A systematic review of the effect of red blood cell transfusion on mortality: evidence from large-scale observational studies published between 2006 and 2010

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ABSTRACT

Objective: To carry out a systematic review of recently published large-scale observational studies assessing the effects of red blood cell transfusion (RBCT) on mortality, with particular emphasis on the statistical methods used to adjust for confounding. Given the limited number of randomised trials of the efficacy of RBCT, clinicians often use evidence from observational studies. However, confounding factors, for example, individuals receiving blood generally being sicker than those who do not, make their interpretation challenging.

Design: Systematic review.

Information sources: We searched MEDLINE and EMBASE for studies published from 1 January 2006 to 31 December 2010.

Eligibility criteria for included studies: We included prospective cohort, case–control studies or retrospective analyses of databases or disease registers where the effect of risk factors for mortality or survival was examined. Studies must have included more than 1000 participants receiving RBCT for any cause. We assessed the effects of RBCT versus no RBCT and different volumes and age of RBCT.

Results: 32 studies were included in the review; 23 assessed the effects of RBCT versus no RBCT; 5 assessed different volumes and 4 older versus newer RBCT. There was a considerable variability in the patient populations, study designs and levels of statistical adjustment. Overall, most studies showed a higher rate of mortality when comparing patients who received RBCT with those who did not, even when these rates were adjusted for confounding; the majority of these increases were statistically significant. The same pattern was observed in studies where protection from bias was likely to be greatest.

Conclusions: Recent observational studies do show a consistently adverse effect of RBCT on mortality. Whether this is a true effect remains uncertain as it is possible that even the best conducted adjustments cannot completely eliminate the impact of confounding.

INTRODUCTION

Randomised controlled trials are considered the gold standard with which to evaluate the efficacy of a particular healthcare intervention. In 2005, Blajchman published a study that explored the impact that 10 landmark randomised controlled trials had on the practice of transfusion medicine. The use of
randomised trials to evaluate transfusion medicine has only been established since the 1980s. Given the limited number of high-quality randomised trials of the efficacy of blood transfusion and the challenges of conducting new trials, clinicians often rely on evidence from observational studies. In a randomised trial, patients are allocated to compare groups at random, so the level of disease is likely to be similar in each group and differences in disease severity unlikely to be the explanation for any differences in outcome seen. In an observational study, the groups of patients being compared are not likely to be comparable and the differences in prognostic factors may of themselves lead to differences in outcome. The impact of such ‘confounding’ can be reduced by adjustment in the statistical analysis, but the success of this is dependent on the technique used, complete identification of the factors which might influence outcome and their accurate measurement in the patients in the study. As all the factors influencing outcome may never be known, adjustment is unlikely to ever completely account for the confounding occurring in observational studies. The unknown interdependence of multiple factors is also a major challenge.

There is increasing implementation of restrictive policies for transfusion, and evidence of reduction in blood use in several countries such as the UK and the USA with no evidence of poorer clinical outcomes. However, there remains considerable variation between hospitals in blood reduction in the UK and elsewhere, suggesting that overall blood usage could be further reduced without compromising patient safety. Observational studies may have influenced these changes in transfusion practice along with evidence from randomised controlled trials, national guidelines and process-driven initiatives, but the impact that the contribution of data from observational studies has made to the practice of transfusion medicine has not been systematically explored. The aims of this systematic review were therefore to identify recently published (2006–2010), large-scale observational studies assessing the effects of red blood cell transfusion (RBCT) on mortality. In particular, we aimed to critique the statistical methods and the assumptions made in the analyses of the observational data, and to consider the validity of these data as an evidence base for the practice of transfusion medicine and to inform future research in this field.

METHODS
Criteria for selecting studies
Type of participants
We included both adults and children receiving RBCT for any cause. We also included studies which stated that patients received red blood cells and other blood products. When reported by the primary studies, we assessed the effects of RBCT separately from other blood products.

Type of intervention and comparator
We included the following risk factors:
- RBCT versus no RBCT;
- Volume ‘A’ of RBCT versus volume ‘B’ of RBCT (as defined by the primary studies);
- ‘Older’ RBCT versus ‘newer’ RBCT (as defined by the primary studies).

Type of outcome measure
Our primary outcome measure was death, mortality or survival measured at any time point.

Type of studies
We included prospective cohort, case–control studies or retrospective analyses of databases or disease registers where the effect of the above risk factors on death, mortality or survival is examined. Studies must have included more than 1000 participants. This was a pragmatic limit designed to focus attention on studies most likely to have had an impact and least likely to have been affected by chance.

Search strategy
We carried out a comprehensive search (21 January 2011) of MEDLINE and EMBASE for studies published from 1 January 2006 to 31 December 2010 using the strategies in online supplementary appendix 1. Again, we chose to use a pragmatic approach and limited our search to studies published in the last 5 years. We also excluded conference abstracts unless they had subsequently been published as full articles.

Data collection and analysis
One review author (CD) initially screened all search results for relevance against the eligibility criteria and discarded all those that were clearly irrelevant. Thereafter, another author (SH) independently screened all remaining hits. We retrieved full text articles for all those references where we are unable to decide on eligibility based on the title and abstract alone. All full text articles were independently screened by two review authors (SH and MFM) to ensure that they met the eligibility criteria.

Data extraction and management
Two review authors (SH and OO) independently extracted data from all included studies. Any disagreements were resolved by discussion or by consulting a third author if there was still uncertainty. We extracted data on the following study characteristics: the study design, how patients were recruited, the country where the study was conducted, the source of funding, the type of participants, their age, disease area, setting, the type of intervention/comparator and nature of the exposure, the number of participants in each group, whether any formal prescribing guidance was reported, the type of outcome measure (ie, mortality) and the time point at which it was measured.

We also extracted information on the statistical methods used to adjust for differences between study
groups, in particular the number of study covariates measured, whether important covariates relating to red cell transfusion were assessed (ie, age, sex, comorbidity and haemoglobin) and whether these were incorporated in the analysis, whether the choice of covariates were prespecified or data driven and the statistical model used for the statistical adjustment. We also assessed the effects of smoking as a study covariate in relation to blood transfusion and its effect on mortality. In terms of the study results, we extracted data on the presentation of both the unadjusted and adjusted results for the effect of red cell transfusion on mortality as reported by each study. If not reported by the primary study, we calculated (where there were sufficient data) the OR for the effects of blood transfusion on mortality for unadjusted analyses using STATA (V11). We assessed, for the unadjusted and adjusted results, whether the study reported summary statistics for each comparison group, the treatment effect, CI, p value and whether the result was statistically significant. If a study reported more than one adjusted analysis, we selected in order of preference (1) the main adjusted analysis mentioned in the abstract, (2) the main adjusted analysis mentioned in the conclusions and (3) the main adjusted analysis mentioned in the results section. If mortality was assessed for more than one time point (ie, at 30 days and 1 year), then we used the shorter time point in our analysis.

Assessment of methodological quality

We also assessed whether studies met important methodological criteria for the reporting of observational studies: whether the samples were representative of those to whom the results might be generalised, whether important covariates in relation to RBCT and mortality (eg, sex, age, smoking, comorbidity, haemoglobin level) were measured and incorporated in the analysis, whether the method of dealing with confounding between patient groups was adequate, whether a statistician was listed as an author of the study and whether the data were collected prospectively following an agreed study design. We included smoking as a covariate as, while not directly correlated with transfusion, it is considered to be important when assessing mortality.

Method of analysis

We have presented the results separately for the three different types of comparisons. Within each, owing to the varied nature of the clinical conditions, study designs and level of statistical adjustment, we decided a priori not to combine the results of individual studies in a meta-analysis and instead present the results of the individual studies descriptively in the text, tables and figures.

RESULTS

Searches of MEDLINE and EMBASE identified 4318 possible records. In total, 4272 did not meet the eligibility criteria for this study. Full articles were retrieved for 45 studies; 13 further studies were excluded as they did not fulfil our eligibility criteria (see online supplementary appendix 2 for list of excluded studies). Thirty-two studies were included in the review; 29 studies assessed the effects of RBCT versus no RBCT; 5 studies assessed different volumes of RBCT and 36 assessed giving older versus newer RBCT (see figure 1).

**RBCT versus no RBCT**

Twenty-three studies assessed the effects of RBCT versus no RBCT on mortality. Four of these studies included both red cell transfusion and other blood products (eg, platelets, plasma and cryoprecipitate); for one study, data were available separately for RBCT and mortality. For three studies, it was unclear if other blood products were transfused along with red blood cells.

**Study characteristics**

Eight studies were prospective cohort studies following up a planned group of patients, whereas the other 15 studies assessed data from a retrospective patient registry or database (table 1). Fourteen studies were conducted in the USA, two in the UK, two in Israel and the remainder in Belgium, the Netherlands, Iran and Denmark; one study was conducted in multiple countries. The time period assessed was between 1989 and 2008. Twelve of the studies specifically looked at adults undergoing cardiac surgery, five were on patients in the intensive care unit, two were on adults trauma patients, two were on patients following hip fracture/replacement, one was on oncology patients and the other on paediatric trauma patients. Three of the studies specifically looked at the effects of RBCT in older adults (eg, >60 years). The size of the studies varied from 1624 participants to 504 208 participants with an overall median sample size of 4344 (IQR 2085–11963); median 1068 (IQR 450–5812) for patients undergoing RBCT compared with median 2252 (IQR 1636–6151) for patients with no RBCT. The time period at which mortality was assessed also varied across studies from in-hospital to mortality at 7 years; the most common time point being mortality at or within 30 days. Several studies reported mortality for more than one time period. Only 7 of the 23 studies provided any mention of guidelines for the prescription of RBCT; 2 stated that no formal protocol was used.

**Statistical methods**

All 23 studies provided information on the statistical methods used to adjust for differences in the baseline characteristics of patients who received RBCT and those.
who did not (table 2). However, the amount of detail and appropriateness of the method used varied across studies. In 13 studies, the choices of covariates measured were reported as prespecified and not data driven, but this was unclear for the remaining 10 studies. The number of covariates measured and incorporated in the analysis also varied across studies with half the studies reported to assess more than 20 different covariates. Despite the high number of covariates assessed in these studies, not all measured covariates which appeared to be of specific importance in relation to RBCT. All of the 23 studies did report measuring the age and sex of the patients and 21 reported measuring patient comorbidity. Overall, only eight studies measured and incorporated the covariates age, sex, smoking, comorbidity and haemoglobin level in the adjusted analysis. Fourteen of the 23 studies reported using logistic regression (ie, mortality was reported as a binary outcome) as the method of adjusting for differences in the baseline characteristics between the two patient groups; six studies reported using the Cox proportional hazard (ie, mortality was reported as a time to event outcome) and three studies reporting using both methods; in these three studies, mortality was assessed for more than one time period. Nine studies reported using propensity scores prior to adjusting for confounders; however, sometimes this matching was done only by using a much smaller subgroup of patients. For full details of the statistical methods, see online supplementary appendix 4.

Presentation of adjusted and unadjusted results

There were marked differences in the presentation and reporting of the unadjusted and adjusted results when comparing the effects of RBCT versus no RBCT on mortality (table 3). Seven of the 23 studies reported summary statistics for each group for both the unadjusted and adjusted analyses. Five studies reported summary statistics for only the unadjusted analysis and one study for the adjusted analysis only. Where possible, we calculated the OR for the effect of RBCT on mortality for unadjusted analyses if it was not reported in the published article.

Seventeen of the 23 studies reported a statistically significant result for the unadjusted analysis, and 15 for the adjusted analysis (figure 2), when comparing the effect of RBCT versus no RBCT on mortality, with more deaths occurring in patients receiving transfusion. This effect was statistically non-significant in seven studies based on the
result of the adjusted analysis. Prospective studies were more likely to show a statistically significant effect for blood transfusion on mortality compared with retrospective studies for both the unadjusted and adjusted analyses. For full details, see online supplementary appendix 5.

**Volume ‘A’ red blood versus volume ‘B’ red blood cells**

Five studies\(^3\)–\(^5\) assessed the effects of different volumes of RBCT on mortality. One of these studies\(^5\) included both RBCT and other blood products.

**Study characteristics**

One study assessed a prospective cohort and followed up a planned group of patients,\(^\) whereas the other four studies assessed data from a retrospective patient registry or database. Two of the studies\(^3\)–\(^5\) specifically looked at adults undergoing cardiac surgery, one was on trauma patients,\(^\) one was on patients undergoing major surgery and one on patients in the intensive care unit.\(^\) The size of the studies varied from 1841 participants to 125 177 participants, with an overall median sample size of 8215 (IQR 3037–8799). The volume of RBCT varied considerably across studies from 1 to 2 units to more than eight units. The time period at which mortality was assessed also varied across studies from in-hospital to mortality at 8 years. Three of the five studies provided any mention of guidelines for the prescription of red blood cells; however, only one gave any specific requirement stating a haemoglobin of <8 g/dl\(^\) (see table 1; online supplementary appendix 3).

### Table 1 Summary of characteristics of included studies

| Type of comparison | RBCT vs no RBCT (n=23) | Volume ‘A’ vs Volume ‘B’ (n=5) | Old RBC vs new RBC (n=4) |
|--------------------|-------------------------|-------------------------------|--------------------------|
| **Design**         |                         |                               |                          |
| Prospective        | 8 (35%)                 | 1 (20%)                       |                          |
| Retrospective      | 15 (65%)                | 4 (80%)                       | 4 (100%)                 |
| **Country**        |                         |                               |                          |
| Australia          | 1 (20%)                 |                               |                          |
| Belgium            | 1 (4%)                  |                               |                          |
| Denmark            | 1 (4%)                  |                               |                          |
| Germany            | 1 (20%)                 |                               |                          |
| Iran               | 1 (4%)                  |                               |                          |
| Israel             | 2 (9%)                  |                               |                          |
| The Netherlands    | 1 (4%)                  | 1 (25%)                       |                          |
| Sweden             | 1 (25%)                 |                               |                          |
| The USA            | 14 (61%)                | 3 (60%)                       | 2 (50%)                  |
| The UK             | 2 (9%)                  |                               |                          |
| (multiple sites)   | 1 (4%)                  |                               |                          |
| **Time period assessed** |                 |                               |                          |
| All patients       | 1989–2008               | 1993–2007                     | 1993–2007                |
| RBC transfusion    | 4344 (IQR 2085–11963)   | 8215 (IQR 3037–8799)          | 4358 (2264–185019)       |
| No RBC transfusion | 2375 (IQR 1636–6151)    |                               |                          |
| **Disease area**   |                         |                               |                          |
| Cardiac surgery    | 12 (52%)                | 2 (40%)                       | 2 (50%)                  |
| Hip fracture/replacement | 2 (9%)               |                               |                          |
| Intensive care     | 5 (22%)                 | 1 (20%)                       |                          |
| Oncology           | 1 (4%)                  |                               |                          |
| Surgery            | 1 (20%)                 |                               |                          |
| Trauma adults      | 2 (9%)                  | 1 (20%)                       | 1 (25%)                  |
| Trauma paediatrics | 1 (4%)                  |                               |                          |
| Not reported       |                         |                               |                          |
| **Prescribing guidance** |                   |                               |                          |
| Reported           | 7 (30%)                 | 3 (60%)                       | 1 (25%)                  |
| Not reported       | 16 (70%)                | 2 (40%)                       | 4 (100%)                 |
| **Mortality assessed** |                     |                               |                          |
| In hospital        | 8                       | 2                             | 2                        |
| 30 days            | 10                      | 2                             | 1                        |
| 3 months           | 3                       |                               |                          |
| 6 months           | 3                       |                               |                          |
| >1 year            | 4                       | 1                             | 2                        |
| Time period not specified | 2                 |                               |                          |

*Studies reported mortality for >1 time point based on binary only and/or time-to-event outcome.

RBCT, red blood cell transfusion.
All five studies provided information on the statistical methods used to adjust for differences in the baseline characteristics of patients who received different volumes of red blood transfusion, however, as with the studies of RBCT versus no RBCT, the amount of detail and appropriateness of the method used varied across studies. In all five studies, the choices of covariates measured were reported as prespecified. The number of covariates measured and incorporated in the analysis varied across studies with two of the studies reported to assess more than 20 different covariates. Once again, despite the high number of covariates assessed in these studies, not all measured covariates seem to be of specific importance in relation to RBCT. All five studies reported measuring age and sex and patient comorbidity; however, one measured and incorporated the covariates age, sex, smoking, comorbidity and haemoglobin level in the adjusted analysis (table 2).

### Table 2 Method of adjusted analysis

| Type of comparison          | RBCT vs no RBCT (n=23) | Volume ‘A’ vs Volume ‘B’ (n=5) | Old RBC vs new RBC (n=4) |
|-----------------------------|------------------------|-------------------------------|-------------------------|
| Choice of covariates        |                        |                               |                         |
| Prespecified                 | 13 (57%)               | 5 (100%)                      | 4 (100%)                |
| Post hoc                     |                        |                               |                         |
| Unclear                     | 10 (43%)               |                               |                         |
| No. of covariates measured  |                        |                               |                         |
| 1–5                         | 2 (9%)                 | 1 (20%)                       | 2 (50%)                 |
| 6–10                        | 4 (17%)                | 1 (20%)                       | 2 (40%)                 |
| 11–20                       | 3 (13%)                | 2 (40%)                       | 1 (25%)                 |
| >20                         | 12 (52%)               | 2 (40%)                       | 1 (25%)                 |
| Unclear                     | 2 (9%)                 |                               | 1 (25%)                 |
| Important covariates assessed |                       |                               |                         |
| Age                         | 23 (100%)              | 5 (100%)                      | 4 (100%)                |
| Sex                         | 23 (100%)              | 5 (100%)                      | 4 (100%)                |
| Smoking                     | 8 (35%)                | 1 (20%)                       | 1 (25%)                 |
| Comorbidity                 | 21 (91%)               | 5 (100%)                      | 3 (75%)                 |
| Hb level                    | 14 (61%)               | 4 (80%)                       | 2 (50%)                 |
| Important covariates incorporated into analysis |   |                               |                         |
| Yes                         | 8 (35%)                | 1 (20%)                       | 1 (25%)                 |
| No                          | 15 (65%)               | 4 (80%)                       | 3 (75%)                 |
| Method of adjustment        |                        |                               |                         |
| Cox proportional hazard     | 6 (26%)                | 1 (20%)                       | 1 (25%)                 |
| Logistic regression         | 14 (61%)               | 4 (80%)                       | 2 (50%)                 |
| Both*                       | 3 (13%)                |                               |                         |
| Not reported                |                        |                               | 1 (25%)                 |

*Studies reported >1 method of adjustment when mortality was assessed for >1 time point.

RBCT, red blood cell transfusion.

**Statistical methods**

All five studies provided information on the statistical methods used to adjust for differences in the baseline characteristics of patients who received different volumes of red blood transfusion, however, as with the studies of RBCT versus no RBCT, the amount of detail and appropriateness of the method used varied across studies. In all five studies, the choices of covariates measured were reported as prespecified. The number of covariates measured and incorporated in the analysis varied across studies with two of the studies reported to assess more than 20 different covariates. Once again, despite the high number of covariates assessed in these studies, not all measured covariates seem to be of specific importance in relation to RBCT. All five studies reported measuring age and sex and patient comorbidity; however, one measured and incorporated the covariates age, sex, smoking, comorbidity and haemoglobin level in the adjusted analysis (table 2).

**Presentation of adjusted and unadjusted results**

As with the studies of RBCT versus no RBCT, there were marked differences in the presentation and reporting of the unadjusted and adjusted results when comparing the effects of different volumes of RBCT on mortality. Two studies reported a statistically significant result for the adjusted analysis with more deaths occurring in patients receiving larger volumes of RBCT. This effect was statistically non-significant in two studies based on the result for adjusted analysis and was not reported for the remaining one study (table 3). No studies reported on the statistical significance of the result of the unadjusted analysis (see online supplementary appendices 4 and 5).

**‘Older’ red blood cells versus ‘newer’ red blood cells**

Four studies assessed the effects of age of RBCT on mortality, one of which specifically looked at leukodepleted RBCT.

**Study characteristics**

All four studies assessed data from a retrospective patient registry or database. Two of the studies specifically looked at adults undergoing cardiac surgery, one was on trauma patients, while the other did not mention a specific patient group. The size of the studies varied from 1813 participants to 364 037 participants, with an overall median sample size of 4358 (IQR 2264–185 019). The period of time in which the blood was stored varied considerably across studies. Two studies assessed RBCT stored for less than 14 days compared with those stored for more than 14 days, one study compared blood stored for less than 18 days and with blood stored for more than 18 days and one study.
looked at multiple storage periods ranging from 1 to 42 days (table 1). None of the studies provided any mention of guidelines for the prescription of red blood cells (see online supplementary appendix 2).

### Statistical methods

All four studies provided information on the statistical methods used to adjust for differences in the baseline characteristics of patients who received RBCT stored for...
different time periods; however, once again, the amount of detail and appropriateness of the method used varied across studies. The number of covariates measured and incorporated in the analysis also varied across studies. All of the four studies reported measuring the age and sex of the participants. Only two studies reported measuring patient haemoglobin levels and three studies reported assessing patient comorbidities. Only one of the four studies measured and incorporated the covariates age, sex, smoking, comorbidity and haemoglobin level in the adjusted analysis (table 2).

Presentation of adjusted and unadjusted results
As with the studies of RBCT versus no RBCT and of volume ‘A’ red blood cells versus volume ‘B’ RBCT, there were marked differences in the presentation and reporting of the unadjusted and adjusted results when comparing the effects of RBCT stored for different time periods on mortality. Two studies reported a statistically significant result for the unadjusted analysis and one study reported a statistically significant result for the adjusted analysis. In two of these three studies, there were more deaths occurring in patients receiving older blood and in one study, there were more deaths in patients receiving newer blood. This effect was statistically non-significant in three studies based on the result for adjusted analysis (see table 3; online supplementary appendices 4 and 5).

Assessment of methodological quality
Overall, the assessment of methodological quality varied across studies and by study group with only 10 of the 32 included studies assessing a prospective cohort following up a planned group of patients over time; the remaining two-third of the studies assessed data from a retrospective patient registry or database. In most studies, the sample of patients included in the study was considered to be representative of those to whom the results might be generalised. Four studies specifically focused on older adults (>60 years) and one study on children, so the findings from these studies should only be interpreted in relation to these specific patient groups. The baseline characteristics of patients who received RBCT compared with those patients who did not receive RBCT (or patients who received different volumes or age of blood) were often very different, and so we wanted to assess whether the studies had adjusted for these differences when carrying out their statistical analysis. Only 10 studies measured and incorporated in the analysis covariates which we deemed to be of specific importance in relation to RBCT (ie, age, sex, smoking, comorbidity and haemoglobin level); thus, we deemed the method of dealing with confounding between patient groups as adequate in only 31% of studies. Critically however, when we restricted our analysis of results to studies with adequate methods, the pattern of an increase in mortality associated with RBCT remained unchanged (table 4).

DISCUSSION
Summary of main findings
We identified 32 observational studies of more than 1000 participants published between 2006 and 2010 assessing the effect of RBCT on mortality. Twenty-three studies compared RBCT versus no RBCT, five compared different volumes and four compared different storage times. Overall, there was considerable variability in the characteristics of the observational studies. However, the majority of studies were retrospective designs assessing patients from an existing patient register or database. We also identified considerable variability in the statistical methods used to adjust for differences in the baseline characteristics of patients who received RBCT and those who did not. It was often unclear if the choices of the covariates measured and used in the adjusted analyses were prespecified at the start of the study or were driven by the underlying data. Perhaps most importantly, around half of the 32 studies did not measure and adjust for covariates which we deemed to be of specific importance to blood transfusion.

Overall, more studies found a higher rate of mortality in patients receiving RBCT compared with those who did not, and this effect was seen in both the adjusted and unadjusted results. In general, where measured equivalently within the same study, the unadjusted estimate of risk was greater than the adjusted risk, emphasising that adverse prognostic factors are more common in patients receiving RBCT and that adjusting for them leads to a smaller estimate of risk. Considering the adjusted risks, although the size of the effect was not consistent across all studies, the direction of the effect was. Most studies suggest an increased risk of mortality associated with RBCT. Further, those studies which were designed prospectively and which used better methods of adjusting for differences in the baseline characteristics between groups were more likely to show an increase in the risk of mortality compared with studies which were based on retrospective registries or databases, although, again, the size of the effect was not consistent across all studies. However, it is important to remember that even with the best methods of adjustment, it cannot completely eliminate the impact of confounding as the sicker the patients (thus, an increased risk of mortality), the more likely they are to have received RBCT.

Comparison with other studies
We are aware of one other systematic review of observational studies looking at the effects of RBCT on mortality, which focused specifically on critically ill adults in intensive care units and adult trauma and surgical patients. This systematic review by Mark and colleagues included more studies (n=45) than our review as it did not restrict its inclusion criteria to studies with >1000 patients; the median number of patients analysed was 687. They also found that RBCT was associated with an increased risk of mortality based on a meta-analysis of 12 studies (OR 1.7; 95% CI 1.4 to 1.9). However, there was
considerable heterogeneity in the meta-analysis, suggesting that it might not have been appropriate to combine the results of the individual studies and supports our decision not to conduct a meta-analysis.

In an overview of evidence from randomised controlled trials, Wilkinson et al.\(^4\) identified 142 trials in RBCT. The majority compared the effects of leucoreduced RBCT or different transfusion triggers (n=71). However, they did identify 12 trials comparing the effects of RBCT versus no transfusion, 7 looking at different volumes of RBCT and 11 different ages of red blood cells. The size of the trials was very small (median 30–40 patients) and the overview did not specifically examine the effect of RBCT on mortality. Currently, we are aware of at least 14 ongoing or recently completed randomised controlled trials examining the effects of the age of RBCT on clinical outcomes including the ARIPi (Age of Red blood cells In Premature Infants)\(^4\) ABLE, (Age of Blood Evaluation trial in the resuscitation of critically ill patients),\(^4\) RECESS (REd CEll Storage duration Study)\(^4\) and INFORM (Effects of transfusing fresh versus standard-issue red cells on in-hospital mortality) trials, for which mortality or survival is a specified outcome measure.

Limitations

Our study has several limitations. First, we only included studies published in the last 5 years and which included more than 1000 patients. This was because we took a pragmatic approach as we hypothesised that more recent studies were more likely to use better statistical methods and also that studies with a larger sample size were more likely to show a truer effect of the intervention.\(^4\) Thus, we aimed to provide a ‘snapshot’ of current practice rather than provide a comprehensive review of all available evidence. It is possible, therefore, that the overall effect seen here might be different in older studies and/or in those carried out in smaller samples.

### Table 4

| Study ID | Data collected prospectively | Sample representative | Important covariates measured | Important covariates incorporated in analysis | Method of dealing with confounding adequate* |
|----------|-------------------------------|-----------------------|------------------------------|---------------------------------------------|--------------------------------------------|
| Aronson 2008\(^7\) | Yes                           | Unclear               | Yes                          | Yes                                         | Yes                                        |
| Bernard 2009\(^3\) | No                            | Yes                   | Yes                          | Yes                                         | Yes                                        |
| Bochicchio 2008\(^8\) | Yes                           | Yes                   | No                           | No                                          | No                                         |
| Charles 2007\(^2\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Edgren 2010\(^3\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Engoren 2009\(^9\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Engoren 2009\(^10\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Garty 2009\(^11\) | No                            | Yes                   | Yes                          | Yes                                         | Yes                                        |
| Jani 2007\(^12\) | No                            | Yes (>60 years)       | Yes                          | Yes                                         | Yes                                        |
| Johnson 2006\(^13\) | No                            | Unclear               | No                           | No                                          | No                                         |
| Khorana 2008\(^14\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Koch 2006\(^15\) | Yes                           | Yes                   | Yes                          | Yes                                         | Yes                                        |
| Koch 2008\(^37\) | No                            | Yes                   | Yes                          | Yes                                         | Yes                                        |
| Murphy 2007\(^17\) | No                            | Yes                   | Yes                          | Yes                                         | Yes                                        |
| Nikolsky 2009\(^18\) | Yes                           | Yes                   | Yes                          | Yes                                         | Yes                                        |
| O’Keeffe 2010\(^23\) | Yes                           | Yes                   | No                           | No                                          | No                                         |
| Pederson 2009\(^19\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Rogers 2006\(^20\) | No                            | Yes (>65 years)       | No                           | No                                          | No                                         |
| Rogers 2009\(^21\) | No                            | Yes (>65 years)       | No                           | No                                          | No                                         |
| Ruttinger 2007\(^24\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Salehmanran 2009\(^22\) | No                            | Yes                   | Yes                          | Yes                                         | Yes                                        |
| Stone 2008\(^23\) | No                            | No (<16 years)        | No                           | No                                          | No                                         |
| Surgenor 2009\(^24\) | Yes                           | Yes                   | No                           | No                                          | No                                         |
| Taylor 2006\(^25\) | Yes                           | Yes                   | No                           | No                                          | No                                         |
| van de Watering 2006\(^3\) | Yes                           | Yes                   | No                           | No                                          | No                                         |
| van Straten 2010\(^26\) | Yes                           | Yes                   | No                           | No                                          | No                                         |
| Vincent 2008\(^27\) | Yes                           | Yes                   | No                           | No                                          | No                                         |
| Weightman 2009\(^28\) | Yes                           | Yes                   | No                           | No                                          | No                                         |
| Weinburg 2008\(^29\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Weinburg 2008\(^30\) | No                            | Yes                   | No                           | No                                          | No                                         |
| Wu 2010\(^29\) | No                            | Yes (>65 years)       | Yes                          | Yes                                         | Yes                                        |
| Zilberberg 2008\(^31\) | No                            | Yes                   | No                           | No                                          | No                                         |

*The method of dealing with confounding was deemed adequate if important covariates were measured and adjusted for in the analysis.*

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numbers of patients. Second, we decided not to combine the results of individual studies because of the variability in clinical settings and study methods, and instead presented the results of individual studies descriptively in the text and in tables and figures. More formal statistical analysis might have given a more precise indication of the overall effect of red cell transfusion on mortality, but would have ignored the significant amount of clinical and methodological heterogeneity between studies which we identified a priori and which was very apparent in the analysis done by Marik and Corwin.40 However, in the absence of a more formal statistical analysis, we have inevitably had to rely on a vote-counting approach which also has great dangers, particularly the assumption that each included study has equal weight. Our main protection against this is the very pronounced nature of the pattern we have observed and the fact that we have limited our conclusions to the direction of effect.

Finally, we limited our inclusion criteria to published articles and excluded unpublished studies or those published only as conference abstracts; thus, our study could be subject to publication bias, as studies which did not show a significant effect of red cell transfusion on mortality might have been less likely to be published in full.46 Outcome reporting bias may also be a problem, although difficult to combat, in the case where a risk has been measured at different time points but only those time points which are ‘positive’ are reported. However, in the case of both publication and outcome reporting bias, the extreme nature of the pattern makes it relatively implausible that there are sufficient unpublished studies or time points to reverse it.

**Implications for clinical practice**

In recent years, many developed countries including the UK, the USA and Australia have developed national initiatives for better blood transfusion practice, sometimes called ‘patient blood management’.4, 5 These include the development of guidelines on blood usage promoting restrictive transfusion strategies and initiatives for using alternatives to transfusion (eg, cell salvage techniques; improvements in the education and training of clinical staff prescribing blood; the provision of mechanisms for reviewing blood use with feedback of data to clinicians). National data on blood usage in the USA suggest an estimated decline of 3% over each of the last two years (2009–2010),4 and similar data are available in the UK where the demand for red cell units, which steadily increased during the 1990s, has decreased by about 20% in the last 10 years. However, there remains considerable variation between hospitals in blood reduction, and national audits of blood components in the UK and elsewhere suggest that overall blood usage could be further reduced without compromising patient safety.5

It is difficult to assess how observational studies may have influenced these changes in transfusion practice in comparison to evidence from randomised controlled trials, national guidelines and process driven initiatives. The most likely answer is that they have all played a role in changing practice. Randomised controlled trials have found that ‘restrictive’ transfusion strategies are associated with similar or improved clinical outcomes compared with ‘liberal’ transfusion strategies.47 Many national guidelines have adopted restrictive transfusion strategies,47 while needing to make assumptions about the generalisability of the findings of randomised controlled trials in specific clinical groups of patients. There have been many smaller observational studies of process initiatives to reduce transfusion that also indicate reductions in the use of blood without any significant impact on clinical outcomes.48–50

**CONCLUSION**

The findings from this systematic review of recent large-scale observational studies show considerable variability in the patient populations and study methods when comparing the effects of RBCT on mortality. Overall, the observational studies do show a consistent adverse effect of RBCT on mortality. Although it seems unlikely that this can be entirely explained by selective sampling or a predominance of poorer quality observational studies, it remains possible that even the best conducted adjustments cannot completely eliminate the impact of confounding.

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