Restrictive versus Liberal Transfusion Strategy in the Perioperative and Acute Care Setting. A Context-specific Systematic Review and Meta-analysis of Randomized Controlled Trials

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ABSTRACT

Background: Blood transfusions are associated with morbidity and mortality. However, restrictive thresholds could harm patients less able to tolerate anemia. Using a *context-specific* approach (according to patient characteristics and clinical settings), the authors conducted a systematic review to quantify the effects of transfusion strategies.

Methods: The authors searched MEDLINE, EMBASE, CENTRAL, and grey literature sources to November 2015 for randomized controlled trials comparing restrictive *versus* liberal transfusion strategies applied more than 24 h in adult surgical or critically ill patients. Data were independently extracted. Risk ratios were calculated for 30-day complications, defined as inadequate oxygen supply (myocardial, cerebral, renal, mesenteric, and peripheral ischemic injury; arrhythmia; and unstable angina), mortality, composite of both, and infections. Statistical combination followed a *context-specific* approach. Additional analyses explored transfusion protocol heterogeneity and cointerventions effects.

Results: Thirty-one trials were regrouped into five *context-specific* risk strata. In patients undergoing cardiac/vascular procedures, restrictive strategies seemed to increase the risk of events reflecting inadequate oxygen supply (risk ratio [RR], 1.09; 95% CI, 0.97 to 1.22), mortality (RR, 1.39; 95% CI, 0.95 to 2.04), and composite events (RR, 1.12; 95% CI, 1.01 to 1.24—3322, 3245, and 3322 patients, respectively). Similar results were found in elderly orthopedic patients (inadequate oxygen supply: RR, 1.41; 95% CI, 1.03 to 1.92; mortality: RR, 1.09; 95% CI, 0.80 to 1.49; composite outcome: RR, 1.24; 95% CI, 1.00 to 1.54—3465, 3546, and 3749 patients, respectively), but not in critically ill patients. No difference was found for infections, although a protective effect may exist. Risk estimates varied with successful/unsuccessful transfusion protocol implementation.

Conclusions: Restrictive transfusion strategies should be applied with caution in high-risk patients undergoing major surgery. **(ANESTHESIOLOGY 2016; 125:00-00)**

D ESPITE studies suggesting unfavorable outcomes after the administration of erythrocytes,¹⁻⁵ the optimal transfusion strategy in surgical and critically ill patients remains unclear. Concerns have been raised about harmful effects of low hemoglobin transfusion thresholds in individuals less able to tolerate anemia, such as the elderly and patients with cardiovascular disease or cancer.⁶⁻⁹ Previously published meta-analyses were inconclusive: minimizing exposure to allogeneic blood reduced the risk of infection, but patients assigned to these restrictive transfusion strategies seemed also at a higher risk of myocardial infarction (MI).^{10,11}

Since variability among studies is inevitable, undertaking meta-analyses generally entails some degree of heterogeneity, of which three different subtypes have been described¹²: (1) *clinical heterogeneity*, which results from

What We Already Know about This Topic

 Although many studies and some systematic reviews have examined the role of transfusion strategies in patient morbidity and mortality, these have not included the role for contextspecific (patient characteristics and clinical setting) conditions

What This Article Tells Us That Is New

 In a review of 31 trials grouped into 5 context-specific strata, restrictive transfusion strategies increased the risk of mortality and composite morbidity in patients undergoing cardiac/vascular procedures and in elderly orthopedic patients

variability in participants, interventions, or outcomes; (2) *methodologic heterogeneity*, a consequence of variability in study design and risk of bias; (3) *statistical heterogeneity*,

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This article is featured in "This Month in Anesthesiology," page 1A. Corresponding article on page XXX. James C. Eisenach, M.D., served as Editor-in-Chief for this article. Supplemental Digital Content is available for this article. Direct URL citations appear in the printed text and are available in both the HTML and PDF versions of this article. Links to the digital files are provided in the HTML text of this article on the Journal's Web site (www.anesthesiology.org).

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which results from clinical or methodologic heterogeneity, or both. Identifying and addressing each type of heterogeneity remains a key step in undertaking meta-analyses. To date, however, most systematic reviews on transfusion strategies failed to address *clinical heterogeneity*, thereby limiting their interpretation.^{13–16}

Therefore, we conducted a *context-specific* systematic review and meta-analysis of randomized controlled trials (RCTs) investigating the effects of restrictive transfusion strategies in the perioperative and acute care settings. The rationale for a *contextual* approach (*i.e.*, stratification of the analysis according to patient characteristics and clinical settings) was based on the prespecified assumption that a high degree of clinical heterogeneity may hinder the identification of group-specific effects: pooling data from various patient populations (young and elderly patients, for instance) or from various settings (such as cardiac surgery and postpartum setting) may result in a dilution of the intervention effects. Since clinical diversity may also result from variability in study interventions, we were also interested in the effects of different transfusion protocols and in the contributing role of cointerventions (*i.e.*, administration of non-erythrocyte blood products, hemostatic agents, or intravenous fluids) in complication rates.

Materials and Methods

We followed the Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines.¹⁷ Eligibility criteria, outcomes, and methods of analysis were prespecified (study protocol available at: http://alfredanaesthesia.org.au/ research).

Eligibility Criteria

Only fully published reports of RCTs were included. For duplicates or follow-up or ancillary studies, the first published article was considered the main study.¹⁸ Crossover designs or studies not adequately controlled were excluded. Trials evaluating a multiinterventional protocol were excluded if the effect of transfusion strategies could not be distinguished from the effect of other interventions. Cluster randomized trials were included only if methods of analysis allowed for clustering.¹⁹

Only trials conducted among adult patients (more than 18 yr old) in the perioperative, emergency, or intensive care settings were considered. We excluded studies conducted in patients with sickle cell disease.

We searched for studies comparing two different laboratory values (or using symptoms of anemia) to guide erythrocyte administration. We excluded studies applying transfusion strategies 24 h or less, trials using a hemodilution protocol, and interventions relying on preoperative autologous blood donation, since some effects (such as immunomodulation) were unlikely to develop,²⁰ and this approach is no longer recommended.²¹

We were interested in studies reporting events associated with, or worsened by, anemia.^{5,22–28} To fully capture all the effects of transfusion strategies, 29,30 individual outcome events were combined into the following categories: "inadequate oxygen supply" (myocardial, cerebral, renal, mesenteric, and peripheral ischemic injury; arrhythmia; unstable angina), "mortality," and a composite category "inadequate oxygen supply + mortality" (see description provided in Supplemental Digital Content 1, http://links.lww.com/ALN/ B275, which describes outcome categories). Only events occurring within 30 days were retrieved because substantial hemoglobin recovery seems to occur within 2 months after surgery/intensive care unit stay.^{31,32} Our aim was also to explore the immunomodulatory effects of allogeneic blood: since transfusions have been associated with impaired host defenses,^{33,34} we searched for studies reporting new infections occurring within 30 days.

Data Sources and Searches

We performed a systematic electronic search in the MED-LINE (Ovid), EMBASE (Ovid), and Cochrane CENTRAL databases. Both MeSH terms and keywords combined with Boolean operators were used (see strategy provided in Supplemental Digital Content 2, http://links.lww.com/ALN/ B276, which provides the full search strategy used in this systematic review). The following sources of grey literature were screened: OpenGrey, International Clinical Trials Registry Platform, ClinicalTrials.gov. Additional reports were identified by hand-searching bibliographies. No language or date restrictions were applied. The last electronic search was done on November 17, 2015.

Study Selection

Titles and abstracts were assessed for eligibility by two independent reviewers (Drs. Hovaguimian and Myles). Duplicate publications were identified through comparison of reports for author names, enrolment date, setting, intervention, participant number, or baseline data. Disagreements were resolved through discussion.

Data Extraction

Data were extracted from original reports by one reviewer (Dr. Hovaguimian) and entered in a form specifically designed for this review (see description provided in Supplemental Digital Content 3, http://links.lww.com/ALN/B277, which details which information was extracted). The second reviewer (Dr. Myles) verified these data, and queries were resolved through discussion. Missing, unclear, or incomplete data in the original report were clarified by contacting authors (two provided additional data).^{35,36} Outcome data were not considered for analysis if no clarification could be obtained. Data from duplicates were extracted and merged under a unique study identification name. Data were subsequently entered into the Cochrane Review Manager software (RevMan, version 5.3.3—The Cochrane Collaboration, The

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Nordic Cochrane Centre, Denmark, 2014) by one reviewer (Dr. Hovaguimian) and checked by the second reviewer (Dr. Myles).

Risk of Bias in Individual Studies

The risk of bias was assessed using the Cochrane "Risk of bias" tool, which evaluates randomization method, concealment of treatment allocation, blinding of participants and personnel, blinding of outcome assessor, risk of incomplete outcome data, risk of selective reporting, and other sources of bias (ethics approval, informed consent, funding, and conflict of interest).³⁷ Each item was rated at "low," "unclear," or "high" risk of bias. The effects of detection and attrition bias were specifically explored, since this may affect studies evaluating adverse events (AEs).³⁸ For cluster randomized trials, we used specific items as recommended elsewhere.¹⁹ Disagreements were resolved through discussion.

Measures of Effect, Data Handling, and Transformation

Dichotomous outcomes were reported as risk ratios with 95% CIs, while continuous data were expressed as weighted mean differences with 95% CI. All statistical analyses were performed with the Cochrane Review Manager software. Data handling and transformation were necessary for some endpoints. Composite outcome categories were obtained by combining individual outcome data, as performed in previous reports (see description provided in Supplemental Digital Content 4, http://links.lww.com/ALN/ B278, which outlines data handling, transformation, and combination).^{10,13}

Data Synthesis and Analysis

Differences between studies in terms of patient characteristics and/or clinical settings (*i.e.*, clinical diversity) may affect effect estimates.¹² To control for these sources of heterogeneity, we used a prespecified *context-specific* approach and stratified the analysis by (1) patient-specific risk of developing complications (according to age, comorbidities, and concomitant medication); (2) setting-related risk of complications (type of surgery). Studies conducted in similar populations and settings were regrouped into *risk strata* (see description provided in Supplemental Digital Content 5, http://links.lww.com/ALN/B279, which explains the rationale and methods used for strata generation). Meta-analyses were performed only if data were obtained from at least two studies.

To assess if our context-specific approach was sensible, we conducted for each outcome category a nonstratified analysis (*i.e.*, data pooling without controlling for clinical diversity) and performed a test of interaction using the Cochran Q and Higgins l^2 . We considered that different population parameters were represented within each risk stratum when the Cochran Q *P*-value was less than 0.05 or when l^2 was greater than 50%.^{12,39} When data combination was deemed inappropriate, a qualitative assessment was performed.

Statistical heterogeneity was assessed by visual inspection of forest plots and by using the chi-square test and the I² statistic. When data were heterogeneous (P < 0.1, I² greater than 50%), we searched for methodologic sources of heterogeneity.¹² We used a fixed-effect model, unless overt clinical or residual statistical heterogeneity was present (see fig., Supplemental Digital Content 6, http://links.lww.com/ ALN/B280, which outlines how heterogeneity assessment was performed).^{39,40}

Additional Analyses

Effect of "Successful Studies." We assumed that only studies demonstrating successful transfusion protocol implementation would reflect true intervention effects. Success was arbitrarily defined as a statistically significant difference between transfusion groups in two performance indicators: (1) hemoglobin levels over time *and* (2) mean erythrocyte units/group. *P* values were assessed from original reports. We also explored other possible determinants of success, such as hemoglobin thresholds, transfusion-sparing effect, and adherence rates (see description provided in Supplemental Digital Content 7, http://links.lww.com/ALN/B281, which outlines which indicators were used to assess successful protocol implementation).

Effect of "Cointerventions." Several routinely administered drugs or other management measures may participate in the occurrence of AE. Thus, we assessed the effects of nonerythrocyte blood products (cryoprecipitate, fresh frozen plasma, and platelets), antifibrinolytics, clotting factor concentrates, and fluids administration.

Results

Study Selection

We identified 4,684 records from MEDLINE, 858 records from EMBASE, and 1741 records from CENTRAL (fig. 1). Other sources retrieved no additional records. Of these 7,283 records, 7,193 were excluded after preliminary screening. Of the 90 remaining reports, 53 were discarded because the study population was ineligible, the intervention was inappropriate, the design was problematic, or the study presented other issues (see description provided in Supplemental Digital Content 8, http://links.lww.com/ALN/ B282, which lists excluded studies). Of 37 relevant reports, 6 were excluded after more thorough examination: 2 were duplicate publications, 1 was a preliminary analysis, and 3 were ancillary or follow-up studies of included studies.⁴¹⁻⁴⁶ Thus, we included 31 RCTs comparing restrictive with liberal transfusion strategies in the perioperative or acute care setting.6-9,35,36,47-71

Study Characteristics

Study Design, Participants, and Setting (table 1). Included studies were published between 1956 and 2015, and all were reported in the English language. A two-arm parallel design

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Fig. 1. Flow diagram illustrating the study selection process. ICU = intensive care unit; RCT = randomized controlled trial.

was used in all trials except in a factorial 2×2 design investigating transfusion strategies and erythropoietin administration.⁶⁵ One study was a cluster randomized trial.³⁵

Trials with similar patient populations and clinical settings were regrouped into five risk strata: (1) patients with cardiovascular disease undergoing cardiac or vascular procedures (surgery or catheterization-8 trials, 3,323 patients)^{6,47,48,51,55,60,66,67}; elderly population with varying cardiovascular dis-(2)undergoing orthopedic surgery (9 trials, 3,777 ease patients)^{8,9,49,50,52,54,62,63,68}; (3) mixed surgical and medical patients with varying comorbidities admitted to an acute care facility (emergency or intensive care unit-10 trials, 4,129 patients)7,35,36,53,56-59,69,70; (4) younger, less comorbid population admitted for acute traumatic brain injury or subarachnoid hemorrhage (2 trials, 244 patients)^{61,65}; (5) other patients or settings: one conducted among anemic women in the postpartum phase and one including thrombocytopenic middle-aged patients with hematologic cancer (2 trials, 579 patients).^{64,71}

Intervention: Transfusion Protocol (table 2). In the first group, four studies failed in implementing their protocol: in two, the absolute difference in hemoglobin thresholds was only 10 g/l, the transfusion-sparing effect was less than 20%, and nonadherence rates in the restrictive group were more than or equal to 15%;^{47,48} in one, transfusion thresholds were higher in some patients of the restrictive group compared to liberal⁶⁷; in one, nonadherence rates, transfusion-sparing effects, and transfusion thresholds seemed adequate, but the sample size was small.⁵¹

In group 2, data to assess success were lacking in one trial,⁶³ and three studies showed unsuccessful implementation: in

one, the transfusion-sparing effect was less than 20%, and nonadherence rates in the restrictive group were more than or equal to 15%⁶²; in one, the lack of standardized protocol in the liberal group resulted in a negative transfusion-sparing effect⁶⁸; and in one, surgical procedures were not adequately balanced between groups (liberal patients were more likely to bleed).⁸

In group 3, success was unclear in two studies ("mean erythrocyte units" were reported *per transfused patients* instead of *per strategy group*).^{7,53} In one unsuccessful study, the transfusion-sparing effect was less than 20%, and nonadherence rates in the liberal group were more than or equal to 15%.³⁵

Group 4 included two studies that demonstrated successful implementation but the protocols were heterogeneous: one study used particularly high thresholds (restrictive: hemoglobin 100 g/l; liberal: hemoglobin 115 g/l), while the other compared 70 with 100 g/l. 61,65

In the last group, success was unclear in one study reporting negligible transfusion-sparing effects and high nonadherence rates in the restrictive group.⁷¹

Risk of Bias in Included Studies

The risk of selection bias was deemed low in only 7 of 31 included studies (see fig., Supplemental Digital Content 9, http://links.lww.com/ALN/B283, which summarizes the risk of bias in included studies).^{7,56,58,66,68,70,71} All studies were at high or unclear risk of performance bias. Outcome assessors were blinded in 13 studies.^{6–8,36,49,50,54,55,58,60,61,65,71} The risk of attrition bias was low in all studies except three: in two,^{54,67} it was not possible to assess if attrition was balanced between groups, and in one,⁶⁴ attrition rates exceeded 20%. The risk of selective reporting was low in all studies, but in one trial, study findings were reported in three different publications without mention of other existing reports.^{9,41,42}

Results of Individual Studies and Data Syntheses Events Associated to or Worsened by Anemia.

Inadequate Oxygen Supply. (See fig., Supplemental Digital Content 10, http://links.lww.com/ALN/B284, which illustrates the risk of events reflecting inadequate oxygen supply.) In group 1, early MI was reported in seven studies, arrhythmia in five, angina in two, stroke or transient ischemic attack (TIA) in five, acute kidney injury in five, and mesenteric ischemia in one (see table, Supplemental Digital Content 11, http://links.lww.com/ALN/B285, which outlines outcome reporting across studies). In one study, stroke/TIA was reported in combination with delirium and could not be extracted.⁵⁵ Thus, data from 8 studies (3,322 patients) were combined^{6,47,48,51,55,60,66,67}: in patients with cardiovascular disease assigned to a restrictive strategy and undergoing high-risk surgery, there was a possible increase in events reflecting inadequate oxygen supply (risk ratio [RR], 1.09; 95% CI, 0.97 to 1.22).

In group 2, early MI was reported in five studies, arrhythmia in three, stroke/TIA in five, and AKI in two (see table,

Study Characteristics				Ра	atient Charac	teristics		
					Mean Age	(yr) Partio	cular Medicat	ion n (%)
Study ID	Design	Total Participants (n)	Setting or Reason for Admission	Particular Population	Restrictive Li	beral Namo	e Restri	ctive Liberal
Group 1: CV disease, carc Bracey <i>et al.</i> , 1999⁴7	liac/vascular surgen Two-arm RCT	/ or interventional - 428	catheterization Elective cardiac (CABG)	No comorbidities reported	61 6	62 Aspirin Oral AC	151 (7 4 (2	(1) 147 (68) () 4 (2)
Bush <i>et al.</i> , 1997 ⁴⁸	Two-arm RCT	66	Elective vascular /activeinfrainquinal/	Variety of CV diseases	66	Other ant	i-PLT 11 (5 NR	11 (5)
Carson <i>et al.</i> , 2013 ⁶	Two-arm RCT	110	catheterization	Variety of CV diseases, hemoglobin < 100 g/l	74 6	7 Aspirin Other ant Chronic A	48 (8 i-PLT 44 (8	(7) 51 (93) (0) 45 (82)
Cooper <i>et al</i> ., 2011 ⁵¹	Two-arm RCT	45	Interventional catheterization or cardiac surgery	Variety of CV diseases, hematocrit < 30%	202	6 Aspirin Other ant Heparin Gn antaoi	24 (1 i-PLT 21 (8 10 (4 10 (4	00) 21 (100 88) 17 (81) 1) 7 (33) 1 (5)
Hajjar <i>et al.</i> , 2010 ⁵⁵	Two-arm RCT	502	Elective cardiac (CABG, valve)	Variety of CV diseases; low-risk of bleeding*	59 6	1 Aspirin	94 (3	() (41) (41) (41)
Murphy <i>et al.</i> , 2015 ⁶⁰	Two-arm RCT	2003	Elective cardiac (CABG, valve, aortic)	Variety of CV diseases; low-risk of bleeding*	202	1 Aspirin Other ant	277 (2 i-PLT 41 (4	(28) 284 (28) (3.7) 37 (3.7)
Shehata <i>et al.</i> , 2012 ⁶⁶	Two-arm RCT	50	Elective cardiac (CABG, valve)	High risk of transfusion†	67 6	9 Other ant Oral AC	i-PLT 18 (7 7 (2	2) 17 (68) 8) 8 (32)
Slight <i>et al.</i> , 2008 ⁶⁷	Two-arm RCT	86	Elective cardiac (CABG, valve)	Variety of CV diseases	65 6	90	NR	• •
Group 2: Elderly, orthopedi Carson <i>et al.</i> , 1998⁴ ⁹	c surgery Two-arm RCT	84	Hip fracture	Mostly ASA III with CV disease (45% IHD), mostly community dwellind	83	11 Aspirin Chronic⊿	12 (2 C 14 (3	.8.6) 15 (35.7 .3.3) 15 (35.7
Carson <i>et al.</i> , 2011 ⁵⁰	Two-arm RCT	2016	Hip fracture	Mostly ASA III with CV disease (40% IHD). mostly community dwelling	82	52	NR	
Fan <i>et al.</i> , 2014 ⁵²	Two-arm RCT	186	Elective lower limb joint replacement (hip)	Mostly ASA II, some with CV disease (only 10% IHD)	73 7	5	NR	
Foss <i>et al.</i> , 2009 ⁸	Two-arm RCT	120	Hip fracture	Mostly ASA III with CV disease (12% IHD), community dwelling	81	1	NR	
Gregersen <i>et al.</i> , 2015 ⁹	Two-arm RCT	284	Hip fracture	21% with CV disease, nursing home or sheltered housing	86	37 Iron suppler	123 (8 nent	8) 118 (87)
Grover et al., 2006 ⁵⁴	Two-arm RCT	218	Elective lower limb joint replacement	Some with CV disease (only 12% IHD)	71 7	5	NR	
Nielsen <i>et al.</i> , 2014 ⁶² Parker, 2013 ⁶³	Two-arm RCT Two-arm RCT	66 200	Elective hip revision Hip fracture	Mostly ASA II, only NYHA I Mostly ASA III with CV disease (only 15% IHD), community dwelling	68 84 8	2 4	AN AN AN	
So-Osman <i>et al.</i> , 2010 ⁶⁸	Two-arm RCT	603	Elective lower limb joint replacement	Mostly "high-risk" patients‡	17	'0 No signifi and AC	cant differenc . No values n	ce for NSAID eported.
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Study ID		- - -	: : :		Mean Ag	e (yr)	Particular M	1edication n ((%)
	Design	Iotal Participants (n)	Setting or Heason for Admission	Particular Population	Restrictive	Liberal	Name	Restrictive	Liberal
Group 3: Mixed medical/surgical	cases, acute ca	Ð							
de Almeida <i>et al.</i> , 2015 ⁷ Tv	wo-arm RCT	198	Surveillance postmajor abdominal surgery	Low risk of bleeding*; some with CV disease (7% IHD)	64	64		NR	
Fortune <i>et al.</i> , 1987 ⁵³ Tv	wo-arm RCT	25	Trauma or surgical bleeding	Hemorrhagic shock class III or IV, all intubated; no history of MI	47	47		NR	
Hébert et al., 1995 ⁵⁷ Tv	wo-arm RCT	69	Various diagnoses	Mixed surgical and medical population with hemoglobin < 90 g/l; some with CV disease	59	59		NR	
Hébert <i>et al.</i> , 1999 ⁵⁶ Tv	wo-arm RCT	838	Various diagnoses	Mixed surgical and medical population with hemoglobin < 90 g/l	57	58		NR	
Holst <i>et al.</i> , 2014 ⁵⁸ Tv	wo-arm RCT	1,000	Septic shock	Mixed surgical and medical population with hemoglobin < 90 g/l; some with CV disease	67	67		NR	
Jairath <i>et a</i> /., 2015 ³⁵ Tv	wo-arm cluster RCT	6 clusters, 936 patients	Upper-GI bleeding	No exsanguinating bleeding; some with CV disease (15% IHD)	58	60 Iroi LN	n supplement //WH	43 (11) 9 (2)	47 (9) 5 (1)
Markatou <i>et al.</i> , 2012 ⁵⁹ Tv	wo-arm RCT	52	Surveillance postmajor abdominal surgery	Low risk of bleeding*	58	63		NR	
Topley and Fisher, 1956 ⁶⁹ Tv	wo-arm RCT	22	Trauma	Actively bleeding, no elderly patient	NR	NR 50		NR 47 (44)	60 (10 E)
VIIIarlueva <i>et al.</i> , ∠U13°		0000	upper-ai pieeairig	Excluded It exsarigumating predung or major cardiovascular disease	P C	28 011		4/ (11) 4	(0.61) 00
Walsh <i>et al.</i> , 2013 ³⁶ Tv	wo-arm RCT	100	Various diagnoses	Mixed surgical and medical popula- tion with hemoglobin < 90 g/l, mechanically ventilated; some with CV disease (32% IHD)	67	68		NR	
Group 4: Younger, fitter, brain inj	jury/intracranial	bleeding							
Naidech <i>et al.</i> , 2010 ⁶¹ Tv	wo-arm RCT	44	Neuro-ICU	Subarachnoid hemorrhage, some with CV disease	59	54		NR	
Robertson <i>et al.</i> , 2014 ⁶⁵ F ₆	actorial 2×2	200	Neuro-ICU	Closed traumatic brain injury, GCS motor ≤ 5, no major chronic disease or AC	28	31 Iro	n supplement oO	99 (100) 1 49 (50)	101 (100) 53 (52)
Group 5: Other patients and set	tings								
Prick <i>et al.</i> , 2014 ⁶⁴ Tv	wo-arm RCT	519	Postpartum hemorrhage	Hemodynamically stable ASA I women with hemoglobin 48–79 g/l; some had surgery	31	31		NR	
Webert <i>et al.</i> , 2008 ⁷¹ Tv	wo-arm RCT	60	Hematooncology	Thrombocytopenic patients, no IHD in past 6 months, no coagulation disorders	51	45		NR	
"Patients with thrombocytopenia, cc ‡Defined as nonsinus rhythm, unstal ischemic attack, left ventricular hype Anti-PLT = antiplatelet therapy; ASA = A	pagulopathy, or ch ble ischemic heard artrophy (electroca American Society of	rronic anticoagulat disease (IHD), or r rdiogram/transthor Anesthesiologists (s	ion (AC) therapy were exclude mycoardial infarction (MI) < 6 r acic echocardiography), chror score); CABG = coronary artery k	ed. †Risk assessment based on comorbiditie mo, heart failure, valvular disease, age > 70 y nic pulmonary disease with polyglobulism, an sypass graft, CV = cardiovascular, EPO = enythro	es and comp //r, peripheral a nd insulin-dep opoietin; GCS	lexity of surg artery diseas bendent diab = Glasgow O	gery (as reporte se, large vessel s betes mellitus. toma Scale; GI = (d in the origin surgery, stroke gastrointestinal;	al article). #/transient ; Gp = gly-

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Volume, Mear Hematocrit)	NR	NR	Mainly LeukoR	NR	Non-LeukoR, 250-320 ml, hematocrit	80% NR	e NR
Follow-up	Discharge	Discharge	6 mo	30 d	30 d	3 mo	Discharge
/ Protocol Application	РОР	IOP + POP	Hospital stay (maximum	30 d) Discharge	IOP + POP (ICU only)	РОР	10P + POP

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	Study ID	Setting or Reason for Admission	Restrictive	Liberal	∇	Restrictive	Liberal 2	∆ Re	strictive Libe	Mea Erythro UI Diffe Significa betwe ral Group	n cyte Hemo red Lev antly Diffe en Signifiu s? over T	globin els ⊮red cantly ïme? A	Protocol	Follow-up	Type of Erythrocyte (Leukoreduction: Volume, Mean Hematocrit)	, Restrictive	Libe
Bush et al. 1997*Excitor viscular intraviound)00100201267Ves100-PCPDischargeNNN1997*intraviound) intraviound)3 (s0)10020736 6 727YesHospital intraviound)6 motication intraviound)20736 6 727YesHospital intraviound)6 motication intraviound)207020207020207020207020	Group 1: CV Bracey <i>et al.</i> 1999 ⁴⁷	' disease, cardiac/v , Elective cardiac (CABG)	ʻascular surgt 80	ery or interv 90	ention: 10	al cathete 65	rization 52 1	13	15 16	Yes	Ż	о В	ЧC	Discharge	RN	NR	ЧN
Casesnet et al. 2013 ⁻¹ Interventional catheterization S (8) 100 20 73 6 6.7 2 7 Ves Hespital fragminun surgery 30 d) Emol NR Mainly reaction 233 M	Bush <i>et al.</i> , 1997 ⁴⁸	Elective vascular (aortic + infrainguinal)	06	100	10	20	12	œ	20 NI	°N N	¥	OI Sé	POP + POP	Discharge	NR	NR	RN
$ \begin{array}{l l l l l l l l l l l l l l l l l l l $	Carson <i>et al.</i> , 2013 ⁶	Interventional catheterization	S (80)	100	20	73	0	22	2	Yes	¥	ж	ospital stay (maximum 30 d)	6 mo	Mainly LeukoR	23	25
Hajlar et al., 2010 ⁵⁶ Elective cardiac (CABG, valve)80100207036470830Non-LeukoR, anos32010 ⁵⁶ (CABG, valve)79015478301055300Non-LeukoR, anos306Non-LeukoR, anos3Non-LeukoR, anos33Non-LeukoR, anosNo	Cooper <i>et al</i> 2011 ⁵¹	 Interventional catheteriza- tion or cardiac surgerv 	80	100	20	46	0	9	4	No	¥	SS Di	scharge	30 d	R	RN	RN
	Hajjar <i>et al.</i> , 2010 ⁵⁵	Elective cardiac (CABG, valve)	80	100	20	20	36 3	34	5	Yes	¥	O Se	ICU only)	30 d	Non-LeukoR, 250–320 ml, hematocrit 80%	ო	က
Shehata et al., Elective cardiac 70 lOP, 95 lOP, 100 POP, 100	Murphy <i>et al.</i> , 2015 ⁶⁰	Elective cardiac (CABG, valve, aortic)	75	06	15	47	က ထ	6	10	, Yes	¥	se P(AC	3 mo	NN	NR	ЯN
Slight <i>et al.</i> , Elective cardiac RCV 80–90 NA 67 47 20 9 5 Yes No POP (48h 3 mo NR NR 20087 (CABG, valve) schema CaBG, valve) schema Group 2: Elderly, orthopedic surgery Group 2: Elderly, orthopedic surgery Carson <i>et al.</i> , Hip fracture S (80) 100 (20) 55 2 53 10 2 Yes Yes POP 60 d NR NR $^{1908^{40}}$ Carson <i>et al.</i> , Hip fracture S (80) 100 (20) 55 2 53 10 2 Yes Yes POP 60 d LeukoR 220150 Carson <i>et al.</i> , Elective lower 80 100 20 56 44 12 NR NR Yes Yes POP 105charge? NR NR $^{2014^{52}}$ Imb joint replacement (hip)	Shehata <i>et al.</i> 2012 ⁶⁶	., Elective cardiac (CABG, valve)	70 IOP, 75 POP	95 IOP, 100 POP	25	48	12 3	36	NA AN	A Yes	¥	Se O	POP + POP	Discharge	» NR	NR	ЧN
Carson et al., Hip fracture S (80) 100 (20) 55 2 53 10 2 Yes POP 60 d NR N	Slight <i>et al.</i> , 2008 ⁶⁷ Group 2: Eld	Elective cardiac (CABG, valve) erly, orthopedic sui	RCV schema rgery	80-90	NA	67	47 2	0	u) O	Yes	z	о Д	DP (48h only)	3 mo	RN	NR	N
Carson <i>et al.</i> , Hip fracture S (80) 100 (20) 59 3 56 6 9 Yes Yes POP 60 d LeukoR 22 201 ⁵⁰ 201 ¹⁵⁰ Elective lower 80 100 20 56 44 12 NR NR Yes Yes IOP + POP Discharge? NR NR NR Pop 2014 ⁵² limb joint replacement (hip)	Carson <i>et al.</i> , 1998 ⁴⁹	Hip fracture	S (80)	100	(20)	55	25	33	10 2	Yes	¥	sé P	ЧC	60 d	NR	NR	ЧN
Fan <i>et al.</i> , Elective lower 80 100 20 56 44 12 NR NR Yes Yes IOP + POP Discharge? NR NR 2014 ⁵² limb joint replacement (hip)	Carson <i>et al.</i> , 2011 ⁵⁰	Hip fracture	S (80)	100	(20)	59	35	99	9) Yes	¥	sé P(ЧС	60 d	LeukoR	22	20
	Fan e <i>t al.</i> , 2014 ⁵²	Elective lower limb joint replacement (hip)	80	100	20	56	44 1	N	NR	R Yes	≯	Se IO	404 + 90P	Discharge?	RN	N	ЧN

Study Charact	eristics							Interve	ntion Charac	cteristics					
		Hemo Thresh	oglobin (l/g) blor		Not Expr Allogeneic f	sed to 3lood (%)	Nonadh (%	erence* 6)	Succ	Sess				Mean St Time	orage (d)
Study ID	Setting or Reason for Admission	Restrictive	Liberal	⊲	Restrictive	Liberal ∆	Restrictiv	e Liberal	Mean Erythrocyte UI Differed Significantly between Groups?	Hemoglobin Levels Differed Significantly over Time?	 Protocol Application 	Follow-up	Type of Erythrocyte (Leukoreduction, Volume, Mean Hematocrit) F	testrictive	Liberal
Foss <i>et al.</i> , 2009 ⁸	Hip fracture	80	100	20	63	27 36	NR	NR	Yes	No	РОР	30 d?	Non-LeukoR	RR	NR
Gregersen <i>et al.</i> , 2015 ⁹	Hip fracture	97	113	16	25	0 25	Q	Q	Yes	Yes	РОР	90 d	LeukoR, 300ml, erythrocyte > 1.65a/dl	Up to 5 wk	Up to 5 wk
Grover <i>et al.</i> , 2006 ⁵⁴	Elective lower limb joint replacement	80	100	20	66	58 8	NR	N	Yes	Yes	РОР	Discharge	LeukoR	RN	NR
Nielsen <i>et al.</i> , 2014 ⁶²	Elective hip revision	73	89	16	67	52 15	18	NR	No	No	10P + POP	30 d	NR	NR	NR
Parker, 2013 ⁶³	Hip fracture	ი	100	ΝA	89	NR NA	NR	NR	NR	NR	РОР	1 yr	NR	NR	NR
So-Osman et al., 2010 ⁶⁸	Elective lower limb joint replacement	Standardized schema st	Non- tandardizec	NA	65	69 -4	NA	NR	No	No	IOP + POP	Discharge	LeukoR	RN	NR
Group 3: Mixe	d medical/surgic	al cases, acute	e care												
de Almeida et <i>a</i> l., 2015 ⁷	Surveillance post-major abdominal surgery	70	06	20	79	58 21	7	13	Unclear	Yes	POP (ICU only)	30 d	LeukoR, 250–350ml, hematocrit 70%	10	13
Fortune <i>et al</i> ., 1987 ⁵³	Trauma or surgical bleeding	100*	133*	33	NR	NR NA	NR	NR	Unclear	Yes	ICU stay	3 d	NR	NR	NR
Hébert <i>et al.</i> , 1995 ⁵⁷	Various diagnoses	70–75	100–105	30	46	3 43	9	9	Yes	Yes	ICU stay	30 d	NR	RN	NR
Hébert <i>et al.</i> , 1999 ⁵⁶	Various diagnoses	70	100	30	33	0 33	-	ო	Yes	Yes	ICU stay	60 d	LeukoR, 240–340ml, hematocrit 80%	RN	RN
Holst et al., 2014 ⁵⁸	Septic shock	20	06	20	36	1 35	g	21	Yes	Yes	ICU stay (max 90 d)	356 d	LeukoR	RN	RN
Jairath <i>et al.</i> , 2015 ³⁵	Upper-GI bleeding	80	100	20	67	54 13	N	24	No	No	Discharge	28 d	NR	RN	NR
														(Co	ntinued)

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Study Charac	teristics							Inte	rvention Char	acteristics					
		Herr Thres	noglobin hold (g/l)		Not Exp Allogeneic	osed to Blood (%	s) Noi	nadherence (%)	°*	scess				Mean S [.] Time	torage (d)
Study ID	Setting or Reason for Admission	Restrictive	Liberal		Restrictive	Liberal ∠	∖ Rest	rictive Libe	Mean Erythrocyt- UI Differec Significantl between ral Groups?	e Hemoglobii I Levels y Differed Significanth over Time?	Protocol Application	Follow-up	Type of Erythrocyte (Leukoreduction, Volume, Mean Hematocrit)	Restrictive	Liberal
Markatou <i>et al.</i> , 2012 ⁵⁹	Surveillance post major abdominal surgerv	77	6 6	22	64	30 3 [,]	4	NR NF	Yes	Yes	1 404 + 901	Discharge	NR	22	28
Topley and Fisher 1956 ⁶⁹	Trauma	70-80% of RCV	Normal RCV	AN	33	0	~ ~	80	Yes	Yes	NR	NR	NR	NR	RN
Villanueva et al., 2013 ⁷⁰	Upper-GI bleeding	20	06	20	51	41 1	0	ຕ ດ	Yes	Yes	Until discharge	45 d	LeukoR, 250-320 ml, hematocrit 60%	15	15
Walsh <i>et al.</i> , 2013 ³⁶	Various diagnoses	70	06	20	22	0	2	14 37	7 Yes	Yes	ICU stay (maximum 14 d)	180 d	LeukoR, 220-340ml, hematocrit 50-70%†	21	21
Group 4: You Naidech <i>et al.</i> 2010 ⁶¹	nger, fitter, brain , Neuro-ICU	injury/intracra 100	nial bleedir 115	ןנ 15	19	5 1	4	0 10) Yes	Yes	14 d	3 mo	Leukodepleted	24	26
Robertson e <i>t al.</i> , 2014 ⁶⁵	Neuro-ICU	70	100	30	48	28	0	4	Yes	Yes	End of ventilation/ ICP monitoring	6 mo	Leukodepleted	ЯN	RN
Group 5: Oth Prick <i>et al.</i> , 2014 ⁶⁴	er patients and s Postpartum hemorrhade	ettings S	89	NA	87	3 8	74	1.3	Yes	Yes	Discharge	6 wk	NR	NR	RN
Webert <i>et al.</i> , 2008 ⁷¹	Hematologic cancer	80	120	40	10	~	n	36 NI	R Unclear	Yes	RN	25 d	Leukodepleted, 240–340 ml, hematocrit 55–65%	N	R
Studies with sig difference in tra Variables in bo l *Nonadherence	prificant differences insfusion thresholds dindicate why tran was defined as any	s in mean erythr s between group isfusion protocol y violation result	ocyte units/ s (∆ hemogl l implementi ing in a dilut	group al lobin thr ation me ion of th	nd in hemo(eshold), the ₃y have faile ie protocol €	jlobin leve transfusio d. Variable	els over on-sparit es in <i>ital</i> .	time were d∉ ng effect (∆ e <i>ïc</i> correspon	emed successfi xposure to allog d to hematocrit patients in the l	ll, <i>i.e.</i> , demons eneic blood), a values convert restrictive grou	trating successfind nd nonadherence ed to hemoglobin p were transfuse	ul transfusic e rates were n values. ed above the	n protocol impleme considered as dete prescribed thresho	intation. The rminants fo Id or situati	e absolute r success. ons where

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CABG = coronary artery bypass graft; CV = cardiovascular; GI = gastrointestinal; ICP = intracranial pressure measurement; ICU = intensive care unit; IOP = intraoperative; LeukoR = leukoreduced; NA = not applicable; NR = not reported; POP = postoperative; RCV = red cell volume; UI = units; S = symptoms of anemia.

Supplemental Digital Content 11, http://links.lww.com/ ALN/B285, which outlines outcome reporting across studies). Outcomes could not be extracted in one study reporting "neuropsychiatric complications."⁶⁸ Thus, data from 7 studies (3,465 patients) were combined^{8,49,50,52,54,63,68}: in an elderly population undergoing orthopedic surgery, events reflecting inadequate oxygen supply were significantly increased in the restrictive group (RR, 1.41; 95% CI, 1.03 to 1.92).

In group 3, six studies reported "inadequate oxygen" events (see table, Supplemental Digital Content 11, http://links.lww. com/ALN/B285, which outlines outcome reporting across studies).^{7,56,58,70} Data from one study could not be extracted, since events were reported as composite outcomes.⁵⁶ When data were combined, there was no difference between groups (3,590 patients; RR, 0.89; 95% CI, 0.72 to 1.09).

Among group 4 and 5 studies, only one study reported the incidence of stroke.⁶¹ In this trial, patients with subarachnoid hemorrhage at risk of cerebral vasospasm were allocated to high transfusion thresholds. There was no difference between groups (RR, 1.36; 95% CI, 0.59 to 3.15).

Early Mortality. (See fig., Supplemental Digital Content 12, http://links.lww.com/ALN/B286, which illustrates the risk of early mortality.) Most group 1 to 3 studies reported early mortality (see table, Supplemental Digital Content 11, http://links.lww.com/ALN/B285, which outlines outcome reporting across studies). When data were combined, there was a possible increase in events when a restrictive strategy was applied in group 1 (RR, 1.39; 95% CI, 0.95 to 2.04—7 studies, 3,245 patients)^{6,47,48,51,55,60,66} but not in group 2 (RR, 1.09; 95% CI, 0.80 to 1.49—7 studies, 3,546 patients)^{8,9,49,50,54,63,68} or group 3 (RR, 0.94; 95% CI, 0.73 to 1.20—7 studies, 2,894 patients).^{7,35,36,56–59} Mortality was not reported in groups 4 and 5.

Composite Events "Inadequate Oxygen Supply + Mortality" (fig. 2)

The risk of events was significantly increased when patients were assigned to a restrictive transfusion strategy in group 1 (RR, 1.12; 95% CI, 1.01 to 1.24—8 studies, 3,322 patients)^{6,47,48,51,55,60,66,67} and group 2 (RR, 1.24; 95% CI, 1.00 to 1.54—8 studies, 3,749 patients)^{8,9,49,50,52,54,63,68} but not in group 3 (RR, 0.90; 95% CI, 0.74 to 1.10—8 studies, 3,762 patients).^{7,35,36,56–59,70} Data in groups 4 and 5 were too scarce to allow statistical combination.

Immunomodulatory Effects of Allogeneic Blood Transfusions (fig. 3)

The incidence of infections was reported in the majority of included studies (see table, Supplemental Digital Content 11, http://links.lww.com/ALN/B285, which outlines outcome reporting across studies). In groups 1 and 3, no difference was found (group 1: RR, 1.11; 95% CI, 0.94 to 1.31—6 studies, 3,141 patients^{6,47,55,60,66,67}; group 3: RR, 0.99; 95% CI, 0.85 to 1.17—5 studies, 2,616 patients^{7,35,56,59,70}). In group

2, patients assigned to a restrictive policy seemed to have less septic events (RR, 0.75; 95% CI, 0.53 to 1.04—9 studies, 3,815 patients).^{8,9,49,50,52,54,62,63,68} In group 4, both studies reported the incidence of infection, but transfusion protocols were deemed too heterogeneous for statistical combination. When individually assessed, no difference was found (RR, 0.77; 95% CI, 0.51 to 1.16⁶⁵; RR, 0.91; 95% CI, 0.14 to 5.92⁶¹). Finally, in a study conducted in the postpartum setting, there was no difference between groups (RR, 1.08; 95% CI, 0.63 to 1.87).⁶⁴

We found significant interaction between risk strata, thereby indicating that our context-specific approach was appropriate (inadequate oxygen supply: Cochran Q P = 0.003, I² = 82.7%; early mortality: P = 0.11 but I² = 54.2%; composite outcome: P = 0.0007, I² = 86.1%; infections: P = 0.04, I² = 69.1%). Data pooling without controlling for clinical heterogeneity (*i.e.*, no context-specific approach) resulted in a dilution of the intervention effect (inadequate oxygen supply: RR, 1.02; 95% CI, 0.94 to 1.11; early mortality: RR, 1.00; 95% CI, 0.89 to 1.12; composite outcome: RR, 1.01; 95% CI, 0.95 to 1.08; infections: RR, 1.97; 95% CI, 0.89 to 1.07).

Additional Analyses

The effect of successful protocol implementation on the risk of AE was explored by excluding unsuccessful studies from each analysis (see table, Supplemental Digital Content 13, http://links.lww.com/ALN/B287, which outlines how risk estimates varied according to successful protocol implementation). In group 1, risk estimates increased further away from the null: patients in the restrictive group seemed to have more events reflecting inadequate oxygen supply (RR, 1.12; 95% CI, 0.99 to 1.27), a 59% increase in mortality (RR, 1.59; 95% CI, 1.04 to 2.44) and a significant increase in the composite outcome (RR, 1.16; 95% CI, 1.03 to 1.31). However, these findings were not reproducible for groups 2 and 3: risk estimates decreased toward the null or further in favor of a restrictive strategy. The effect of successful implementation on infections was inconsistent.

We also explored the effect of cointerventions on the risk of AE, but data were scarce and comparison across studies was difficult (see table, Supplemental Digital Content 14, http://links.lww.com/ALN/B288, which illustrates cointerventions across studies). The use of clotting factor concentrates or antifibrinolytics was reported in three cardiac and one orthopedic surgery studies.^{55,60,62,66} Their administration was well balanced between transfusion groups. The use of cryoprecipitate, fresh frozen plasma, and platelets was reported in four,^{35,55,60,66} eight,^{35,52,55,58,60,62,66,70} and seven studies, 35,55,58,60,66,70,71 respectively. Overall, the administration of blood products was similar between transfusion groups (cryoprecipitate: RR, 0.99; 95% CI, 0.77 to 1.27; fresh frozen plasma: RR, 0.89; 95% CI, 0.75 to 1.04); platelet therapy: RR, 0.95; 95% CI, 0.81 to 1.12), but compared to others, patients undergoing cardiac surgery were more

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Fig. 2. Forest plots illustrating the risk of composite events. For Jairath *et al.*³⁵ and Walsh *et al.*³⁶ data were obtained by contacting the authors. Composite events: myocardial infarction, arrhythmia, unstable angina, stroke, acute kidney injury, mesenteric ischemia, peripheral ischemia, and mortality (occurring within 30 days); group 1 studies: patients with cardiovascular disease undergoing cardiac or vascular procedures (surgery or catheterization); group 2 studies: elderly patients undergoing orthopedic surgery; group 3 studies: mixed surgical/medical patient population admitted to an acute care facility (emergency or intensive care unit). M-H = Mantel-Haenszel data analysis.

systematically exposed to nonerythrocyte blood products. Finally, 10 studies reported intravenous fluid usage: in only three, unbalanced administration was found.^{52,57,62}

Exploring the effect of studies at high or unclear risk of detection and attrition bias did not yield clinically meaningful results: because of the small remaining number of studies, 95% CI was large and risk estimates varied only mildly (data not shown).

Discussion

In this context-specific systematic review, we found that restrictive transfusion strategies were associated with an increased risk of complications in situations combining highrisk patients with major surgery. Those with cardiovascular disease undergoing cardiac or vascular procedures seemed to have more events reflecting inadequate oxygen supply, higher mortality rates, or both. In the elderly orthopedic

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Fig. 3. Forest plots illustrating the risk of infection. Group 1 studies: patients with cardiovascular disease undergoing cardiac or vascular procedures (surgery or catheterization); group 2 studies: elderly patients undergoing orthopedic surgery; group 3 studies: mixed surgical/medical patient population admitted to an acute care facility (emergency or intensive care unit). M-H = Mantel-Haenszel data analysis.

population, a <u>restrictive</u> policy led to a <u>40% increase</u> in <u>ischemic</u> events or <u>AKI.</u>

These findings were consistent with physiologic studies suggesting that multiple perioperative factors may undermine normal compensatory responses to anemia^{72,73}: cardiac dysfunction (induced by anesthetic drugs or surgical trauma), vasoconstriction (due to endogenous or exogenous catecholamines), or postoperative hypoventilation (due to pain or residual effects of anesthetics) may compromise adequate oxygen delivery to vital organs. In normal conditions, systemic oxygen delivery largely exceeds oxygen consumption, resulting in a positive *oxygen reserve.*⁷³ In patients having a preexisting low reserve, however, the combination of acute anemia with impaired compensatory responses may induce a state of *oxygen supply dependency*, resulting in acidosis and organ failure. In this particular situation, administrating erythrocyte could restore the oxygen reserve by increasing blood oxygen content and tissue oxygenation.⁷³

Surprisingly, no evidence of harm was found when restrictive strategies were applied in critically ill patients, although similar impairment of compensatory responses was expected. One explanation might be the heterogeneity in oxygen reserve among this mixed population: medical patients might be at lower risk of oxygen supply dependency than their surgical counterparts, who have the additional burden of surgery, pain, and recovery from anesthesia.

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When we excluded studies showing unsuccessful protocol implementation, effect estimates differed across risk strata. While harm was more pronounced in patients undergoing cardiac or vascular procedures, it decreased toward the null in those undergoing orthopedic surgery. Two reasons may account for this: first, cardiac patients were probably more likely to enter a state of oxygen supply dependency (due to the combination of advanced cardiovascular disease and high-risk surgery) than the orthopedic population, where ischemic heart disease ranged from 10 to 45%. Second, cardiac patients might have been exposed to context-specific factors increasing their risk of AE: antifibrinolytics and clotting factor concentrates were more systematically administered in this population, while this was rarely reported in orthopedic patients. Although tranexamic acid seems safe in trauma patients,⁷⁴ the thrombogenic potential of other hemostatic agents in nontrauma settings remains unclear.^{75–77} Thus, the combination of a restrictive transfusion strategy and highly thrombogenic drugs in patients with advanced cardiovascular disease might be particularly unfavorable.

Our analysis of the <u>risk of infection</u> remained <u>inconclu</u>sive. Although there was a possible reduction in septic events in orthopedic patients assigned to a restrictive strategy, effect estimates differed widely across risk strata: the benefit of reduced exposure to allogeneic blood was less clear in cardiac patients, which is consistent with a previously published meta-analysis.¹⁰ One reason might be that immune response impairment was more pronounced in these patients: in our analysis, the cardiac surgery population was more likely to receive nonerythrocyte allogeneic blood products (such as platelets or fresh frozen plasma), which are also known to have immunomodulatory effects.^{33,78–80} Alternatively, cardiac surgery itself might induce particularly high levels of perioperative stress, which has also been found to interfere with immune responses.^{22,81–83}

Our systematic review differs from others in many aspects. First, in most previous meta-analyses, data were combined despite high clinical heterogeneity,^{10,13,14,16} which may hinder the identification of group-specific effects. We addressed this methodologic limitation by performing a *context-specific* analysis, using strict criteria for risk-strata generation and subsequent data pooling. We were eventually able to show that indiscriminate data combination (*i.e.*, performing analyses without controlling for clinical diversity) resulted in a dilution of the intervention effects. Furthermore, to fully explore the impact of clinical diversity, the issue of transfusion protocol variability was examined using two performance indicators and exploring three different determinants of success. One other meta-analysis investigated protocol diversity, but only hemoglobin thresholds were explored.¹⁰ Additionally, our review was the first to assess the role of cointerventions: we were able to show that in some patients, the combination of restrictive transfusion policies and thrombogenic drugs could be particularly

detrimental; we also identified a possible effect of nonerythrocyte blood products and of perioperative stressors on the risk of infection.

This review has some limitations. First, although strict criteria were used to handle unclear/missing outcome data, outcome reporting and definitions varied across studies, and so this may result in residual clinical diversity. However, a certain degree of heterogeneity is desirable to ensure wide applicability of the findings. Second, in order to capture the full spectrum of effects related to transfusion strategies and to improve statistical precision, we used arbitrarily defined outcome categories. However, although endpoints combination might be biologically well founded, individual components may differ in clinical importance, and our categories may have failed to reflect endpoints truly relevant for patients.³⁰ Third, in some risk strata, large studies having high event rates appeared to dominate the analysis, but the risk of a small-study effect was deemed low: we used the Mantel-Haenszel method to account for smaller studies and addressed thoroughly all sources of clinical and methodologic diversity. It seems therefore unlikely that our findings derive solely from the effect of larger studies. Fourth, our assessment of methodologic heterogeneity (detection and attrition bias) was hampered by the scarcity of data. The same problem was encountered with data on cointerventions, which reduced our ability to fully explore their role in the occurrence of complications. Finally, indicators of successful protocol implementation were arbitrarily defined; using a different model might have yielded other results.

This analysis provided <u>clear evidence that the decision</u> to transfuse (or not transfuse) requires more than a "onesize-fits-all" approach. As highlighted recently,^{84–86} the identification of populations at higher risk of oxygen supply dependency who might particularly benefit from erythrocyte administration remains a real challenge. New transfusion algorithms should aim to integrate additional clinical parameters, such as patient comorbidities, particular settings, or oxygen reserve estimates.⁷³ We recommend that future trials systematically collect and report data regarding the use of nonerythrocyte blood products, antifibrinolytics, and clotting factor concentrates, since their role in the risk of AE remains unclear.

Conclusion

This meta-analysis suggests that the use of restrictive transfusion strategies might be detrimental in high-risk patients undergoing major surgery. Further research is needed to evaluate the contributing role of cointerventions in the occurrence of complications.

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Competing Interests

The authors declare no competing interests. The funding organizations had no role in the design and conduct of the study; in the collection, management, analysis, or interpretation of the data; or in the preparation, review, or approval of the manuscript.

Reproducible Science

Full protocol available from Dr. Hovaguimian: frederique. hovaguimian@usz.ch. Raw data available from Dr. Hovaguimian: frederique.hovaguimian@usz.ch.

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