Developmental outcome in newborn infants treated for acute respiratory failure with extracorporeal membrane oxygenation: present experience

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Objective: To describe the later health status of newborn infants who received extracorporeal membrane oxygenation (ECMO) for acute respiratory failure in the era after the UK ECMO trial.

Design: Prospective follow up study of newborn infants who received ECMO at a single centre between January 1997 and January 2001.

Setting: Departments of ECMO and Paediatric Intensive Care, University Hospitals of Leicester.

Patients: All babies who received ECMO within 14 days of birth.

Interventions: Neurodevelopment screening using the schedule for growing skills-II (SGS-II) assessment tool.

Main outcome measures: Survival at 12 months of age by disease and functional development at follow up.

Results: A total of 145 neonates received ECMO for treatment of respiratory failure. Of these, 108 (75%) were alive at 1 year of age. There were no deaths in children treated for respiratory failure secondary to meconium aspiration syndrome (73/145). Ninety three (86% of survivors) infants attended a follow up visit at 11–19 months postnatal age. Eighty two were classed as normal, seven as having ‘impairment’, and four as having ‘severe disability’.

Conclusions: Most newborn infants with acute respiratory failure treated with ECMO will have a normal neurodevelopmental screening assessment at 11–19 months of postnatal age. There is no evidence to suggest that changes in neonatal practice since the UK ECMO trial have led to changes in outcome of infants undergoing ECMO therapy.
disability’’, whereas children with overall DQ scores of 50–69% predicted, or a locomotor DQ of less than 70% predicted, or were receiving regular anticonvulsants, or needed tube feeding, or required oxygen were concluded to have “impairment”. Children with an overall DQ greater than 70% predicted were classed as “normal”. Comparisons of the present data with UK ECMO trial data were made using a $\chi^2$ test for dichotomous data. $p < 0.05$ was considered significant.

RESULTS

Between January 1997 and January 2001, 152 newborn infants aged less than 14 days received ECMO therapy in our unit (fig 1). Seven infants with heart lesions were excluded. Between January 1997 and January 2001 inclusive at Glenfield Hospital, Leicester. PICU, Paediatric intensive care unit.

Twelve month survival for the cohort was 75%. Although no deaths were recorded in infants with MAS, 36 infants died in hospital and one child with congenital diaphragmatic hernia (CDH) died after discharge. A third of non-survivors had CDH. Six other babies had untreated respiratory conditions. Figure 1 and table 2 give the details.

Tables 3 and 4 and fig 2 give the health status of the 93/108 children assessed. The mean overall DQ score for the group was 105. Mean group scores for individual skill domains varied between 92 (cognitive) and 119 (locomotor). Data for active and passive posture domains are not shown, as nearly all children received maximum scores in these skills. Eighty two children were classed as normal, seven as having impairment, and four as having severe disability. No infant was blind, totally deaf, or receiving oxygen therapy. All four children with severe disability had overall and locomotor DQ scores of less than 50 and additional health problems such as squint and tube feeding. One had trisomy 21. Four of seven children with impairment had DQ scores <70 in the locomotor skill domain. One of them also required a hearing aid. The three other children with impairment were receiving supplemental feeding. Figure 2 is a scatter plot showing individual locomotor skill domain DQ scores. Four children with DQ scores in the normal range had squints. Although the status of the 15 infants who did not receive follow up is uncertain, contact with families and correspondence from referring units suggest that none of these children have severe health problems.

![Figure 1](https://www.archdischild.com)

**Figure 1** A flow diagram detailing the outcome of neonates referred for and treated with extracorporeal membrane oxygenation (ECMO) between 1997 and 2001 inclusive at Glenfield Hospital, Leicester. PICU, Paediatric intensive care unit.
Outcomes in neonates referred for ECMO

DISCUSSION

The main findings of this report are that survival for newborn infants with severe acute respiratory failure treated with ECMO is excellent and that most survivors are normal on developmental screening at 11–19 months. However, data from other follow up programmes, including that for the UK collaborative ECMO trial, indicate that a full picture regarding the health status of these children will not emerge until later.4

Survival (75%) to 1 year of age for newborns receiving ECMO in the study period is in line with other ECMO programmes and better than reported by the UK collaborative ECMO trial group.1–3 Infants who died could be divided into a group with congenitally abnormal lungs and a group with normal lungs at birth. Idiopathic persistent pulmonary hypertension of the newborn (PPHN) and sepsis were the only two diagnostic categories in the latter. The group with congenitally abnormal lungs included infants with CDH, alveolar capillary dysplasia, and hereditary surfactant protein B deficiency. All infants with CDH were referred for ECMO therapy before surgical repair. Only a few could be weaned from ECMO support on to mechanical ventilators despite early surgical repair of the hernia and a prolonged (>2 weeks) ECMO run. The disappointing survival for infants with CDH (3/15) is indicative of the severe lung dysplasia present in some children with this disorder, and it is our view that outcome for this subgroup of infants with CDH is unlikely to improve in the absence of an effective “lung growth” strategy, with or without ECMO.

Survival for infants with MAS was 100%, a finding that has been reported by other ECMO programmes.10–12 These observations show that ECMO is a very effective treatment for MAS. Moreover, the data show that zero or near zero mortality is an achievable goal for this disorder. One reason for the excellent survival statistics for MAS may be a general improvement in the care of infants with hypoxaemic respiratory failure. For example, compared with the UK ECMO trial, present day infants were referred earlier in the course of their illness and were more likely to have received treatments known to improve oxygenation, such as surfactant, inO, and HFOV. It is therefore possible that these treatments, in conjunction with early institution of ECMO, resulted in present day infants having less prolonged and/or profound episodes of hypoxaemia-ischaemia than infants in the UK ECMO trial. The net result is the excellent survival figures for MAS. It is noteworthy that, despite the greater use of surfactant, inO, and HFOV before ECMO, there was no significant difference in the time of referral for ECMO. This is also encouraging as there had been concern that a strategy of ECMO avoidance, where possible, might result in late referral of a number of infants. The only evidence of such an effect is the 21% of babies whose referral still occurs after 48 hours.

Comparison of the current outcome data with UK ECMO trial data2 shows more children in the severely disabled category in the present day cohort (even after accounting for the child with trisomy 21 who would have been excluded from the UK ECMO trial), but fewer children with impairment. There are several possible explanations for these differences. Firstly, the better survival rate for the present day cohort compared with the UK ECMO trial equates to about 10 more infants being alive at follow up. These “extra” survivors may have been those most likely to have disability. Although it could be argued that compared with the UK ECMO trial, infants in the present era were less sick (lower OIs) and had a shorter period of postnatal hypoxaemia-ischaemia (earlier referral), neither the time to ECMO referral nor a lower OI correlate with better outcomes.1–4 Differences in assessment methodology may also explain the disparity in developmental outcomes. More specifically, as SGS-II is a screening tool designed to detect definite “normality” and definite “abnormality”, it has little sensitivity for detecting subtle abnormalities. Thus DQ scores above 100 do not indicate above average development, and few children are likely to have DQ scores in the “impairment” range. A third possible explanation is that the missing data on 15 infants have distorted the pattern of results. Finally, the age of assessment may also have distorted the findings, as the UK ECMO trial data show that it is important that follow up continues at least until school age.

<table>
<thead>
<tr>
<th>Skill domain</th>
<th>Table 3</th>
<th>Development quotients (DQs) for eight of the 10 skill categories in the schedule for growing skills-II (SGS-II) developmental assessment tool for infants who received follow up after extracorporeal membrane oxygenation (ECMO) at Glenfield Hospital</th>
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<tbody>
<tr>
<td>Overall</td>
<td>105 (29)</td>
<td>Values are mean (SD) (n = 93).</td>
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<tr>
<td>By type of ECMO</td>
<td></td>
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<tr>
<td>VenOAerial</td>
<td>110 (30)</td>
<td></td>
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<tr>
<td>Venovenous</td>
<td>105 (22)</td>
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<tr>
<th>Cognitive</th>
<th>Speech and language</th>
<th>Hearing and language</th>
<th>Manipulative</th>
<th>Visual</th>
<th>Locomotor</th>
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<tr>
<td>92 (31)</td>
<td>97 (26)</td>
<td>101 (25)</td>
<td>109 (21)</td>
<td>100 (21)</td>
<td>119 (36)</td>
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<tr>
<th>Self care social</th>
<th>Interactive social</th>
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<td>108 (34)</td>
<td>112 (36)</td>
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<th>Tables 4</th>
<th>Developmental outcome for infants who received follow up (n = 93) by disease and type of extracorporeal membrane oxygenation (ECMO) run</th>
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<tr>
<td>All infants</td>
<td>93</td>
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<td>CDH</td>
<td>3</td>
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<tr>
<td>MAS</td>
<td>65</td>
</tr>
<tr>
<td>Idiopathic PPHN</td>
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<tr>
<td>Sepsis</td>
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<tr>
<td>Venovenous</td>
<td>81</td>
</tr>
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<td>Venoarterial</td>
<td>11</td>
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<td>UK ECMO trial</td>
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<tr>
<th>Table 5</th>
<th>Developmental outcome for infants in the UK ECMO trial are listed for comparative purposes. Numbers in parentheses are percentages. CDH, Congenital diaphragmatic hernia; MAS, meconium aspiration syndrome; PPHN, persistent pulmonary hypertension of the newborn.</th>
</tr>
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SUMMARY AND CONCLUSIONS

In the current era, we have found that most term and near term newborn infants with hypoxaemic respiratory failure treated with ECMO have normal neurodevelopment. The zero mortality for MAS indicates that "normal physical and neurodevelopmental outcome" is a realistic aim for all neonates with severe respiratory failure, but survivors are at high risk of adverse neurodevelopment.

A strategy of early institution of ECMO therapy is likely to result in the best outcomes for newborns with hypoxaemic respiratory failure. However, at present, there are few markers that predict which infants need ECMO therapy and which can be safely managed with non-ECMO therapies. At present, our in house criteria for newborns with acute respiratory failure include an OI of >25, systemic hypotension requiring treatment with moderate to large doses of inotropes, the presence of air leaks, and cardiorespiratory instability. These criteria may require modification as more outcome data for specific treatments become available. Hence, as recommended by the American Academy of Paediatrics,14 we would encourage the collection and publication of data on short and long term outcomes for infants with severe acute respiratory failure on a centre and treatment specific basis. We acknowledge the care provided by all the staff of the Paediatric Intensive Care Unit, Glenfield Hospital, Leicester to babies referred to the ECMO service.

What is already known on this topic

- ECMO is highly effective at improving survival in neonates with severe respiratory failure, but survivors are at high risk of adverse neurodevelopment
- Alternative means of respiratory support are available for use before ECMO, potentially altering the pattern of ECMO referrals and results

What this study adds

- New treatments are being used before referral; there is no evidence that this is delaying the time that otherwise eligible infants will receive ECMO therapy
- There is no evidence of an adverse impact on neurodevelopmental outcome during the second year of life

REFERENCES


IMAGES IN NEONATAL MEDICINE

Neonate with staphylococcal scalded skin syndrome

This boy was born after an uncomplicated pregnancy and delivery. On the 5th day of life, a small blister was seen on the right buttock and thigh. During the course of a few hours, erythema developed on the trunk and blisters in the napkin area. The patient was admitted to our burn centre, and the diagnosis of staphylococcal scalded skin syndrome (SSSS) was made. His vital functions and temperature remained normal. The skin was reddened and painful, with 50% of the total body area covered by blisters. The Nikolsky sign (separation of the superficial skin from the deeper layers on application of light pressure) was positive. The mucous membranes of the mouth and anus were unaffected. During the next few hours the blisters became more extensive, until they involved 90% of the total body area (fig 1).

Skin cultures showed a growth of Staphylococcus aureus, which produces exfoliative endotoxin B.1 Treatment included administration of intravenous antibiotics and fluid.2 The fluid amount was adapted to compensate for evaporation through the large wound surface. Normal body temperature was maintained with extra heat in an incubator, and pain was relieved with intravenous morphine and rectal acetaminophen. The wound areas were covered with a polyurethane film (Omniderm). This semipermeable wound cover reduces water permeability and thereby fluid loss. It also creates a favourable environment for epithelialisation. On the third day after admission, epithelialisation began and was complete on the 5th day. The patient was discharged home with intact skin, without scars, seven days after admission (fig 2).

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Competing interests: none declared

REFERENCES

Figure 1 The patient a few hours after the diagnosis of staphylococcal scalded skin syndrome, when 90% of the total body area was blistered. The child’s parents have consented to the publication of this picture.

Figure 2 The patient seven days after admission, with intact skin and no scars. The child’s parents have consented to the publication of this picture.