First experiences in the Netherlands with a new single catheter-based veno-venous extracorporeal carbon dioxide removal system

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Abstract · A 34-year-old woman suffering from severe systemic lupus erythematosus was admitted to the ICU with respiratory failure due to a pneumocystis jiroveci infection resulting in severe hypercapnia which could not be controlled with conventional strategies. A new single catheter-based CO₂ removal system was initiated in order to control severe hypercapnia. The case and relevant literature are discussed.

Keywords · Artificial respiration, Pulmonary fibrosis, Extracorporeal membrane oxygenation/instrumentation, heparin

Introduction
Extracorporeal carbon dioxide (CO₂) removal has been used in the treatment of patients with acute respiratory distress syndrome (ARDS). As a consequence, tidal volumes and plateau pressure could be reduced in these patients [1,2].

Previous reports have demonstrated the safety and feasibility of a pumpless arteriovenous CO₂ removal system [2,3]. However, this system requires cannulation of the femoral artery and vein with large bore catheters and may induce a considerable left-to-right shunt [4-7].

Recently, a new technique has been launched, allowing a pump-driven system with a veno-venous approach using a standard 13 French single double lumen CVVH-catheter. Results of an experimental study in adult sheep were promising [8].

Case Report
A 34 year old woman was admitted to the Internal Medicine department of Erasmus Medical Center Rotterdam with fever. Her past medical history revealed SLE, diagnosed at the age of 26, leprosy, genital herpes, hypertension and steroid myopathy. Cerebral manifestations had been present since Autumn 2008, for which she had been treated with cyclofosfamide. Because of a high suspicion of a pneumocystis infection, she was started on trimethoprim/sulfamethoxazole 1920 mg tid iv and prednisolone 40 mg bid iv therapy. Two days after admission, she developed respiratory failure requiring intubation and mechanical ventilation. Broncho-alveolar lavage (BAL) showed pneumocystis jiroveci and computerized tomography of the chest revealed extensive bilateral infiltrates with hardly any aerated areas left. High positive end-expiratory pressures were required to maintain adequate oxygenation. Peak inspiratory pressure rose to 52 cmH₂O.

In the first 3 weeks, her clinical condition improved with a PaO₂/FiO₂ ratio increasing from 80 up to 230, and declining peak end-expiratory pressures (from 19 cm H₂O to 8 cm H₂O). A percutaneous tracheostomy was performed to facilitate weaning from mechanical ventilation.

On day 37, her condition suddenly deteriorated due to a sepsis with positive blood cultures for enterococcus. Despite treatment with vancomycin, her respiratory condition also deteriorated with increasing requirements for oxygen. The chest X-ray showed increasing bilateral infiltrates, suggestive of an intercurrent pneumonia. Furthermore, possible signs of pulmonary fibrosis were present.

At this time it was felt that despite the severe underlying disease, the clinical condition might still be partially reversible. High inspiratory peak pressures (up to 52 cmH₂O) were needed to control hypercapnia. Permissive hypercapnia was used, however, despite all measures taken, pCO₂ levels rose to high levels (11 kPa) and oxygenation and haemodynamic problems were encountered when trying to decrease the pCO₂ level. Full extracorporeal membrane oxygenation was not an option due to marginal arterial access in this patient.

Because of failure to control the progressive hypercapnia while limiting inspiratory pressures, the patient was placed on a extracorporeal veno-venous carbon dioxide removal system.

This veno-venous carbon dioxide removal system (Decap®, Hemodec srl, Salerno, Italy) is a low-flow system using a percutaneous single venous access with a double lumen catheter. The machine has two circuits. In the first circuit, blood flows to the decapneizator, containing a CO₂ filter. In this circuit, oxygen
is used as a sweep gas at a flow rate of 5 litres per minute. This device allows a maximum oxygen flow rate of 10 l/min resulting in 8-10% oxygen increase in the patient’s blood. The second circuit is a pre-dilution circuit removing CO₂ from the plasmatic water, improving the decapneizator efficiency (Figure 1). The maximum blood flow is 350-400 ml/min allowing maximal CO₂ extraction of approximately 30%. In the present case, the blood flow averaged 300 ml/min. Heparin was needed as a coagulant in order to prevent thrombus formation. In order to reach adequate anticoagulation, an activated partial thromboplastin time (APTT) between 60-80 seconds is required. Figure 2 shows the PaCO₂, pH and peak pressures after starting the veno-venous removal system. The inspiratory peak pressures could be decreased from 52 cmH₂O to 37 cmH₂O.

At day 41, an accidental bolus of intravenous heparin was administered, leading to heparin overshoot with an APTT over 240 seconds. Within a couple of hours, the patient suffered an uncontrolled bleeding from the groin at the insertion site of the double-lumen catheter. After multiple transfusions of erythrocyte concentration and fresh frozen plasma, the decision was made to remove the catheter in order to control the bleeding more optimally with a pressure-device. After a few hours of fluid resuscitation, the patient’s condition stabilized. Considering the marginal vascular access in this patient, we decided not to restart the CO₂ removal therapy at this point.

A few days after termination of the CO₂ removal therapy, again a progressive hypercapnia developed. The diagnosis of progressive pulmonary fibrosis was made. In agreement with her husband and her family, treatment was stopped.

### Discussion

Our case describes a young female suffering from pneumocystis infection leading to severe pulmonary fibrosis. Conventional therapeutic strategies were not effective, resulting in very high arterial PaCO₂ with corresponding respiratory acidosis and low, but not yet life-threatening PaO₂ levels. When PaCO₂ rose to levels as high as 11 kPa (82.5 mmHg), and oxygenation deteriorated further, we decided to apply an extracorporeal gas exchange device.

It was felt that extracorporeal membrane oxygenation or NovaLung™ in this patient, requiring optimal arterial access was not an option due to the very poor arterial status. For this reason, a new veno-venous CO₂ removal system was the only available option. This new technique is attractive because of its simplicity permitting a single venous access. The rationale for using this device was primarily to reduce PaCO₂ levels and thus reduce inspiratory peak pressures. At the time of starting the treatment with Decap, the patient had already required high peak pressures for more than a week and PaCO₂ was progressively rising.

Several studies have shown the safety and feasibility of using pumpless CO₂ removal systems in patients with acute lung injury exhibiting hypercapnia and respiratory acidosis [3,4,9-12]. Systems used in these studies were either arterio-venous or veno-venous systems. An experimental study in sheep has demonstrated that a significant reduction in PaCO₂ using the Decap system can be achieved [8].

In our case, suboptimal anticoagulation led to clotting of the system. Due to an accidental bolus of heparin, a severe bleeding of the groin at the insertion site occurred requiring multiple...
transfusions. It was necessary to terminate the CO₂ removal and remove the venous catheter in order to control this bleeding.

After removal of the catheter, the PaCO₂ level remained acceptable for three more days, without the need to increase ventilatory pressures to control PaCO₂. For this, we do not have a satisfactory explanation. Unfortunately, our patient died five days after discontinuing Decap due to progressive respiratory failure caused by severe pulmonary fibrosis.

**Conclusion**

We show that this new veno-venous CO₂ removal technique is an effective method in controlling PaCO₂ in a patient with severe respiratory failure due to pneumocystis infection, complicated by pulmonary fibrosis. More experience and clinical trials are needed to evaluate this technique.

**Figure 2.** PaCO₂, pH and peak pressures at start, during and after cessation of the treatment.

**References**