ambulance service trust involved. By law the trust was liable for employees involved in accidents while on duty, but nevertheless the recourse to obtain compensation was through the usual delays.

As a result of our experiences we have undertaken a comprehensive review of our neonatal transport operations. Initial problems of incubator security have been temporarily addressed with ring bolts set in the floor of ambulances which allow secure anchorage with cargo straps. Incubators are loaded using portable rams. In the long term, in cooperation with our ambulance service, we have redesigned our transport system with a reduced payload around a no lift platform that will be compatible with all types of stretcher. The ambulance service has also undertaken to modify ambulance anchorages to accommodate newer stretcher types with transport incubators on board.

The standards of safety for neonatal transport incubators need to be reviewed. Attention needs to be given to reducing payload, and to a universal mounting platform for loading and securing incubators within ambulances, whether road vehicle or aircraft. The problems of different incubators, ambulances, and stretcher designs can be avoided by using a universal mounting platform, thus each unit needs to be aware of its responsibilities to its staff regarding liability in the event of an accident.

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Unfriendly incubators

Editor.—Fractures in infants of very low birth weight (<1500 g) not related to birth trauma are being recognised with increasing frequency.¹ These fractures are usually associated with bone demineralisation, which is frequently present in very low birthweight infants, and almost two thirds of them involve the extremities.¹ Fractures in these infants can occur as the result of ‘trauma’ during physiotherapy or other procedures such as placement of intravenous lines.¹ ² ³

We have observed four infants with birth weights under 1000 g who sustained traumatic fractures of the extremities (one arm in three cases, one leg in the other), and we believe that the contributing factor was accidental trapping of the extremities by or under the plastic tray on which the mattress of the incubator lies. The fractures, diagnosed radiologically between 4 and 6 weeks of age, were courts with significant dislocation. Two of these babies also had well documented bone demineralisation. Subsequent to these independent observations in Athens and Montreal, we conducted a survey of incubators used in neonatal intensive care units and found that in many (even in some of the latest models) there was sufficient gap between the plastic tray and the incubator wall for an arm or a leg of the baby to slip between the incubator’s wall and the plastic tray (figure). In many instances the nursing staff had tried to cover this gap with rolled sheets. In some incubators the tilting mechanism could not be securely locked at the desired tilted position

![Incubator showing gap between plastic tray and wall allowing arm to be trapped.](image)

and the plastic tray could therefore accidentally fall or jolt on the trapped arm or leg. Injury might also occur if the baby is pulled or turned by a nurse or a doctor unaware that a limb is trapped. The possibility that a fracture may be caused in this manner is enhanced if the bones are significantly demineralised.

The design of an incubator should be such that the plastic tray extends to the wall of the incubator such that any gap is too small for a baby’s extremity to slip through and the tilting mechanism should be securely locked at the desired tilt. Incubators currently in use should be checked for this potential hazard.

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Excessive rates of asphyxia – accedect or fact?

Editor.—It is well known that hypoxic ischaemic encephalopathy (HIE) in neonates is associated with adverse neurological outcomes.¹ ² The incidence of HIE can be a useful indicator of quality of intrapartum care provided to term infants,³ while the severity relates to subsequent neurological outcome.² The Doncaster perinatal service has received much adverse publicity suggesting that their incidence of HIE was excessive (‘Trial of labour’, World in Action, Granada Television, 2 Nov 1992). We analysed data from the Trent Regional Neonatal Survey in order to establish whether these criticisms were valid.

The Trent Regional Neonatal Survey collects data on all high risk babies admitted to perinatal units within the region. All babies greater than 35 weeks gestation admitted because of HIE were included in this analysis. HIE was diagnosed using modified criteria of Levene et al.¹ Grade II HIE was defined as the infant having a history suggestive of asphyxia plus convulsions, while grade III was defined as grade II plus the need for respiratory support. Data was collected by two independent visiting observers. The last three years of complete data were analysed (1 April 1990–30 March 1993).

There were a total of 168 435 live births in 16 perinatal centres (five subregional units and 11 smaller units) and 263 babies with HIE; 141 grade II and 122 grade III (table). The incidence of HIE for the region was 1.56 per 1000 live births (range 0.58–5.1 per 1000 live births). The Doncaster unit (number 6 in the table) had an incidence of 1.53 per 1000 live births – that is, similar to the region as a whole.

It is clear that criticisms levied at this perinatal unit were unfounded and based on anecdotal evidence rather than fact. Data relating to HIE is not routinely collected by most perinatal units, however, we believe such data provide a valuable method of improving perinatal care. Only if the rate of HIE falls outside the ‘normal range’ is concern warranted.

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Diagnosis and management of non-immune hydrops in the newborn

Editor.—We read with interest the article by Stephenson et al on the diagnosis of non-immune hydrops in the newborn¹ and we would like to emphasise that a wide range of inborn errors of metabolism (IMD) have been reported to be associated with non-immune hydrops and in many cases a feasible pathogenetic mechanism can be hypothesised.² As a cause of hydrops they are rare
but collectively they are significant and hence appropriate investigations should be considered. It is particularly important to consider IMD when more than one case of hydrops occurs in the same family or in twins. Hydrops is a history of spontaneous abortion or unexplained deaths in close family members. We would like to describe our protocol for the investigation of fetal and neonatal hydrops for IMD. Investigations are considered when the more common causes of hydrops have been excluded.

From the dead fetus a skin biopsy can be taken for collagen and this should be taken as soon as possible. The cultured cells can be assayed for β-galactosidase (GM1 gangliosidosis, Morquio’s disease type B, galactosialidosis), β-glucuronidase (Sly’s disease), β-glucosidase (Gaucher’s disease), α-neuraminidase (muco- lipidosis I), N-acetylgalactosamine-6-sulphate sulphatase (Morquio’s disease type A), sialic acid (sialic aciduria) and cholesterol esterification (Niemann-Pick C). DNA from the cultured cells can be investigated for expanded trinucleotide repeat sequence found in myotonic dystrophy.

In cases where hydrops has been diagnosed antenatally amniotic fluid can be tested for I-cell disease and this should be taken as soon as possible. Other assays can be undertaken on cultured amniotic fluid cells or chorionic villus tissue (direct or cultured). In the live neonate investigations which can be performed on blood include enzyme kinetic, plaque cell proliferative, and tissue culture investigations. Liver tissue may be of value for liver iron quantitation and possibly also for enzyme assays. If liver is to be taken, it should be frozen immediately and stored below –20°C.

While not all these tests will be indicated on every case of hydrops it is important to consider informed consent of parents as a cause of non-immune hydrops and to collect appropriate specimens for investigations where indicated. Hence discussion with a specialist centre for inherited metabolic disorders is recommended.

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Do maternal β-sympathomimetics influence the development of retinopathy in the premature infant?

Editor.—Recent studies suggest that a number of factors other than the injudicious delivery of oxygen contribute to the development of retinopathy in the premature infant.1,2 One hundred and fifty infants on our neonatal unit who required oxygen treatment for more than 60 days were reviewed retrospectively in order to determine the range of factors contributing to this pathology. All infants requiring oxygen for more than two weeks were examined at two weekly intervals by an ophthalmic surgeon (ES). Fourteen cases of retinopathy greater than stage 3 were identified. These patients were matched for gestation (+1 week), birth weight (±50 g), sex, and ethnic group with infants who had required oxygen for more than 60 days, but did not have retinopathy. The two sets of cases showed no significant differences in their Apgar scores, requirement for exchange transfusions, or the incidence of significantly patent ductus arteriosus.

Mothers of eight of the retinopathy group had received infusions of β-sympathomimetics (seven ritodrine, one salbutamol) to arrest premature labour: only one of the non-retinopathy group received ritodrine (p<0.008). This observation may be of physiological importance, as animal experiments have suggested that the retinal circulation is strongly influenced by these pharmacological agents acting jointly through the sympathetic nervous system and endothelial factors.3,4 The use of β-sympathomimetics is known to induce transient ischaemic changes in the myocardium of the newborn if given over long periods by intravenous routes.4,5 Maternal ritodrine has been associated with reduced blood pressure in the newborn.6,7 In the premature infant in whom antenatal retinal perfusion is influenced by β-sympathomimetics, these compounds possibly predispose to retinopathy. This observation merits review in a larger series of infants. Further, it casts doubt as to the safety of β-sympathomimetics which are widely used in the therapy of premature labour with little evidence to support their efficacy.

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Immunisation and C reactive protein in infants on neonatal intensive care units

Editor.—C reactive protein estimation is a well known indirect method of detecting bacterial infection in neonates.1 Premature infants on our neonatal unit receive their primary course of immunisation for diphtheria, pertussis, tetanus, and Haemophilus influenzae b as per the immunisation schedule at two, three, and four months postnatally.2 Plasma C reactive protein estimation is performed as part of a weekly or daily infection surveillance. Serial estimations are continued in infants with an increased concentration (normal <4 mg/l). We noticed, as an incidental finding, that values increased within 24 hours of immunisation.

During a period of three months (November 1993 to January 1994) 12 babies had C reactive protein estimations before and after immunisation. The median value preimmunisation was 2.3 mg/l (range 0.3–4.8) and postimmunisation ranged from 0.5 to 14.7 mg/l (median 4.8). The median duration of increased C reactive protein was two days. During the initial period of the study one of the babies had a full infection screen based on the raised C reactive protein and clinical findings (systemically unwell, postimmunisation). Plasma C reactive protein is known to increase after immunisation with diphtheria, pertussis, and tetanus vaccines in malnourished children.3,4 To the best of our knowledge it has not been reported in premature infants who have been immunised. It is important to realise that immunisation leads to an increase in the plasma C reactive protein concentrations so that unnecessary investigations may be avoided.

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Diagnosis and management of non-immune hydrops in the newborn

Editor.—The article by Stephenson and colleagues concerning the diagnosis and management of non-immune fetal hydrops was both informative and practical. However, information regarding the reporting of associations was particularly thorough.1 I would like to make a minor point regarding the inclusion of asplenia syndrome among the cardiovascular associations.

Asplenia syndrome (right atrial isomerism) is characterised by complex cardiac defects including primitive atrioventricular septal defects and pulmonary atresia. This too is listed in Stephe son’s table.1 Nevertheless, it is far more common to see polysplenia syndrome (left atrial isomerism) in association with fetal hydrops.2,3 Left atrial isomerism is well associated with the atrioventricular block, as well as other structural cardiac lesions.2 Complete heart block can occur, rarely, in right atrial isomerism, as in one case in Schmidt’s study.