B-type natriuretic peptide levels predict outcomes for children on extracorporeal life support after cardiac surgery

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Objective: Extracorporeal life support is used in 3% to 8% of infants and children after cardiac surgery. B-type natriuretic peptide may have utility as a biomarker in these patients. The objective of this study was to investigate potential associations between changes in B-type natriuretic peptide during trials off extracorporeal life support and clinical outcome.

Methods: Ten infants and children requiring extracorporeal life support after cardiac surgery were studied prospectively. Before separation from extracorporeal life support, a shunt was placed in the circuit, allowing for temporary trials off life support. Serum lactate, arterial–venous oxyhemoglobin saturation difference, and B-type natriuretic peptide levels were determined before each trial off life support and at the end of each trial off life support, and the ability to predict postoperative outcome from these data was evaluated.

Results: During trials off extracorporeal life support, lactate, the arterial–venous oxyhemoglobin saturation difference, and B-type natriuretic peptide levels increased above pre-trial values (P < .05). Only the arterial–venous oxyhemoglobin saturation difference predicted successful separation from extracorporeal life support after a trial (P < .05). There were no associations between long-term outcome and alterations in lactate and the arterial–venous oxyhemoglobin saturation difference during the final trials off life support. However, an increase in B-type natriuretic peptide levels during the final trial off life support (trial/pre-trial ratio of >1) had a sensitivity of 80% and a specificity of 100% for predicting the need for an unplanned operation or death within 3 months (P < .05).

Conclusion: B-type natriuretic peptide determinations may be a useful tool for clinicians caring for infants and children requiring extracorporeal life support after cardiac surgery.

Extracorporeal life support (ECLS) is used in 3% to 8% of infants and children after corrective or palliative surgery for congenital cardiac defects.1–5 Although historically a rescue therapy, technologic advances and accumulated clinical experience have expanded the indications for ECLS to permit earlier initiation and even prophylactic treatment of low cardiac output syndrome.6–9 Consequently, patients supported with ECLS are an increasingly heterogeneous group, with some who require only time for recovery and others who require further interventions. Distinguishing between these types of patients is important to minimize the duration of ECLS and to ensure that patients requiring further investigations and interventions are identified. Thus, much of the clinical focus during ECLS is aimed at assessing recovery of cardiovascular function. One common method involves trials off ECLS that are achieved by placing a shunt in the ECLS circuit, allowing mechanical support to be temporarily stopped, but keeping the cannulas...
and circuit components free of thrombosis and ready for immediate resumption. During these trials off ECLS, cardiovascular function is assessed by observing the hemodynamic response and through the measurement of various markers of cardiac output, such as lactate, central venous hemoglobin saturation, acid–base status, and urine output. Unfortunately, none of these factors has been shown to adequately predict either a readiness to separate from ECLS or the need for further intervention. Thus, novel biomarkers that might aid in these assessments are needed.

B-type natriuretic peptide (BNP) is a 32-amino acid polypeptide hormone with diuretic, natriuretic, and vasoactive properties. As a cardiac hormone produced by the atria and ventricles in response to myocyte stretch, it has been shown to have unique clinical utility as a biomarker and is thus widely used in the management of adult cardiac disease, with an emerging use in pediatric cardiac disease. In fact, BNP has been studied in adult and pediatric patients requiring mechanical cardiac support. However, the value of BNP levels in predicting successful separation from ECLS or in predicting outcome after surgery, including the need for further interventions, has not been established.

Thus, the objectives of this study were (1) to determine whether alterations in BNP during trials off ECLS were associated with our ability to separate patients from ECLS and (2) to investigate potential associations between changes in BNP during trials off ECLS and outcome. We prospectively studied 9 infants and children who required ECLS after cardiac surgery. Systemic arterial plasma BNP determinations were made before and during each trial off ECLS, and whether a change in BNP could predict separation from ECLS, and/or postoperative outcome was evaluated.

Materials and Methods
This prospective cohort study was conducted in the Pediatric Cardiac Intensive Care Unit (PCICU) at the University of California, San Francisco, between March 2005 and June 2006. Patients with congenital cardiac defects undergoing surgical repair were enrolled and patients requiring postoperative ECLS were included in the study.

The patients were followed up during their entire course in the PCICU. The perioperative anesthesia management, cardiopulmonary bypass (CPB) strategy, and subsequent PCICU management followed standard institutional practices. An on-service team that was blinded to the BNP values made all decisions regarding patient management.

Written informed consent was obtained from the patients’ parents or guardians before enrollment of the patients into the study. The Institutional Review Board at the University of California, San Francisco, reviewed and approved this study.

Indications for ECLS
The indications for ECLS included (1) the inability to separate a patient from CPB, necessitating the initiation of ECLS in the operating room, (2) the elective initiation of ECLS in the operating room for low cardiac output after discontinuing CPB, (3) the development of refractory low cardiac output syndrome in PCICU, or (4) postoperative cardiac arrest, with ECLS used as a part of cardiopulmonary resuscitation. Signs of low cardiac output in the operating room or intensive care unit that resulted in the initiation of ECLS included sustained systemic hypotension and/or persistent metabolic acidosis refractory to medical management.

ECLS
Our techniques for circulatory support have been described in detail previously. The ECLS circuit was a Medtronic BioMedicus Portable Cardiopulmonary Bypass System 1000, consisting of a centrifugal pump Bio-Console model 550 (BP-50 Bio-Pump; Medtronic Inc, Anaheim, Calif) and a hollow-fiber membrane oxygenator (Mini-max Plus or Affinity NT, Medtronic), in which the whole surface was heparin-bonded. Arterial (left atrial or aortic) and venous (right or common atrial) cannulas were placed through either a transcervical or transsternal approach. Left ventricular decompression with an additional left atrial cannula was used selectively for patients with biventricular hearts and severe left ventricular dysfunction. Systemic–pulmonary shunts were kept open during ECLS. For patients with adequate pulmonary function and oxygenation requiring only left ventricular support without an oxygenator, a single venous cannula was placed in the left atrium and the arterial cannula was placed in the ascending aorta. Circuit flow was targeted at 80 to 120 mL · kg⁻¹ · min⁻¹ or 150 mL · kg⁻¹ · min⁻¹ for shunt-dependent patients. Inotropes were discontinued or decreased during ECLS support. Heparin infusions were used in most cases to maintain an activated clotting time of 180 to 220 seconds.

Separation from ECLS
The decision to discontinue ECLS was based on patients recovering adequate cardiovascular function. To assess native cardiovascular function, we placed a shunt or bridge between the arterial and venous cannulas. With the use of 3-way connectors, ECLS flow was diverted away from the patient, but full flow was maintained across the remainder of the circuit, allowing for trials off ECLS without removal of the cannulas. The cannulas were flushed during this time with a continuous infusion of heparin solution. During trials off ECLS, an observer blinded to the BNP data collected hemodynamic and biochemical data. In most trials, echocardiography was performed to assess ventricular function. ECLS was discontinued, with removal of the cannulas, if the cardiac team believed that cardiovascular function was adequate. These assessments were variably based on the hemodynamic, echocardiographic, and circuit components free of thrombosis and ready for immediate resumption. During these trials off ECLS, cardiovascular function is assessed by observing the hemodynamic response and through the measurement of various markers of cardiac output, such as lactate, central venous hemoglobin saturation, acid–base status, and urine output. Unfortunately, none of these factors has been shown to adequately predict either a readiness to separate from ECLS or the need for further intervention. Thus, novel biomarkers that might aid in these assessments are needed.

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graphic, and biochemical data obtained during the trials off ECLS, and the decisions were individualized for each patient. In general, a period of stable hemodynamics with the support of fluid volume loading and inotropic agents, the appearance of improvement in cardiac function by echocardiography as compared with the period before ECLS initiation, and minimal changes in venous oxygen saturation and acid–base status were sought. All members of the cardiac team making these decisions were blinded to the BNP data.

Outcomes
A good short-term outcome was defined as separation from ECLS after a trial off ECLS. Poor short-term outcome was defined as the need for resumption of ECLS after a trial off ECLS. Good long-term outcome was defined as the absence of death or unplanned operation within 3 months of separation from ECLS. Poor long-term outcome was defined as an unplanned operation within 3 months or death within 3 months of separation from ECLS. BNP levels, hemodynamic data, and other biochemical values were evaluated for their association with these end points.

Data Collection
Blood samples were obtained from an arterial catheter preoperatively, before each trial off ECLS, and at the end of each trial off ECLS. The samples were immediately placed on ice in chilled ethylenediaminetetraacetic acid tubes and centrifuged at 3000 rpm for 15 minutes at 4°C. Separated plasma was stored at −20°C. Within 7 days of obtaining the sample, the plasma was thawed to room temperature, and BNP levels were measured with a commercially available fluorescence immunoassay (Triage Meter Plus; Biosite Diagnostics, San Diego, Calif). The measurable range of BNP on this device is between 5 and 5000 pg/mL. The estimated coefficient of variation for the assay is 9.2% to 11.4%.

Clinical and biochemical data were prospectively collected at each sampling point and once daily thereafter by an observer blinded to the BNP data. The clinical data collected included demographics, CPB time, myocardial ischemic time during the operation, duration of ECLS, duration of each weaning trial, duration of mechanical ventilation, inotrope dose, mean systemic arterial pressure, central venous pressure, urine output, and fluid balance. Biochemical data included arterial and venous blood gases, base deficit, and serum lactate levels. The duration of mechanical ventilation was quantified as the number of ventilator-free days within the first 28 days after the operation, so as to avoid confounding the data with patients who died while receiving mechanical ventilation.

Calculations
Inotrope use was quantified by a score adapted from Wernovsky and associates. The score was calculated from the level of inotropic support the patients were receiving (in micrograms per kilogram per minute) at each sampling point according to the following equation: dopamine + dobutamine + (epinephrine + norepinephrine) × 100) + (milrinone × 20). The arterial–venous oxyhemoglobin saturation difference (AVdO2) was calculated as the co-oximetric arterial oxyhemoglobin saturation minus the central venous oxyhemoglobin saturation.

Analysis of the Data
Data are described as mean ± standard deviation, range, or percentage, as appropriate. Differences in the continuous variables between good outcome and poor outcome subgroups were tested with the Student t test or Mann–Whitney U test. Differences of variables between subgroups within category were tested with the Fisher exact test. Contingency tables were formulated to determine the sensitivity value, specificity value, positive predictive value, and negative predictive value of changes in serum BNP, lactate, and the AVdO2. Differences of variables within groups at two different time points were compared by paired t test. Correlations between variables were performed by the Spearman rank correlation method. Statistical analyses were performed with the use of Prism (GraphPad Software, Inc, San Diego, Calif).

Results
Patients
A total of 11 patients required ECLS after cardiac surgery and were enrolled in the study. Two patients died without undergoing a trial off ECLS and thus were excluded from the analysis. One patient underwent 2 separate operations, 3 months apart, and required ECLS after each operation. For the analysis, this patient is considered separately on each admission (Patients 6 and 7, Table 1). The patients’ demographic data, cardiac defects, surgical repairs, indications for ECLS, types of ECLS, ECLS durations, and long-term outcomes are outlined in Table 1. The mean intraoperative bypass time was 123 minutes ± 43 (range 69–188 minutes), with a mean crossclamp time of 61 minutes ± 30 (range 26–116 minutes). Four (40%) patients had single ventricle physiology. In 3 (30%) patients, ECLS consisted of mechanical ventricular support without an oxygenator; the remaining 7 patients received mechanical ventricular support with an oxygenator. There was no association between the use of an oxygenator and outcome.

Trials of Patients off ECLS
For the 10 patients studied, there were 18 trials off ECLS. BNP levels were obtained during 16 of these trials. Five (50%) patients underwent only 1 trial off ECLS. The mean duration of the trials was 94 minutes ± 102 (range 9–340 minutes). Hemodynamic and biochemical data before (pretrial) and during trials off ECLS (trial) are shown in Table 2. When all trials were taken together, heart rate and central venous pressure increased, and mean arterial pressure decreased from pre-trial values (P < .05). In addition, the inotrope score increased during trials off ECLS (P < .05). Furthermore, serum lactate, the AVdO2, and serum BNP values increased above pre-trial values (P < .05). There was no correlation between the duration of the trials off ECLS and any of the biochemical values or hemodynamic indices.

Separation of Patients From ECLS
Nine (90%) patients were ultimately able to separate from ECLS. One patient who underwent trials off ECLS died
without separating from ECLS. In both the poor and good
short-term outcome groups, mean arterial pressure de-
creased, and central venous pressure, heart rate, and inotro-
pic score increased during the trials ($P < .05$).

Biochemical changes during the trials for each group are
shown in Figure 1. Serum BNP and lactate levels tended to
increase in the poor short-term outcome group, but these
changes did not reach significance (Figure 1). However, the
AVdO₂ was significantly higher in the poor short-term out-
come group than in the good short-term outcome group
(Figure 1; $P < .05$). In fact, an AVdO₂ of greater than 40
during a trial off ECLS had a sensitivity of 100% and a
specificity of 70% for predicting a poor short-term outcome
($P < .05$).

Long-term Outcomes
The long-term outcome for each patient can be seen in
Table 1. There were 5 (50%) poor long-term outcomes and
5 (50%) good long-term outcomes. One patient (patient 1,
Table 1) required an unplanned operation within 3 months
after ECLS and died within 6 months of ECLS. The un-
planned operation is considered the single poor outcome for
this patient. Thus, 3 (30%) patients died within 3 months
after ECLS, and 2 (20%) patients required an unplanned
operation within 3 months after ECLS.

Patients with poor long-term outcomes had a longer
aortic crossclamp time (43 minutes ± 14 vs 78 minutes ±
29; $P < .05$) and more trials off ECLS (1.4 ± 0.5 vs 2.8 ±
1.1; $P < .05$) than patients with good long-term outcomes. There were no significant differences between the two
groups in age, weight, type of cardiac defect, preoperative
BNP levels, bypass time, ECLS duration, or hospital days.
In addition, there were no significant differences between
the groups in pre-trial or trial mean arterial pressures, cen-
tral venous pressures, heart rates, urine output, or inotrope
scores.

Days in the PCICU tended to be longer in the poor
outcome group, but the difference did not reach significance
(25 days ± 13.6 vs 13.4 days ± 6.6; $P = .12$). Patients in
the poor outcome group had fewer mechanical ventilation–
free days than patients in the good outcome group (8.4 days ±
8.1 vs 20 days ± 5.7; $P < .05$).

Biochemical changes during the final trial off ECLS
for patients with good and poor long-term outcomes are
shown in Figure 2. There were no differences in the
AVdO₂ or in the change in lactate during the trials be-
tween groups (Figure 2). Pre-trial lactate levels were
greater in the poor long-term outcome group than in the
good long-term outcome group (1.8 mmol/L ± 1.0 vs 1.0
mmol/L ± 0.21; $P = .11$), but the values were within the
normal range and were not predictive of outcome. Abso-
lute lactate levels during the trials were also not predic-
tive of outcome.

<table>
<thead>
<tr>
<th>Table 1. Patient demographics</th>
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<tbody>
<tr>
<td>Patient No.</td>
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<td>3</td>
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<td>4</td>
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<td>5</td>
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<td>6*</td>
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<td>7*</td>
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<tr>
<td>8</td>
</tr>
<tr>
<td>9</td>
</tr>
<tr>
<td>10</td>
</tr>
</tbody>
</table>

ECLS, Extracorporeal life support; DORV, double-outlet right ventricle; VSD, ventricular septal defect; RVOT, right ventricular outflow tract obstruction; HLHS, hypoplastic left heart syndrome; AVC, atrioventricular canal; AV, atrioventricular; TGA, transposition of great arteries; CPB, cardiopulmonary bypass; ASD, atrial septal defect; AS, aortic valve stenosis; AI, aortic insufficiency; RV, right ventricle; OR, operating room; ALCAPA, anomalous left coronary artery from the pulmonary artery; TAPVR, total anomalous pulmonary venous return. *The same patient with two different admissions 3 months apart.
The change in BNP during a trial, as defined by the ratio of the trial to pre-trial BNP level, was significantly higher in the poor long-term outcome group than in the good long-term outcome group (Figure 2; $P<.05$). BNP levels decreased from pre-trial levels during the final trials off ECLS in all 5 patients with a good long-term outcome (from 388 pg/mL to 306 pg/mL; $P<.05$) and increased during the final trials in 4 of the 5 patients with a poor long-term outcome (from 638 pg/mL to 953 pg/mL; $P=.07$). The 1 patient (patient 2, Table 1) with a poor long-term outcome who had a decrease in BNP during the final trial off ECLS was separated from ECLS but 48 hours later had support withdrawn owing to multisystem organ failure. In fact, an increase in BNP during the final trial off ECLS (trial/pre-trial ratio of $>1$) had a sensitivity of 80% and a specificity of 100% for predicting a poor long-term outcome ($P<.05$).

**Discussion**

These data indicate that BNP may have clinical utility as a biomarker for infants and children requiring ECLS after cardiac surgery. Although BNP levels during trials off ECLS did not predict subsequent separation from ECLS, an increase in BNP during the final trial off ECLS was associated with the need for an unplanned operation or death within 3 months. Interestingly, no other changes during trials off ECLS were associated with these long-term outcomes, and thus BNP may reflect myocardial derangements in a manner not captured by other physiologic markers.

We chose to investigate BNP because it is a cardiac hormone integrally related to the myocardium, as opposed to other biomarkers that reflect more global processes. BNP is a member of a structurally related group of peptide hormones, termed natriuretic peptides. Biosynthesis of BNP begins with a 134-amino acid precursor, which is cleaved in steps to form the active C-terminal 32-amino acid hormone, BNP, and the inactive N-terminal proBNP. Active BNP is stored in the atria in specific storage organelles. Basal BNP levels result from continuous secretion from the atria. With acute myocardial distention, BNP release increases greatly from this depletable storage pool, in a manner independent of BNP synthesis. However, under chronic conditions of increased cardiac volume loading, such as congestive heart failure, increases in circulating BNP are

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<th>ECLS duration (d)</th>
<th>Long-term outcome</th>
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<tr>
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<td>Yes</td>
<td>2</td>
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<td>Yes</td>
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<td>Yes</td>
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**TABLE 2.** Hemodynamic and biochemical data in all patients before (pre-trial) and during (trial) trials off ECLS

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<th>$P$ value</th>
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<tr>
<td>Lactate (mmol/L)</td>
<td>1.44 ± 0.71</td>
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<td>Inotrop score</td>
<td>15.6 ± 5.2</td>
<td>21.1 ± 9.1</td>
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<tr>
<td>HR (beats/min)</td>
<td>136 ± 17</td>
<td>152 ± 21</td>
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<tr>
<td>MAP (mm Hg)</td>
<td>62 ± 11</td>
<td>50 ± 10</td>
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<tr>
<td>CVP (mm Hg)</td>
<td>10 ± 4</td>
<td>15 ± 4</td>
</tr>
<tr>
<td>ScvO$_2$ (%)</td>
<td>72 ± 14</td>
<td>50 ± 22</td>
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<tr>
<td>AVdO$_2$ (%)</td>
<td>28.5 ± 14.2</td>
<td>44.2 ± 20.2</td>
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<td>BNP (pg/mL)</td>
<td>581 ± 382</td>
<td>777 ± 637</td>
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$HR$, Heart rate; $MAP$, mean arterial pressure; $CVP$, central venous pressure; $ScvO$_2, central venous oxygen saturation; $AVdO$_2, arterial–venous oxyhemoglobin saturation difference; $BNP$, B-type natriuretic peptide.
maintained owing to ventricular re-expression of the fetal gene program. In addition, acute myocardial ischemia and inflammatory mediators, such as tumor necrosis factor-α and interleukin-1β, result in rapid ventricular ex-
pression of BNP. The primary actions of BNP, which are mediated by the secondary messenger cGMP, are vascular smooth muscle relaxation, diuresis, and natriuresis. Clearance of BNP, which has a half-life of approximately

Figure 1. Biochemical changes during trials off extracorporeal life support (ECLS) in patients with good and poor short-term outcomes. A good short-term outcome was defined as the separation from ECLS after a trial, and a poor short-term outcome was defined as the resumption of ECLS after a trial. A, The arterial–venous oxyhemoglobin saturation difference \(\Delta VdO_2\) was significantly greater in the poor short-term outcome group than the good short-term outcome group. B, The change in lactate during trials is expressed as the ratio of the level during the trial (on-trial) to the level before the trial (pre-trial). The change in lactate during trials was not significantly different between groups. C, The change in B-type natriuretic peptide (BNP) during trials is expressed as the ratio of the level during the trial to the level before the trial. The change in BNP during trials was not significantly different between groups.

Figure 2. Biochemical changes during the final trials off extracorporeal life support (ECLS) in patients with good and poor long-term outcomes. Good and poor long-term outcomes were defined as the absence and presence, respectively, of an unplanned cardiac operation or death with 3 months after the operation. A, The arterial–venous oxyhemoglobin saturation difference \(\Delta VdO_2\) was not different between groups. B, The change in lactate during trials is expressed as the ratio of the level during the trial (on-trial) to the level before the trial (pre-trial). The change in lactate during the final trials off ECLS was not significantly different between groups. C, The change in BNP during trials is expressed as the ratio of the level during the trial to the level before the trial. BNP levels during the final trials off ECLS increased to a greater extent in patients with a poor long-term outcome than patients with a good long-term outcome.
20 minutes, occurs through degradation by an endopeptidase, and by endocytosis through a specific natriuretic peptide receptor, termed NPR-C. Owing to these characteristics, BNP has been successfully used as a biomarker in adult and, to a lesser extent, pediatric cardiac disease.11-15

In fact, alterations in BNP during mechanical myocardial support, including support for pediatric patients after cardiac surgery, have been investigated previously18-20,28 In adult patients with cardiac failure, the use of ventricular assist devices has been shown to decrease BNP levels and gene expression.18,19,28 In a recent study, Huang and colleagues30 studied 15 pediatric patients requiring ECLS for cardiogenic shock. Shock developed after cardiac surgery in 11 of the 15 patients. These investigators did not find an association between BNP levels during the course of ECLS and survival after ECLS. However, they did find that BNP levels on the first and fourth days after separation from ECLS were significantly higher in nonsurvivors than survivors.20 In contrast to the present study, these investigators determined BNP levels on ECLS before weaning, when the heart was fully supported. This support may have confounded any association between BNP and the underlying condition of the myocardium.

Cardiovascular hemodynamics may be significantly altered during mechanical support of the heart, particularly ventricular loading conditions. Thus, we believed that BNP determinations off ECLS would have the greatest potential clinical utility. However, several factors obviate the use of absolute BNP values in pediatric patients after cardiac surgery. Perhaps most important, the available data on BNP in patients with congenital cardiac defects suggest that BNP levels vary substantially depending on the specific defect.29-32 Furthermore, BNP levels are altered after operations with CPB in a manner that also appears in part dependent on the underlying defect, and BNP levels vary with age with levels highest at birth, followed by a fall in the first 2 weeks after birth to levels generally below that of adults.32-34 Therefore, we chose to assess the degree of change in BNP during a trial to control for variability in baseline BNP levels between patients. Although we expected BNP levels to increase in all trials off ECLS, owing to increased cardiac volume loading, we hypothesized that the magnitude of the change would differ significantly between patients with persistent myocardial dysfunction and patients with adequate recovery. In fact, when all trials off ECLS were analyzed together, mean BNP levels did increase (from 581 pg/mL ± 382 to 777 pg/mL ± 637; P < .05); however, quite surprisingly, in all patients with a good long-term outcome, BNP levels actually decreased during trials off ECLS. The mechanism for this decrease is unclear and warrants further study.

Clinically, the recovery of adequate cardiovascular function is the key requirement to separate patients from ECLS. Therefore, accurate clinical assessments of cardiovascular function during weaning from ECLS are critical. Our data exemplify the difficulties inherent in these assessments. During trials off ECLS, we found significant increases in lactate, inotrope score, central venous pressure, heart rate, BNP, and the AVdO₂. In addition, we found significant decreases in mean arterial pressure and central venous oxygen saturations. Importantly, we did not find an association between the duration of the trials with any of the biochemical or hemodynamic indices. However, only the AVdO₂ correlated with the ability to separate from ECLS after a trial. We did not use a defined protocol to guide decisions about separating patients from ECLS, and thus it is in fact more accurate to state that the AVdO₂ was correlated with the decision to separate patients from ECLS, whereas the other factors were not. That is, patients not separated from ECLS after a trial may have survived separation. Furthermore, because the cardiac team was not blinded to any of the hemodynamic or biochemical data, other than BNP, in reaching a decision to separate from ECLS we likely considered some or all of these factors. Nevertheless, assessments of the AVdO₂ are known to improve the management of pediatric patients after cardiac surgery, and thus our data suggest that this may also apply to the subset of these patients that requires ECLS postoperatively.35

Despite an increase in the use of ECLS for pediatric patients after cardiac surgery, mortality has remained at approximately 60% to 70%.5,9,36 In our study, 3 (30%) patients died within 3 months of ECLS and an additional patient died within 6 months. Previous factors that have been reported to predict death in pediatric patients on ECLS include age, gender, lactate, type of cardiac defect, inotrope score, renal failure, hepatic failure, multisystem organ failure, duration of ECLS, sepsis, arrhythmia before ECLS, and the duration of mechanical ventilation before ECLS.1,3,5,36,37 None of these associations was identified in the current study, which may relate to its small sample size. However, in our study only the change in BNP during the final trial off ECLS, predicted poor long-term outcomes.

Two patients required an unplanned operation within 3 months of ECLS. One patient (patient 1, Table 1) required pulmonary artery banding for excessive pulmonary blood flow owing to left–to–right shunting, and the second patient (patient 6, Table 1) required a Ross–Konno procedure and aortic arch augmentation for aortic stenosis, aortic obstruction, and aortic insufficiency. All patients had transesophageal echocardiography performed at least once while receiving ECLS. In addition, echocardiography was performed during 6 of the 9 final trials off ECLS. The present study design does not allow comparisons between risk stratification based on echocardiography and BNP levels. However, after echocardiography was performed, patients who ulti-
mately died or required an unplanned operation were separated from ECLS. Although echocardiography can precisely identify the cardiac anatomy and function at the time of the study, assessing the long-term impact of a finding (eg, left-to-right shunting) requires subjective judgments. Our data suggest that BNP may provide additional information that might strengthen these judgments.

The management of pediatric patients requiring ECLS after cardiac surgery is daunting. Unfortunately, few studies have put forth potential tools to aid in the management of these patients. This pilot study, although preliminary owing to its sample size and heterogeneous population, suggests that BNP determinations during trials off ECLS allow for risk stratification. The availability of this information before removal of the ECLS cannulas could allow for important decisions to be made at a time when full mechanical support can be readily re-established or increased. If BNP levels increase during a trial off ECLS, physicians might consider re-evaluation of potential surgical options or consider whether the patient would be a transplant candidate. Further studies are warranted to confirm and define a role for BNP in this vulnerable patient population.

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