Prospective study of preoperative autologous blood donation for patients with high risk of allogeneic blood transfusion in lumbar fusion surgery: a study protocol of a randomised controlled trial

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ABSTRACT

Introduction Preoperative autologous blood donation (PABD) can be used to reduce the exposure of allogeneic blood transfusion in patients undergoing elective surgery. Better blood management to avoid anaemia and reduce allogeneic blood transfusion after spine surgery become increasingly important with development of enhanced recovery after surgery. We present here the design of a randomised controlled trial with three groups to verify the clinical effectiveness of PABD in patients at high risk of transfusion for lumbar fusion surgery and explore the optimal timing of autologous blood donation.

Method and analysis Patients (age 18–70 years) who will receive lumbar fusion surgery for degenerative disease with haemoglobin over 110 g/L and ‘high risk’ of allogeneic blood transfusion are eligible, unless they refuse participation or are diagnosed with malignant metastases, infection, cardiovascular and cerebrovascular diseases, haematological disorders or relevant drug history and critical illnesses. A total of 1200 patients will be recruited and randomised into three groups. Patients in group A will not receive PABD and be regarded as control group. PABD will be performed for patients in groups B and C. Blood donation will be finished at 1 week (±3 day) before surgery in group B and 2 weeks (±3 day) before surgery in group C. Primary outcome measures will include haemoglobin decline, incidence and amount of allogeneic blood transfusion. Secondary outcome measures will include days of hospitalisation after surgery, haematocrit level and incidence of complications. This study is a single-centre and open-label randomised controlled trial. The sample size is calculated with reference to the retrospective data and previous studies.

Ethics and dissemination This trial has been approved by the Peking University Third Hospital Medical Science Research Ethic Committee (no: 2020-262-02). Results of the trial will be submitted for publication in a peer-reviewed journal and as conference presentations. Trial registration number ChiCTR2000039824, preregistration.

INTRODUCTION

Preoperative autologous blood donation (PABD) can be used to reduce the exposure of allogeneic blood transfusion in patients undergoing elective surgery.1 2 Allogeneic blood transfusion is safe in our opinion currently but usually limited in clinical practice due to increasing blood shortage,3 and there are still inherent risks for allogeneic blood transfusion.4-6 Autologous blood transfusion techniques involve the collection and reinfusion of the patient’s own blood with PABD being the common used form.7-9

Elective surgery in orthopaedics like spine or hip procedures is often associated with massive blood loss and high risk of anaemia after surgery.6 10-12 Delayed wound healing and infection could be related to anaemia after surgery, and those are more susceptible and severe in orthopaedics due to the implant. Additionally, anaemia contributes to higher cardiac burden and may affect the functional exercise in the early postoperative period, which is essential for patients who underwent orthopaedic surgery. With the development of the concept of enhanced recovery...
after surgery in recent years, better blood management to avoid anaemia after surgery becomes increasingly important. In addition to an alternative in allogeneic blood shortage, there are also other advantages about PABD. It can be used for patients with rare blood groups, multiple allo-antibodies or religious objections to allogeneic transfusion. However, we should also be careful about this technique especially for the necessity, indication and cost-effectiveness in the clinical practice. PABD programme should be a multidisciplinary issue and based on a discussion between doctor and patient regarding the procedure’s risks and benefits.

Surgical procedures with high risk of allogeneic blood transfusion should be considered for PABD. The less likely the transfusion, the more likely donated blood will not be used. Lumbar fusion surgery for patients with degenerative lumbar spine diseases often results in massive blood loss due to long time of operation, large wound and high difficulty to stop bleeding in the spinal canal area. All levels of anaemia were reported to be significantly associated with prolonged length of hospitalisation and poorer operative or 30-day outcomes in patients undergoing elective spine surgery. Blood management could be essential especially for patients with spine deformity and long segments fusion. Kennedy et al found that PABD was more efficient in patients who underwent instrumentation fusion but not all spine surgery. Solves et al also reported that PABD significantly decreased the allogeneic blood transfusion for spine instrumentation fusion in young patients. However, it is still controversial about the appliance and effectiveness of PABD in spine surgery. Brookfield et al reported that it is not beneficial for patients who underwent short lumbar spine fusion with normal blood coagulation. Moreover, evidence about the timing of blood donation in PABD programme still remains insufficient. More evidence about the appliance and effectiveness of PABD in spine surgery is warranted.

In this randomised clinical trial, we aim to verify the clinical effectiveness of PABD in patients at high risk of allogeneic blood transfusion for lumbar fusion surgery with respect to the incidence and number of allogeneic blood transfusion, haemoglobin (Hgb) decline, days of hospitalisation after surgery, haematocrit level and incidence of complications. Study design of different time interval between blood donation and surgery help us to explore the optimal timing of autologous blood donation simultaneously.

METHOD

Study design

This study is planned to be a prospective and open-label randomised controlled trial with three groups (figure 1).

Recruitment and informed consent

This single-centre study will be conducted in the Peking University Third Hospital (PUTH). Eligible participants will be recruited from the patients who are going to receive lumbar fusion surgery in a 3 years period from 1 January 2022 to 31 December 2024 by our researchers. Recruitment, assessment and randomisation will be finished in the Inpatient Management Centre (IMC) that is in charge of preoperative evaluation before patients’ admission. Researchers will discuss with eligible patients when they have decided to receive lumbar fusion surgery and then finish the informed consent. After enrolment, participants will be coded as a unique number and general information will be recorded.

Eligibility

Inclusion criteria

1. Patients who will receive elective lumbar fusion surgery for lumbar degenerative disease.
2. Age between 18 and 70 years.
3. Hgb over 110 g/L.
4. ‘High risk’ for the risk score of allogeneic blood transfusion for lumbar fusion surgery.

Exclusion criteria

1. Diagnosed with malignant metastases.
2. Infectious diseases.
3. Cardiovascular and cerebrovascular diseases such as coronary heart disease and severe aortic stenosis.
4. Haematological disorders or drug history which are not suitable for blood donation.
5. Critically ill patients.
6. Refuse to participate for any reason.

Randomisation
Eligible participants recruited from the IMC in PUTH will be randomised to three groups via random number method by researcher who is blinded for outcome collection. Patients in group A will not receive PABD and be regarded as control group. PABD will be performed for patients in groups B and C. Blood donation will be finished at 1 week (±3 day) before surgery in group B and 2 weeks (±3 day) before surgery in group C. Randomisation of the three groups will be on a 1:1:1 basis.

Blinding
Participants and surgeons will not be blinded to the interventions. As the PABD plans should be informed to patients clearly. The assessment after surgery will be performed by research assistants who are blinded to the recruitment and randomisation.

Interventions
Blood donation and transfusion
All eligible participants will be randomised to three groups after preoperative evaluation. Participants in group A will not receive PABD and regarded as control group. Donation of 400 mL autologous blood will be finished once at 1 week (±3 day) before surgery for participants in group B and 2 weeks (±3 day) before surgery for group C. We also allow participants to finish the donation 3 days before or after the time points in consideration of the feasibility in practice. All donated blood in our study will be transfused back during surgery unless the Hgb level is still above 125 g/L for man and 115 g/L for woman before wound closure, then the autologous blood will not be transfused and be stored until the discharge of patients. Tranexamic acid and intraoperative blood salvage will be applied in three groups as usual. Autologous blood donation, preservation at 2–6°C in dedicated refrigerator and transfusion during surgery will be assisted and finished by department of Blood Transfusion and Anesthesiology. Patients in groups B and C will be given subcutaneous injection of 10 000U erythropoietin (EPO) every other day (quaque altera die (qod)) until the date of surgery. Time schedule of interventions can be followed in table 1.

Assessment and management
We have established a risk score of allogeneic blood transfusion for lumbar fusion surgery in the preliminary study. This score system consist of six parameters including age, body mass index, number of fusion and fixation segments, spine deformity and Hgb level (table 2). The risk score of allogeneic blood transfusion for lumbar fusion surgery was established based on the retrospective data of 5101 cases of lumbar spine surgery in the past 2 years from 2018 to 2019. We have performed preliminary validation for the risk score system in the patients from January to June 2020 prospectively. The effectiveness was acceptable with sensitivity of 76% (area under curve, AUC=0.83).

All participants will be assessed at outpatient and IMC, including demographic information, height and weight, blood test and radiographic examination. Spine surgeons

### Table 1 Time schedule of interventions

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Admission</th>
<th>1–2 weeks before surgery</th>
<th>Perioperation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preoperative evaluation</td>
<td>√</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blood donation</td>
<td>√</td>
<td></td>
<td></td>
</tr>
<tr>
<td>EPO injection</td>
<td>√</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Autologous blood transfusion</td>
<td>√</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

EPO, erythropoietin; .

### Table 2 Risk score of allogeneic blood transfusion for lumbar fusion surgery

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td></td>
</tr>
<tr>
<td>&lt;60 years</td>
<td>0</td>
</tr>
<tr>
<td>≥60 years</td>
<td>1</td>
</tr>
<tr>
<td>BMI</td>
<td></td>
</tr>
<tr>
<td>≥18.5</td>
<td>0</td>
</tr>
<tr>
<td>&lt;18.5</td>
<td>1</td>
</tr>
<tr>
<td>No. of fusion segments</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>≥4</td>
<td>4</td>
</tr>
<tr>
<td>No. of fixation segments</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>≥5</td>
<td>4</td>
</tr>
<tr>
<td>Spine deformity</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>0</td>
</tr>
<tr>
<td>Yes</td>
<td>1</td>
</tr>
<tr>
<td>Haemoglobin (g/L)</td>
<td></td>
</tr>
<tr>
<td>≥140</td>
<td>0</td>
</tr>
<tr>
<td>125–140</td>
<td>1</td>
</tr>
<tr>
<td>&lt;125</td>
<td>2</td>
</tr>
</tbody>
</table>

Total score 0–4=low risk; total score 5–13=high risk. BMI, body mass index.
will make a general surgical plan including fusion and fixation segments. Patients with low risk of allogeneic blood transfusion will be excluded from our study. Baseline iron metabolism and vitamin will be screened for all participants. Oral iron supplements and vitamin drugs will be initialed and continued until the day before surgery for patients with iron and vitamin deficiencies. All patients will receive one unit of allogeneic blood transfusion if their Hgb level drop below 8.0 g/dL and the patient display clinical symptom of anaemia (tachycardia and/or hypotension) despite intravenous fluid boluses.

Outcome measurements

Primary outcome

Incidence of allogeneic blood transfusion
Intraoperative or postoperative allogeneic blood transfusion will be recorded as binary outcome. Incidence of allogeneic blood transfusion will be statistically compared as primary outcome.

Amount of allogeneic blood transfusion
Total number (mL) of allogeneic blood transfusion for each patient in three groups will be recorded as continuous outcome and analysed as the primary outcome.

Hgb decline
Hgb decline (g/L) from 1 day before surgery to 1 day before discharge will be recorded as continuous outcome and analysed as one of the primary outcomes.

Secondary outcome

Days of hospitalisation after surgery
Length of hospitalisation after surgery could be an indicator for recovery and will be recorded as continuous outcome in this study. Days of hospitalisation after surgery will be compared and statistically analysed as secondary outcome.

Haemotocrit level
Haemotocrit check will be scheduled weekly for patients in groups B and C. Preoperative haemotocrit level will be assessed in group A regularly. Initial and preoperative haemotocrit level will be compared between two groups receiving EPO administration and recorded as secondary outcome.

Incidence of complications
Wound infection and complications associated with blood transfusion will be recorded as binary outcome and analysed as secondary outcome.

Data management
Each patient will receive a unique number and all data will be recorded with this number. An attending spine surgeon and a research assistant will be in charge of the examination and assessment in the perioperative period. Research assistant will maintain the follow-up 1 month after surgery. Data entry and transfer will be performed by two staff and two computers. All data including baseline information, risk score, Hgb result, allogeneic blood transfusion and ability assessments will be secured in PUTH and the access will be restricted to the research team. Database will be established after finishing all the data collection and back-ups will be made in multiple disks. All raw data will be kept in the Medical record department.

Patient and public involvement
Patients or public were not involved in the design, conduct, reporting or dissemination plans of our research.

Safety monitoring and adverse events
Participants in this study will receive PABD before lumbar spine surgery. All participants will be observed for 40 min after blood donation. Patients with Hgb less than 110 g/L, infectious diseases, cardiovascular or cerebrovascular diseases will be excluded from our study in consideration of safety. Strictly standard collection and storage processes will be performed and monitored by department of IMC and Blood Transfusion. All expected or unexpected adverse events from this study will be recorded and monitored. Patients suffered from any adverse events related to the interventions in research will receive free treatment.

Sample size calculation
There were 5101 cases of lumbar fusion surgery from January 2018 to November 2019 and 817 cases received allogeneic blood transfusion. The incidence of allogeneic blood transfusion was 16%. The preliminary study of PABD from August 2020 to September 2020 demonstrated that the incidence of allogeneic blood transfusion decreased by 18% compared with the same time period in 2019. We hypothesise that the incidence of allogeneic blood transfusion decreased by 18% via PABD for patients at high risk score for lumbar fusion surgery. We should recruit 400 patients for both groups PABD and non-PABD. This is based on \( \alpha \) at 0.025 and power at 80% considering a 1:1 allocation rate and accuracy rate of 80% for risk score system. On the other hand, a single injection of EPO was reported to result in Hgb increase of 2.9 g/L in adolescence. We hypothesise that a single injection of EPO could attain Hgb increase of 2.5 g/L in adults who are going to receive lumbar fusion surgery, the Hgb level in group C before surgery should be 10 g/L more than patients in group B and we assume to be 120 g/L and 110 g/L, respectively. Then we should recruit 330 patients for both groups B and C. This is based on \( \alpha \) at 0.025 and power at 80% considering a 1:1 allocation rate. To sum up, we will recruit a total of 1200 patients for three groups.

Statistical analysis
The baseline characteristics of all participants will be summarised by group and presented as means (SD) for continuous variables, and count (%) for categorical variables. All the confounding variables which may influence the primary outcome will be recorded and compared among three groups. Incidence of allogeneic
blood transfusion and complications will be measured as binary outcome. Amount of allogeneic blood transfusion (mL), Hgb decline, days of hospitalisation after surgery and haematocrit level will be measured as continuous outcome. χ² test and logistical regression will be used for the binary outcome. Non-parametric test or t test will be performed for continuous outcome according to the distribution. A value of p<0.05 will be considered as statistically significant. All analysis will be performed using SPSS V.17.0 by a researcher who is blinded to recruitment and data collection.

Ethic and dissemination
This trial has been approved by the PUTH Medical Science Research Ethic Committee (no.: 2020-262-02) and registered on ChiCTR.org (registration number: ChiCTR2000039824). Informed consent will be obtained for all participants. Results of the trial will be submitted for publication in a peer-reviewed journal and as conference presentations.

DISCUSSION
We have presented the rationale and design of a prospective randomised controlled trial to compare the outcomes of PABD in patients at high risk of allogeneic blood transfusion for lumbar fusion surgery. The randomised controlled trial (RCT) will compare the outcome among three groups to verify the clinical effectiveness of PABD and explore the optimal timing of blood donation with adjuvant EPO injection in lumbar fusion surgery.

In the previous studies of PABD, the time interval between first blood donation and surgery varied from less than 2 weeks to more than 4 weeks.14-20 Adequate time interval was thought to be crucial for the red blood cell (RBC) regeneration of donated blood.22 But it should also be noted that there is an outdate for the donated blood in plastic bags. An appropriate time interval to balance the regeneration and preservation of donated blood is significant for the efficiency of PABD programme. We choose an interval of 1–2 weeks to reduce the storage time of donated blood in autologous blood bank. Meanwhile, adjuvant EPO injection after blood donation will be used for all patients in groups B and C to accelerate the RBC regeneration. EPO injection was reported to be useful for increasing the RBC before hip surgery and avoiding allogeneic transfusion during spinal deformity surgery in PABD programme.21-23 To explore the optimal timing of autologous blood donation with EPO injection is also one of the main goals in this prospective study.

In this study, a validated risk score system of allogeneic blood transfusion for lumbar fusion surgery based on retrospective study with large sample size will be implemented to ensure the necessity for PABD programme. Both this technique and EPO injection should be more efficient in the patients with higher risk of blood transfusion.20-24 Waste of donated blood was an inherent risk for PABD programme.25-26 The review by Singbartl reported that the wastage of unneeded PABD units varied from 18% to above 50%.27 All donated blood in our study will be storage for a relatively shorter time period and transfused back during operation unless the Hgb level is still above 125 g/L for man and 115 g/L for woman before wound closure, then the autologous blood will not be transfused and be storage until the discharge of patients. This design aims to eliminate the waste of donated blood and simultaneously decrease the risk of allogeneic blood transfusion in perioperative period for targeted patients who will receive lumbar fusion surgery. A multidisciplinary cooperation including department of IMC, Blood Transfusion, Anesthesiology and Orthopedics in the hospital will ensure the safety and feasibility of this prospective trial in clinical practice.

REFERENCES