Veno-arterial extracorporeal membrane oxygenation without therapeutic anticoagulation for intra-operative cardiopulmonary support during lung transplantation

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Abstract: Cardiopulmonary support is frequently required during lung transplantation. Veno-arterial extracorporeal membrane oxygenation (VA-ECMO) has been recently shown to be an acceptable and, perhaps, safer alternative to cardiopulmonary bypass machine (CBM) for lung transplantation. However, therapeutic anticoagulation used during both these techniques can cause significant bleeding diathesis, especially in the setting of re-transplantation or when severe adhesions are encountered. Technological advances have reduced the thrombogenic potential of VA-ECMO circuits and unlike CBM, VA-ECMO does not include a reservoir containing a stagnant column of blood. Here, we demonstrate that VA-ECMO can be used without therapeutic anticoagulation for short-term intraoperative support.

Keywords: Anticoagulation; extracorporeal membrane oxygenation (ECMO); Lung transplantation; heparin

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Introduction

Anticoagulation for intraoperative mechanical cardiopulmonary support during lung transplantation is associated with significantly increased bleeding complications especially when pleural adhesions are encountered or during re-transplantation (1). Cardiopulmonary bypass machine (CBM) has a reservoir containing a stagnant column of blood that necessitates therapeutic anticoagulation to avoid thrombosis. However, veno-arterial (VA) extracorporeal membranous oxygenation (ECMO) does not have such a blood reservoir. Furthermore, technological advancements including heparin-bonded tubing, heparin-coated polymethylpentene oxygenators, and centrifugal pumps have made the components in contemporary ECMO circuits significantly less thrombogenic (2,3). Hence, we postulated that it might be possible to use VA-ECMO without therapeutic anticoagulation. We recently reported the successful anticoagulation-free veno-venous ECMO support for respiratory failure and bridge to lung transplantation (4). Here, we used VA-ECMO without therapeutic anticoagulation for intraoperative support during lung transplantation.

Case presentation

A 65-year-old male with end-stage lung disease and severe pulmonary hypertension from pneumoconiosis underwent left lung transplantation through posterolateral thoracotomy. A prior right-sided Clagett window (Figure 1A) for chronic empyema associated with significant pleural plaques (Figure 1B) precluded double lung transplantation. Intraoperatively, the patient was intolerant to right single lung ventilation and developed hypoxemia. Dense pleural adhesions and hilar scarring were encountered. The patient was, therefore, placed on VA-ECMO. A 6 mm Hemashield graft (vendor) was sewn onto the left femoral artery and used as arterial inflow. The femoral vein was directly cannulated using a 25 F Biomedicus multi-port cannula for venous drainage. The...
VA-ECMO was initiated and blood flows maintained at >4 liters/min. Oxygen saturations were monitored on the right hand to ensure adequate oxygenation of arch vessels. Left pneumonectomy was performed after extensive extrapleural mobilization, pleurolysis and tedious dissection of hilar structures. Subsequently, implantation of left single lung transplantation was performed with a total ischemic time of 210 minutes. The activated clotting time (ACT) was less than 160 seconds during the procedure. To de-air the allograft, pump flows were reduced to obtain right ventricular cardiac output of 1.5–2 L/min. Since we reduced the flows, we administered 5,000 U of unfractionated heparin. The allograft was ventilated after which the arterial and venous cannulas were removed, after clamping them for 5 minutes. The flow in the ECMO circuit was kept flowing using a bridge while we established hemodynamic and respiratory stability prior to removing the cannula. Circulation using an ECMO bridge was performed to facilitate going back on ECMO in case there was development of primary graft dysfunction. Total VA-ECMO time was 107 minutes. The patient had an eventful post-operative course and was discharged breathing room air on day 10 without evidence of arterial and venous thromboembolism. The patient received routine prophylaxis using unfractionated heparin for deep vein thrombosis.

**Discussion**

Several centers including Toronto General Hospital, Vienna Medical Center, and Columbia University have published the feasibility of VA-ECMO during lung transplantation (5-7). The current guidelines of the Extracorporeal Life Support Organization (www.elso.org) recommend ACT levels of 180–220 for VA-ECMO. We postulate that VA-ECMO can be used safely without therapeutic anticoagulation particularly for short-term intra-operative support. To reduce thrombogenicity, we believe that the flows should be kept high and therefore adequate sized venous and arterial cannulas should be used. If there is reduction in circuit flow, for example due to peripheral vasodilation, extra fluid or vasopressors need to be administered intravenously since ECMO is a closed circuit. Alternatively, if flows over 3.5–4 L/min cannot be achieved consistently we recommend administering unfractionated heparin to reduce thrombogenicity. Our practice is to give 5,000 U of unfractionated heparin. The generally accepted guidelines for anticoagulation during VA-ECMO are based on limited or anecdotal experience (8). This case reflects equipoise in the conventional use of therapeutic anticoagulation for VA-ECMO. Prospective studies are required to evaluate use of VA-ECMO without heparin, especially in patients with high bleeding risk. Long-term use of VA-ECMO without systemic anticoagulation should also be further investigated. Avoiding or minimizing anticoagulation in selected patients may greatly improve the safety of ECMO. However, in patients with a pro-coagulant state, anticoagulation-free VA-ECMO is not advised.

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Footnote

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