Extracorporeal membrane oxygenation
Warwick Butt¹-³ and Graeme MacLaren¹,²,⁴ *

Addresses: ¹Paediatric Intensive Care Unit, Royal Children’s Hospital, 50 Flemington Road, VIC 3052, Australia; ²Department of Paediatrics, The University of Melbourne, VIC 3010, Australia; ³Murdoch Children’s Research Institute, Clinical Sciences, 50 Flemington Road, VIC 3052, Australia; ⁴Cardiothoracic Intensive Care Unit, National University Health System, 5 Lower Kent Ridge Road, Singapore 119074

* Corresponding author: Graeme MacLaren (gmaclaren@iinet.net.au)

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Abstract
Extracorporeal membrane oxygenation (ECMO) is an advanced form of life support technology whereby venous blood is oxygenated outside of the body and returned to the patient. ECMO was initially used as last-resort rescue therapy for patients with severe respiratory failure. Over the last four decades, it has developed into a safe, standard therapy for newborns with progressive cardiorespiratory failure, as a resuscitation therapy after cardiac arrest, and in combination with other treatments such as hypothermia and various blood filtration therapies. ECMO has also become routine for children and adults with all forms of cardiogenic shock and is also routine in early graft failure after transplantation. The one area of ongoing debate is the role of ECMO in adults with hypoxemic respiratory failure. As ECMO equipment becomes safer, earlier use improves patient outcomes. Several modifications of the two basic venovenous and venoarterial ECMO systems are now occurring, as are many minor variations in cannulation strategies and systems of care for patients receiving ECMO. The indications and situations in which ECMO have been tried continue to change, and ECMO for sub-acute and chronic illnesses is now commonplace, as is the use of ECMO in patients with clinical problems previously regarded as contraindications, such as sepsis, malignancy, and immunosuppression.

Introduction
ECMO is an advanced form of life support technology, whereby venous blood is removed from the patient and passed through a membrane lung where oxygen is added and carbon dioxide removed. The blood is then returned to the patient [1]. Both circulatory and respiratory support may be provided if oxygenated blood is returned to an artery (venoarterial ECMO). If oxygenated blood is returned to a vein (venovenous ECMO), then only respiratory support is provided. ECMO can keep patients alive for days to weeks even when the heart or lungs are completely non-functional [2]. Other forms of extracorporeal technology can be added onto the ECMO circuit to provide support for other organ systems [3]. For example, connecting a hemofiltration circuit to the ECMO system to serve as an artificial kidney is frequently done in many centers.

ECMO began in the early 1970s as a prolonged form of cardiopulmonary bypass to treat severe respiratory failure. The initial approach with ECMO was to siphon blood from a vein and return it to an artery (venoarterial ECMO), and the duration was fixed at 5 days. Treatment was started when death was viewed as near-certain, because of concerns about the severity of complications at the time. The first randomized controlled trial [4], published in 1979, reflected this, and patient outcomes were poor in both conventional and ECMO groups. One of the authors (Robert Bartlett) decided that the therapy held promise but what was required were patients with a disease that would recover quickly within 5 days. He successfully applied ECMO in newborn infants with acute respiratory failure and persistent pulmonary hypertension of the newborn, which usually resolves within a few days of birth.
During the 1980s, the use of ECMO rapidly proliferated as early successes with newborn respiratory failure were reported. Trials by committed clinicians used novel statistical methods to try to establish the benefits of ECMO, but controversy remained as to the efficacy and safety of the technology. A randomized trial of inverse ratio ventilation and extracorporeal carbon dioxide removal in adult respiratory distress syndrome [5] showed no benefit, further dissuading clinicians in adult medicine from adopting the technology. However, ECMO was used in children with cardiorespiratory failure and sepsis and adapted to children with cardiogenic shock following cardiac surgery. As results improved, ECMO gradually gained credibility as a useful treatment in children, and a randomized controlled trial of ECMO in severe neonatal respiratory failure, published in 1996, showed improved survival and better long-term neurodevelopmental outcomes in the ECMO group [6,7].

Refinements in clinical management and ECMO equipment continued, and ECMO gradually changed from a “rescue therapy of last resort” to a standard therapy with clear indications and contraindications. Results continued to improve, and innovations in circuit design resulted in wider applications to more diverse clinical situations. The term ECMO was changed to extracorporeal life support (ECLS) and this acronym is used instead of ECMO in many parts of the world. Resuscitation and transport of critically ill patients on ECMO has become routine. There has also been increasing use of associated adjunctive therapies, such as hypothermia or various types of filtration therapies, as part of the ECMO circuit. The structure and nature of ECMO programs and models of care for patients on ECMO are also varied and depend on local skills and expertise rather than any one definitive or “best practice” model. This leads to variations in practice in different parts of the world.

Goals of support
The fundamental goal of ECMO is to enhance systemic oxygen delivery. The oxygen saturation goals vary from patient to patient and from center to center. The circuit flow rate chosen may also vary substantially and consequently alters much of how ECMO is done. For example, in pediatric sepsis guidelines, a maximum blood flow rate of 110 mL/kg/min has been recommended [8], although there are reports of improved survival with 150 to 200 mL/kg/min [9]. For adult ECMO patients with acute respiratory distress syndrome, some centers aim for an arterial oxygen saturation of more than 80%, while others prefer more than 90%. These differences in management goals have an impact on the way ECMO is conducted, with regards to catheter size and cannulation strategies.

Cannulation strategies and organ support
The type of ECMO applied is determined by the type of organ support required, the type of patient being treated, and the desired outcome. Support of respiratory function is usually done with venovenous cannulation; oxygen delivery is determined by the proportion of the patient’s cardiac output being diverted through the extracorporeal circuit (i.e. circuit flow rate), whilst carbon dioxide clearance is principally determined by the amount of fresh gas flow through the oxygenator. If there is a large amount of blood flow required for the circuit or if the ECMO catheters are in close proximity to each other, then recirculation is possible. This is the term for the recycling of already oxygenated blood back into the circuit, with a consequent decrease in overall patient oxygen delivery. Patient diagnoses (acute or chronic disease), mobility and need for physiotherapy (e.g. chronic obstructive pulmonary disease, cystic fibrosis), and ultimate destination (recovery or transplantation) all subtly affect management strategies. Thus many different venovenous strategies exist, depending on the goals of respiratory support. These include single-catheter tidal flow, double-lumen systems, or two- and three-catheter systems.

Support of cardiac function (either single or both ventricles) will require venoarterial cannulation, and the degree of support is determined by the amount of circuit blood flow. Blood may or may not be ejected by the failing ventricle. The diagnosis, age of the patient, and the ultimate destination are important in determining the type of cannulation performed. In infants who weigh less than 10kg, these catheters are generally placed in the neck, and in the groin in older children and adults. After cardiac surgery, some centers use central or transthoracic catheter placement.

Complications of different cannulation strategies
Catheter placement is either by surgical cut-down and insertion under vision or percutaneous with the use of ultrasound. In small infants and children, surgery is generally preferred, and vessel repair is performed at removal; in adults, percutaneous placement is standard, with surgery occasionally needed to help with difficult access or to repair arterial vessels.

Central cannulation requires the surgical insertion of catheters into major vessels or directly into either atria and therefore carries an increased risk of bleeding and infection. The sternum and skin may or may not be closed, and further major surgery is required to remove these catheters. This strategy is common in small infants but increasingly uncommon in adults.

Venoarterial ECMO in the groin vessels can be associated with cerebral hypoxemia if ventricular function recovers.
whilst severe respiratory failure is present [10]. The right ventricle propels blood through the poorly functioning lungs, which leads to desaturated blood coming into the left ventricle and then being ejected into the ascending aorta. This is known as differential cyanosis. It can be treated with a number of different strategies, including increasing ventilatory support of the lungs (by increasing the fraction of inspired oxygen or increasing the amount of positive end-expiratory pressure), adding an additional venous catheter to return a proportion of oxygenated blood into the venous system (venoarteriovenous ECMO), moving the arterial return catheter more proximally into the right subclavian or carotid artery, or changing to central ECMO. Femoral cannulation is associated with substantial vascular complications in 10% of cases [9] and these may include ischemia and the need for amputation.

Complications of extracorporeal membrane oxygenation
The complications of all extracorporeal technology are similar and include bleeding, thromboembolism, access difficulties, and infection. Mechanical equipment failure is now very uncommon with modern systems. The incidence of major complications that impact on long-term outcomes is also lower and varies depending on the type of patient, severity of illness, and indication for ECMO. As indications for ECMO and treatment protocols change, so do the complication rates. Use during cardiac arrest increases the likelihood of cerebral hemorrhage; use after trauma, in severe multiorgan failure, or initiation after seven days of mechanical ventilation is complicated and more frequently associated with bleeding.

Circuit design and equipment
The basic ECMO circuit is shown in Figure 1. The circuit consists of vascular access catheters, a pump, an oxygenator, a temperature control system, monitors, and access points.

Many different catheters can be employed, but catheter selection is based on local preference and little debate occurs about these. However, much debate exists about the relative merits of the roller pump (North America) and the centrifugal pump (Europe, Asia, and Australasia). Both pumps have potential problems. The roller pump uses gravity drainage and a bladder as a reservoir and thus clotting is possible. Tubing rupture has also occurred due to compression. The centrifugal pump generates suction at the inlet, and hemolysis is possible. A recent study highlighted this as a potentially serious issue for newborn infants [11].

A number of different oxygenators are used worldwide. Heparin bonding and newer materials are used to minimize the inflammatory response and platelet adhesiveness to these materials. It is likely that a truly biocompatible plastic will eventually be developed (possibly with an endothelial coating) and obviate the need for anticoagulation. The different oxygenators have different resistances and gas exchange properties, but in practice these do not substantially alter patient outcome and thus local preferences determine which is used.

In-line saturation monitors are being replaced by non-contact external devices that monitor temperature, blood saturation, and flow. Standards for the use of pressure monitors vary, but those pediatric programs that use centrifugal pumps tend to monitor access and return pressures, while programs that use roller pumps tend to monitor pre- and post-oxygenator pressures. This latter approach is common in adult centers irrespective of the type of pump used.

Access points to the circuit have obvious benefits for circuit sampling, monitoring, and connection of other devices (such as dialysis) but these are also points of turbulent blood flow and potential thrombus formation. Circuit bridges (connections between access and return parts of the circuit adjacent to the cannulas) are points of stasis and blood clotting. These were useful to allow circuit changes with minimal disruption of patient support; with modern equipment, many pediatric centers and most adult centers are increasingly eliminating these from the ECMO circuit.
Anticoagulation is essential for the safe use of ECMO at present. A number of key factors influence the likelihood of blood clots forming in the circuit, in particular the patient’s inherent coagulability and the rate of circuit blood flow. Newborn infants, patients with a large systemic inflammatory response after cardiopulmonary bypass, or patients with disseminated intravascular coagulation from sepsis are likely to have circulating pro-coagulants and develop clots within a circuit. Stasis of blood will also lead to clots and therefore low circuit blood flow rates require increased anticoagulation. The commonest anticoagulant is heparin, but enoxaparin, bivalirudin, argatroban, and antiplatelet agents such as prostacycline have been used.

Institutions all use different protocols of anticoagulation and monitor anticoagulation in different ways. Blood tests of clotting effectiveness, such as activated partial thromboplastin time, prothrombin time, anti-Xa levels, activated clotting time, and thromboelastography, have all been used to assess the balance between bleeding and thrombosis. Pediatric programs tend to initiate anticoagulation substantially earlier and aim for higher targets than adult programs because of the lower blood flow rates, larger circuit volumes relative to patient blood volume, and differences in infant coagulation physiology.

Indications and contraindications
The indications for ECMO are continually changing. Initial indications were acute reversible heart or lung disease in patients likely to die. However, high doses of inotropes and high pressure/volume ventilation are dangerous; prolonged hypotension or hypoxia also leads to multiorgan failure. Coupled with the increasing safety of ECMO, this realization has led to the earlier use of ECMO with consequent patient benefits.

ECMO has now become established as a means of resuscitating patients from acute cardiogenic shock, cardiac arrest, and acute fulminant respiratory failure; this gives time to identify other treatments that may be effective. Bridging to long-term mechanical support (such as ventricular assist devices) and transplantation is routine [12-14]. The role of ECMO in adults with severe respiratory failure continues to be debated, but increasing numbers of patients are receiving the treatment. Much of the debate revolves around timing and therapeutic strategies rather than whether it works [15-19].

ECMO was historically contraindicated in patients with sepsis and immunosuppression, but now survival of these patients can be as high as 50% to 75%. Similarly, patients with trauma or bleeding were not offered ECMO, but there are now reports of successful outcomes in these patients.

Outcomes
Survival after ECMO is determined by many factors, including the nature and severity of disease, age, size, pre-existing patient co-morbidities, and attitudes towards quality of life after critical illness. These factors are independent of any ECMO complications; any complications that affect patient outcome such as cerebral hemorrhage or systemic thromboembolism are additive.

Evidence of benefit and current role of extracorporeal membrane oxygenation
There are no trials that demonstrate the absolute benefit of ECMO in improving survival or quality of life. One reason for the paucity of controlled trials is that it is extremely difficult to design a randomized controlled study of life support that is ethically acceptable and yet has the utmost scientific rigor [20,21]. There can be no question that further trials of ECMO are needed, but controversy exists as to how these should be conducted [22]. Nonetheless, over 55,000 patients have been reported with a survival to discharge of more than 60% [23], with outcomes varying with age and diagnosis.

Sufficient experience with ECMO has accumulated worldwide that it is regarded as a standard of care for cardiogenic shock in both pediatric and adult medicine [24,25]. It is also regarded as a standard of care in refractory neonatal and pediatric respiratory failure [6]. However, the debate continues over adult respiratory failure in many centers [19]. Two recent reports illustrated the difficulties of studying ECMO for this indication and both involved adults with hypoxic respiratory failure from influenza A (H1N1). The first study used a number of different statistical techniques, including propensity matching to compare two similar groups of patients in the United Kingdom; one group received ECMO and the other did not [26]. Transfer to an ECMO-capable hospital was associated with lower mortality irrespective of the statistical technique used.

The second study used similar statistical methods to analyze a comparable French cohort but was unable to demonstrate that ECMO was associated with improved survival [27]. However, 51 patients in this series received ECMO but were not able to be matched to non-ECMO controls. Despite having significantly worse gas exchange before ECMO than the matched cohort, they had significantly lower mortality (22% vs. 50%, \(P < 0.01\)). The presence of this group made interpretation of the study’s results more difficult. Although a large randomized controlled trial showed that referral to an ECMO-capable center was associated with reduced mortality [28], the methodology was heavily criticized. Complex and potentially conflicting results such as these continue to fuel the controversy. Another multinational trial of
Extracorporeal Life Support Organization

The Extracorporeal Life Support Organization (ELSO) was founded in 1989. The group maintains an international registry that contained details of 55,668 patients by July 2013. The results of this registry are published biannually and distributed to members. The complications in patients who receive ECMO are recorded by ELSO; this list is very comprehensive and includes all events whether due to ECMO or not.

Conclusions

ECMO is a standard therapy in critical care. It is used as a treatment for acute severe cardiorespiratory failure and as a rescue strategy in many clinical scenarios. It is also used as a “bridge” to other treatments and transplantation. It continues to be applied to more complex and chronic situations. It is being integrated into multiple-organ support therapies. Substantial improvements in biotechnology and clinical practices over the last 40 years have allowed ECMO to provide a vital role in acute organ support in patients of all ages. It is likely that further such advances will diminish complication rates, facilitate more widespread adoption of the technology in middle- and high-income countries, and improve outcomes from refractory heart, lung, and multiorgan failure.

Abbreviations

ECMO, extracorporeal membrane oxygenation; ELSO, Extracorporeal Life Support Organization.

Disclosures

The authors declare that they have no disclosures.

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