Clinical biochemistry of the neonatal period: immaturity, hypoxia, and metabolic disease

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SUMMARY This review attempts to provide practical information on common problems in the laboratory medicine of newborn infants and also considers unresolved problems in achieving neonatal diagnoses. A common cause of upset in the newborn—intrapartum asphyxia—can now be positively diagnosed. This leaves a small group whom it is necessary to investigate because they may have metabolic disease. The initial investigation of metabolic disease at the district general hospital should be limited to the commoner conditions.

This review aims to present everyday practice of clinical biochemistry in the neonatal period and to consider how cases which need more detailed consideration by a laboratory medicine team might be selected. The neonatal period is from birth to four weeks of age. Newborn infants require special care, generally for immaturity. For those few whose condition does not improve with time, a system of investigation can be devised. Fig 1 shows the basis for such an approach.

For a baby born after a 40 week gestation the most likely cause for upset is hypoxia during birth; the non-specific but extremely variable clinical changes produced by hypoxia decrease with time after birth. On discharge from hospital at about 7–10 days no obvious ill effects may be detectable, but the infant is on total bed rest and constantly nursed. Later in life increased demands on the central nervous system made by reading and writing or the necessity of secreting more growth hormone may show up defects. Infections, like congenital malformations and metabolic disease, tend to get worse. The pattern of events over time is of major diagnostic importance. In a preterm infant time also allows the clusters of adaptive changes, described as maturation, to take place and in all infants permits the healing of birth trauma. The success of many of the supportive procedures like increased oxygen concentration and even ventilation in preterm infants depends on such maturation.

When the infant does not mature normally then a wide variety of conditions are caused. Anatomical defects can be detected clinically or by various imaging techniques, failing which, they may eventually be shown at necropsy. Congenital disease of the heart, gut, or renal tract is often clinically apparent. Such clinically obvious disease is generally detected using a systematic clinical approach. Serious defects which are not obvious require conscious search with a specific screening procedure. Congenital dislocation of the hip and phenylketonuria are examples which are not fatal but later produce obvious and serious “disease”.

Much has been written about the neonatal period. Books selected or written by local clinical staff are obviously the best guide to local clinical practice, but many of them are long. There are also a few extremely practical guides to clinical practice, but little help is available for those in laboratory medicine. The physiological principles entailed in the care of the newborn, especially preterm infants, have been reviewed in addition to the relevant clinical biochemistry procedures. Both these reviews focus on the principal workload of special care baby units that is generated by the problems of immaturity: “normal” values for conventional clinical biochemical results in newborn infants have been compiled.

The available reference values in the fetus and the newborn need to be interpreted with caution as these are often generated at or after delivery when there might be undiagnosed asphyxia present. Asphyxia can raise fetal inorganic phosphate and sulphate concentrations, thus generating an apparent fetal-maternal gradient which may not have existed during intrauterine life. Such a gradient has already been noted for sulphate concentration at delivery. The inclusion of values from infants with undiagnosed intrapartum asphyxia may also be responsible for the high upper limits of reference ranges for newborn in-
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Fig 1 Changes seen in the patterns of disease intensity with age. Improvement with age occurs after trauma and in the common effects of immaturity and hypoxia. Deterioration with age occurs in the effects of infection, congenital malformations, and inherited metabolic disease.

Infants. The values used at Northwick Park Hospital, for example, are uric acid concentration 90–580 μmol/l, serum aspartate transferase activity 30–75 IU/l, creatine kinase 40–500 IU/l and γ glutamyl transferase activity 12–200 IU/l; the urate is a by-product of the increased purine output associated with adenosine triphosphate (ATP) depletion and the leakage of enzymes from cells depleted of ATP causes the high upper limits for the enzyme activities.

This paper aims to help those workers in laboratory medicine in hospitals which deliver between 1000 and 4000 babies each year. Some responsibilities for newborn infants will probably be a part of their routine work and a relatively large part of their emergency work. Units with smaller numbers of deliveries will probably refer their perinatal problems, preferably in utero, or failing that, in ambulances.

Attempts are made later in this review to integrate the every day care of the newborn with the problems of diagnosis of metabolic disease. The small range of diagnoses made in special care baby units contrasts with the long lists of possibilities and the success of some screening programmes. Antenatal diagnosis must also be mentioned as it is one factor that affects the incidence of open neural tube defects and chromosomal anomalies.

Routine laboratory care of the newborn

General considerations

Most newborn infants require no special care. About 10% are admitted to a special or intensive care unit from which many emergency requests can arise. Newborn infants change rapidly, and specimens for analysis should be transported rapidly and biochemistry reports should be available twice daily to reduce the errors and interruptions inherent in telephone reports. There is also a place for coordinated ward laboratories used by medical and nursing staff to carry out simple tests that are urgently needed.

Local experience is used extensively as the basis of this review. Such personal practice, however, may not always be suitable for a different environment. Furthermore, accepted practice may not always have been adequately evaluated and may reflect the past rather than the present. Accepted practices like the extensive use of measurements of sodium and potassium concentrations are difficult to evaluate and it is even more difficult to change their use. If practice is not to become frozen into existing rituals it is advisable to evaluate the uses and limitations of all procedures, including those performed by nurses. Some laboratories have stopped measuring chloride without ill effects, and this indicates that large changes can be made safely. Safe reductions in emergency requests can also be achieved by medical laboratory staff.

Antenatal care

Biochemical care of the newborn begins in the antenatal ward: routine clinical biochemistry is of little benefit. The obstetrician wants to know the fetal age, size, presentation, and fetal risk. Clinical biochemistry can be fittingly used to measure fetal risk.

Pulmonary maturity

In preterm infants pulmonary immaturity is a major problem. Treatment by prolonged ventilation and
increased oxygen concentrations is expensive and not entirely safe. In a pregnancy of less than 36 weeks the assessment of pulmonary maturity by amniotic fluid phospholipid concentration—either by thin layer chromatography relative to the internal standard of endogenous sphingomyelin concentration or by more quantitative methods—provides information on the risk of respiratory distress. It is true that such investigations are not necessary if the baby must be delivered for obstetric reasons. Evaluation of obstetric risk, however, is usually imprecise, and delays in delivery are often justifiable because clinical experience shows that the stimulation of pulmonary development for 48 hours before delivery by glucocorticosteroids is effective.

**Rhesus isoimmunisation**

The extent of Rhesus isoimmunisation can be estimated by spectrophotometric estimations of bilirubin in filtered amniotic fluid. These results offer a precise guide to treatment, including intrauterine transfusion.

**Oestrogen analyses and time intervals sampled by different procedures**

Measurements of fetal size by ultrasound scanning may detect fetal growth retardation over an interval of one or more weeks. The biochemical function of the growth retarded or at risk fetoplacental unit can be estimated from oestrogen excretion over 24 hours. If three or more results show a consistent fall to concentrations below normal the risk to the fetus is high. If oestrogen excretion is consistently <20 µmol/24 hours but with normal fetal growth, steroid sulphatase deficiency is probable and presents little or no risk except at birth. Measurement of blood oestrogen concentrations can be informative over a shorter time interval of about three hours; monitoring of fetal heart rate gives an instant record. The usefulness of oestrogen analyses in diagnosing specific defects of the fetal hypothalamo-pituitary-adrenocortical-placental system and the low usefulness of these results in indicating general metabolic failure has been discussed, as have the differences between blood and urinary oestrogens.

To avoid spurious conflicts a coordinated approach is required for the allocation of resources between methods which provide different information or study different time intervals. The routine application of any of the above monitoring methods to an unselected population can provide little or no benefit. A combined systematic approach using a preliminary screening method is difficult to evaluate as developments, especially in ultrasound scanning techniques, are changing this area. It is often difficult to control the quality of ultrasound observations and the interpretation of fetal heart rate patterns in routine clinical work.

**Rationalisation**

Requests for analyses of amniotic fluid for phospholipids and bilirubin, as well as urinary oestrogen excretion, are decreasing to a point where it is difficult for any district hospital laboratory to produce reliable results. As these analyses can be useful, regional centres for these measurements should now be agreed.

**Clinical biochemistry in special and intensive care neonatal units**

Much of this can be done in the ward by clinical and nursing staff using side room equipment.

Small volumes of blood are all that can be spared in infants weighing about 1 kg. For the central laboratory the provision of a service using very small quantities of blood is difficult. “Top up” transfusions are to be avoided because of their attendant risks of viral infection, especially hepatitis. Junior medical staff are probably the most practical and skilled source of blood samples, using indwelling catheters, peripheral veins, or capillaries. In this way disturbance can be minimised and the volume taken can be easily monitored. It is necessary for laboratory staff to spend considerable time and effort in using as much as possible of the samples. It is surprisingly simple to scale down a modern routine method so that only micro-litre samples will be needed. Small specimen tubes sampled carefully using thin pipette tips and more staff time are all that may be needed to provide an improved “micro” service.

**Anoxia**

The first need of a newborn infant is for oxygen. The monitoring of ventilatory efficiency in preterm infants with pulmonary immaturity or in infants with asphyxial lung damage is efficiently and continuously performed by transcutaneous pO₂ monitoring with periodic checks of blood acid base and gas variables—pH, pO₂, pCO₂, base excess, and standard bicarbonate.

**Salt and water balance and renal function**

The second major need is for water and minerals, although some regimens for the treatment of intrapartum hypoxic damage restrict fluid intake “to reduce cerebral oedema”. In addition to measuring urine volume, nurses use refractometers and dipstick methods to measure the specific gravity of urine. The dipstick methods can also detect renal damage manifest as proteinuria and haematuria. Accepted clinical practice is to use serum urea concentration as an index of renal failure but experience in paediatric nephrology suggests that measurement of serum creatinine is better. Renal failure in the first two days of life is a feature of the damage caused by intrapartum asphyxia.
and is seen as oliguria or anuria.

When oral feeding is difficult or impossible, or in cases with excessive gut or even renal losses, monitoring serum sodium and potassium concentrations is useful especially if combined with measurements of serum creatinine. Renal immaturity in preterm infants can also result in excessive loss of renal sodium which can be treated with supplements. Diagnostically confusing amounts of lactose and galactose as well as amino acids are also found in urine from preterm infants.

**Hypoglycaemia** The third major need is for an energy source. Hypoglycaemia can occur in the newborn, particularly in preterm infants who have inadequate glycogen stores and an inability to perform gluconeogenesis, and in the infants of diabetic mothers. Hypoglycaemic infants may have used their glycogen stores during intrapartum asphyxia. Regular ward monitoring of blood glucose with dipsticks or tapes is standard practice. The role of the laboratory is to confirm the dipstick results for which a fluoride blood sample is advisable.19

**Hypocalcaemia** Another example of physiological immaturity is hypocalcaemia. This can also follow intrapartum hypoxia, possible due to defective renal hydroxylation of 25-hydroxycholecalciferol. Experienced clinicians will recognise slight “jitteriness” as a good guide to this condition. Measurement of serum or plasma calcium is therefore often requested and the results of treatment are good.20 Only if there is no response to a calcium supplement for about three days is it justifiable to measure serum magnesium. Hypomagnesaemia complicating hypocalcaemia has been reported and both respond well to supplementary magnesium. Hypomagnesaemia alone is virtually unknown: emergency magnesium estimations cannot therefore be justified. It is rare for any infants to require further investigation for any of the diseases associated with incorrect calcium metabolism despite clear and even recurrent evidence of neonatal hypocalcaemia. Such investigations are only practicable in a metabolically stable and mature infant.

**Jaundice** The immaturity of detoxication systems is seen in the “physiological” jaundice of the newborn, in which rapid breakdown of fetal haemoglobin and its further metabolism can not be adequately performed by the liver. Unconjugated bilirubin concentration rises to a peak at 3–5 days of age, and this may be measured by various side room bilirubinometers that simply measure the yellow colour of plasma or serum, or even skin. These are adequate at bilirubin concentrations of less than 200 μmol/l and can monitor the effects of phototherapy as well as hepatic maturation, thus saving the laboratory a large load of emergency and routine bilirubin estimations. These instruments are less reliable, however, at high concentrations, and laboratory methods are needed for clinical decisions such as exchange transfusion.17

Jaundice is the clearest example of a condition which is, initially, nearly always “physiological immaturity.” If it persists and especially if a pronounced rise in conjugated bilirubin occurs investigation is required.

The “problem oriented” case record has been used in the systematic investigation of neonatal jaundice by Mathew and Wharton.21 Their chart showing serial plots of plasma bilirubin against age in days, with an action limit for phototherapy at about 250 μmol/l and that for exchange transfusion at about 350 μmol/l, is similar to charts used in many units. Their careful documentation of experience with infants born in 1976 (n = 1988) suggested that their action limits may need changing and that the selection of tests which are going to yield useful diagnoses for their population should be changed. Possible ABO incompatibility was common (34 cases) but unsuspected infection was rare (four cases), and transient and questionable hypothyroidism occurred in four preterm infants. Congenital hyperbilirubinaemia was found in three cases with probable neonatal “hepatitis”. There were two infants with chronic illness, one with a “pulmonary syndrome”, and another had propionic academia that was unresponsive to biotin. Neither glucose-6-phosphate dehydrogenase deficiency nor galactosaemia were detected in any infant. A systematic review of the investigation of jaundice will produce an effective local protocol.

**Breast milk jaundice** The incidence of breast milk jaundice is between 1 and 2%.22 As the milk of a species is adapted to the nutrition of that species23 breast feeding is encouraged. Breast feeding has again become more popular in some areas of the United Kingdom.24 Detailed consideration of breast milk jaundice is thus justified. The mean rise in bilirubin concentration in breast fed infants may be no larger than in formula fed term but not preterm infants.25 In 1–2% of mother and infant pairs, however, there is an iatrogenic effect of human but not cow’s milk.

The often cited association between 5β-pregnane-3α,20β-diol in human milk and jaundice came from New York. It has been confirmed that 5β-pregnane-3α,20β-diol was present in lactating mothers with jaundiced infants26 but disappeared after the end of lactation. These mothers probably had some abnormality of the steroid metabolism of the lactating mammary gland. The predominantly 5α structure for steroids in milk and studies of mammary gland metabolism in vitro suggest that the breast changes the pattern of steroids in milk from that in plasma.27 28 Large doses of steroids can raise unconjugated bilirubin concentrations in plasma in the new-
born, and milk causing jaundice was inhibitory in vitro to glucuronol transferase activity. Such inhibition, however, was not explained by the effects of 5β-pregnane-3α,20β-diol.

General consideration of bilirubin metabolism suggested that a wide variety of medium molecular weight compounds could be responsible for breast milk jaundice. Many steroids inhibit steroid glucuronol transferase activity with similar micromolar inhibition constants and might therefore inhibit bilirubin detoxication by these enzymes. A survey method, suitable for the identification of peaks using gas chromatography-mass spectrometry (GC-MS), was applied to milk. The method showed a low concentration of pregnanediones from pregnanediols and pregnanolones in human milk, with no increases found in milk causing breast milk jaundice. Another sensitive GC-MS method did not detect pregnanediols in human milk.

Many studies on the transmission of steroid analogues, other drugs, and insecticides in milk have shown low concentrations. Any such compound found in milk has been of a low and usually biologically unimportant dose; the exceptions have been listed in advice to prescribers.

### Table 1: Steroids in human milk from mothers with jaundiced (n = 9) and normal (n = 6) infants

<table>
<thead>
<tr>
<th>Normal infants*</th>
<th>Jaundiced infants*</th>
<th>Quantity in final fraction (µg/100ml milk)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (days)</td>
<td>Steroid detected in GC survey</td>
<td></td>
</tr>
<tr>
<td></td>
<td>5α androstane-3,17-dione</td>
<td>0.57</td>
</tr>
<tr>
<td>1</td>
<td>5β androstane-3,17-dione</td>
<td>0.08</td>
</tr>
<tr>
<td>2</td>
<td>5α androstane-3,17-dione</td>
<td>0.12 and 0.12</td>
</tr>
<tr>
<td>3</td>
<td>5α androstane-3,17-dione</td>
<td>0.12</td>
</tr>
<tr>
<td>4</td>
<td>5α androstane-3,17-dione</td>
<td>0.29</td>
</tr>
<tr>
<td></td>
<td>5α androstane-3,17-dione</td>
<td>0.46</td>
</tr>
<tr>
<td>5</td>
<td>5β androstane-3,17-dione</td>
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</tr>
<tr>
<td>6</td>
<td>5α androstane-3,17-dione</td>
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</tr>
<tr>
<td>7</td>
<td>5α androstane-3,17-dione</td>
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</tr>
<tr>
<td>8</td>
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<td>0.17</td>
</tr>
<tr>
<td>9</td>
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<td>0.21</td>
</tr>
<tr>
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</tr>
<tr>
<td>14</td>
<td>5α androstane-3,17-dione</td>
<td>0.29</td>
</tr>
</tbody>
</table>

*Milk from three mothers with jaundiced infants and from two mothers with normal infants contained no detectable steroids. 10-50 µg/100ml = 14 nmol/l. Data from JAB Darling, the Department of Paediatric Biochemistry, Royal Hospital for Sick Children, Edinburgh.
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It was clear from gas chromatography (figs 1 and 2)\(^6\) and from thin layer chromatography evidence that human milk contained more lipid soluble medium molecular weight compounds than cows's milk. There was, however, no component on thin layer chromatography or gas chromatography that was associated with jaundice in the infants receiving the milk (Darling and Harkness, unpublished observations, 1975). There were, therefore, more potential inhibitors in human milk\(^3\) but no one compound and certainly no pregnanediol could account for the jaundice. The evidence suggests that some breast milk jaundice is due to cumulative inhibitory effects from small amounts of many compounds on bilirubin detoxication and excretion. Many different compounds are detoxicated by the same mechanisms as bilirubin; specifically, the glucuronyl transferases are "receptors" of wide specificity.\(^3\)

Care should be exercised in checking the specificity of analytical methods used on milk; it has not proved possible to confirm reports of high vitamin D sulphate concentrations in milk based on a colourimetric method.\(^4\)

The large numbers of compounds in human milk may arise from a highly varied diet compared with that of dairy cattle. Specifically, it seems pointless to analyse milk for pregnanediols.

MAINTENANCE OF WARD AND SIDE ROOM METHODS

Mention has been made of the techniques whereby ward blood glucose monitoring can be checked and its quality maintained. The laboratory can also usefully train medical and nursing staff and check their performance.\(^17\) A simple technique to reduce anomalous results is for the performance of the actual bottle of sticks used to be checked. A poor batch may result from changes that have occurred during storage, or from operator errors. Quality control for the bilirubinometer can in practice depend on comparison of results from ward and laboratory from the same patient at about the same time.\(^17\) The increasing use of dry stick methods is clinically desirable and will increase.

A simple and robust instrument like a refractometer used for measurements of urine specific gravity or a bilirubinometer in a side room may only need to be cleaned after use. A stock of commonly needed spares must also be kept. These requirements can be efficiently handled at ward level.

Complex machinery like existing blood gas assemblies requires regular quality control and service checks by laboratory staff. The machine must therefore be in continuous use to compensate for the work needed to maintain it in good order, otherwise, it should be placed in the general laboratory medicine area.

A careful and agreed definition of responsibilities in the area of ward and side room methods, with a continuing communication network among designated staff can avoid unnecessary conflicts. In the NHS this is difficult except at consultant level. The recommended guidelines on the performance of chemical pathology assays outside the laboratory\(^18\) provides a useful guide to harmonious working practices.

NEWER DEVELOPMENTS: DRUG MONITORING

Expensive developments in laboratory methods are an ongoing problem for those assigning resources. After enthusiasts show that their procedure can be useful there is a need for critical assessment of cost and of benefit and a clear description of where and how such methods can solve or prevent clinical problems.

Drug monitoring is potentially one of the most expensive areas of laboratory medicine and may be of particular importance in the newborn period because the neonate lacks the necessary detoxication systems. The vulnerability of developing systems is also high: examples are the sensitivity of multiplying mitochondria to chloramphenicol\(^44\) and of myelin to hexachlorophene.\(^45\)

The prescription of antibiotics on suspicion or risk of infection is common in special care baby units, although less than 40% of neonates have any definite evidence of infection. Gentamicin is often used which requires regular monitoring to avoid toxicity. Another widely used drug, which is regularly monitored in the newborn, is theophylline, a stimulant to the respiratory centre used in preterm infants who have periodic apnoea. Apnoea is usually well tolerated and occurs about five to seven days after delivery when the baby has excreted the caffeine derived from its mother. Methylated xanthines have very much longer half lives in newborn infants than in adults. Although good cost benefit assessments are not available, the use of theophylline in the newborn has been reviewed.\(^46\)

The use of drugs only on specific indications is advisable but this is difficult in practice. The intensive efforts made to save newborn infants, especially preterm infants, require the administration of many potent drugs. To avoid unnecessary side effects the safest course would be to improve the standards of diagnosis, an area in which laboratory medicine has a major role.

THE CHRONIC CARE OF PRETERM INFANTS

Once the problems of pulmonary immaturity are over, the care of preterm infants focuses on nutrition, which makes great demands on mothers and nurses.
Harkness

Fig 3 Convergence of processes to produce any final product allows homeostasis to mask a defect five stages away from its initial cause (line 1). Unless variables at or near the primary defect are measured, overlapping values will be found (lines 3 and 4). Lines 1–5 represent values of different variables in a chain of causal events. The values of the different variables are plotted as standard deviations of the normal population from the mean. The relative amounts of the normal "population" encompassed by 1, 2, and 3 standard deviations, 68.3, 95.5, and 99.7% respectively, are indicated below the normal distribution.

The excess loss of sodium caused by the failure of the kidney to respond to aldosterone can be detected by regular estimations of serum sodium and potassium. Calcium deficiency can also occur and cause rickets; radiological diagnosis is unequivocal but not soon enough. Gross increases of alkaline phosphatase activity in plasma can be shown but the wide variation in normal values still renders interpretation of early changes difficult.

Diagnosis in the newborn

The need for transient "homeostatic" care conceals many diagnostic problems. A widely adopted solution is to conduct diagnostic investigations only if the state of the baby does not improve with time (fig 1). Many major and eventually fatal anomalies compatible with live birth also allow four to five days of extrauterine life. In the first two to three days of life the assumption that the major problems are transient products of immaturity and do not require a diagnosis is probably justifiable, but severe infections and intrapartum hypoxia must be diagnosed. At present both infections and intrapartum hypoxia are diagnosed on non-specific signs similar to those listed in table 2 and fig 2, indicating metabolic disease.

Diagnosis and Measurement of Hypoxic Damage

Intrapartum asphyxia is common, being severe enough to cause clinical problems in 1% of all births. Infection can usually be successfully treated by antibiotics. Anatomical abnormalities are detectable by detailed clinical examination and imaging techniques. The positive laboratory diagnosis of intrapartum hypoxia would help obstetricians and allow a small group of children to be selected who show signs attributable to hypoxia, but in whom inherited or congenital metabolic disease are considered more likely because they show no specific evidence of hypoxic damage. It is probable that in this group are found many of the permanent neurological defects previously ascribed to asphyxia. A variety of evidence
suggests that we may be overdiagnosing asphyxia as a cause of long term handicap.

The diagnostic use of non-specific variables and the related overlapping distribution of variables Why are we continually trying to measure risk and even achieve diagnoses using overlapping distributions of variables? Because we use commonly disturbed variables which are the end products of many rarer, initial defects. Many different initial defects converge to produce a final effect like growth failure. Fig 3 gives an explanation of the overlapping distribution of variables, which represents values of several associated variables.

Most variables used in monitoring show poor discrimination between normal and abnormal (fig 3). This arises because the initial defects causing the disturbances are several stages removed, and the different linking stages are all subject to homeostatic mechanisms that tend to restore values to normal. In other words, the further away a measurement is made from the site of a primary defect, the greater the overlap produced by homeostasis between normal and abnormal distributions of values. Good discrimination is probably the reason for the consistent progress that is being made in the diagnosis and treatment of rare conditions. This progress contrasts with the negative or equivocal results of trials of methods detecting final common products, like intrauterine growth failure.

Table 2 Features suggesting metabolic disease

| Clinical features and findings of routine investigations: |
|-------------------------------|-----------------|
| Persistent vomiting           | Poor feeding, failure to thrive |
| Neurological abnormalities    | Depressed consciousness from somnolence and lethargy to coma, convulsions, hyperventilation and hyperventilation, absence or loss of functions expected in the newborn, such as ability to suck and swallow |

| Liver abnormalities:          | Acute liver failure, jaundice, bleeding, depressed conscious state, large or small liver depending on stage of disease, Pronounced or prolonged jaundice, Hepatomegaly with or without splenomegaly, Metabolic acidosis, Abnormal smell, Unexplained hypoglycaemia, Neutropenia, thrombocytopenia, particularly with metabolic acidosis, Hyponatraemia/hyperkalaemia with or without ambiguous genitalia, Miscellaneous, and less commonly, diarrhoea, hypothermia, abnormal hair, cardiomegaly and heart failure, catacar 

2 Important supporting factors: 

| Family history of similar incidents or unexplained death |
| Familial consanguinity |
| Onset of illness corresponds to a change of feeding practice, or the timing of symptoms related to feeds |
| Symptoms improve on glucose/saline fluids |
| Onset of illness associated with infection, fasting, or surgery |


Fig 4 Amniotic fluid and cord blood plasma concentrations of hypoxanthine, xanthine, and urate from mother and child monitored during labour, showing criteria of fetal distress, fetal heart rate abnormalities, and meconium stained amniotic fluid.

Because several initial or primary defects converge to produce changes in growth, for example, growth failure is more common than any one individual primary or initial defect and is therefore a tempting target. Clinically, it is rarely practicable to measure each different primary or initial defect, but the routine detection of intrauterine growth failure by ultrasound examination is only useful in a high risk group because of the overlapping distributions of values. Clinical assessment Clinical diagnosis and assessment of damage due to intrapartum hypoxia depend on non-specific signs, many of which are ranked to produce a cumulative Apgar score. The Apgar score, however, is too insensitive to measure the efficacy of obstetric procedures which are now beginning to lower the stillbirth rate. If obstetric intervention is successful infants are born with high Apgar scores after showing signs of intrauterine hypoxia, such as slowing of the fetal heart following uterine contractions and meconium staining of amniotic fluid.

pH and blood gas variables The use of pH and blood gas variables in cord blood and even from fetal scalp to assess hypoxia are limited in their application. The period of time sampled is limited to about 30 minutes or less. The widespread evidence of a lack of correlation of cord blood pH or hypoxanthine (fig 4) or newborn pH after resuscitation with fetal damage is largely due to the rapid changes of pH and other variables in blood. As labour generally lasts more than four hours a more "cumulative" record of hypoxic damage is needed. Moreover, most changes
Hypoxanthine
Carbon
dioxide
pCO2

CELL

Hypoxanthine

pH Lactate

ATP

Glycolysis

Glucose

Oxygen

Organic acids

Nucleus

Mitochondrion

Fig 5 Energy generation in the cell by the mitochondria and cytosol using oxygen and glucose to produce the energy currency of the cell ATP. During depletion ATP is broken down to hypoxanthine. Glycolysis is reflected by lactate concentrations which can affect blood pH. The sites of defects in ATP generation are indicated by asterisks and include hypoxia, hypoglycaemia, and ischaemia, as well as intracellular defects in glycolysis, organic acid metabolism, other mitochondrial functions and a variety of genetic defects in nuclear DNA.

in pH are due to metabolic acidosis and reflect lactate concentrations which are produced by anaerobic glycolysis. Anaerobic glycolysis can maintain ATP concentrations in many tissues, especially in the fetus. Measurement of pH does not reflect a fall in ATP concentrations, which is the critical initial event in metabolic damage caused by hypoxia.

ATP METABOLISM IN HYPOXIA

New methods of assessment

It is possible to define asphyxia at the molecular level as a failure of energy supply sufficient to cause cellular damage. As the energy currency of cells the purine nucleotide ATP the definition is, “ATP depletion sufficient to cause irreversible or partially reversible cellular damage,” (which also covers damage by hypoglycaemia). ATP is confined to living cells but ATP depletion can be measured in extracellular fluids because such depletion increases output of hypoxanthine, the central intermediate product in purine metabolism, from tissues into blood, urine, amniotic fluid and cerebrospinal fluid. The mechanisms and quantitative associations between concentrations of hypoxanthine and tissue energy supply have been established.55-60 Fig 5 represents the information provided by lactate on the rate of glycolysis and by hypoxanthine on the extent of ATP depletion.

The relation between ATP depletion and subsequent tissue damage is more difficult to establish.61,62 Unexpectedly, there is a close positive correlation between neurological damage, assessed clinically, and persistently excessive hypoxanthine excretion occurring 24 hours after ATP depletion and lasting for several days.54,63

Anatomical origin of the changes in ATP metabolites and methods of sampling for clinical analysis

Evidence from studies of progesterone and human

Fig 6 Plasma concentrations at elective operative deliveries of hypoxanthine, xanthine, and uridine in samples from maternal artery, peripheral vein, and uterine vein, as well as fetal umbilical vein and artery. Individual patients and their infants are shown by the symbols ○, ●, and □.
placental lactogen production shows that the fetus, and especially its brain, is more easily damaged than the placenta. Systematic studies on this area have been made in sheep. In man the vulnerability of the fetus was shown from studies of concentrations of ATP metabolites and the related pyrimidine nucleoside uridine present in maternal artery, peripheral and uterine veins, and fetal umbilical vein and artery samples at operative delivery. Fig 6 shows the results from elective operative deliveries. There is no steep gradient, suggesting that there is no excess ATP breakdown in the normal fetus. In contrast, emergency operative deliveries without evidence of fetal distress show a trend for higher hypoxanthine concentrations in the uterine vein (fig 7), consistent with the active myometrium consuming ATP and releasing hypoxanthine. In cases with two or more clinical criteria of fetal distress, however, the increases in metabolites in uterine vein and umbilical artery compared with umbilical vein (fig 8) are consistent with a sharp increase in hypoxanthine output due to ATP breakdown by the fetus but not the placenta during fetal distress. The increases in uridine concentrations show a somewhat more consistent pattern and are compatible with the hypoxanthine concentrations (figs 6–8).

The slower breakdown of ATP in the placenta compared with that of the fetus has been confirmed in the rat and guinea pig. Their tissue concentrations of hypoxanthine, xanthine, and uridine are also significantly negatively correlated with the relative ATP concentrations expressed as energy charges. Similar correlations also exist in human placenta.

The changes occurring in amniotic fluid over a considerable time period, possibly of about 12 to 24 hours, however (fig 4), and later in neonatal urine give more diagnostic information than the rapid changes in blood and tissues. Fig 4 shows an abnormally high concentration of hypoxanthine in amniotic fluid in association with meconium staining, and fetal heart rate patterns justifying emergency caesarean section. The umbilical artery and vein concentrations and those in maternal peripheral vein returned to normal. The study in fig 4 thus illustrates the weak correlation of the rapidly responsive cord blood variables with antecedent state. It is also clear that obstetric intervention removes evidence of asphyxia. Amniotic fluid concentrations of hypoxanthine may thus become a means of validating and measuring the common obstetric diagnosis of fetal distress.

Although the placenta can be a source of tissue for enzymological confirmation of a diagnosis, there is no good biochemical evidence that placental insufficiency exists apart from arylsulphatase C or steroid sulphatase deficiency. Many routine procedures which assess the placenta are therefore of dubious value. In the more vulnerable fetus cerebrospinal fluid can be analysed to obtain a neurological prognosis from the hypoxanthine concentration. Hypoxanthine output in urine and amniotic fluid are a record of hypoxia over about the previous 24–48 hours. Analyses of these fluids in varying combinations may therefore help diagnose and assess the effects of intrapartum asphyxia and cardiac and ventilatory arrest.
Tissue damage after ATP depletion
A wealth of clinical and experimental evidence from drowning \(^70\) shows that it is the duration of anoxia relative to the irreducible energy requirements associated with an organ’s temperature which determine the reversibility of the changes. After 10 minutes of extrauterine anoxia the prognosis is poor. In perfused rat hearts the duration of low ATP concentrations seems to be related to the ability of the tissue to recover.\(^71\) If glycolysis succeeds in maintaining ATP concentrations then no damage will occur; a case of intrapartum hypoxia has been reported with a pH of 6.6 and a bicarbonate concentration of 12.9 mmol/l at one hour with good recovery.\(^72\)

Evidence from exercise and muscle physiology suggests that there are coordinated muscle and neural mechanisms of fatigue that normally limit the energy drain and fall in ATP concentrations, so that they are easily reversible.\(^73\) Forcing the heart to work during anoxia increases the amount of damage. It could be that the adverse effect of pronounced hyperglycaemia on anoxic brain damage\(^74\) is caused by the increased effort required to metabolise high glucose concentrations.

Clinically, diabetes mellitus predisposes to intrapartum asphyxia. High fructose concentrations deplete liver adenosine phosphates,\(^75\) which reconciles the experimental observations of Myers et al\(^74\) with a variety of well established clinical observations. A normal glucose concentration seems to preserve brain function in the fetus, and in extrauterine life hypoglycaemia damages the brain. Glycogen stores are rapidly used by tissues to maintain ATP concentrations and their absence in the fetus leads to damage.\(^64\)

Recovery from damage
Vogt and Farber\(^76\) showed that the reversibility of change was a reliable prognostic index in the rat kidney. This also seems to be true for the rat heart.\(^71\) It is the degree of function in EEG records that can be maintained after the episode which relates to prognosis.\(^70\)\(^77\)

Diagnosis of metabolic disease at the district general hospital
Two approaches have been used in the diagnosis of inherited metabolic diseases—stepwise analysis and population screening. Both have limitations and neither is of value in conditions which are rapidly fatal.

**Conventional clinical investigation**
The classic stepwise approach of clinical medicine, which progressively limits the possibilities, can be highly effective. In addition to the infant’s history, a family history may suggest an inherited disease. Repeated obstetric problems suggest a maternal cause, which is often ill defined. Diagnostic success for the stepwise approach involving several groups and reference laboratories probably depends on a severe progressive illness in which an infant stays alive only for months or years.

Cystic fibrosis can be diagnosed in this way. It is the

---

**Fig 8** Plasma concentrations at emergency operative deliveries of hypoxanthine, xanthine, and uridine in samples from maternal artery, peripheral vein, and uterine vein, as well as fetal umbilical vein and artery. The infants showed two or more criteria of fetal distress and are shown by the symbols ○, ●, □ and ■.
Clinical biochemistry of the neonatal period: immaturity, hypoxia, and metabolic disease

Table 3  Approximate incidence of some metabolic diseases in populations of western European origin

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Approximate No of diagnoses per 100 000 births</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cystic fibrosis</td>
<td>50</td>
</tr>
<tr>
<td>Neonatal hypothyroidism*</td>
<td>25</td>
</tr>
<tr>
<td>Other endocrine disorders†</td>
<td>30</td>
</tr>
<tr>
<td>Phenylketonuria</td>
<td>5–25</td>
</tr>
<tr>
<td>(Histidinemia)</td>
<td>6</td>
</tr>
<tr>
<td>Homocystinuria</td>
<td>1</td>
</tr>
<tr>
<td>Maple syrup urine disease*</td>
<td>1</td>
</tr>
<tr>
<td>Urea cycle defects*</td>
<td>4</td>
</tr>
<tr>
<td>Cystinuria</td>
<td>8</td>
</tr>
<tr>
<td>Non-ketotic hyperglycaemia*</td>
<td>1</td>
</tr>
<tr>
<td>Organic acidoses*</td>
<td>3</td>
</tr>
<tr>
<td>Galactosaemia</td>
<td>2</td>
</tr>
<tr>
<td>Other carbohydrate disorders*</td>
<td>9</td>
</tr>
<tr>
<td>Mucopolysaccharidoses</td>
<td>8</td>
</tr>
<tr>
<td>Other storage disorders</td>
<td>19</td>
</tr>
</tbody>
</table>

*Can cause acute disorder in the perinatal period.
†Mainly steroid 21-hydroxylase defects and arylsulphatase C or steroid sulphatase defects.

Data modified from Association of Clinical Biochemists Broadsheet, No 237.

most common lethal genetic disease of caucasian populations. In newborn infants it presents as meconium ileus and the usual pattern of lung disease develops later. The diagnosis depends on the classic sweat test which shows the characteristic abnormally high sodium concentrations. This test combined with those for meconium albumin and pancreatic function requires care and experience with good clinical-laboratory liaison.

The high detection rate for endocrine disease may be due to the early arousal of suspicion by the clinically obvious effects of hormones, like virilisation of external genitalia. Similarly, many infants deficient in adenosine deaminase activity (ADA) manifest in severe combined immunodeficiency, have been diagnosed, although this condition is very rare—about 1000 times rarer than phenylketonuria. An extensive screening programme to detect ADA deficiency in infants in New York was not helpful and was abandoned.

The known incidence of the storage disorders caused by acid hydrolase deficiencies has been estimated at 27 in 100 000 births, which is high (table 3). In older children the protracted and characteristic clinical course of the storage disorders has been associated with many successful diagnoses in large numbers of index cases. As a consequence of successful diagnosis of index cases, antenatal diagnoses have become highly developed.

Table 4  Incidence of diagnosed inherited metabolic disorders over two years in the United Kingdom

<table>
<thead>
<tr>
<th>Group diagnosis</th>
<th>Manchester (n = 110000)</th>
<th>Sheffield (n = 60000)</th>
<th>Bristol (n = 70000)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amino acids</td>
<td>5</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Organic acids</td>
<td>5</td>
<td>2</td>
<td>Not tested</td>
</tr>
<tr>
<td>Urea cycle</td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Galactosaemia</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Data modified from Association of Clinical Biochemists Newsheet No 237.

Therefore seems practical to suggest that once the team is clear that a baby is probably ill, an investigation protocol should be initially restricted to diseases with an incidence of about 1 in 10–20 000. This proposal is in line with current screening practice for phenylketonuria, which has an incidence of 1 in 10 000 and hypothyroidism, which has an incidence of 2 in 10 000. Of the several thousand possible defects, the number which should be considered by a non-specialist can probably be limited to between 10 and 30. Considering the incidence of diagnosable and, as a first priority, treatable diseases in the population in one's care generally yields a small group of diseases in the United Kingdom (tables 3 and 4) and for Switzerland.

The introduction of methods capable of detecting organic acidoses may increase the detection rate of these diseases. Three groups are commonly encountered: methylmalonic acidemias, some of which are B₁₂ responsive; multiple carboxylase deficiencies, which may be biotin responsive; and the generally fatal propionic acidemias. The total incidence may be as high as 1 in 10 000. From this evidence the necessary methods for preliminary diagnosis can be defined.

It might be helpful to indicate a condition which is impracticable to detect usefully by whole population screening. Maple syrup urine disease is too rare and too rapidly fatal for the index case to be saved in most cases. It is practicable, however, to diagnose a second case in an affected family. Newborn infants in whom serious metabolic disease is anticipated can be fed glucose and thoroughly and rapidly investigated. Such a procedure organised from a reference laboratory caring for 75 000 births a year detected only six babies at risk, of whom four were normal. One had massive hepatic necrosis of unknown cause and the other citrullinaemia.

Population screening

Phenylketonuria and congenital hypothyroidism are common enough—1 and 2 in 10 000—respectively, to produce a considerable number of live born irre-
versibly damaged infants who require continued institutional care. These two conditions are not common enough, however, to be an effective part of a paediatrician's clinical response to their very vague and often late clinical signs. The unexpected success of the laboratory screening programmes for phenylketonuria and later for hypothyroidism mitigates against a policy of dismissing all such metabolic disease as too rare to justify diagnostic consideration.

With so many possible diseases it is hardly surprising that such insidious and vague early signs are missed in newborn infants. Reference laboratories often care for patients who have shown early signs (table 2) that were not investigated, but it is impractical to investigate all patients with such signs. A major problem is the selection of a high risk group. This needs positive diagnoses for the common causes of the ill defined clinical signs that provide the initial clues. Before it can be justified to adopt screening procedures for specific disease most centres regard a good cheap overall screening method with a minimum of false positive or negative results and an effective treatment as vital.

Screening programmes also have limitations. Samples are generally taken on the fifth to eighth day of life when rapid changes are taking place. The analytical problems of measuring large numbers of compounds are considerable. In addition, substrate loading by milk feeding is also necessary in galactosaemia. The success of adapting the hypothyroid and steroid 21-hydroxylation deficiency screening programme to the dried blood sample used for phenylalanine analyses suggests that many more compounds could be included if abnormalities were common enough. Various methods are already available.

Histidinaemia can be reduced with an appropriate diet, but a recent careful and extensive prospective study of histidinaemia detected by whole population screening in North America and Japan has shown that there seems to be no associated disease. Previous associations were due to selective investigation of an abnormal group. Histidinaemia is not a "dis-adaptive" phenotype or disease. As well as abnormality being treatable when detected in the second week of life it is also necessary for there to be an associated clinical abnormality. Antenatal treatment for most defects is at the experimental stage.

In summary, the present pattern seems to be firstly, national population screening by central laboratories for common genuinely treatable disorders with an insidious clinically indistinguishable onset and course. The latest candidate for population screening is steroid 21-hydroxylase defects. Secondly, stepwise investigation of chronic, progressive severe disease is needed, in which a conventional clinical approach of:

1. A severely ill infant aged four to five days or more, without gross congenital anatomical defects or infection, probably has one of more than a 1000 rare diseases. The only practicable mechanism for the diagnosis and management of these diseases has been the creation of reference centres. In this review I have indicated how patients should be selected for referral. Widening the criteria for acceptance of patients for detailed investigation of organic acidemias, however, has not increased the number of cases detected. Preliminary local investigation by a well informed paediatric and laboratory team can result in a higher proportion of positive diagnoses.

2. The investigative flow diagram in fig 2 is similar to that agreed by a United Kingdom working party. It was emphasised that the district general hospital should take investigation to the second line; the reference laboratory then takes over at the third and final line. It was also emphasised that the reference laboratory is available for consultation, and to accept material it must have been initially consulted. Many problems, when defined clearly on the telephone or in a summary letter, are soluble with the clinical and biochemical experience of the reference laboratory and require no further investigation. The reference
centre can guide initial investigation and interim
treatment. Many endocrine diseases are amenable to
diagnosis by a team which can recognise the clinical
effects of hormones and systematically test even rare
possibilities.95

If the newborn period alone is considered, published
tables of possible causes of metabolic disease are lengthy. Haan and Danks96 listed 38 conditions;
these tables have been condensed by Holton.97 Con-
sideration of the actual United Kingdom incidences
listed in tables 3 and 4 shows that most of these can be
initially disregarded. The evidence in fig 2 and tables
3 and 4 suggest that the following tests should be
available at a district general hospital.
1 Careful check for sodium loss due to steroid
21-hydroxylation deficiency which has an inci-
dence of about 1 in 10 000.84-89 Urinary concen-
trations of sodium greater than 20 mmol/l should
be regarded as high in a normally fed term infant.
2 An amino acid chromatogram of blood and urine
will show various disorders of amino acid metabo-
lism.
3 Check for persistent metabolic acidosis (at least
plasma bicarbonate) as a reduced concentration is
usually seen in organic acidemia. Simple chro-
matographic methods for organic acids are useful.
4 Possibly check plasma ammonia. This is very high
in urea cycle defects and hepatic failure.
5 Do a test for galactose. This is a reducing sub-
stance that, if shown not to be glucose in the urine
of a mature milk fed infant, indicates galac-
tosaemia.
6 Perform a sweat test for cystic fibrosis.
These tests cover most of the conditions which are
at present recognised and emphasise those which are
treatable. Mucopolysaccharidoses and other storage
disorders do not usually present as acute metabolic
disorders in the newborn.

If there are doubts about the results of these tests
then the reference laboratory should be consulted and
the results checked by those with more experience,
especially in interpreting amino acid patterns. If all
the test results are negative and the infant is still de-
teriorating transfer of the patient to the care of a spe-
cialised clinical and laboratory team is advisable.
Incurable damage due to failure to detect a treatable
condition should thus be avoided.

It is the role of the reference laboratory to reach
systematically one of the many possible diagnos-
es.99 Current evidence suggests that this is an un-
solved problem.100

The use of a conventional 12 channel clinical chem-
istry analyser to generate an “admission” profile at a
paediatric referral centre made only a small con-
tribution to overall care.101 Most conventional clin-
ical chemistry analyses are of limited use in diagnosis
despite their value in the problems of immaturity. Ex-
tensive chromatographic skills can show a variety of
defects—the profiling of organic and amino acids and
of steroids, for example. For confirming diagnoses,
the ability to estimate enzyme activities in tissues gen-
erally on a micro scale may be required by reference
centres, which will also need extensive clinical experi-
ence.

Conclusions

The field of neonatal diagnosis and treatment is con-
tinually developing. The acquisition of diagnostic
skills is shown in the clustering of case reports of new
diseases.

Management of suspected metabolic disease is to
feed the patient only glucose and water for about
12–48 hours, when preliminary results should be
available.82 100 In cases in which death is imminent
the aim should still be to secure a diagnosis. Diagnos-
sis after death of an index case of maple syrup disease
can help the affected family. For such diagnosis Bur-
ton and Nadler102 recommended that measurements
of urine, plasma, and sterile skin kept at 37°C or at
least at room temperature in tissue culture medium or
even 5% glucose in physiological saline should be se-
cured. The preliminary use of an ethyl chloride spray
sterilises skin; alternatively, fascia can be used. Fi-
broblasts grow if obtained up to about 48 hours after
death. If a necropsy is performed a specimen of fresh
liver stored at −20°C is also recommended. If the
necropsy shows grossly affected tissues these should
be retained. Vitreous humour can be taken at
necropsy103 and gives clearer amino acid patterns
than other available extracellular fluids. Every effort
should be made, however, to take specimens of blood,
urine, and even cerebrospinal fluid before death. In
general, analyses of necropsy specimens are extremely
difficult to interpret.97 104 105 In severely ill infants a
high index of suspicion for inherited metabolic dis-
ease and a readiness to do some tests can produce
useful results.106

This article could not have been written without the
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Committee.
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