Oxygen delivery during cardiopulmonary bypass (and renal outcome) using two systems of extracorporeal circulation: a retrospective review

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Abstract

OBJECTIVES: To investigate the combined influence of blood flow and haemodilution with either a miniaturized (Mini-CPB) or a conventional cardiopulmonary bypass (C-CPB) circuit on average oxygen delivery during bypass. The influence of this on clinical outcome, particularly renal dysfunction after routine coronary artery bypass surgery (CABG), was measured.

METHODS: Retrospective analysis in two groups of 160 patients based on the surgeon’s preference for bypass circuit. We compared consecutive patients undergoing isolated CABG surgery by two surgeons using Mini-CPB with a matched cohort of patients, from the same period, undergoing isolated CABG surgery by four other surgeons using a C-CPB. No trial-related intervention occurred. Data on bypass circuit parameters and clinical outcomes were acquired from routinely collected data sources.

RESULTS: Average cardiopulmonary bypass pump flow was significantly lower with Mini-CPB compared with C-CPB. Mini-CPB resulted in significantly less haemodilution. The resultant calculated average oxygen delivery provided by the two systems was the same. Percentage change in plasma creatinine was significantly and inversely related to the oxygen delivery during CPB. There was no difference in percentage change in plasma creatinine between groups. The risk of having Acute Kidney Injury Network (AKIN) score ≥1 increased 1% for every 1 ml min⁻¹ m⁻² decrease in oxygen delivery (P = 0.0001, OR 0.990, 95% CI 0.984–0.995).

CONCLUSIONS: Despite aiming for the same target pump flow, periodic limitations of venous return to the pump resulted in a significant reduction in average flow delivered to the patient by Mini-CPB. Less haemodilution compensated for this reduction, so that the average oxygen delivery was the same. The association between oxygen delivery and postoperative change in plasma creatinine was evident in both groups. Further work to understand whether there is a particular cohort of patients who benefit (or are put at risk) by one method of CPB vs the other is warranted.

Keywords: Oxygen delivery • Extracorporeal circulation • Miniaturized cardiopulmonary bypass • Cardiac surgery

INTRODUCTION

The factors related to the process of cardiopulmonary bypass (CPB) that contribute most significantly to ‘optimal perfusion’ remain an important topic of debate. Randomized data on best practice in managing haemodilution, perfusion pressure, haematocrit and pump flow are extremely limited, but in an extensive review of the literature [1], the authors conclude that oxygen delivery remains ‘one of the most important determinants of optimal perfusion’.

Organ dysfunction after cardiac surgery has been linked to a decrease in oxygen delivery during CPB [2, 3]. Conventional CPB (C-CPB) requires an initial crystalloid prime of 1500–2000 ml, resulting in dilutional anaemia at the onset of bypass. Autologous priming of the circuit after cannulation reduces the prime, but is incomplete, and haemodilution still occurs. Acute haemodilution during cardiac surgery is associated with an increased risks of renal failure [4–6], stroke [7] and mortality [8–10].

A recent development, miniaturized cardiopulmonary bypass (Mini-CPB), has appeared to offer theoretical advantages. These include the absence of a venous reservoir, considerably lowering the priming volume to 200–500 ml after the circuit is retrogradely primed, resulting in minimal haemodilution. This should increase the oxygen delivery (DO₂) during bypass, but there are suggestions that the pump flow achieved with Mini-CPB is lower than with C-CPB [11], and that periods of ‘low flow’ are more frequent and last longer [12].

We evaluated the combined influence of blood flow during bypass together with any change in haematocrit concentration...
on average oxygen delivery. Because of the frequently reported association between oxygen delivery and subsequent renal dysfunction [2, 13], we used this as the main marker of differences in clinical outcome.

**MATERIALS AND METHODS**

Over a 1-year period, a total of 160 patients underwent coronary artery bypass grafting (CABG) with the Mini-CPB circuit (two surgeons). After approval from the Cornwall and Plymouth Research Ethics Committee, we compared these patients with the first 160 who had undergone CABG with C-CPB during the same period (four other surgeons). We used data from the cardiac surgery and perfusion database.

**Conventional and miniaturized cardiopulmonary bypass**

The C-CPB circuit was a standard open circuit with a centrifugal arterial pump, membrane oxygenator and hard-shell reservoir (Quadrox-I, Maquet Cardiopulmonary). Venous return was gravity-dependent with ½ in. tubing. The circuit was primed with 1000 ml Hartmann’s solution, 700 ml gelofusine, 100 ml 20% mannitol and 5000 IU heparin. No retrograde autologous priming was used. Cardiotomy suction was used and aspirated blood returned into the reservoir. A cell-saver device was used depending on surgeon preference.

The Mini-CPB circuit (Maquet Cardiopulmonary) was a preconnected closed-loop circuit incorporating a RotaFlow centrifugal pump, a Quadrox-I diffusion membrane oxygenator and a Quart arterial blood filter. There was both an arterial and a venous bubble detector and a venous bubble trap (Maquet Cardiopulmonary). Pericardial blood was aspirated into a cell-saver device (Electa, Sorin). If sufficient volume was aspirated, it had been processed and re-transfused at the end of the operation. In cases where large volumes returned to the cell saver, the processed blood was returned to the circuit via the bubble trap. The circuit was heparin coated (Bioline Coating, Maquet Cardiopulmonary) and primed with 800 ml Gelofusine that was reduced to between 200 and 500 ml by retrograde autologous priming following aortic cannulation.

**Anaesthesia and surgical management**

All patients were anaesthetized with a low-dose opioid technique, using 10 µg kg⁻¹ of fentanyl. Maintenance of anaesthesia was done with a combination of isoflurane and propofol. Surgical technique did not differ between groups. Arterial access was the same in all patients and was achieved through an ascending aorta cannulation [24-fr angled (72 524, Medtronic)], and venous access was done through a two-stage venous cannula inserted through the right atrial appendage (91251C, Medtronic).

**Perfusion management**

Moderate hypothermia to 34°C, alpha-stat pH management and target flow rates of 2.5 l min⁻¹ m⁻² were used for both groups. Mean arterial pressure target was 50–60 mmHg. Anticoagulation to achieve an activated clotting time >400 s was achieved with heparin 300 IU kg⁻¹. Final cardioplegia concentrations were the same (K⁺ 20 mmol l⁻¹ induction and 10 mmol l⁻¹ maintenance). In the Mini-CPB group, retrograde autologous priming of the circuit was undertaken over ≤1 min prior to the commencement of bypass. Target haematocrit (Hct) during bypass was maintained at 21% or higher, with red blood cell transfusion given as necessary.

**Intensive care unit**

A unit-based transfusion policy was used for both packed red blood cell transfusion, to maintain the haematocrit >24%, and for the transfusion of platelets and fresh-frozen plasma depending on the results obtained with thrombo-elastography. Standard protocols were used for weaning from the ventilator, discharge from the Intensive Care Unit (ICU) and High-Dependency Unit (HDU).

**Study end-points**

Average pump flow over the whole period of bypass, arterial PaO₂, and arterial oxygen saturation (SaO₂) (Sorin, Datamaster) were automatically collected by the JocapXL data acquisition system on the bypass machine. The JocabXL has a sample frequency every 15 s. Arterial blood gas measurements were taken every 30 min during bypass, measured with the ABL815 blood gas analyser (Radiometer) and entered onto the electronic database management system. An average haemoglobin (Hb) concentration was calculated from these measurements, together with the peak lactate concentration. Average indexed oxygen delivery (DO₂I) was calculated:

$$DO₂I \, (ml \, min^{-1} \, m^{-2}) = \frac{\text{average pump flow} \, \text{(ml \, min \, m}^2\text{)}}{\text{arterial blood flow} \, \text{(ml \, min \, m}^2\text{)}} \times \left[ \left( Hb \, (g \, dl^{-1}) \times 1.36 \times SaO₂(\%) \right) + 0.003 \times PaO₂ \, (mmHg) \right]$$

Routine blood counts, biochemistry and number and type of blood transfusion given were recorded by the hospital pathology system. Time of discharge from the critical care and hospital was recorded on the hospital patient manager system. Postoperative information, including requirement for renal replacement therapy or reoperation, was recorded on the cardiac surgery database.

**Data acquisition and statistical analysis**

Continuous variables were compared using a Student t-test to test for a difference in means between the two patient groups. Two-sample tests of proportions were used to test for differences in proportions of patients with given characteristics or outcomes in the two groups. Comparisons of dichotomous data were made using Fisher’s exact test. A Bonferroni adjustment was applied to allow for multiple testing.

For all the tests, a P-value of ≤0.05 was deemed statistically significant. Statistical analysis was performed using the software package R (R Development Core Team, Vienna, Austria).
Summary statistics for patient demographics, comorbidities, intraoperative and ICU variables and clinical outcomes are reported as Bonferroni adjusted mean ± standard deviation or n (%), [95% confidence interval (Mini-CPB - C-CPB)].

RESULTS

Patients’ demographics, comorbidities and procedural characteristics are presented in Table 1, and show a well-matched patient population, but patients in the Mini-CPB group started at a slightly higher Hb and had a marginally lower left-ventricular ejection fraction. One patient in the C-CPB group had an incomplete dataset and was excluded from the final analysis.

Intraoperative variables

Table 2 shows the intraoperative variable between the two groups. Initiation of bypass diluted patients on Mini-CPB 30.5% compared with 39.3% with C-CPB (P < 0.0005). This resulted in a greater percentage of C-CPB patients requiring a perioperative packed red blood cell (PRBC) transfusion to maintain Hb above target, compared with Mini-CPB. Patients in the C-CPB group were also more likely to require a platelet transfusion, while the fresh-frozen plasma (FFP) requirement, based on coagulation parameters, was the same (Table 2). Despite the higher transfusion rate on the day of surgery, patients in the C-CPB had a significantly lower Hb and percentage fall in Hb from their preoperative level by Day 1.

Average CPB pump flow index with Mini-CPB was significantly lower. When this figure was combined with average Hb during bypass, average PaO2 and arterial saturation, this resulted in the same DO2/l with Mini-CPB compared with C-CPB (Fig. 1A–C). The peak lactate measured during bypass was higher with C-CPB compared with Mini-CPB.

Postoperative management and morbidity

Table 3 shows estimated differences in ICU variables between the two groups, in terms of proportions of patients with a given characteristic. By the time of discharge, similar numbers of patients in each group had received PRBC transfusion.

Daily measurement of serum creatinine demonstrated the same peak rise and maximum percentage rise in plasma creatinine irrespective of group. We classified individual patients by the criteria described by the Acute Kidney Injury Network (AKIN) [14]. Similar numbers of patients in each group developed AKIN scores 1–3. Ten patients (Mini-CPB) and 9 patients (C-CPB) developed AKIN 2 or 3, with 6 patients in the Mini-CPB group and 7 in the C-CPB group requiring postoperative renal replacement therapy. Using a Fisher’s exact test, there was no evidence of an association between group and AKIN scores (P = 0.21).

Both groups had an association between average oxygen delivery and postoperative percentage change in plasma creatinine (ΔCr). The difference in this effect between groups was not significant. For every 1 ml min$^{-1}$ m$^{-2}$ increase in oxygen delivery, ΔCr decreased by 0.0012 (P = 0.006).

A logistic regression model revealed a strong risk of developing AKIN 1 or more, compared with a score of 0, with a decreasing oxygen delivery. There was no evidence of a similar effect with flow index. The risk of having AKIN ≥1 increased by ~1% for every 1 ml min$^{-1}$ m$^{-2}$ decrease in oxygen delivery (P = 0.0001, OR 0.99, 95% CI 0.984–0.995).

Length of stay

Length of stay in critical care and in hospital was the same between groups. We used a multiple regression model fitted to the log ICU stay, incorporating the covariates flow during bypass, oxygen delivery and AKIN score 2 or 3. Backward elimination, to include only the AKIN score, demonstrated that compared with those with an AKIN score of 0, patients with AKIN 1 are estimated

### Table 1: Patient demographics

<table>
<thead>
<tr>
<th>Variables</th>
<th>Mini-CPB (n = 160)</th>
<th>C-CPB (n = 159)</th>
<th>Statistics P [95% CI (Mini-CPB to C-CPB)]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>67.1 ± 10</td>
<td>66.3 ± 10</td>
<td>NS (−1.40 to 3.00)</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>171.9 ± 8</td>
<td>172.1 ± 8</td>
<td>NS (−1.96 to 1.56)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>84.1 ± 16</td>
<td>84.7 ± 14</td>
<td>NS (−3.91 to 2.71)</td>
</tr>
<tr>
<td>Body surface area (m$^2$)</td>
<td>2.0 ± 0.2</td>
<td>2.0 ± 0.2</td>
<td>NS (−0.04 to 0.04)</td>
</tr>
<tr>
<td>Logistic EuroSCORE</td>
<td>4.4 ± 6.0</td>
<td>3.7 ± 5.0</td>
<td>NS (−0.52 to 1.92)</td>
</tr>
<tr>
<td>'Urgent' case (n)</td>
<td>45</td>
<td>37</td>
<td>NS (P = 0.37)</td>
</tr>
<tr>
<td>Ejection fraction (%)</td>
<td>55.1 ± 10.5</td>
<td>60.7 ± 14.2</td>
<td>P &lt; 0.0005 (−8.35 to −2.85)</td>
</tr>
<tr>
<td>Diabetes (n)</td>
<td>32</td>
<td>26</td>
<td>NS (P = 0.39)</td>
</tr>
<tr>
<td>NYHA grade 3/4 (n)</td>
<td>32</td>
<td>40</td>
<td>NS (P = 0.29)</td>
</tr>
<tr>
<td>Hypertension (n)</td>
<td>97</td>
<td>106</td>
<td>NS (P = 0.30)</td>
</tr>
<tr>
<td>Preop Hb (g dl$^{-1}$)</td>
<td>14.0 ± 1.5</td>
<td>13.6 ± 1.3</td>
<td>P &lt; 0.011 (0.091 to 0.71)</td>
</tr>
<tr>
<td>Preop Cr (µmol l$^{-1}$)</td>
<td>95 ± 25</td>
<td>99 ± 43</td>
<td>NS (−11.7 to 3.74)</td>
</tr>
<tr>
<td>CPB time (min)</td>
<td>68 ± 20</td>
<td>68 ± 21</td>
<td>NS (−4.52 to 4.52)</td>
</tr>
<tr>
<td>Aortic cross-clamp (min)</td>
<td>42 ± 14</td>
<td>40 ± 14</td>
<td>NS (−1.08 to 5.08)</td>
</tr>
</tbody>
</table>

Differences in preoperative demographic variables, co-morbidities and bypass times between patients before either Mini-CPB or C-CPB. Each variable is shown as either mean ± SD, or absolute number.
to stay in ICU for approximately twice as long (95% CI 1.69–2.53), while those with a score of 2 or 3 are estimated to stay in ICU for approximately three times as long (95% CI 2.31–4.30).

**Major morbidity and mortality**

Four patients in the C-CPB group and 3 in the Mini-CPB group died before discharge from hospital.

**DISCUSSION**

We have shown that despite the same target flow indexed flow is limited at some point, and for long enough, to significantly reduce the average flow during perfusion with the Mini-CPB compared with C-CPB. We have also demonstrated that, despite this reduction in flow, the effects of minimal haemodilution with Mini-CPB results in the preservation of average oxygen delivery.

This study concurs with previous studies, and with the conclusion of a best-evidence review in this journal [15], by demonstrating the benefits of the Mini-CPB circuit, together...
CONCLUSIONS

Miniaturized cardiopulmonary bypass is associated with lower average indexed flows compared with conventional circuits despite aiming for the same target flows. The reduction in haemodilution with Mini-CPB results in the preservation of oxygen delivery despite lower net flows. In both groups, a postoperative rise in plasma creatinine and risk of developing AKIN 1 are strongly associated with any fall in oxygen delivery.

Conflict of interest: none declared

REFERENCES