An extracorporeal carbon dioxide removal (ECCO$_2$R) device operating at hemodialysis blood flow rates

R. Garrett Jeffries$^{1,2}$, Laura Lund$^3$, Brian Frankowski$^2$ and William J. Federspiel$^{1,2,4,5,*}$

Abstract

Background: Extracorporeal carbon dioxide removal (ECCO$_2$R) systems have gained clinical appeal as supplemental therapy in the treatment of acute and chronic respiratory injuries with low tidal volume or non-invasive ventilation. We have developed an ultra-low-flow ECCO$_2$R device (ULFED) capable of operating at blood flows comparable to renal hemodialysis (250 mL/min). Comparable operating conditions allow use of minimally invasive dialysis cannulation strategies with potential for direct integration to existing dialysis circuitry.

Methods: A carbon dioxide (CO$_2$) removal device was fabricated with rotating impellers inside an annular hollow fiber membrane bundle to disrupt blood flow patterns and enhance gas exchange. In vitro gas exchange and hemolysis testing was conducted at hemodialysis blood flows (250 mL/min).

Results: In vitro carbon dioxide removal rates up to 75 mL/min were achieved in blood at normocapnia (pCO$_2$ = 45 mmHg). In vitro hemolysis (including cannula and blood pump) was comparable to a Medtronic Minimax oxygenator control loop using a time-of-therapy normalized index of hemolysis (0.19 ± 0.04 g/100 min versus 0.12 ± 0.01 g/100 min, $p = 0.169$).

Conclusions: In vitro performance suggests a new ultra-low-flow extracorporeal CO$_2$ removal device could be utilized for safe and effective CO$_2$ removal at hemodialysis flow rates using simplified and minimally invasive connection strategies.

Keywords: Extracorporeal carbon dioxide removal, CO$_2$ removal, Artificial lung, Gas exchange, Chronic obstructive pulmonary disease, Acute respiratory distress syndrome

Background

Mechanical ventilation has long been the standard of care for severe lung failure. A major and paradoxical complication of mechanical ventilation is direct trauma to already ailing lungs caused by over-distention and damage of alveolar tissue due to excessive positive pressures or volumes in the lung [1]. Extracorporeal membrane oxygenation, or ECMO, has more recently become recognized as a last resort option for severe lung failure when mechanical ventilation is failing or is not an alternative. Unlike mechanical ventilation, ECMO performs the function of blood oxygenation and carbon dioxide (CO$_2$) removal independently of the lungs, allowing injured tissue to rest and heal [2]. ECMO is associated, however, with a higher risk of severe complications compared...
to mechanical ventilation because it requires full circulatory diversion of venous blood to
achieve its intended function [3].

The primary complication risks of ECMO are associated with cannulation, exposure of
blood to foreign materials, the concomitant requirement for systemic anticoagulation, and
the stresses induced by mechanical pumping [4]. The degree of risk associated with these
factors correlates with the extracorporeal blood flow rate necessary for treatment [4–6].
To provide full extracorporeal oxygenation of venous blood requires circuit flows up to
the full cardiac output (4000–7000 mL/min) [5, 7]. In contrast, full metabolic CO₂ re-
moval can be achieved at much lower extracorporeal blood flows. CO₂ is predominantly
carried in the form of highly soluble bicarbonate ion that rapidly restores depleting CO₂
as it is eliminated, and the CO₂ dissociation curve is essentially linear and does not satu-
rate like the oxyhemoglobin dissociation curve [8–10]. These differences also provide the
opportunity to augment CO₂ removal efficiency with gas exchanger design features aimed
at reducing the thickness of the diffusive boundary layer at the gas exchange surface,
where gas transport through blood is limited to diffusion [11].

The degree of risk associated with extracorporeal lung support is reduced when lower
blood flows are needed to provide clinically meaningful benefit [5]. The ability to effi-
ciently remove CO₂ at lower blood flows has motivated use of extracorporeal CO₂
removal, or ECCO₂R, as an alternative or supplement to mechanical ventilation. The
two primary clinical indications where this objective is feasible are acute exacerbations
of chronic obstructive pulmonary disease (ae-COPD) and moderate to severe ARDS,
where lung protective ventilation strategies are necessary but are unable to maintain
safe levels of CO₂ removal [12, 13]. ECCO₂R was shown to reduce intubation rates in
ae-COPD patients failing less-invasive ventilation and assisted in weaning from ventila-
tion [14–19]. Hypercapnia was also managed in moderate ARDS patients using
ECCO₂R to facilitate more protective ventilation strategies by enabling reduction of
tidal volumes to ≤ 4 mL/kg without complications [20, 21]. Associated risk remains the
primary obstacle of ECCO₂R adoption in these indications however. Currently
approved ECCO₂R systems can operate at blood flows around 500 mL/min, but still
require cannula with size greater than 15 Fr [4]. The ability to provide the same levels
of CO₂ removal at even lower flows will enable the use of smaller catheters that are
similar in size to commonly used dialysis catheters that are 9–14 Fr.

We are developing a next-generation ECCO₂R device that operates at lower
blood flows (250 mL/min) with minimal surface area and clinically significant CO₂
removal rates. CO₂ removal of 70–160 mL/min has been shown to benefit patients
with hypercapnia, which translates to a target CO₂ removal rate of ≥ 25–35% meta-
bolic CO₂ production (~200–250 mL/min) at normocapnia (pCO₂ = 45 mmHg)
[22, 23]. The device also should maintain an optimal degree of fluid washing
around the hollow fiber membranes to eliminate regions of stagnation but without
causing unacceptable levels of blood cell trauma. We have adapted technology from
our intravenous respiratory assist catheter to accomplish these objectives [24, 25]. Using
an array of rotating impellers within an annular hollow fiber membrane bundle,
high fluid velocities, and improved blood flow distribution enhances gas exchange.
This paper reports on the design and bench testing of an ultra-low-flow ECCO₂R
device (ULFED) utilizing the rotating impeller concept. In vitro gas exchange and
hemolysis were evaluated.
Methods

Device description

The ultra-low-flow ECCO₂R device (ULFED) (Fig. 1) contains six rotating impellers fixed on a rigid stainless steel driveshaft (3/16 in. diameter [4.76 mm]). A stainless steel safety coil surrounding impellers (diameter 14.4 mm) protects a 30-cm-long polypropylene (PP) fiber bundle (300 μm diameter, x30-240; Membrana Celgard, Wuppertal, Germany) with total surface area 0.42 m². The bundle/impeller assembly is housed in cylindrical acrylic tubing (inner diameter 1.375 in. [34.9 mm]) with 1/4 in [6.35 mm] inflow/outflow ports for a total priming volume of 240 mL. Impellers (Fig. 2) were fabricated from a hydrophobic epoxy resin (Watershed XC11122; DSM Somos, Sittard) using stereolithography (SLA). Impellers measured 4 mm in length with a maximum outer diameter of 11.7 mm. Impellers are designed to only generate flow radially in/out of the surrounding fiber bundle to avoid perturbing circuit blood flow rates.

The impeller drive shaft extends out of the blood pathway and is sealed (400054; SKF, Gothenburg, Sweden) and supported by bearings (Ceramic R3; Ortech, Inc., Sacramento, CA). An external DC brushless servomotor (4490 H 048B; MicroMo Electronics, Inc., Clearwater, FL) drives shaft rotation. Saline is continuously infused along the shaft at 30 mL/h to lubricate and protect the seal and bearing from blood backflow up the drive-shaft. Heparin was added to the saline infusion (20 U/mL) to maintain anticoagulation levels in the blood for consistency across test circuits. The shaft is supported distally using a custom pivot bearing (ceramic pin (MSC Industrial Supply, Melville, NY) nested in an ultra-high molecular weight polyethylene cup (UHMWPE; Orthoplastics, Lancashire, UK)) shown in Fig. 2.

Gas exchange

CO₂ removal performance of the ULFED prototype was evaluated in a single-pass flow loop (Fig. 3) at a hemodialysis blood flow rate of 250 mL/min. The evaluations followed ISO 7199:2009 standards for gas exchange testing in blood oxygenators [26]. Filtered and heparinized bovine blood (20 U/mL) was collected fresh from the slaughterhouse the day of testing. The fluid circuit consisted of a centrifugal blood pump (BPX-80; Medtronic, Minneapolis, MN), a commercial oxygenator (Affinity NT; Medtronic, Minneapolis, MN), two blood reservoirs connected in parallel, and the ULFED. Blood continuously recirculated at 4500–5500 mL/min while gas tensions were balanced by the commercial
oxygenator to normocapnic venous conditions (pCO₂ = 45 ± 5 mmHg) using a N₂/CO₂/O₂ gas mixture. Blood temperature was maintained at 37 ± 1 °C with a heat exchanger integrated into the commercial oxygenator. Flow recirculated only to and from the primary reservoir during balancing, while secondary reservoir tubing remained clamped. Gas levels in the recirculating loop were monitored with a blood gas analyzer (RapidPoint 405; Siemens, Erlangen, Germany) until venous conditions were reached. The loop was then converted to single-pass mode for data collection by diverting flow to the secondary reservoir and clamping the bypass tubing in parallel to the ULFED. Measurements were collected at rotation speeds from 0 to 5000 RPM once all measured parameters remained stable for ≥ 2 min. The order of tested rotation speeds was randomized, and a minimum of two measurements were collected at each rotation speed. Blood
flow rate was continuously monitored with an ultrasonic flow probe (Transonic Systems, Ithaca, NY).

Pure O₂ sweep gas was pulled through fibers counter-current to blood flow at 8.0 L/min by a sealed vacuum pump (N811 KV.45P; KNF Neuberger, Trenton, NJ) and was regulated with a thermal mass flow controller (GR-116-1-A-PV-O₂; Fathom Technologies, Georgetown, TX). The fraction of CO₂ in outlet sweep gas \( F_{CO_2} \) was measured by a gaseous CO₂ analyzer (WMA-4; PP Systems, Amesbury, MA) and was used to calculate total CO₂ removal \( V_{CO_2} \) together with the STP-corrected sweep gas flow rate \( Q_{OUT}^{STP} \) according to Eq. 1.

\[ V_{CO_2} = Q_{OUT}^{STP} F_{CO_2} \quad (1) \]

\( V_{CO_2} \) was normalized to our target inlet pCO₂ of 45 mmHg to reduce variability in measurements associated with small fluctuations in gas inlet conditions (± 5 mmHg) according to Eq. 2.

\[ V'_{CO_2} = V_{CO_2} \times \frac{45 \text{ mmHg}}{pCO_2^{INLET}} \quad (2) \]

\( pCO_2^{INLET} \) was measured from a fresh blood sample immediately prior to each data point. Three identical ULFED prototypes were fabricated for repeatability testing, but gas pathway failure in one device limited gas exchange testing to two devices. Total ULFED gas exchange is reported as average and standard deviation of the CO₂ removal rates of both prototypes at each rotation speed.

**In vitro hemolysis testing**

Filtered and heparinized bovine blood (20 U/mL) was collected fresh from the slaughterhouse the day of testing per ASTM standards (F1841–97) [27]. The gas exchange loop was modified for hemolysis testing by removing the bypass tubing parallel to the ULFED, the commercial oxygenator, the secondary reservoir, and the ULFED gas pathway components. The ULFED was evaluated in two circuits so that overall hemolysis reflected that of clinical setups. A reasonable cannula for ECCO₂R at the target 250 mL/min of blood flow (13 Fr Avalon Elite DLC 10013; Maquet, Rastatt, Germany) and a pediatric centrifugal pump (PediMag; Thoratec, Pleasanton, CA) were selected for the ULFED “standard circuit”. Blood (1000 mL) was continuously recirculated for 3 h. The reservoir was submerged in a heated water bath to maintain a circuit temperature of 37° ± 1 °C. ULFED rotation was set to the minimum speed necessary where CO₂ removal did not differ significantly from the maximum rate achieved. The second ULFED circuit (“dialysis configuration”) evaluated performance using a hemodialysis controller roller pump (Prisma; Baxter, Deerfield, IL) and cannula. A larger bore 14-Fr, 15-cm dialysis cannula (AK-22142-F; Teleflex, Morrisville, NC) was used in the second circuit due to availability of parts recommended for the target blood flows.

A control circuit (“Minimax”) was tested to evaluate ULFED hemolysis against an approved low-flow blood oxygenator (Minimax Plus; Medtronic, Minneapolis, MN). Blood flow in the control loop was maintained at the minimum rate necessary (1250 mL/min) to match ULFED CO₂ removal performance according to the manufacturer [28]. Pump (BP-50; Medtronic, Minneapolis, MN) rotation speed in the loop was maintained at 2100–2200 RPM against 180 mmHg to simulate inclusion of cannula recommended for
use at the target blood flows (14 Fr Biomedicus 96820-014 venous, 12 Fr Biomedicus 96820-012 arterial) [29, 30]. Pressure against the pump was adjusted using a Hoffman clamp on ULFED outlet tubing and was continuously monitored with a differential fluid pressure transducer (PX771-025DI; Omega Engineering, Inc., Stamford, CT) across the pump. All other components and conditions were consistent between circuits. All three ULFED prototypes fabricated for gas exchange testing were evaluated for hemolysis in both circuit configurations, as the gas pathway failure observed in one prototype did not interfere with hemolysis testing.

Samples were drawn every 30 min to measure hematocrit (HCT) and plasma-free hemoglobin (pHb). Plasma was isolated from whole blood in two centrifuge spins (15 min at 0.8g, 10 min at 7.2g), and absorbance at 540 nm was measured spectrophotometrically (Genesys 10S UV-Vis; Thermo Scientific, Waltham, MA). pHb concentration was calculated from absorbance using a standard curve developed from a linear-fit of serially diluted whole blood with 100% hemolysis versus absorbance [31].

The normalized index of hemolysis (NIH) was calculated for circuit comparisons:

\[
\text{NIH (g/100L)} = \frac{\Delta \text{pfHb} \times V \times \frac{100 - \text{HCT}}{100} \times \frac{100}{\Delta t \times Q}}{100 - \text{HCT}}
\]

Where NIH = normalized index of hemolysis in grams of hemoglobin released into the blood per 100 L of flow through the circuit (g/100 L); \(\Delta \text{pfHb}\) = increase in pHb over the sampling time interval (g/L); \(V\) = circuit volume (L); \(\text{HCT}\) = hematocrit (%); \(\Delta t\) = sampling time interval (min); \(Q\) = average blood flow rate (L/min). A time-of-therapy normalized index was also calculated, since the NIH equation does not reflect total hemolysis returned to a patient in the context of treatment duration. Flow rate normalization in the NIH equation is eliminated in the new therapeutic index of hemolysis (TIH) calculation to indicate the total grams of hemoglobin released to the blood per 100 min of therapy (g/100 min):

\[
\text{TIH (g/100 min)} = \frac{\Delta \text{pfHb} \times V \times \frac{100 - \text{HCT}}{100} \times \frac{100}{\Delta t}}{100 - \text{HCT}}
\]

Statistics
All statistical comparisons were conducted in SPSS (IBM, Armonk, NY). A one-way ANOVA with Tukey HSD post hoc testing was used to compare removal rates at each RPM after data satisfied assumptions of homogeneity of variance, normality, and independence. Comparisons were used to identify the minimum speed necessary to achieve statistically equivalent performance to the maximum CO\(_2\) removal rate. The determined rotation speed was used for subsequent hemolysis testing. Mean NIH values were compared using a one-way ANOVA with Tukey HSD post hoc test after satisfying relevant assumptions. TIH data violated the assumption of homogeneity of variance via Levene’s test, and means were compared with Welch’s F test. Subsequent Games-Howell post hoc tests were used for between-group comparisons of means. All comparisons of means were considered significant at the level \(p < 0.05\).
**Results**

**In vitro gas exchange**

Figure 4 shows the raw and normalized CO₂ removal rates of the ultra-low-flow CO₂ removal device (ULFED) as a function of impeller rotational speed. A sharp increase in CO₂ removal occurred between 0 and 2000 RPM before subsequently leveling off at higher rotation speeds. A maximum normalized CO₂ removal rate of 75.1 ± 1.1 mL/min was achieved at 5000 RPM. Normalized performance at 4000 RPM did not differ significantly from the maximum rate however (74.1 ± 3.3 mL/min, p = 0.99).

**In vitro hemolysis**

Measured rates of pfHb accumulation were highly linear over testing periods as shown in Fig. 5 (ΔpfHb versus elapsed time $R^2 > 0.95$ in all tests). Table 1 shows the calculated hemolysis indices for the ULFED (at 4000 PRM) and the control device. NIH values for the standard-ULFED circuit (0.78 ± 0.19 g/100 L), dialysis-ULFED circuit (1.55 ± 0.03 g/100 L), and the control circuit (0.11 ± 0.01 g/100 L) each differed significantly from one another (ANOVA $p < 0.001$, all group-wise comparisons $p < 0.001$). The TIH value of the standard-ULFED (0.190 ± 0.041 g/100 min) did not differ significantly from the control circuit (0.123 ± 0.013 g/100 min; Welch's test $p < 0.001$, group-wise $p = 0.169$). The hemolysis using dialysis circuit components (0.386 ± 0.010 g/100 min) was significantly greater than both other test groups (each $p < 0.05$). Average hematocrit at each sampling interval is shown in Fig. 5 (right). Hematocrit decreased by ~1.5–2.5% from baseline over test periods, which is consistent with dilution due to saline infusion.

**Discussion**

Supplementing respiration by removing CO₂ independent of the lungs can improve outcomes for patients at risk of requiring or already receiving invasive mechanical ventilation. We developed the ultra-low-flow ECCO₂R device (ULFED) to operate at blood flow rates consistent with renal hemodialysis to simplify circuit management and minimize invasiveness of CO₂ removal. In vitro CO₂ removal rates up to 74 mL/min at 4000 RPM were achieved by the ULFED with minimal cell trauma.

![Fig. 4 Raw (V_{CO2}) and normalized (V'_{CO2}) CO₂ removal rates versus rotation speed. Error bars represent one standard deviation of measured removal rates](image-url)
(therapeutic index of hemolysis, TIH = 0.19 g/100 min) at blood flows consistent with dialysis (250 mL/min).

CO₂ removal systems used in conjunction with non-invasive or protective ventilation strategies have been shown to correct pCO₂ and pH in hypercapnic patients with removal rates equivalent to ~25–35% of the metabolic CO₂ production (~200–250 mL/min) [22, 23]. CO₂ removal at these levels prevented intubation in patients with ae-COPD failing or unresponsive to non-invasive ventilation [14, 16, 17, 32]. Partial respiratory assistance has also aided weaning from ventilation [14, 33, 34] and allows reduction of ventilator tidal volumes to ultra-protective levels (3–4 mL/kg) [21, 35, 36]. The ULFED exceeded these CO₂ removal rates by eliminating ~30–37% of the metabolic CO₂ production at normocapnic test conditions (inlet pCO₂ = 45 mmHg). Gas exchange will also increase proportionally with pCO₂ in hypercapnic patients, where CO₂ removal up to 50% or more of metabolic production can be required.

Efficient gas exchange in the ULFED minimizes necessary fiber surface area and enables clinically significant CO₂ removal rates at hemodialysis blood flows. Pump-less arteriovenous CO₂ removal (AVCO₂R) requires dual cannulation (13–19 Fr) for circuit flows of 600–2000 mL/min that is shunted between the femoral artery and vein through a 1.3-m² oxygenator [36–38]. A newer integrated pump-oxygenator system uses a rotating core to generate active mixing to improve gas exchange up to 60% with a 0.59-m² bundle [39]. Comparatively lower flows (350–500 mL/min) are possible in the simplified veno-venous circuit, but CO₂ removal decreases with blood flow (~50 mL/min at 300 mL/min blood flow) and connection requires 15.5-Fr cannulation [4, 39]. Developing systems combine existing oxygenators with dialysis controllers targeting even lower flows (200–300 mL/min) to minimize cannulation invasiveness (≤14 Fr). These systems utilize larger surface area gas exchangers (≥1 m²) to improve performance [32, 40] or target lower CO₂ removal using smaller pediatric oxygenators (40–55 mL/min with a 0.3-m²

**Table 1** Summary of in vitro hemolysis testing

| Test device                  | Blood flow (mL/min) | TIH (g/100 min) | NIH (g/100 L) |
|------------------------------|---------------------|-----------------|---------------|
| ULFED (standard circuit)     | 250                 | 0.190 ±0.041    | 0.775 ±0.186* |
| ULFED (dialysis configuration)| 250                 | 0.386 ±0.010*   | 1.551 ±0.025* |
| Minimax                     | 1250                | 0.123 ±0.013    | 0.105 ±0.012* |

*Significant at p < 0.05 versus all other devices. TIH therapeutic index of hemolysis. NIH normalized index of hemolysis.
bundle in pigs with \( \text{PaCO}_2 > 80 \text{ mmHg} \) [41]. Approaches to enhance \( \text{CO}_2 \) removal such as bicarbonate dialysis [42], blood acidification [43], electrodialysis [44], plasma recirculation [45], and fiber enzyme coatings [46] are also being explored to reduce necessary blood flows for treatment.

The rotating impellers in the ULFED enhance gas transfer by generating an “active mixing” effect in the fiber bundle that improves convective mixing at gas exchange surfaces [24, 25, 47, 48]. Computational simulations have indicated development of continuously recirculating flow pathways in/out of the fiber bundle with impeller mixing [25]. Blood is pumped radially outward through the bundle by impellers, then pulled back into the bundle toward low-pressure regions in the gaps between impellers before converging onto the impeller blade and cycling through the bundle again. Increasing flow velocity past gas exchange surfaces is a well-established mechanism for improving transfer efficiency by diminishing the thickness of the surface diffusive boundary layer [49]. This facilitates replenishment of gases to the membrane surface to maximize the concentration gradients spanning fiber walls. Recirculating flow also maintains a high level of washing in the bundle that eliminates regions of stagnation where thrombus formation may otherwise occur at low blood flows.

The measured rate of hemolysis in the standard-ULFED circuit was comparable to a clinically approved oxygenator circuit. Two indices of red cell trauma are reported here that indicate the rate of \( \text{pfHb} \) accumulation over time, the key difference being how time is reported. A major limitation of the NIH calculation is that hemolysis is normalized for blood flow rate, but operating flow rate is ultimately irrelevant. Two systems intended for use at 5000 mL/min versus 250 mL/min that cause equivalent rates of total cell damage would differ in NIH by a factor of 20, despite returning an equal number of \( \text{pfHb} \) species to a patient. As a result the NIH calculation is bias against low-flow devices. The TIH calculation removes the flow rate normalization and provides a clinically relevant time-of-therapy rate of hemolysis. The limitation of both indices however is that no reliable benchmark threshold values have been validated for low-flow devices against in vivo performance to our knowledge. More information or in vivo testing is therefore necessary to make conclusions regarding acceptability of the dialysis-ULFED performance. No difference in hemolysis was observed between the control and standard-ULFED circuits, so we expect in vivo hemolysis to be acceptable in this configuration.

Anticoagulation of saline infused to the ULFED may have clinical implications with extended use. Heparin levels in the saline infusion line were chosen primarily to avoid dilution of circuit anticoagulation for consistency between tests. Approved blood pumping devices utilizing saline-lubricated seals anticoagulate infusion lines at higher rates [50], while others do not require anticoagulation [51]. Elimination or minimization of local anticoagulation from the ULFED will be investigated in future prototypes utilizing a saline infusion line.

**Conclusions**

Evidence continues to grow that \( \text{ECCO}_2 \text{R} \) can effectively prevent intubation, facilitate earlier extubation, or allow reduction of ventilator settings in hypercapnic respiratory failure. The ULFED eliminates clinically significant levels of \( \text{CO}_2 \) from blood with acceptable hemolysis at hemodialysis blood flows, making minimally invasive dialysis connection
strategies and simplified management possible for ECCO₂R. Future work may focus on in vivo validation of benchtop performance or improvements to the ULFED aimed at simplifying the design, such as sealing the blood compartment with a magnetically coupled driveshaft that would obviate the driveshaft seal and saline infusion.

Abbreviations
ECCO₂R: Extracorporeal carbon dioxide removal; ECMO: Extracorporeal membrane oxygenation; NIH: Normalized index of hemolysis; pfHb: Plasma-free hemoglobin; TIH: Therapeutic index of hemolysis; ULFED: Ultra-low-flow ECCO₂R device

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Availability of data and materials
All relevant data is presented within the text to support findings. No additional data was submitted with the article.

Authors’ contributions
RJ, BF, and WF contributed to the conception and design of the study and analysis and interpretation of data. RJ, LL, and WF contributed to the drafting the manuscript for important intellectual content. All authors read and approved the final manuscript.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
WF is an equity holder and Head of the Scientific Advisory Board, for which he receives compensation, at ALung Technologies, which is commercializing an artifical lung device independent of the device described in this article. LL is a full-time employee at ALung Technologies. RJ and BF have no relevant competing interests to report.

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Author details
1 Department of Bioengineering, University of Pittsburgh, Pittsburgh, PA, USA. 2 McGowan Institute for Regenerative Medicine, University of Pittsburgh, 3025 E Carson St, Suite 226, Pittsburgh, PA 15203, USA. 3 ALung Technologies, Inc., 2500 Jane Street, Suite 1, Pittsburgh, PA 15203, USA. 4 Department of Chemical Engineering, University of Pittsburgh, Pittsburgh, PA, USA. 5 Department of Critical Care Medicine, University of Pittsburgh Medical Center, Pittsburgh, PA, USA.

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