb;17(2):133-7.
- 50. Zhang XF, Dong JM, Gong ML, Shen SM, Zhou Y, Pan YF, Mao JP. [Effectiveness of preoperative autologous plateletpheresis combined with intraoperative autotransfusion on the blood coagulation in orthopaedic patients]. Zhonghua Wai Ke Za Zhi. 2008 Jan 15;46(2):118-21. Chinese.

Appendix 1: Search Strategy

MEDLINE search strategy

- 1. cell\$ sav\$.mp.
- 2. cell\$ salvage.mp.
- 3. blood transfusion, autologous/
- 4. autotransfusion\$.mp.
- 5. auto-transfusion\$.mp.
- 6. blood salvage.mp.
- 8. solcotrans system.mp.
- 9. constavac.mp.

7. autovac.mp.

- 10. solcotrans.mp.
- 44 1
- 11. hemovac.mp.
- 12. BRAT.mp.13. fresenius.mp.
- 14. consta vac.mp.
- 15. cell saver.mp.
- 16. dideco.mp.
- 17. electromedic.mp.
- 18. electromedics.mp.
- 19. gish biomedical.mp.
- 20. haemonetics.mp.
- 21. orth-evac.mp.
- 22. pleur-evac.mp.

- 23. sorenson.mp.
- 24. reinfusion system.mp.
- 25. sorin biomedical.mp.
- 26. or/1-25
- 27. exp blood transfusion/
- 28. exp hemorrhage/
- 29. exp anesthesia/
- 30. transfusion\$.mp.
- 31. bleed\$.mp.
- 32. blood loss\$.mp.
- 33. hemorrhag\$.mp.
- 34. haemorrhag\$.mp.
- 35. or/27-34
- 36. 26 and 35
- 37. randomized controlled trial.pt.
- 38. controlled clinical trial.pt.
- 39. randomized controlled
- trials.sh.
- 40. random allocation.sh.
- 41. double blind method.sh.
- 42. single blind method.sh.
- 43. or/37-42

- 44. clinical trial.pt.
- 45, exp Clinical trials/
- 46. (clin\$ adj25 trial\$).ti,ab.
- 47. ((singl\$ or doubl\$ or trebl\$ or
- tripl\$) adj25 (blind\$ or
- mask\$)).ti,ab.
- 48. placebos.sh.
- 49. placebo\$.ti,ab.
- 50. random\$.ti,ab.51. research design.sh.
- 52. or/44-51
- 53. comparative study.sh.
- 54. exp Evaluation studies/
- 55. follow up studies.sh.
- 56. prospective studies.sh.
- 57. (control\$ or prospectiv\$ or
- volunteer\$).ti,ab.
- 58. or/53-57 59. 43 or 52 or 58
- 60. 36 and 59
- 61. animal/ not human/
- 62. 60 not 6

Appendix 2: Study characteristics and risk of bias assessment of included studies

	Blinding		High risk	High risk	High risk	High risk	High risk
risk of bias	Allocation B concealment		Unclear risk H	High risk	Unclear risk H	High risk H	Unclear risk H
Assessment of risk of bias	Random sequence generation		Low risk	High risk	Unclear risk	High risk	Unclear risk
	Tourniquet		Yes	Yes	Yes	Yes	Yes
	Treatme nt policy in control group ⁴		Control 1	Control 0	Control 1	Control 1	Control 1
		Subgroup ³	Trigger 2	Trigger 2	Trigger 2	Trigger 1	Trigger 1
	Transfusion trigger	Transfusion threshold	Hb < 9.0 g/dl	Hb < 9.0 g/dl	Hb < 9.0 g/dl	Hb < 8.0 g/dl	Hb < 8.0 g/dl or Hct < 25% and clinical symptoms of anaemia
	Transfus	Yes/ No	Yes	Yes	Yes	Yes	Yes
		Timing ²	POST	POST	POST	POST	POST
stics	Interventions	Description: Intervention (I) and Control (C)	I: autotransfusion (Bellovac ABT autotransfusion system), n=52; C: standard suction drain (Redivac), n=52	l: autotransfusion (Solcotrans - Solco Basle UK Itd.)), n=24; Control: no drain, n=24	I: autotransfusion (ConstaVac CBCII system), n=16; C: standard care (2 drains for shed blood drainage), n=16	I: autotransfusion (Bellovac ABT autotransfusion system), n=92; C: standard vacuum drain, n=86	I: autotransfusion (Transolog), n=17 (hip) and n=20 (knee); C: routine hemovac drain, n=19 (hip) and n=21 (knee)
characteri	Type¹		Knee	Knee	Knee	Knee	Hip and knee
Summary of study characteristics	Participants: Patients undergoing		Primary cemented total knee arthroplasty	primary total knee arthroplasty	bi- or tri- compartmental total knee arthroplasties	total knee replacement	hip and knee arthroplasty
	0.	Auth Year	Abuzakuk Abuzakuk	AthedlabA 8601	lanitlA 7002	nimA 800S	yetA 0102
<u> </u>							

	1						
High risk	High risk	High risk	High risk	High risk	High risk	Unclear risk	High risk
High risk	High risk	High risk	Unclear risk	Low risk	Unclear risk	Low risk	Unclear risk
High risk	High risk	Unclear risk	Low risk	Low risk	Unclear risk	Low risk	Unclear risk
ς. Υ	O _N	O _N	N.A.	o Z	Yes	Yes	Z.A.
Control 1	Control 0	Control 0	Control 0	Control 1	Control 1	Control 0	Control 1
None	Trigger 2	Trigger 2	None	Trigger 2	Trigger 2	None	Trigger 2
	Hb 9-10 g/dl: 1 unit; Hb 8-9 g/dl: 2 units; Hb 7-8 g/dl: 3 units	Hb < 9.0 g/dl		Hb < 8.0 g/dl or signs of anaemia or tachycardia	Hb < 9.0 g/dl or clinical symptoms of anaemia		EVF < 27% (i.e. Hb < 9.2 g/dl)
<u>0</u>	Yes	Yes	ON.	Yes	Yes	No	Yes
POST	PERI/ POST	POST	POST	PERI	POST	POST	PERI
I: autotransfusion (Autovac postoperative orthopaedic autotransfusion canister), n=67; C: closed suction drainage system, n=89	I: 1. autotransfusion (Dideco Compact Advanced and ConstaVac CBCII), n=92; 2. Autotransfusion (ConstaVaC CBCII), n=71; C. no drain, n=85	I: autotransfusion (DONOR system), n=26; C: no drain, n=34	I: autotransfusion (Bellovac ABT autotransfusion systemt), n=53; C: no drain, n=48	I: autotransfusion (OrthoPAT), n=70; C: no retransfusion system, n=70	I: autotransfusion (CellTrans system), n=32; C: Standard vacuum drain, n=17	I: autotransfusion (Bellovac ABT autotransfusion system), n=23; C: no drain, n=25)	I: Autotransfusion (Haemonetics CellSaver 4, Althin model AT 1000 or Shiley/Dideco STAT), n=15, C. no autotransfusion, n=15
qiH	Knee	Knee	Hip	Knee	Knee	Knee	Hip
primary total hip arthroplasty	unilateral total knee replacement	unilateral total knee arthroplasty	primary total hip replacement	total knee arthroplasty	primary unilateral total knee arthroplasty	total knee arthroplasty	total hip arthroplasty
1995 1995	2010	2002	2010	2012	9007	2012	1995 1995
Ауегѕ	Blatsoukas	Suədə	guəny	qiD	Dramis	Dutton	Екраск

High risk	High risk	High risk	High risk	Low risk	High risk	High risk
High risk	Unclear risk	Unclear risk	Unclear risk	Low risk	Unclear risk	High risk
Unclear risk	Low risk	Unclear risk	Unclear risk	Unclear risk	High risk	Unclear risk
N.A.	Yes	Unknown	Yes	N.A.	Yes	N.A.
Control 0	Control 1	Control 1	Control 1	Control 0	Control 1	Control 1
Trigger 2	Trigger 2	None	Trigger 1	Trigger 1	Trigger 2	Trigger 2
Hb < 8.5 g/dl	Hb < 9.0 g/dl or by internist based on patients' condition		Hb 8.0-8.9 g/dl: 1 unit; Hb 7.0-7.9 g/dl: 2 units; Hb 6.0-6.9 g/dl: 3 units, Hb 5.0-5.9 g/dl: 4 units	Hb < 6.4/ 8.0/ 9.6 g/dl dependent on ASA classification	Hb < 10.0 g/dl	Hct at 30% (i.e. Hb <10.2 g/dl)
Yes	Yes	ON	Yes	Yes	Yes	Yes
INTRA	POST	POST	POST	POST	POST	PERI
I: Autotransfusion (Electromedic Autotrans AT100) autotransfusion system), n=20; C: no drain, n=20	I: Autotransfusion (Solcotrans), n=124; C: standard suction canister, n=115	I: autotransfusion (Ortho-Evac system or Solcotrans), n=75; C: standard wound drainage system, n=43	I: autotransfusion (Solcotrans), n=39; C: standard care (drained blood collected by a Davol suction unit and discarded), n=40	I: autotransfusion (Bellovac ABT autotransfusion system), n=50; C: no drainage, n=50)	I: autotransfusion, n=78; C: standard vacuum drain, n=77	I: autotransfusion (Haemonetics Haemolite-2 system), n=29; C: no autotransfusion, n=30
qiH	NAS	NAS	Knee	Hip	Knee	NAS
primary total hip arthroplasty	total hip or total knee replacement	hip arthroplasty, total knee arthroplasty or spine fusion	elective knee arthroplasty	total hip arthroplasty	total knee arthroplasty	total hip arthroplasty or dorsal lumbo- sacral spinal fusion
bewel3 1991	nonns2 1991	1994 1994	əlppəH	Horstmann 2012	Kirkos 2006	Koopman Koopman
				l		

High risk	High risk	High risk	High risk	High risk	High risk	High risk
Unclear risk	Unclear risk	Unclear risk	Unclear risk	Unclear risk	High risk	Unclear risk
Unclear risk	Low risk	Unclear risk	Low risk	Unclear risk	Low risk	Low risk
N.A.	Yes	Yes	Yes	N.A.	Yes	Yes
Unknow	Active	Control 1	Control 1	Active	Control 1	Control 1
Trigger 2	Trigger 2	Trigger 2	None	Trigger 2	Trigger 1	None
Hb < 9.0 g/dl (operating room, IC); Hb < 10.0 g/dl (other)	Hb < 10.0 g/dl	Hb < 9.5 g/dl or if indicated hemodynami cally		Hb < 9.0 g/dl or Hct < 28% (i.e. Hb < 9.5)	Hb < 8.1/ 8.9/ 9.7 g/dl dependent on ASA classification	
Yes	Yes	Yes	No	Yes	Yes	NO N
PERI	POST	POST	POST	POST	POST	POST
I: Autotransfusion, n=16; C: standard care, n=15	I: autotransfusion (Electromedics BT-795), n=44; C: standard care, n=55	I: autotransfusion (Solcotrans), n=20; C: three standard Redivac drains	I: autotransfusion (CBC ConstaVac), n=57; C: standard post-operative collection system, n=54	I: autotransfusion (Autotrans BT 795 P, Dideco system), n=14; C: No autotransfusion, n=12 (both groups also received crystalloids and colloids)	I: autotransfusion (Bellovac ABT autotransfusion system), n=35 (hip) and n=45 (knee); C: regular post- operative low-vacuum drainage, n=48 (hip) and n=32 (knee)	I: autotransfusion (Dideco 797 transfusion system), n=35, C: standard Hemovac suction drain, n=35
H d	Knee	Knee	NAS	Н Ф	Hip and knee	Knee
total hip artrhoplasty	elective primary total knee replacement surgery	primary unilateral total knee arthroplasty	elective primary total hip arthroplasty and total knee arthroplasty		primary total knee arthroplasty or total hip arthroplasty	unilateral total knee replacement
T66T	deM 2991	Majowski 1991	Mauerhan 1993	1992 Tees	Moonen 7007	1997
2400001	7~14	MarroloM		-525574	-3000V4	

Low risk	High risk	High risk	High risk	High risk	High risk	High risk
Unclear risk	Unclear risk	High risk	Unclear risk	Unclear risk	Unclear risk	Unclear risk
Low risk	Unclear risk	Unclear risk	Unclear risk	Unclear risk	Low risk	Unclear risk
Ä.	Yes	N.A.	Yes	Yes	Yes	Unknown
Control 1	Control 0	Active	Control 0	Control 1	Control 0	Control 1
Trigger 2	Trigger 2	None	Trigger 2	None	Trigger 2	Trigger 2
Htt < 25% (i.e. Hb < 8.5 g/dl)	Hb < 9.0 g/dl	Based on clinical condition of patient	Hct at 30% (i.e. Hb <10.2 g/dl)		Hb < 9.0 g/dl	Hb < 10 g/dl or Hct < 30% (i.e. Hb < 10.2 g/dl)
Yes	Yes	NO	Yes	No	Yes	Yes
POST	POST	PERI/PO ST	POST	POST	POST	POST
I: autotransfusion (Solcotrans), n=25; C: postoperative drained blood collected into Solcotrans Orthopedic Plus system but salvaged blood was not considered for reinfusion, n=25	I: autotransfusion (Solcotrans), n=78 (hip) and n=137 (knee); C: no drainage system, n=62 (hip) and n=138 (knee)	I: 1. autotransfusion (Haemonetics), n=35; 2. autotransfusion (Solcotrans), n=40; C: no drain, n=38	I: autotransfusion (Ortho-Evac system or Solcotrans), n=20; C: no drain, n=10	I: autotransfusion, n=60; C: standard care without autotransfusion, n=60	I: autotransfusion (Haemonetics Cell Saver 3), n=50; C: no drain, n=50	I: autotransfusion (Solcotrans), n=12; C: closede suction drain, n=12
Other ortho paedic	Hip and knee	Hip	Knee	Knee	Knee	NAS
elective, non- emergency spinal surgery	primary total hip or total knee replacement	primary total hip arthroplasty	knee replacement surgery	total knee arthroplasty	total knee replacement	elective primary total joint arthroplasty
Jeer 1994	Ritter 1994	T995	January Januar	Jis2 1999	Shenolikar 1997	nosqmi2
ia	Ap++iq	olioa	Rocencher	+102	acdiloned2	aosami2

High risk	High risk	High risk	High risk	High risk	Low risk
Unclear risk	High risk	High risk	Low risk	Unclear risk	Low risk
Unclear risk	Low risk	Low risk	Low risk	Unclear risk	Low risk
ON.	N.A.	Yes	Yes	Yes	Z.A.
Active	Control 1	Control 1	Control 1	Control 1	Control 1
None	Trigger 1	None	Trigger 1	Trigger 2	Trigger 2
	Hb < 8.0 g/dl and in symptomatic patients with Hb of 8.0- 10.0 g/dl: 2 units		Hb < 6.4 for age < 60 years; Hb < 8.1 g/dl for age > 60 years; Hb < 9.6 g/dl in high risk	Hb < 9.0 g/dl	Hb < 8.5 g/dl or clinical symptoms of anaemia
O _N	Yes	ON	Yes	Yes	Yes
POST	POST	POST	PERI/	POST	PERI
I: autotransfusion (Hemolite cell salver), n=24 (hip) and n=27 (knee); C: Hemovac standard drainage system, n=26 (hip) and n=25 (knee)	I: autotransfusion (ABTrans autologous re- transfusion system), n=76, C: two standard Medinorm vacuum drains, n=82	I: autotransfusion (DONOR or Bellovac ABT autotransfusion system), n=35 (hip) and n=12 (knee); C: standard closed suction wound drainage, n=11 (hip) and n=11 (knee)	I: 1. autotransfusion (OrthoPat), n= 412 (hip); 2. Autotransfusion (Donor or Bellovac ABT autotransfusion system), n=419 (hip) and n=436 (knee); C. low vaccum wound drain, n=419 (hip) and n=417 (knee)	I: autotransfusion (Haemonetics Cell Saver 5), n=115, C: all drainaged blood was discarded, n=116	I: autotransfusion (Sangvia Blood Management System), n=106; C: regular postoperative low vacuum drain, n=110
Hip and knee	Нір	Hip and knee	Hip and knee	Knee	Hip
total hip or knee replacement	primary total hip replacement	primary or revision total hip or knee replacement	primary or revision total hip or knee replacement	total knee replacement	primary or revision total hip arthroplasty
sigel2 1991	Z007	nsm20-o2	nsm2O-o2	Thomas Thomas	Thomassen

High risk	High risk	High risk
Unclear risk	Unclear risk	Unclear risk
Unclear risk	Unclear risk	Unclear risk
N.A.	Yes	N.A.
Active	Control 1	Control 1
Trigger 2	Trigger 2	None
Hb < 10 g/dl or Hct < 30% (i.e. Hb < 10.2 g/dl)	Hb < 9.0 g/dl Trigger 2	
Yes	Yes	No
POST	POST	INTRA
I: autotransfusion (BIODREN system), n=30; C: no autotransfusion, n=30	I: autotransfusion (Gish Orthofuser system), n=30; C: standard wound suction drainage system, n=30	I: autotransfusion (Haemonetics Cell Saver 5 system), n=20; C: standard care, n=20
Hip	Knee	NAS
primary total hip replacement	unilateral total knee replacement	orthopaedic procedures
2008	2007	2008
Tripkovic	Zacharapoulos	BnsdZ

¹ Type: Hip, knee, hip and knee or not able to split (NAS).

² Timing Cell Salvage: POST = postoperative, INTRA = intraoperative, PERI = both intra- and postoperative.

⁴ Treatment policy in control group: Control 0 = in control groups no drain is used; Control 1 = in control group standard suction or vacuum drain is used; Active = in control group active treatment (active plus cell salvage versus active comparisons). ³ Subgroup: Trigger 1 Hb=<8.0 g/dl; Trigger 2 Hb> 8.0 g/dl

