Serial Determinations of Blood Lactate in Respiratory Distress Syndrome

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Serial determinations of blood lactate in respiratory distress syndrome. Blood lactate was measured 4-hourly in 21 newborn babies with respiratory distress syndrome, of whom 13 survived and 8 died. In general, lactate levels were higher in babies who died than in survivors, but there were inconsistencies which were uninterpretable if only a single estimation were made in a baby. Analysis of serial determinations showed that all patients in whom the lactate level never exceeded 35 mg/100 ml survived, and babies with high but falling values also did well. Only those who had rising lactate values, even if initially normal, died.

In most cases a high or normal Pao\textsubscript{2} was associated with normal or decreasing lactate levels, but babies with Pao\textsubscript{2} below 60 mmHg had often also normal or decreasing lactate levels. Some babies had high and increasing lactate levels despite having normal Pao\textsubscript{2}.

In order to use lactate levels for prognosis in respiratory distress syndrome (RDS) serial determinations are required.

It is concluded that there may be a wide range of hypoxaemia without oxygen deficit in body tissues, so that it is not possible to define a 'lower acceptable Pao\textsubscript{2}', which will define adequate tissue oxygenation.

The respiratory distress syndrome (hyaline membrane disease) is an example of a condition where the baby has difficulty in taking up adequate amounts of oxygen and in transporting oxygen to sites where it is needed, so that oxygen therapy is an important and much discussed problem in the care of such babies. Because of risks of producing retrolental fibroplasia or lung damage oxygen therapy must be carefully controlled. The practice at the Hammersmith Hospital is to try to maintain Pao\textsubscript{2} between 60 and 90 mmHg in aortic blood. The optimum Pao\textsubscript{2} is very difficult to establish; there is some agreement as to the maximum safe limits (Boston, Geller, and Smith, 1966; Robertson et al., 1968; Scopes, 1970), but there are no well-established values for the minimum Pao\textsubscript{2} levels to achieve acceptable tissue oxygenation (Brumley, 1971; Auld, 1971). The minimum Pao\textsubscript{2} levels have been experimentally studied by Takano (1968) in dogs and by Vaughan and Eitzman (1969) in puppies. Both authors agree that there is evidence of anaerobic metabolism when the Pao\textsubscript{2} is below 25–35 mmHg. But even in their normal healthy animals the Pao\textsubscript{2} levels at which evidence of anaerobiosis appeared varied greatly.

Tissue oxygenation depends not only on arterial Pao\textsubscript{2} but also on other factors as haemoglobin concentration, cardiac output, degree of shunts, and peripheral circulation. Clinically, babies are seen who are apparently well and others who are very ill, though having the same Pao\textsubscript{2}. The most rational evaluation of oxygen therapy in the treatment of respiratory distress syndrome (RDS) would be to measure mitochondrial oxygen uptake which is clearly not practicable. Even the measurement and interpretation of tissue Pao\textsubscript{2} (Rodger et al., 1968) has considerable difficulties. However, measurement of blood lactate might provide an indirect evaluation of tissue oxygenation, because tissue hypoxia produces anaerobic metabolism with a large increase in lactate concentration. This has been shown experimentally by Neill et al. (1969) and Huckabee (1961); in adult circulatory failure by Weil and Affi (1970); and in intrapartum asphyxia by Derom (1965), Yu et al. (1965), and Daniel, Adamsom, and James (1966).

Another important problem in RDS is that of establishing an early prognosis in order to evaluate different therapeutic regimens. The Pao\textsubscript{2} achieved

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by a newborn baby breathing 100% O₂ (N₂ washout) seems the best prognostic sign available (Boston et al., 1966; Roberton et al., 1968). Other factors such as pH, serum K, P<P₀₂, and buffer base have been evaluated and scored (Stahlman et al., 1967). Initial blood lactate levels as a prognostic index in RDS were studied by Stahlman et al. (1967) and regarded as not adding more information than pH alone. Matthieu, Gautier, and Prod’hom (1971) studied blood lactate levels before 3 hours of age and found a significant difference between the values of those babies who died and those who survived from RDS, claiming that lactate measurements were of some prognostic value.

There are reports of lactate levels in cord blood related to birth asphyxia (Payne and Acharya, 1965; Marx and Greene, 1965; Kornacki et al., 1967), in normal babies (Koch and Wendel, 1968; Yu et al., 1965), and of single determinations in babies with RDS (Stahlman et al., 1967; Matthieu et al., 1971) where the levels were related to the prognosis. There is one report in abstract form (Wang et al., 1963) of serial determinations of lactic acid in RDS.

Our purpose was to study serially blood lactate levels in newborn babies with RDS, firstly in order to seek evidence of adequate tissue oxygenation in babies who would inevitably have a low P<sub>O₂</sub> at some stage of their illness, and secondly to assess the prognostic value of these measurements.

**Clinical material**

The patients studied were those admitted to the Neonatal Unit at the Hammersmith Hospital with clinical manifestations of RDS defined as having two out of the following three signs at 4 hours of age: grunting, recession, and a respiratory rate over 60 per minute. All had umbilical catheters inserted, for monitoring pH and blood gas levels, and chest x-rays. They were treated according to the usual regimen of the Unit (Scopes, 1971).

In all, 21 patients were studied, 7 born at the Hammersmith Hospital and 14 referred from other hospitals. 13 patients survived and 8 died. Their gestational age, birthweight, sex, type of delivery, result of hyperoxia test, age at the onset of study, and necropsy findings are shown in Table I.

**TABLE I**

Details of 21 Cases of Respiratory Distress Syndrome (RDS)

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Gestational Age (wk)</th>
<th>Birthweight (g)</th>
<th>Sex</th>
<th>Delivery</th>
<th>Birth Asphyxia</th>
<th>N₂ Washout P&lt;sub&gt;O₂&lt;/sub&gt; (mmHg)</th>
<th>pH</th>
<th>Age at Onset of Study (hr)</th>
<th>Other Diagnosis or Necropsy</th>
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<tr>
<td>Survived</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>30</td>
<td>1190</td>
<td>F</td>
<td>Vertex</td>
<td>Yes</td>
<td>143</td>
<td>7.18</td>
<td>9</td>
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<td>33</td>
<td>1990</td>
<td>M</td>
<td>Vertex</td>
<td>No</td>
<td>121</td>
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<td>13</td>
<td>Persistent ductus arteriosus</td>
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<tr>
<td>5</td>
<td>27</td>
<td>1060</td>
<td>M</td>
<td>Vertex</td>
<td>No</td>
<td>153</td>
<td>7.26</td>
<td>6</td>
<td>Subarachnoid haemorrhage; apnoeic attacks</td>
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<tr>
<td>7</td>
<td>34</td>
<td>1980</td>
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<td>Vertex</td>
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<td>7.40</td>
<td>4</td>
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<tr>
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<td>Vertex</td>
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<td>26</td>
<td>800</td>
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<td>2600</td>
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<td>Vertex</td>
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<td>7.08</td>
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<td>2120</td>
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<td>1760</td>
<td>F</td>
<td>Breech</td>
<td>No</td>
<td>32</td>
<td>7.08</td>
<td>4</td>
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<tr>
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<td>35</td>
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<td>LSCS</td>
<td>Yes</td>
<td>102</td>
<td>7.10</td>
<td>6</td>
<td>Subarachnoid haemorrhage; IVH; HMD</td>
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</tbody>
</table>

IVH = intraventricular haemorrhage. HMD = hyaline membrane disease. LSCS = lower segment caesarean section.
Fig. 1.—Scattergram of lactate measurements related to postnatal age of patients. ▲, babies who died; ●, babies who survived.

Methods

In all patients routine clinical observations were recorded and blood sugar, arterial pH, arterial blood gases, and blood lactate were measured 4-hourly. Blood sugar was estimated using Dextrostix (Chantler, Baum, and Norman, 1967). Arterial pH and blood gases were measured with an IL pH and blood gas analyser 313. When 0·3 ml blood was withdrawn from the umbilical arterial catheter for pH and blood gases determination, an extra 0·2 ml was taken for lactate measurement. This sample was immediately deproteinized with 0·2 ml 0·6 N perchloric acid and the supernatant kept at 4 °C. Lactate was measured within the next 24 to 48 hours using the Boehringer modification of the Hohorst enzymatic method (Hohorst, 1957, 1963). A calibration line was drawn from standards on each day and optical densities read at a wavelength of 366 mµ. The standard deviation of the measurement was ± 0·48 mg/100 ml at a concentration of 33 mg/100 ml and ± 0·94 mg/100 ml at a concentration of 112 mg/100 ml.

In these 21 patients there were 150 determinations of lactate levels, with a minimum of 2 and a maximum of 14 in any single baby.

Results

Fig. 1 shows all individual estimations of lactate in 21 patients with RDS, indicating whether the baby survived or died. It is clear that in general terms very high lactate levels were found in babies who subsequently died, and levels less than 35 mg/100 ml in those who survived. There were however obvious exceptions which could be more correctly interpreted when serial measurements had been made. Babies with a falling level of blood lactate, even if high, were likely to survive, and babies with a rising level were likely to die.

Every baby whose blood lactate level remained below 35 mg/100 ml survived. Fig. 2 plots the serial measurements of all babies who at any stage had a blood lactate level higher than 35 mg/100 ml. This group includes all babies who died and three survivors. They are best described individually.

Cases 1, 9, and 11 were preterm babies with severe RDS, low P\textsubscript{a}O\textsubscript{2} in N\textsubscript{2} washout and severe acidosis, in whom there was no clinical response to
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FIG. 2.—Lactate curves in babies who at any stage had blood lactate higher than 35 mg/100 ml. The number at the beginning of each curve is the case number, see Table I. (△-△) 8 cases who died; (.....) 3 cases who survived.

treatment, and in whom it was impossible to achieve an acceptable P\textsubscript{a}O\textsubscript{2}. In all these three babies blood lactate levels had an extremely steeply rising curve, even though initially less than 35 mg/100 ml in Cases 9 and 11. All died before 14 hours of age and had hyaline membrane disease (HMD) and intracranial haemorrhage at necropsy.

Cases 8 and 16 were preterm babies with severe birth asphyxia who had as their main clinical problems apnoeic attacks and haemorrhages from different sites. Both were extremely acidic and required the use of a ventilator because of repeated apnoeic episodes. P\textsubscript{a}O\textsubscript{2} in Case 8 was between 52 and 67 mmHg. Their lactate levels were very high initially and continued to rise until their deaths at 32 and 20 hours of age, respectively. At necropsy there were haemorrhages in the respiratory tract and lungs in Case 8, intracranial bleeding in both, and in Case 16 also gastrointestinal haemorrhage. Though both patients had clinical RDS, hyaline membranes were not found at necropsy in Case 8.

Cases 4 and 6 had rising blood lactate levels though in each there was a period of temporary fall. Case 4 was a preterm and small-for-dates baby with severe RDS; his lactate concentration was below 35 mg/100 ml initially and increased progressively up to 96 mg/100 ml at 23 hours. At 26 hours he received a blood transfusion and his blood lactate fell to 65 mg/100 ml one hour later; this was not followed by clinical improvement and the baby died at 30 hours of age. Case 6 was a preterm baby with severe RDS who had very low blood gases during the hyperoxia test, was severely acidic, and required the use of a ventilator during the whole period of study. His P\textsubscript{a}O\textsubscript{2}, initially low (range 14 to 49 mmHg), became satisfactory (range 57 to 82 mmHg) between 23 and 36 hours age and deteriorated thereafter until his death at 47 hours. His blood lactate value was initially very high, decreased somewhat during the period of acceptable P\textsubscript{a}O\textsubscript{2} and then increased again. The main necropsy finding was HMD in these two cases.
Case 10 was a preterm and small-for-dates baby who had multiple congenital abnormalities—hypoplastic lungs, thoracic kyphosis with defect in vertebral bodies T6 and 7, large enterogenous cyst in the abdomen, small enterogenous cyst in posterior mediastinum with extension into the vertebral canal, and undescended testes. Besides all these problems he had clinical RDS and died at 23 hours of age. His lactate level was initially 51 mg/100 ml, fell to 16 mg, and then rose progressively up to 40 mg. His Pao2 was over 100 mmHg during the period of decreasing lactate and between 36 and 40 mmHg after this. This was the only baby in the study where lactate level fell from above to below 35 mg/100 ml and who died. Necropsy examination showed the congenital abnormalities but no HMD.

Cases 2, 17, and 18 were survivors who at some stage had a lactate concentration higher than 35 mg/100 ml. Case 18 had an initial lactate of 42 mg at 4 hours age, having had severe birth asphyxia and apnoeic episodes before this age. Lactate decreased after this initial measurement, despite severe respiratory distress that required the use of a ventilator and high concentrations of O2. Even with this therapy his Pao2 was often below 60 mmHg. Case 2 had an increase in blood lactate between 9 and 13 hours of age reaching a concentration of 61 mg/100 ml after three prolonged apnoeic attacks. After this temporary increase, levels of lactate fell and he recovered. Case 17 was exceptional, as he had the highest initial lactate level (139 mg/100 ml) in the series and yet survived. Before this measurement made at 6 hours of age the baby had had 'terminal birth asphyxia' (Gupta and Tizard, 1967), two prolonged apnoeic episodes, and at age 4 hours a cardiac arrest; he required artificial ventilation from 9 hours to 10 days of age and finally recovered. Lactate levels fell continuously after this exceptionally high initial value.

In most cases a high or normal Pao2 coincided with lactate levels that were either normal or decreasing. When however Pao2 was less than 60 mmHg, that is within the range generally quoted as meaning unacceptably hypoxaemic, lactate levels were often also normal or decreasing. All but one child (Case 6) in whom this set of circumstances arose recovered and survived. Thus, either falling or normal lactate levels gave evidence of adequate tissue oxygenation despite the low Pao2.

For example Case 3 (Fig. 3) was a very ill baby who despite inspired oxygen concentration of near 100% had Pao2 between 25 and 40 mmHg in the first 60 hours. None the less the lactate levels remained stable between 14 and 21 mg/100 ml and the child survived. In Case 17 (Fig. 3) there was a dramatic fall in lactate from 139 to 21 mg/100 ml during a period in which the Pao2 was mostly below 60 mmHg, and she recovered.

The opposite of these circumstances was finding a rising lactate level despite acceptable Pao2 measurements. In Case 16 (Fig. 2) high and rising lactate levels occurred despite Pao2 of over 100 mmHg and the child rapidly succumbed. In Case 8 (Fig. 2) Pao2 measurements were marginally low (52 to 67 mmHg) but lactate levels rose sharply and again the baby died.

Discussion

Normal values of blood lactate were not determined in this study as we did not consider it justifiable to insert catheters in healthy babies, but these have been established by others. Koch and Wendel (1968) established lactate values for normal newborn babies, studying systematically 79 infants from birth to 7 days of age, using umbilical arterial blood and using the same method we used. Their normal values are shown in Table II. Lactate levels in healthy preterm infants were studied by Yu et al. (1965) and did not differ significantly from those of term infants. It is obvious from these two studies that measurements made before 3 hours of age would reflect what had happened at birth rather than what the situation might be at the time of sampling. This is in keeping with the fact that the mean life of lactate in newborn babies is longer than 60 minutes (Ciampolini and Franchini, 1966).

Blood lactate concentration may be increased by causes other than tissue hypoxia, mainly exercise, hypocapnia, or liver dysfunction. When tissue hypoxia occurs, the lactate dehydrogenase system shifts towards a more reduced state, and lactic
acid production exceeds that of pyruvic acid, producing an increase in the lactate/pyruvate ratio (L/P). This led Huckabee (1958a) to propose the calculation of excess lactate (XL) as an unequivocal sign of anaerobic metabolism. However, the value of XL as an invariable indicator of tissue hypoxia has been questioned (Olson, 1963; Alpert, 1965). L/P and XL also vary with hypocapnia (Cain, 1968) and lactate levels alone correlate with signs of tissue oxygen deficit in the same way as XL or L/P (Daniel et al., 1966; Weil and Afifi, 1970). XL is a measurement requiring previous determinations of lactate and pyruvate levels in the non-hypoxic state and is not feasible in clinical situations. Pyruvate determinations require undesirably large samples of blood for a systematic study in the newborn. Therefore, we did not measure pyruvic acid and we interpret the large increases in blood lactate found as qualitative but not quantitative evidence of tissue hypoxia. This interpretation has been established experimentally (Huckabee, 1958b; Takano, 1970), and in several clinical conditions (Weil and Afifi, 1970), including some cases of RDS (Wang et al., 1963; Payne and Acharya, 1965).

The other major causes of increased lactate are exercise, hypocapnia, and liver dysfunction. Muscular activity of patients studied in this series was observed and recorded and no correlation was found between high lactate level and hyperactivity; on the contrary, those babies with high lactate concentrations were usually the very sick and inactive babies. The rise in lactate that occurs with hypocapnia is small compared with the levels described here, and we found no correlation between high lactate levels and the occasionally low Pco₂. We did no liver function tests in these babies; it is probable that a baby who has tissue hypoxia has a degree of liver dysfunction which would in part explain the lactate rise observed, an added reason for not drawing quantitative conclusions from these results.

The relation between pH and lactate concentration is difficult to establish. A high blood lactate
is usually accompanied by a low pH. Alkali therapy corrects pH but does not necessarily change the lactate concentration. Therefore there might be, as we found on several occasions, high lactate concentration coincident with a normal pH, or a normal lactate in the presence of low pH. Increasing lactate levels were not accompanied by decreasing pH curves for the latter was modified by alkali therapy, and therefore pH curves did not have the prognostic value that lactate curves had. Nevertheless lactate is one of the determinant factors of a low pH in metabolic acidosis, and therefore much but not all the information obtainable from its measurements is also obtained by monitoring pH, especially in situations where alkali therapy is not needed.

No attempt was made in this study to standardize the age at which the initial measurement of lactate was made, as umbilical arterial catheters were inserted for clinical reasons at the age patients entered our unit. It was however clear that initial lactate levels were as a whole lower in survivors from RDS than in those who died, but variations were too large in each group to establish any conclusions. Stahman et al. (1967) found the same in their series and left lactate out of their proposed score for prognostication in RDS. Gautier (1971) has recently reported high lactate measurements to be correlated with low Apgar score and bad prognosis in RDS. These measurements were done before 3 hours of age, when they probably say more about the past than about the future.

Our results show that a single measurement of blood lactate is of little prognostic value, only serial measurements to show either an increasing or a decreasing curve being helpful in establishing a good or a bad prognosis in RDS. An early prognosis cannot therefore be established on this basis. The value of N2 washout as an early and practical test was once more corroborated by our results, as can be seen in Table I.

These results give support to the concept that there may be a wide range of hypoxaemia not associated with deficiency of oxygen in body tissues. This makes it impossible to define a 'minimum acceptable P\textsubscript{a}O\textsubscript{2}', for a different figure would apply for each individual baby and for each stage in his disease. Thus a high and increasing lactate concentration gives better evidence of inadequate tissue oxygenation than any arbitrary level of 'low' P\textsubscript{a}O\textsubscript{2}. A baby with a P\textsubscript{a}O\textsubscript{2} below 60 mmHg and a normal or decreasing lactate level has no evidence of lack of oxygen and should not be subjected to the known risks of increasing his ambient oxygen, to achieve an ‘acceptable’ P\textsubscript{a}O\textsubscript{2} of more than 60 mmHg.

In the reverse situation, there may be a failure in oxygen transport to the tissues though the P\textsubscript{a}O\textsubscript{2} is within the accepted normal range. This situation was evident in two of our patients. Therefore, a rising blood lactate level, even in the presence of normal P\textsubscript{a}O\textsubscript{2}, means that the baby needs further support, either by increasing ambient inspired oxygen using continuous airway pressure or by supporting circulation.

We conclude that in RDS systematic measurements of blood lactate evaluate tissue oxygenation better than do P\textsubscript{a}O\textsubscript{2} measurements. Lactate levels may provide useful information for therapy, indicating whether a baby needs to have his inspired oxygen increased or his circulation supported, and they give some prognostic information.

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