Case Report

Heart Failure and Hypothermia in an Infant: Pseudocyanide Syndrome?

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Purpose. Mixed or central venous oxygen saturation has not been described during concurrent heart failure and hypothermia in children, both of which may be associated with hyperlactatemia. This report of an infant with heart failure and hypothermia is significant for increased inferior vena cava (IVC) oxygen saturation and hyperlactatemia.

Case Report. A 36-day-old female was fussy for a day and then developed respiratory distress. In the Pediatric ER, she was tachycardic (260 beats/minute) and hypothermic (32.4 degrees C) with prolonged capillary refill and faint distal pulses. She was placed on continuous positive airway pressure via nasal prongs. Adenosine was given twice via an intraosseous line for supraventricular tachycardia, with conversion to sinus rhythm. She was then intubated for worsening respiratory distress and placed on an FiO₂ of 1.0. An echocardiogram was notable for decreased biventricular systolic function and left to right shunting through a patent foramen ovale. A dobutamine infusion was started. Arterial cannulation was unsuccessful; transcutaneous pulse SO₂ was consistently 100% during rewarming. Results from the first blood gas (all blood gas results were reported uncorrected for temperature) drawn from a catheter placed in the IVC (by Seldinger technique; sutured in place after advancing to its full length overlying the L3 vertebral body; Figure 1) were pH 7.01, PCO₂ 33 mm Hg, PO₂ 101 mm Hg, SO₂ 94%.

Conclusions. Concurrent heart failure and hypothermia in an infant were associated with increased IVC oxygen saturation and hyperlactatemia, similar to lab findings associated with a mitochondrial toxin such as cyanide. Improvement of heart failure and hypothermia were associated with resolution of these lab abnormalities, thus helping to rule out mitochondrial toxins. Additional reports may help better define a pseudocyanide syndrome in this setting.

1. Introduction

Mixed or central venous oxygen saturation (SO₂) is decreased with heart failure and increased with hypothermia; though normal limits for mixed or central venous SO₂ require interpretation within a clinical context, levels below 70% and above 90% are concerns [1]. Hyperlactatemia is common to both heart failure and hypothermia. With concurrent heart failure and hypothermia in children, hyperlactatemia is increased in proportion to heart failure severity [2], but mixed or central venous SO₂ data are lacking. This report of an infant with heart failure and hypothermia is notable for increased inferior vena cava (IVC) SO₂ and hyperlactatemia, similar to the findings associated with a mitochondrial toxin, such as cyanide.

2. Case Report

The Columbia University Medical Center IRB (AAAR0802) exempted this retrospective case report from review.
and lactate 18 mmol/L. Early during rewarming, the IVC SO₂ remained above 90% and was as high as 99% (Figure 2); declines in IVC SO₂ and lactate during the first 3 hours of rewarming correlated (Spearman r: 0.94; p = 0.002). An arterial catheter was then placed, and the FiO₂ weaned. B-type natriuretic peptide was increased (4578.5 pg/mL). Capillary refill and distal pulses improved, and dobutamine was discontinued at 8 hours; a repeat echocardiogram the next day showed improved (though still diminished) biventricular systolic function with continued left to right shunting through a patent foramen ovale. Hypothermia and hyperlactatemia resolved over 10 and 12 hours, and IVC SO₂ was 80% on hospital day 2. The patient was diagnosed with supraventricular tachycardia secondary to Wolff-Parkinson-White syndrome. Prior to discharge a week later, serum acylcarnitine profile, lactate, pyruvate, thyroid stimulating hormone, and thyroxine were all normal.

IVC PO₂ values were corrected retrospectively for the patient’s temperature [3] (Table 1).

3. Discussion

Concurrent acute heart failure and mild (>32 degrees Centigrade [4]) hypothermia were associated with an increased SO₂, profound metabolic acidosis, and severe hyperlactatemia in blood from the IVC of an infant. Heart failure alone is associated with a decreased mixed venous or central venous SO₂—as well as a decreased IVC SO₂—in neonates [5] as well as infants and children; mild hypothermia alone is associated with an increased—though still normal—mixed venous [6] or central venous [7] SO₂ and is unlikely to be responsible for profound acidosis [4] or severe hyperlactatemia [8, 9]. Resolution of heart failure and mild hypothermia in this infant was associated with resolution of the lab abnormalities: a decrease in cardiac output and tissue metabolic rate, as well as peripheral vasoconstriction and vascular shunting, all improved with resolution of heart failure and hypothermia. Similar lab abnormalities are associated with cyanide toxicity [10, 11], though the pathophysiology is different: as oxidative phosphorylation is disrupted, oxygen extraction and consumption decrease due to the metabolic block, and venous SO₂ increases with hyperlactatemia. Though a metabolic disease or toxin might have been associated with the clinical presentation of this infant, there was ultimately no evidence to support such a diagnosis or its associated pathophysiology.

A clinical controversy regarding blood gas results may have escaped the notice of some pediatric intensivists: correcting blood gas results for the patient’s temperature is no longer routine, as recommended [12]. As a result, blood gas specimens are routinely analyzed at 37 degrees Centigrade (normothermia) and so reported. Interestingly, correcting the blood gas results retrospectively in this infant did not alter the findings: early during rewarming (on the first 3 blood gas results), the corrected IVC PO₂ was >80 mm Hg, corresponding generally to SO₂ values on the flat upper part of the oxyhemoglobin dissociation curve. As calculations of SO₂ from PO₂ may be associated with inaccuracies [13], specific SO₂ values were not calculated; however, a leftward shift of the oxyhemoglobin dissociation curve (due both to hypothermia and the presence of fetal hemoglobin in a 5 week old infant) supports IVC SO₂ values >90% for PO₂ values >80 mm Hg. It is thus evident that both corrected and uncorrected IVC SO₂ values early during rewarming were increased above normal.

The IVC SO₂ was >90% in 27% (15 of 55) of values from 20 critically ill neonates and infants undergoing cardiopulmonary bypass (mostly with hypothermia), though lactate was only mildly elevated (mean: 2.2 +/- 0.1 mmol/L) [14].

**Figure 1:** Radiograph of infant with heart failure and mild hypothermia showing central venous catheter overlying L3 vertebral body.

**Figure 2:** Inferior vena cava oxygen saturation (uncorrected for temperature) and lactate during rewarming of infant with heart failure and mild hypothermia.
As cardiopulmonary bypass replaces cardiac function, it is possible that heart failure with mild hypothermia in an infant would be associated with more severe hyperlactatemia and an increased IVC SO₂, as noted in this infant. In any case, the trend in lab results from the IVC catheter was consistent during rewarming following cardioversion and suggests that these findings represent an actual association.

Limitations to this report include those related to retrospective case reports generally. Some laboratory data—including arterial blood gas results as well other metabolic testing—were not available due to limited vascular access during acute illness. The FiO₂ was maintained at 1.0 during the first 3 hours of rewarming while the infant's clinical condition stabilized, and this could have contributed to an increased IVC SO₂; however, this effect was not dramatic, as the IVC SO₂ fell to 83% before the FiO₂ was weaned. In any case, pediatric intensivists may benefit from knowing that the SO₂ from a femoral vascular catheter may be as high as 99% in certain clinical conditions, even when the catheter is not arterial. As the IVC catheter was advanced to its full length and sutured in place overlying the L3 vertebral body, proximal catheter migration was not possible, such that the left to right shunt across the patent foramen ovale did not contribute to an increased IVC SO₂.

4. Conclusion

This report offers anecdotal evidence that IVC SO₂ may not always be low in infants with heart failure, while lactic acidosis may be profound in the presence of mild hypothermia: concurrent heart failure and mild hypothermia were likely responsible for the lab abnormalities in this infant, which to some extent mimic the lab abnormalities associated with a mitochondrial toxin such as cyanide. Further study of a pseudocyanide syndrome associated with concurrent heart failure and mild hypothermia in infants may be helpful.

Disclosure

This study was presented in abstract form at the 2017 American Thoracic Society’s Annual meeting [15].

Conflicts of Interest

The author has no conflicts of interest relevant to this article to disclose.

References


Table I: Temperature, uncorrected and corrected (for temperature) IVC PO₂, uncorrected IVC SO₂, and IVC lactate in infant with heart failure and mild hypothermia.

<table>
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<tr>
<th>Time (minutes)</th>
<th>Temperature (degrees Centigrade)</th>
<th>Uncorrected PO₂ (mm Hg)</th>
<th>Corrected PO₂ (mm Hg)</th>
<th>Uncorrected SO₂ (%)</th>
<th>Corrected SO₂ (%)</th>
<th>Lactate (mmol/L)</th>
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*Interpolated temperatures.


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