weight babies. There will be then also a need for an intravenous line. A short non-invasive ophthalmic procedure is changed into a lengthy and invasive anaesthesia technique.

Our previous experience with a nasal ketamine midazolam mixture in babies suggests that it is an attractive alternative for short ophthalmic procedures. A study is under way to use it for different procedures in babies and small children such as ophthalmic examinations, YAG laser treatment, and cryotherapy for ROP.

Nasal administration of drugs is not a new idea and has been used for some time. Pharmacokinetics of intranasal midazolam at a dose as low as 0.2 mg/kg showed that the hypnotic threshold of 100 ng/ml was obtained with 5–10 minutes and this correlated well with sedation.1 Pharmacokinetics of intranasal ketamine were studied by Russel and Aldrete.2 The dose was 4–6 mg/kg and the onset of action within 8–12 minutes. Compared with intramuscular or intravenous routes, intranasal administration is a non-invasive, pain-free method and has a fast onset of action.34 High bioavailability, rapid peak serum levels, and perhaps absorption through the cribriform plate result in a rapid onset of sedation.1

We would like to stress that this technique requires skilful supervision and adequate monitoring. We do not recommend it to be used by untrained staff in outpatient departments or outside the hospital. Low birthweight babies should recover in a high dependency area.

The absence of respiratory depression, arterial oxygen desaturation with satisfactory sedation suggests that this technique is safe and efficient for non-invasive ophthalmic procedures. A large series is required to evaluate potential side effects in this specific indication.

We have presented a novel technique of intranasal administration in a low weight baby requiring cryotherapy for ROP. The side effects were minimal and the operating conditions excellent.

British Journal of Ophthalmology 1993; 77: 530–533

Congenital third nerve palsy associated with mid-trimester amniocentesis

C K Patel, D S I Taylor, I M Russell-Eggitt, A Kriss, P Demaerel

Amniocentesis is the presumed cause of many types of penetrating fetal injuries including skin marks,1 ocular perforation,2,7 limb, chest, and abdominal trauma.8–10 The following report describes how it may also cause neuro-ophthalmic disease and adds to other reports9,11–13 that suggest it is an important cause of congenital intracranial damage.

Case report

A 3 month male infant was referred because his right eye had never opened. He was born at term following uneventful labour and vaginal delivery. Amniocentesis had been performed at 17 weeks of gestation because of advanced maternal age. Ultrasound was used to localise the placenta. Two unguided attempts were then made to aspirate amniotic fluid by stab through one abdominal site. Both times the fluid was stained with blood, which to our knowledge was not analysed to determine its source.

Ocular examination showed a complete ptosis on the right. A diagnosis of total third nerve palsy was made by finding an externally ophthalmoplegic eye that was deviated down and out (Fig 1). Ocular movements on the left were normal. The infant fixated more actively with the

Figure 1 Right ptosis associated with hypotropia and exotropia.
Congenital third nerve palsy associated with mid-trimester amniocentesis

Figure 2. Flash VEPs and ERGs from our patient (left upper trace) and age-matched control (left lower trace) are shown on the left. VEPs to pattern reversal using 100 second checks are shown on the right. VEPs are recorded from left occipital (L Occ), mid-occipital (M Occ), and right occipital (R Occ) locations referred to a mid-frontal reference (MF). The right eye VEPs are better defined compared with the left eye’s confirming the clinical suspicion that visual acuity was better in the paretic eye. The marked occipital asymmetry in the patient’s VEPs suggests right posterior hemisphere dysfunction (see text for fuller explanation).

right eyelid raised suggesting that there was better vision in the paretic eye. There was no relative afferent pupil defect. Cycloplegic retinoscopy was +3.5 D right and +4.0 DS left. The optic discs looked normal. Visual evoked potentials (VEPs) to flash and to pattern stimulation demonstrated left optic nerve dysfunction (Fig 2).

Physical examination showed a skin mark on the right temple (Fig 3). There was an area of pigmentation separating two linear scars which looked like puncture marks. There were no focal neurological limb signs. Magnetic resonance imaging showed right hemiatrophy of the cerebrum, cerebellum, and brain stem (Figs 4 and 5) with hypoplasia of the right middle cerebral circulation (Figs 4(a) and 6).

Comment
Mid-trimester amniocentesis has been linked with cerebral malformations in three previous cases. Naylor et al described an infant with convulsions, a gaze palsy, and a hemiparesis. They detected clinically a homonymous hemianopia in the absence of any consistent asymmetry in the VEPs. Computed tomography, however, showed a cerebral atrophy on the right. Youroukos et al have shown a porencephalic cyst in an infant who presented with vertical jerky eye movements, a probable left visual field defect, and a left hemiparesis. Chong et al reported a neonate with congenital hydrocephalus and a large subarachnoid cyst.

Ultrasound guidance was not used during amniocentesis for these cases although it did predetermine the location of the placenta in the cases described by Naylor and Chong. The use of real time scanning is likely to reduce the risk of inadvertent fetal puncture but does not necessarily eliminate it. Blood staining of the amniotic fluid was a common feature of all three cases. Naylor et al proved fetal vascular disruption by demonstrating that the blood was 100% fetal. When an injury is suspected retrospectively this proof may not be available. A study has, however, shown that fetal blood is usually present when amniotic fluid is grossly stained suggesting that fetal injury may be common. Routine analysis of blood stained amniotic fluid by obstetricians would be valuable. Presumed cerebral puncture can be proved biochemically by examining amniotic fluid for the neural isoenzyme of acetylcholinesterase.
Skin marks, compatible with needle injury, are a more permanent feature of injury during amniocentesis. In the case described by Chong et al a scar was serendipitously discovered after the infant's head was shaved in preparation for the insertion of a cerebrospinal fluid shunt. Such marks can be inconspicuous if fetal skin heals rapidly and there is delayed traction on the epidermis caused by fibrosis. There should be a thorough examination of the head in all neonates where iatrogenic trauma is suspected even with recourse to shaving the head if the need for diagnosis is sufficiently great. Skin marks are usually dimpled or linear, but may be nodular.
and associated with a discharging sinus.12 The morphology of the pigmented lesion in our patient, the use of a single abdominal site during amniocentesis, and the small size of a 17 week fetal head makes it possible that the head was punctured twice in close proximity. An oblique rather than perpendicularly penetrating injury is suggested by the presence of linear scars rather than dimples and could account for the extensive intracranial damage that occurred.

Congenital third nerve palsy is important to recognise and investigate because the neuromuscular prognosis is not as good10 18 as previously thought.19 20 Trauma during vaginal birth or forceps delivery may damage the peripheral nerve.19 Necropsy, computed tomography, and magnetic resonance imaging have highlighted thromboembolic vascular disease as a cause of nuclear palsy.15 18 Congenital heart disease and infective endocarditis are the most important causes of embolic disease in childhood.20 Mid-brain ischaemia has been reported following vertebral artery trauma during birth.21 The occurrence of pulmonary emboli following umbilical vein catheterisation in neonates may be relevant to our case.22 If the fetal venous circulation is traumatised during amniocentesis, emboli could be diverted into the systemic fetal circulation by the high pulmonary resistance. This could explain the involvement of the internal carotid and verteobasilar territories in our patient. The unilateral distribution of cerebral atrophy, however, favours direct trauma to the anterior and posterior cranial fossae. Magnetic resonance imaging supports this hypothesis by showing accentuated atrophy in the temporal lobe under the skin marks that were observed.

Previous reports have highlighted the finding of better vision in the paretic eye of patients with congenital third nerve palsy.10 23 Where refractive errors or visual pathway defects are clinically inapparent it has been suggested that nystagmus may lead to amblyopia because of the fixational advantage conferred on the paretic eye. Nystagmus was absent in our patient but VEP testing showed its value by identifying subclinical optic nerve dysfunction to account for the reduced vision in the left eye. Figure 2 shows the occipital asymmetry in the VEPs to flash and pattern reversal stimulation. The main positivity of the pattern reversal VEP shows a paradoxical lateralisation as it is mainly distributed ipsilaterally to the half field stimulated. Electrodes over the ipsilateral scalp are well placed for picking up activity generated at the posterior medial aspects of the activated hemisphere. We are surprised that asymmetry to pattern reversal was not detected in the patient described by Naylor1 but technical detail was not given.

We thank S C Patel and A Tank for their secretarial help.

Congenital third nerve palsy associated with mid-trimester amniocentesis.

C K Patel, D S Taylor, I M Russell-Eggitt, A Kriss and P Demaerel

Br J Ophthalmol 1993 77: 530-533
doi: 10.1136/bjo.77.8.530

Updated information and services can be found at:
http://bjo.bmj.com/content/77/8/530.citation

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/