Revised protocol of extracorporeal membrane oxygenation (ECMO) therapy in severe ARDS. Recommendations of the Veno-venous ECMO Expert Panel appointed in February 2016 by the national consultant on anesthesiology and intensive care

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Abstract
Extracorporeal membrane oxygenation (ECMO), which enables effective blood oxygenation and carbon dioxide removal for several weeks, has become a well established technique for the treatment of severe acute respiratory failure (V-V ECMO, veno-venous ECMO) or circulatory failure (veno-arterial ECMO). Veno-venous ECMO is a life-saving treatment in patients in whom severe acute respiratory distress syndrome (ARDS) makes mechanical ventilation unlikely to provide satisfactory blood oxygenation for preventing further vital organ damage and progression to death. The protocol below refers only to V-V ECMO therapy as a measure to support blood gas exchange by means of an extracorporeal circuit in adult patients with severe ARDS. Veno-venous ECMO does not provide treatment for acutely and severely diseased lungs but it enables the patient to survive the critical phase of severe ARDS until recovery of lung function. In addition to preventing death from hypoxemia, this technique can also prevent further progression of lung damage due to mechanical ventilation. Recent experience in ECMO therapy since the outbreak of an influenza A(H1N1) pandemic in 2009, along with technical progress and better understanding of the pathophysiology of ventilator-induced lung injury, have contributed to a significant improvement in ECMO treatment outcomes. Postulated factors related to an increased survival include wider use of ECMO during patient transfer and less intensive anticoagulation protocols. The aim of presenting this revised protocol was to improve ECMO treatment outcomes in patients with severe ARDS, to enhance ECMO accessibility for patients who might possibly benefit from this treatment, to reduce the time until institution of ECMO therapy, and to avoid ECMO therapy in futile cases. The authors believe that this protocol, based on recent papers and their own experience, can provide help and advice both for the centers which develop V-V ECMO program, and for doctors who will refer their patients for treatment in an ECMO center.

Key words: extracorporeal membrane oxygenation, ECMO, ARDS, acute respiratory failure
right atrium, or veno-arterial, when blood is drained from the inferior vena cava, superior vena cava or the right atrium and returned to large artery.

Veno-venous ECMO (V-V ECMO) is used in conditions associated with potentially reversible severe lung dysfunction to a degree that precludes effective gas exchange by mechanical ventilation, while veno-arterial ECMO (V-A ECMO) is used in case of potentially reversible or irreversible heart failure. In the latter situation, V-A ECMO may be used as a bridge to cardiac transplantation or long-term mechanical cardiac support. The present guidelines refer only to V-V ECMO as a method to replace lung function using an extracorporeal circuit in adult patients with severe respiratory failure.

ECMO does not treat lungs but only allows the patient to survive the period of severe lung dysfunction to a degree that precludes effective arterial blood oxygenation and/or carbon dioxide elimination by mechanical ventilation. In addition, V-V ECMO reduces or eliminates the risk of ventilation-related lung damage in patients with severe acute respiratory distress syndrome (ARDS). New-generation ECMO systems provide oxygen at a rate of up to 3 mL kg\(^{-1}\) min\(^{-1}\) and eliminate carbon dioxide at a rate of 3–6 mL kg\(^{-1}\) min\(^{-1}\), which may fully cover the metabolic needs of the patient [1].

Experience with the use of V-V ECMO for the treatment of severe respiratory failure in adults in the recent years, including largely increased use of this technique during an influenza A(H1N1) pandemic in 2009, along with better understanding of the pathophysiology of ventilator induced lung injury (VILI), contributed to a significant improvement in ECMO treatment outcomes [2]. Safe use of an extracorporeal circuit for a longer time has become possible with advances in ECMO technology including use of centrifugal pumps, polymethylpentene oxygenators, and heparin-coated cannulas. Wider use of ECMO during patient transfer and less intensive anticoagulation during this therapy also contributed to reduced complication rates and better treatment outcomes. ECMO is increasingly used in the treatment of life-threatening asthma, and as bridge therapy to lung transplantation [3]. The role of ECMO for the reduction of VILI was highlighted during the Berlin ARDS conference [4]. In these settings, a modification known as partial ECMO assistance is used, with blood flow reduced to values that allow ventilation in accordance with the criteria of lung protective strategy [3].

The goal of the present update of the guidelines on the use of V-V ECMO is to improve treatment outcomes in patients with severe respiratory failure, increase access to ECMO in those patients in whom this therapy is indicated, reduce the duration of mechanical ventilation that induces further lung damage, and define more clearly those patient groups in whom ECMO therapy does not increase the likelihood of recovery. The present guidelines have been developed based on a review of current literature and national experience with the use of V-V ECMO in expert units in Poland. Due to the fact that use of V-V ECMO for the treatment of respiratory failure is a relatively new area of intensive care, subject to further rapid advances, these guidelines should not be treated as definitive, particularly in centers having most expertise with the use of this method.

Clinical studies indicate that ECMO is a relatively safe therapeutic technique which improves outcomes in patients with severe ARDS but is also associated with a risk of serious complications. Thus, ECMO therapy must be undertaken in accordance with specific management principles and algorithms. In the recent years, a number of centers in Poland have been selected that offer adequate organization of care, staff, and equipment to provide ECMO in the treatment of severe respiratory failure. The list of these centers is included in the Appendix 1.

Observational studies suggest improved treatment outcomes if ECMO is initiated in a referring unit and used during patient transport to a specialized tertiary care unit [5]. It the tertiary care unit is equipped with a portable ECMO system, initiation of this therapy in the referring unit by a team from the tertiary care unit should be considered. For this reason, it is desirable to equip ECMO-capable tertiary care units with portable ECMO systems. The basis of appropriate patient selection for ECMO therapy is ineffectiveness of previous guideline-directed conventional therapy.

**OUTCOMES OF VENO-VENOUS ECMO**

Most studies published prior to 2009 indicated that survival in patients with severe and critical hypoxemia in ARDS was comparable in patients treated with ECMO or conventional lung ventilation techniques [6, 7]. The randomized CESAR study that included 180 patients showed somewhat better survival in severe ARDS in patients treated with ECMO compared to conventional treatment [8]. Studies indicate that particularly in severe ARDS complicating influenza, survival with ECMO treatment is higher compared to conventional mechanical lung ventilation. Multiple studies indicate that benefits of ECMO may also be seen in severe ARDS due to other causes but treatment outcomes may be worse compared to severe ARDS secondary to a viral infection.

ECMO is an invasive medical technology which is not free from severe and life-threatening complication, and thus this treatment should be initiated only if reliably defined criteria are met.

**TREATMENT OPTIMIZATION BEFORE PATIENT REFERRAL FOR ECMO**

1. Exclusion or elimination of potentially reversible causes of worsening of lung function and ventilation param-
eters, including pneumothorax, significant pleural effusion, bronchial obstruction with mucus or clots, pulmonary congestion, and increased extravascular lung water volume.

2. Ventilation according to the lung protective strategy: tidal volume ($V_t$) $\leq 6$ mL kg$^{-1}$ for predicted body weight using the ARDSNet calculator, aiming for $P_{\text{plateau}} < 30$ cm H$_2$O, permissive hypercapnia, if $P_{\text{plateau}} > 30$ (max. 35) cm H$_2$O. Recommended ventilation modes include pressure-controlled ventilation (PCV), bilevel positive airway pressure (BiPAP), and BiLevel [9].

3. Adequate sedation. If low volume ventilation is not tolerated and ventilator-patient synchronization is difficult, only in severe ARDS (PaO$_2$/FiO$_2$ < 120, where PaO$_2$ is partial arterial oxygen pressure and FiO$_2$ is oxygen concentration in inspired air) is possible to initiate neuromuscular blockade (the preferred drug is cisatracurium) with boluses/continuous intravenous infusion for up to 48 hours [10].

4. Titration of positive end-expiratory pressure (PEEP) to optimal PaO$_2$/FiO$_2$. Lung compliance with consideration of the effect on hemodynamic parameters in the range of 5–15–20 cm H$_2$O, optimally in the range defined by the derecruitment technique.

5. Performing lung recruitment maneuvers (LRM) every 4–8 hours and after each lung derecruitment. Derecruitment occurs already after 1.5–2.0 seconds following a loss of PEEP in the airways. i.e., after each mucus suctioning from the airways or short-term disconnection of the ventilator circuit. The potential of recruitment may be evaluated using electrical impedance tomography or lung ultrasonography. The following LRM techniques are possible:

   a) PEEP $30$ cm H$_2$O for 20 seconds, peak pressure up to 40 cm H$_2$O; (it is important to maintain respiratory drive at $< 15$ cm H$_2$O)

   b) performing two inspirations prolonged to 20 seconds, with the peak pressure of 40 cm H$_2$O — may be achieved by pressing the „respiratory pause“ key in some ventilator types. Blood pressure and oxygen saturation must be monitored during LRM.

   c) derecruitment technique — LRM combined with determination of optimal PEEP value:

   — BiPAP ventilation;

   — reduction of pressure support (PS) to 0 cm H$_2$O;

   — change of inspiratory positive airway pressure (IPAP) to achieve tidal volume of 6 mL kg$^{-1}$;

   — determination of the IPAP-EPAP (expiratory positive airway pressure) (or IPAP-PEEP) difference needed to maintain the tidal volume of 6 mL kg$^{-1}$;

   — increasing inspiratory pressure by 3–4 steps of 5–7 cm H$_2$O to IPAP of 40–45 cm H$_2$O, maintaining a constant IPAP-EPAP difference determined previously. Duration of ventilation at each pressure level during subsequent steps — 3–4 breathing cycles;

   — if the ventilator cannot change both IPAP and EPAP at the same time, EPAP is increased first, followed by IPAP (to avoid a temporary increase in the tidal volume above 6 mL kg$^{-1}$). The highest pressure level should be maintained for about 3 breathing cycles, with careful monitoring of hemodynamic parameters and reduction of inspiratory pressures in case of instability (hypotension, heart rate changes);

   — gradual reduction of inspiratory pressures while maintaining a constant IPAP-EPAP difference, every 3–4 breathing cycles in steps of 4–5 cm H$_2$O initially (down to EPAP of about 25 cm H$_2$O) followed by 2 cm H$_2$O, while monitoring the tidal volume;

   — if the ventilator cannot change both IPAP and EPAP at the same time, IPAP is decreased first followed by EPAP (to avoid a temporary increase in the tidal volume above 6 mL kg$^{-1}$);

   — tidal volume should be monitored during each step/at each pressure level;

   — during reduction of inspiratory pressures, the tidal volume increases (due to increasing lung compliance) or is kept at the same level until EPAP is reduced below the lower inflection point on the pressure-volume curve (the point when alveolar derecruitment begins);

   — optimal EPAP is 2 cm H$_2$O below the derecruitment point determined as above;

   — as reduction of airway pressure below the derecruitment point resulted in alveolar collapse, repeating the recruitment maneuver described above is needed to reopen the alveoli;

   — during the latter step, inspiratory pressure may be reduced at a faster rate.

   — when EPAP of 2 cm H$_2$O above the determined derecruitment point is achieved, IPAP is adjusted to achieve the tidal volume of about 6 mL kg$^{-1}$.

6. While determining PEEP, use of the stress index may be helpful. Determination of the latter is based on visual representation of the pressure-volume curve during volume controlled ventilation (VCV). Optimal PEEP is associated with a linear pressure-volume curve in its middle segment, also indicating normal elastance [11, 12].

7. Compliance of the respiratory system ($C_{RS}$) is related to the air-filled lung volume. Thus, optimal ventilation should be associated with the driving pressure ($\Delta P$) of $< 7$ cm H$_2$O, where $\Delta P = V_{VT}C_{RS}$ ( $C_{RS}$ — static compliance) ($\Delta P = $ pleural pressure [PPl] − PEEP). A strong correlation is found between the driving pressure and energytrauma and development of VILI [13].
8. Careful and frequent airway toilet (using bronchofiberscope if possible). LRM should be performed after each airway cleaning procedure.

9. Careful consideration of steroid therapy: methylprednisolone 0.5–2.5 mg kg\(^{-1}\) day\(^{-1}\) for 7 days is particularly indicated if lung histology (by transbronchial or fine-needle biopsy) shows evidence of fibrosis.

10. Use of optimal fluid therapy, defined as avoiding volume overload and maintaining a negative fluid balance if possible, using continuous renal replacement therapy techniques if needed. If extravascular lung water (EVLW) volume may be measured, attempts should be made to reduce it below 10 mL kg\(^{-1}\).

11. Aiming to optimize the cardiovascular status based on invasive hemodynamic monitoring in case of hemodynamic instability (techniques that allow EVLW monitoring by transpulmonary termodilution are preferred).

12. If ventilation in the prone position is associated with a significant improvement of the PaO\(_2\)/FIO\(_2\) ratio, the patient should be placed in the prone position for 6 to 8 hours at least twice during 24 hours. During ventilation in the prone position, recruitment maneuvers should be performed and optimal PEEP value should be determined by the derecruitment approach described above.

13. Evaluation of the effectiveness of advanced ventilation techniques (depending on the capabilities and experience of the unit) including pressure-limited ventilation, inverse ratio ventilation (IRV), airway pressure release ventilation (APRV), oscillatory ventilation, independent lung ventilation, and determination of optimal PEEP by esophageal pressure measurement.

14. If available, an attempt may be made to optimize oxygenation using inhaled nitric oxide (NO) or nebulized prostacyclin analog (iloprost). One management algorithm is to administer NO at 10 ppm for 30 minutes. This therapy should be continued if it is associated with a significant increase in PaO\(_2\), and if not — administration of NO should be discontinued [14].

15. Use of techniques to reduce ventilator-associated pneumonia (VAP) including rational antibiotic therapy, avoiding reintubation, tracheal intubation and insertion of an orogastric tube, patient positioning at 30–45\(^{\circ}\), subglottic suctioning, monitoring sedation intensity (e.g., using Richmond Agitation-Sedation Scale or Riker Agitation-Sedation Scale) [15–17], initiation of enteral feeding within 72 hours of treatment, blood glucose level monitoring, and prevention of peptic ulcer disease and deep venous thrombosis.

16. It should be clearly stressed that the selected mode of ventilation should also be used during patient transport for diagnostics and surgical procedures, and to other units. This is possible using an appropriate quality transport ventilator with adequate oxygen supply.

**Before transporting the patient, it is reasonable to ventilate the patient in the intensive care unit with the transport ventilator and evaluate the effectiveness of ventilation.**

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**MONITORING AND LABORATORY TESTS BEFORE PATIENT REFERRAL FOR ECMO**

**Basic:**
- pulse oximetry,
- arterial blood gases — not less than every 3 hours,
- direct blood pressure measurement,
- central venous pressure measurement,
- renal function parameters,
- myocardial necrosis biomarkers,
- ventilation parameters including tidal volume, breathing rate, FIO\(_2\), peak inspiratory pressure (PIP), static lung compliance, and PEEP — recorded not less than every hour,
- chest X-ray,
- pleural ultrasound,
- chest computed tomography should be considered the preferred modality for imaging the lungs,
- blood lactate level.

**Additional (if available in a given unit):**
- echocardiography,
- if influenza A(H\(_1\)N\(_1\)) virus infection is suspected, it is indicated to confirm this etiology by polymerase chain reaction (PCR).

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**MEDICAL INDICATIONS FOR TREATING ACUTE RESPIRATORY FAILURE WITH ECMO**

The indication for V-V ECMO is respiratory failure with hypoxemia and hypercapnia persisting despite use of large oxygen concentrations, advanced ventilator techniques and optimization of the patient’s condition, which is associated with a risk of further worsening of the patient’s condition, ultimately leading to death.

When evaluating the need for patient referral for ECMO, changes in gas exchange parameters following use of advanced ventilator techniques and optimization of the patient’s condition, which is associated with a risk of further worsening of the patient’s condition, ultimately leading to death.

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**Basic criterion:**
- Severe ARDS by the Berlin criteria [4] and at least one of the following criteria:
  - PaO\(_2\)/FIO\(_2\) < 80 for ≥ 3 hours despite V\(_T\) of 6 mL kg\(^{-1}\) and PEEP of ≥ 5 cm H\(_2\)O and performance of LRM as described above;
  - pH < 7.25 for ≥ 3 hours.
Additional criteria:
- pH < 7.2, PaCO₂ > 80 mm Hg,
- static lung compliance < 0.5 mL cm H₂O⁻¹,
- PIP > 40 cm H₂O with Vₕ ≤ 6 mL kg⁻¹,
- oxygenation index OI = (MAP × FIO₂ × 100)/PaO₂ > 60 mm Hg for 30 minutes or > 35 mm Hg for 6 hours (MAP — mean airway pressure);
- chest X-ray showing extensive consolidations in at least two lung quadrants or alternatively:
  - Murray lung injury score (LIS, Lung Injury Score) [18] > 3.0;
  - severity of the patient’s condition should be evaluated twice daily using the Sequential Organ Failure Assessment (SOFA) score. However, the SOFA score does not serve as an inclusion or exclusion criterion for ECMO.

It should be noted that the simplified definition of ARDS (Kigali modification) is used for screening purposes only and should not be used for patient selection for ECMO [19].

**PARAMETERS OF MAJOR IMPORTANCE FOR PREDICTING THE NEED FOR ECMO**

It has been found that outcomes of ECMO are best when this therapy is initiated as soon as possible after respiratory failure develops. Thus, early identification of patients likely to need ECMO is of major importance, so as to refer these patients promptly to the nearest center offering ECMO therapy [20].

**OXYGENATION INDEX**

Compared to PaO₂/FiO₂, a better prognostic parameter indicating the need for ECMO may be the oxygenation index (OI) (see additional criteria above), defined as the product of the reciprocal of PaO₂/FiO₂ and the mean airway pressure (MAP) in cm H₂O [20].

\[
OI = \frac{(\text{FiO}_2 \times \text{MAP} \times 100)}{\text{PaO}_2}
\]

However, the available data are still inadequate to define specific OI index that would serve as a criterion to initiate ECMO.

**ARDS INDEX**

It has been postulated that an important parameter predicting the need for ECMO may also be the ARDS index after 12 hours of ventilation [20].

\[
\text{ARDS index} = \text{OI} \times \left(\frac{\text{MV} \times \text{PaCO}_2}{40}\right)
\]

\[
\text{ARDS index} = \left(\frac{(\text{FiO}_2 \times \text{MAP} \times \text{MV} \times \text{PaCO}_2)}{\text{PaO}_2}\right) \times 40
\]

ARDS index above 300 after 12 hours of ventilation is a significant parameter indicating the need for ECMO [20].

**CHANGE IN PaO₂/FiO₂**

It has been observed that a reduction in PaO₂/FiO₂ by more than 100 during the first 12 hours of ventilation indicates a significant likelihood of the need for ECMO [20].

**ADDITIONAL REMARKS**

In patients with severe lung dysfunction that may soon fulfill the criteria for V-V ECMO, gastric contents aspiration as the cause of impaired gas exchange, body mass index (BMI) > 30 kg m⁻² and conditions associated with immunosuppression decrease the likelihood of the need for life-saving therapy with ECMO [20].

**CONTRAINDICATIONS TO VENO-VENOUS ECMO**

Absolute contraindications:
- previous mechanical ventilation with high peak airway pressure or high oxygen concentration in inhaled air for more than 7 days [5, 20];
- severe systemic disease with unfavorable prognosis, regardless of the effectiveness of therapy for ARDS;
- severe irreversible central nervous system damage, encephalopathy [5];
- cirrhosis with ascites, history of esophageal variceal bleeding [5];
- malignancy associated with unfavorable prognosis [5];
- chronic respiratory disease associated with unfavorable prognosis;
- intracranial bleeding and other absolute contraindications for anticoagulation;
- severe chronic pulmonary hypertension (mean pulmonary arterial pressure [mPAP] > 50 mm Hg) [5];
- severe left ventricular failure (left ventricular ejection fraction [LVEF] < 25%) or right ventricular failure before the occurrence of hypoxemia [5];
- informed patient refusal of ECMO therapy.

Relative contraindications:
- Age > 70 years [20],
- AIDS [5],
- body mass >150 kg [20],
- other factors associated with a significantly reduced likelihood of survival.

**MANAGEMENT TEAM**

Therapy with ECMO involves close cooperation between anesthesiology and intensive care specialists, cardiac surgeons, intensive care unit nursing personnel, and perfusionists. This cooperation should be based on the following principles:

1. An anesthesiology and intensive care specialist familiar with the principles and the management of ECMO therapy is present in the unit for 24 hours a day.
2. If possible, a surgeon familiar with the principles and the management of ECMO therapy, including device operation, is at standby for 24 hours (e.g., a physician on duty in a cardiac surgery unit).
3. Direct patient care is provided by intensive care unit nursing personnel familiar with the principles and the
management of ECMO therapy, including device operation.
4. A perfusionist may be helpful to monitor therapy by checking up the system at least once daily.

**ORGANIZATIONAL ISSUES AND ACTIVITIES ASSOCIATED WITH PATIENT REFERRAL FOR ECMO**

After confirming indications and excluding contraindications, the physician referring a patient for ECMO therapy should undertake the following actions after a consultation with a physician in charge of the unit:
1. If possible, explain the nature of and the risk associated with ECMO to the patient or his/her relative, and obtain the patient’s consent for this therapy.
2. Liaise with the physician on duty in the nearest or organizationally appropriate unit providing ECMO therapy to verify indications and exclude contraindications, and jointly discuss the appropriateness of therapy in case of doubts. The referring physician is obliged to verify accordance of the previous therapy with the above guidelines, particularly in regard to appropriate ventilator therapy.
3. After the patient is selected for ECMO therapy, it is necessary to establish whether the nearest unit providing ECMO is able to accept the patient and whether an ECMO system is available for the patient in question.
4. If an appropriate hospital bed or an ECMO system is not available, a next nearest unit providing ECMO therapy should be contacted and the points 2 and 3 of the procedure should be repeated.
5. If a destination place for further therapy is agreed, patient transport should be arranged using an appropriate quality transport ventilator. The referring unit is responsible for the organization of patient transport. If a portable ECMO system is available in the accepting unit, it is preferable that a team from the accepting unit comes to the referring unit, initiates V–V ECMO, and transports the patient to the accepting unit [20, 21]. However, this approach depends on the organizational and staffing capabilities of the unit providing ECMO therapy and cannot be considered obligatory at the present stage.
6. The referring unit is obliged to accept the patient for further treatment after ECMO therapy is discontinued and the patient’s cardiovascular status is stabilized to a degree that allows safe transport.

**SETTING UP VENO-VENOUS ECMO**
1. Adequate anticoagulation should be instituted before cannulation. The most commonly used approach is to administer unfractionated heparin 5000 units or 100 units kg⁻¹.
2. The device inflow cannula is most commonly inserted to the vena cava inferior through the femoral vein, and the return (outflow) cannula is inserted to the vena cava superior through the internal jugular vein or to the vena cava inferior through the femoral vein. The optimal level for the device inflow cannula insertion may be T10-T11, as deeper insertion may interfere with venous return from the hepatic veins.
3. If inflow through a cannula inserted through the femoral vein is insufficient, an additional cannula may be inserted to the vena cava superior through the internal jugular vein, or to the contralateral femoral vein, depending on the location of the return cannula, and both inflow cannulas are connected with a Y-connector.
4. An alternative approach is insertion of a dedicated ECMO cannula through the inferior jugular vein which allows venous inflow to the ECMO device from the inferior and superior venae cavae and blood return to the right atrium at the level of the tricuspid valve.
5. When setting up the extracorporeal circuit, particular care should be paid to careful air removal from the system.
6. Cannulas should be inserted and connected to the extracorporeal ECMO circuit by a cardiac surgeon, vascular surgeon, or adequately trained intensive care unit physician. Aseptic conditions should be maintained and cannulas should be protected from inadvertent disconnection or sliding out.

**MANAGEMENT DURING ECMO THERAPY**
1. Normal arterial gas partial pressures are obtained mostly by gas exchange in the ECMO oxygenator. Ventilation is only an additional measure and should be conducted in a way that allows as rapid lung regeneration as possible. Moderate or high PEEP values should be maintained (10–18 cm H₂O) with low ventilation rate of 12–14 min⁻¹ and maintenance of a pressure-limited ventilation mode [22].
2. Veno-venous ECMO is the major treatment approach in acute respiratory failure with hypoxemia that cannot be corrected using advanced ventilation techniques. If significant circulatory failure coexists, it is mostly due to hypoxemia, and thus cardiovascular function usually improves rapidly once normal blood oxygenation is restored.
3. In special cases, V-A ECMO should be considered (in patients with heart failure due to myocarditis or pericarditis, previous severe cardiovascular disease, or significantly elevated cardiac biomarkers) if cardiac support with large catecholamine doses is needed. Cases of concomitant ARDS and acute heart failure due to viral myocarditis have been reported, e.g. in influenza A(H1N1) infection.
4. Fresh gas flow rates should be adjusted to PaCO₂, with flow increase in case of hypercapnia, and flow reduction if PaCO₂ is too low.

5. Blood flow rate generated by the centrifugal ECMO pump is adjusted to achieve optimal arterial blood oxygenation (desired PaO₂ 100–150 mm Hg). Initial flow rate is usually set at 3–5 L min⁻¹. Values recommended by the Extracorporeal Life Support Organization (ELSO) are 60–80 mL min⁻¹ [23]. Increasing blood flow through the oxygenator increases PaO₂, and decreasing blood flow reduces PaO₂. Except in patients with ARDS secondary to sepsis, the ratio of ECMO blood flow to cardiac output above 60% allows adequate arterial blood oxygenation (SaO₂ > 90%) [24]. If oxygenation cannot be improved to that level (see Table 1), many centers believe that maintaining SaO₂ of about 80% is safe [25]. Observational studies indicate, however, that persistently reduced PaO₂ correlates with cognitive dysfunction [25].

6. If achieving minimal oxygenation (PaO₂ 70 mm Hg) is not possible due to inadequate venous inflow, correcting the position of the cannula or insertion of an additional inflow cannula should be considered.

7. During ECMO therapy, lung protective ventilation should be used, with tidal volume < 6 mL kg⁻¹, peak inspiratory pressure (PIP) 20–25 cm H₂O, breathing rate 10/min⁻¹, PEEP 10–15 cm H₂O, and FiO₂ 0.3, or ventilator weaning should be aimed for.

8. Use of heparin- or phospholipid-coated systems is beneficial [6].

9. According to the ELSO recommendations, anticoagulation with continuous infusion of unfractionated heparin adjusted to increase the activated clotting time (ACT) to 180–220 seconds or the activated partial thromboplastin time (APTT) to 40–55 seconds is necessary to prevent blood clotting in the extracorporeal circuit during ECMO therapy [23]. In the recent years, a trend to reduce the intensity of anticoagulation has been observed (target ACT 160–200 seconds). Enoxaparin 40 mg daily was also reported to be used to prevent clotting. In an observational study that included 61 patients treated for the overall treatment duration of 56 days, thrombotic complications were observed in 4 patients (3 cases of ECMO circuit failure due to circuit clotting, 1 case of myocardial infarction), and bleeding complications occurred in only 18% of patients [26]. Arguments for reducing the intensity of anticoagulation in patients treated with V-V ECMO seem reasonable but no sufficient evidence from comparative studies are yet available to recommend reduction of anticoagulation to prophylactic low-molecular-weight heparin doses.

10. During ECMO therapy, platelet count should be maintained above 100 G L⁻¹.

11. If possible, invasive procedures should be avoided during ECMO therapy to limit the risk of bleeding complications. Tracheotomy should be considered before initiation of ECMO therapy, or the intensity of anticoagulation should be reduced at the time of tracheotomy [6, 27].

12. Hemoglobin level should be kept above 10–12 g dL⁻¹, depending on the patient’s condition. In practice, this may mean that at least 2–3 units of packed red blood cells may need to be transfused daily [28]. This is also associated with a need for transfusion of fresh frozen plasma. According to the ELSO recommendations, maintaining SaO₂ not lower than 80% with hematocrit of 40% allows adequate oxygen transport [29].

13. If evidence of volume overload is present before or during ECMO therapy, which is often the case, reduction of volume overload should be aimed for but attention should also be paid to maintain appropriate tissue perfusion.

14. Adequate sedation should be continued during the first days of therapy. During subsequent days, attempts should be made to reduce sedation to allow pressure support ventilation (PSV). If allowed by the general condition of the patient, weaning of ventilator therapy, removal of the endotracheal tube and respiratory rehabilitation may be beneficial.

15. All attempts should be made to prevent hypothermia resulting from heat loss by the extracorporeal circuit canulas.

16. Acute kidney injury is a frequent complication in patients with severe respiratory failure who require therapy with V-V ECMO. The Acute Kidney Injury Network (AKIN) II or III criteria are met in 25–67% of patients treated with ECMO, and 21–24% of patients require renal replacement therapy [30]. Available studies do not include a recommendation regarding using the ECMO circuit for renal replacement therapy or performing the latter using a dialysis catheter. The decision regarding the use of the ECMO circuit for renal replacement therapy or using a dialysis catheter should be made based on the experience of the treating unit, characteristics of the renal replacement therapy system in regard to threshold inflow and outflow pressure, and the possibility of inserting a dialysis catheter.

17. The mode of connecting the renal replacement therapy system to the ECMO circuit depends on the relation between pressures in various parts of the ECMO circuit (proximally to the centrifugal pump, between the pump and the oxygenator, and between the oxygenator and the return cannula) and alarm pressures in the inflow and outflow lines in the renal replacement therapy system. In practice, it may be optimal to connect the inflow and outflow lines of the renal replacement therapy system between the centrifugal pump and the oxygenator, or the inflow line between the oxygenator and the return cannula.
cannula and the outflow line between the centrifugal pump and the oxygenator (despite renal replacement therapy system-ECMO circuit recirculation) [31].

**MONITORING DURING ECMO THERAPY**

Basic monitoring:
- pulse oximetry,
- arterial blood gases — not less than every 3 hours,
- direct blood pressure measurement,
- measurement of central venous pressure has a minor role, as interpretation of this parameter is limited by active blood suction by the centrifugal pump,
- renal function parameters,
- ventilation parameters including tidal volume, breathing rate, FiO2, PIP, static lung compliance, and PEEP — recorded not less than twice daily,
- blood lactate level,
- ACT or APTT — not less than every 6 hours,
- international normalized ratio (INR), prothrombin time (PTT), D-dimer, fibrinogen, and antithrombin levels, platelet count — daily,
- chest X-ray — not less than every 3 days,
- device function parameters should be recorded every hour, including blood flow rate and pump speed (revolutions per minute). Decreasing blood oxygenation with constant blood flow indicates device “wear” or an increased clotting risk. In such cases, prepare for replacement of the oxygenator or the whole extracorporeal circuit. PaO2/FiO2 < 200 in a blood sample taken distally to the oxygenator indicates its “wear” and suggests a possible need for prompt device replacement [26]. A proposed nursing observation chart for ECMO therapy is provided in the Appendix 2.

Additional monitoring (if available in the unit):
- transesophageal echocardiography may be helpful to evaluate valvular function and position of the cannula, particularly with the use of a dual-lumen cannula,
- evaluation of extravascular lung water volume using the Pulse Contour Cardiac Output (PICCO) methods. These parameters must be interpreted with caution due to additional variable blood inflow from the ECMO system to the vena cava superior,
- computed tomography of the chest, abdomen, and head depending on clinical indications,
- bedside lung impedance tomography if available.

**TECHNICAL PROBLEMS AND COMPLICATIONS ASSOCIATED WITH ECMO THERAPY**

Patient-related complications:
- bleeding (prevalence approx. 30%, including cardiac tamponade, hemothorax, gastrointestinal bleeding, bronchial bleeding, central nervous system bleeding, genitourinary bleeding [32]), vessel damage related to cannulation,
- hemolysis,
- thrombocytopenia/heparin induced thrombocytopenia (HIT),
- infection (including respiratory infections, catheter-related infections, and infections associated with the ECMO cannula ECMO [32]),
- embolic complications,
- neurological complications and long-term cognitive dysfunction,
- organ failure (kidneys, heart, liver),
- lung barotrauma,
- metabolic disturbances,
- right atrial perforation with a cannula.

Complications related to the ECMO device and circuit (prevalence approx. 5% [3]):
- consequences of incomplete air removal from the circuit,
- air suction to the extracorporeal circuit during therapy,
- cannula dyslocation or sliding out,
- oxygenator dysfunction (wear, thrombosis),
- circuit interruption (disconnection, damage),
- heater-cooler dysfunction,
- pump dysfunction.

Management in case of selected technical problems

Regardless of the mechanism, each device dysfunction leading to reduced blood flow or impaired gas exchange is an immediately life-threatening condition and requires prompt modification of ventilator parameters to allow adequate gas exchange in the lungs. At the same time, a solution of the technical problem should be sought as soon as possible. Most common technical problems and device malfunctions and their suggested management are shown in Table 1.

**DISCONTINUATION OF ECMO THERAPY**

- Extensive cerebral ischemia.
- Massive intracranial bleeding.
- On-treatment diagnosis of another progressive condition precluding lung function recovery.
- No recovery of lung function despite prolonged therapy.
- Brain death declared during ECMO therapy.

**WEANING FROM ECMO THERAPY**

Prerequisites:
- resolving lung abnormalities on chest X-ray,
- improvement of arterial blood oxygenation at FiO2 < 0.6 and improvement of lung compliance (PIP < 30 cm H₂O) during an attempt to reduce ECMO support.
ATTEMPT TO REDUCE ECMO SUPPORT TO EVALUATE THE POSSIBILITY OF WEANING FROM ECMO THERAPY

1. Reduction of V-V ECMO support involves termination of fresh gas supply to the oxygenator, usually for at least 2 hours. Reduction of blood flow rate is associated with a risk of thrombosis in the ECMO circuit and is not justified when evaluating respiratory function during weaning from V-V ECMO [26].
2. Maintenance of normal gas exchange parameters during reduction of V-V ECMO support indicates that ECMO therapy may be safely discontinued. In all cases, this decision should be made with adequate caution, taking into account the whole clinical picture.
3. Cannulas may be removed by a cardiac surgeon, vascular surgeon, or an intensive care unit physician. Cannula removal by the latter is considered safe provided that the procedure is adequately prepared and a surgeon is available to intervene in case of complications [33].

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References:


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e-mail: rlango@gumed.edu.pl
Appendix 1
Centers providing ECMO therapy for the management of severe respiratory failure in Poland as of February 2017

<table>
<thead>
<tr>
<th>Hospital</th>
<th>Unit</th>
<th>Address</th>
<th>Contact phone number (prefix +48)</th>
</tr>
</thead>
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<tr>
<td>Śląskie Centrum Chorób Serca w Zabrzu</td>
<td>Oddział Kliniczny Kardioanestezji i Intensywnej Terapii</td>
<td>Curie-Skłodowskiej 9, 41–800 Zabrze</td>
<td>32 271 27 31, 32 373 38 00</td>
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<tr>
<td>Krakowski Szpital Specjalistyczny im. Jana Pawła II</td>
<td>Oddział Anestezjologii i Intensywnej Terapii</td>
<td>Prądnicka 80, 31–202 Kraków</td>
<td>12 614 33 03</td>
</tr>
<tr>
<td>Szpital Uniwersytecki w Krakowie</td>
<td>Oddział Anestezjologii i Intensywnej Terapii</td>
<td>Skawierska 8, 31–066 Kraków</td>
<td>12 430 53 82</td>
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<td>Uniwersytecki Szpital Kliniczny we Wrocławiu</td>
<td>Klinika Anestezjologii i Intensywnej Terapii</td>
<td>Borowska 213, 50–556 Wrocław</td>
<td>71 733 23 10, 71 733 23 20, 71 733 23 11</td>
</tr>
<tr>
<td>Szpital Uniwersytecki nr 1 w Bydgoszczy</td>
<td>Klinika Anestezjologii i Intensywnej Terapii</td>
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<td>Klinika Anestezjologii i Intensywnej Terapii</td>
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<td>Samodzielny Publiczny Szpital Kliniczny nr 2</td>
<td>Klinika Anestezjologii, Intensywnej Terapii i Ostrzych Zatrąć</td>
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<td>Uniwersytecki Szpital Kliniczny w Białymstoku</td>
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<td>Szpital Wojewódzki w Bielsku–Białej</td>
<td>Oddział Anestezjologii i Intensywnej Terapii</td>
<td>Aleja Armii Krajowej 101, 43–300 Bielsko Biała</td>
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<td>Szpital Kliniczny Przemienienia Paraskiego</td>
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<td>Kliniczny Szpital Wojewódzki nr 2 im. Św. Jadwigi</td>
<td>Klinika Intensywnej Terapii i Anestezjologii z Ośrodkiem Ostrzych Zatrąć</td>
<td>Lwowska 60, 35–301 Rzeszów</td>
<td>17 866 48 59, 17 866 48 60</td>
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Centers providing ECMO therapy in pediatric patients

| Uniwersytecki Szpital Dziecięcy             | Oddział Anestezjologii i Intensywnej Terapii | Wielicka 265, 30–663 Kraków      | 12 658 20 11 ext. 15 22, 15 25   |
| Szpital Kliniczny im. Karola Jonschera     | Klinika Anestezjologii i Intensywnej Terapii Pediatrycznej | Szpitalna 27/33, 60–572 Poznań | 61 849 14 78, 61 849 14 38, 61 849 14 89 |
| Szpital–Pomnik Centrum Zdrowia Dziecka     | Klinika Anestezjologii i Intensywnej Terapii | Aleja Dzieci Polskich 20, 04–730 Warszawa | 22 815 13 35, 22 815 13 32        |
| Uniwersytecki Szpital Kliniczny we Wrocławiu | Oddział Anestezjologii i Intensywnej Terapii Dziecięcej | Borowska 213, 50–556 Wrocław | 71 733 23 10, 71 733 23 20        |
Appendix 2

**Nursing observation chart**

Date: ........................................................................................................

Patient name: ...............................................................................................

PESEL number: ..........................................................................................

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<tr>
<th>Hour (24 hour clock)</th>
<th>ECMO blood flow [L min⁻¹]</th>
<th>ECMO pump speed (revolutions per minute)</th>
<th>SpO₂ (pulse oximetry)</th>
<th>ECMO gas supply flow [L min⁻¹]</th>
<th>Oxygen level in ECMO gas supply</th>
<th>ACT or APTT</th>
<th>Remarks/problems/alarm type</th>
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