**Expression of Mobility Group Box 1 Protein in a Polytrauma Model Treated with ECLS at Ground Level and High Altitude**

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Expression of high mobility group box 1 protein in a polytrauma model treated with ECLS at ground level and high altitude

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Introduction

- Acute respiratory distress syndrome (ARDS) is the most severe form of acute lung injury, characterized by acute onset of hypoxemia, bilateral radiographic pulmonary infiltrates without cardiogenic pulmonary edema, and may lead to sepsis and multi-organ failure.
- Injuries incurred in austere environments, particularly in the combat setting, require immediate evacuation with en-route care. Extracorporeal membrane oxygenation (ECMO) may be used to support ARDS patients during transport, including during aeromedical evacuation.
- High mobility group protein box 1 (HMGB1) is an important indicator of damage-associated molecular pattern (DAMP) expression and disease progression in ARDS.
- HMGB1 has been identified as a mediator of ARDS and is expressed in blood following activation of damaged cells.
- Little is known regarding HMGB1 expression in a pulmonary contusion model of ARDS supported by ECLS at ground level.
- Altitude change effect on HMGB1 expression during air transportation is also unknown.

Hypothesis

We hypothesized that HMGB1 expression in systemic blood increases following chest contusion and that HMGB1 expression is affected by changes in altitude to a greater extent in injured animals supported by ECLS versus healthy animals on ECLS undergoing the same altitude exposure.

Methods

- Female Yorkshire pigs (54.17 ± 1.27 kg) (n=15) were anesthetized and received arterial and venous catheters, followed by tracheostomy.
- Following baseline measurements, animals were cannulated and veno-venous ECMO was initiated (Cardiohelp, Maquet Gmbh, Gettinge Group, Rastatt, Germany), via an Avalon 23 Fr. catheter inserted into the right jugular vein.
- Blood flow was 1.2-3 L/min and sweep gas flow ranged at 4-8L/min. Continuous heparinization was started at cannulation and titrated to 30-50% higher than baseline ACT levels.
- Animals were then transported via a standard NATO litter fitted with a next-generation medical equipment rail kit (MERK, Smed Technologies, Cummings, GA) to an adjacent building housing hypobaric chambers.
- The altitude simulation profile consisted of the multiple levels of simulated atmospheric exposure, and is depicted in Figure 2.
- Altitude exposure occurred in healthy state on Day 1, and injured state on Day 2.
- Injury consisted of bilateral pulmonary contusions using a modified captive-bolt stunner (Model ML, Karl Schermer, Packers Engineering, Omaha, NE) and chest tube placement.
- HMGB1 ELISA (IBL international, STS1011, NC, US) was utilized to analyze the level of HMGB1 in the blood at each time-point.
- Plasma free hemoglobin (pfHb) was measured in real time by spectrophotometer method.
- Plasma total protein concentration (PTPC) was measured by Pierce™ BCA protein assay kit (Thermo scientific, Rockford, IL, US)
- Post-mortem lung tissue samples were fixed by 10% normal buffered formalin and paraffin embedded, thickness 4 µm sliced tissues were stained by Hematoxylin & Eosin or primary antibody immunohistochemistry for HMGB1 and TLR4 (abcam, CA, US).

Results

- Figure 3. Post-mortem Image of Lungs after Bilateral Pulmonary Contusion Treated with ECMO
- Figure 4. Post-Mortem Expression of hmgb1, TLR4 after Bilateral Pulmonary Contusion Treated with ECMO
- Figure 5. Histology image and diffused alveolar damage score in a bilateral pulmonary contusion
- Figure 6. Dramatic change of HMGB1 protein, pfHb and PTPC in a bilateral pulmonary contusion treated with ECMO at ground level and high altitude during en-route care.

Conclusion

- High altitude does not alter HMGB1, pfHb and PTPC to expression in uninjured state on ECLS. Pulmonary contusion causes a transient increase in HMGB1 and pfHb levels. The level of HMGB1 and pfHb of early died animals were significantly higher than survived group. Beside assessment of HMGB1 and pfHb confirms injury and may provide a useful monitoring capability during en-route care, and should be a part of precision medicine lab-on-a-chip type assays in the future.

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