Background and Aims A recent RCT suggested improved neurological outcome at discharge for moderate to severe perinatal asphyxia babies given iv magnesium sulphate. However, this trial was performed in babies who were not cooled.

Methods We present a pilot case series of 3 patients with moderate to severe HIE who satisfied the criteria for cooling and received both cooling and iv magnesium sulphate loading of 200mg/kg. Serum Magnesium levels were monitored at 0, 12, 24, 48, 72 hours of cooling.

The babies were reviewed for adverse effects of magnesium sulphate in terms of hypotension, arrhythmia, feed intolerance, respiratory depression and hypocalcemia.

Results One patient received systemic cooling and two other patients received selective head cooling. In addition to iv magnesium sulphate loading, decision was made to institute continuous infusion of iv magnesium sulphate in one of these patients for 4 days at 20–40 mg/kg/h for PPHN. All babies achieved serum magnesium levels of > 1.2 mmol/l within 24h of the loading dose, which was similar to the level aimed for in the previous RCT.

Magnesium sulphate was well tolerated with only mild hypotension requiring one day of dopamine (max 5 mcg/kg/min) in one patient. No babies had respiratory depression, arrhythmia, feed intolerance or hypocalcemia. Neurodevelopmental outcome to date is also presented.

Conclusions Magnesium sulphate is well tolerated in babies with moderate to severe HIE in the cooling era. A large RCT is required to assess its efficacy, long term impact and further look into adverse effects.

Background Hypoxic Ischaemic Encephalopathy (HIE) affects 1–2 per 1,000 live births in UK. The TOBY study showed that therapeutic hypothermia (TH) is beneficial for babies with moderate HIE. In view of this we established an in-house cooling service.

Aim To review our experiences in establishing an in-house cooling service.

Methods The Badger electronic database was used to identify babies who received TH in the last 2 years (01/01/10 to 31/12/11). The management and outcomes were analysed.

Results In the last two year, 27 babies commenced in-house TH. 3 babies were transferred out to another unit for TH for bed capacity reasons. A further 3 babies died before discharge home. 19 babies received the full 72 hours of cooling in our centre. We did not encounter any major complications with the servo-controlled cooling mattress. Only 50% of babies had their MRI in the defined time period as per the TOBY guidelines. All babies are being followed up by a dedicated consultant Neonatologist and neurodevelopmental physiotherapist to assess their neurodevelopment up to the age of 2 years.

Conclusion We have safely established an in-house cooling service by following the TOBY guidelines. The servo-controlled cooling mattress provides a safe cooling process with a rectal probe. Identifying these babies early and the interpretation of CFAM was an important aspect of training. Our main challenge was to get an MRI post cooling in a timely fashion. This has been resolved with an agreed dedicated slot for these babies at Birmingham Children’s Hospital.
Methods Newborn piglets underwent hypoxia following a standardized model. They were randomly assigned for 30 min resuscitation with air (21% O₂) (n=12) or 2.1% Hydrogen gas mixed into synthetic air, H₂ (n=14) and then observed for 9 hours. One control group (n=6) went through the same procedures and observation time (anesthesia, surgery, ventilation and sample collection). The left hemisphere was used for histopathology. Tissue from prefrontal cortex and liver were snap frozen in liquid nitrogen and stored by −70°C until analysis. The tissue samples were homogenized and the protein extracted. A Quantikine KM 300 immunoassay was used to measure activated caspase-3 protein. Gene expression for Casp-3, BDNF, MMP-2, MMP-9 and VEGFR2 was measured in tissue from prefrontal cortex and liver.

Results The use of 2.1% hydrogen gas mixed into synthetic air decreased activated caspase-3 vs. air. In liver tissue piglets resuscitated with air: 12.6 pg/mg protein SD (9.1) vs. H₂: 5.3 (4.9), p=0.031 whereas in cortex piglets resuscitated with air 26.3 pg/mg protein (14.9) vs. H₂ 15.4 (13.0), p=0.05.

There were no significant changes in gene expression in liver and cortex. Histopathology showed a tendency to less brain damage in the hydrogen group.

Conclusions Hydrogen gas used for newborn resuscitation may reduce apoptosis.

THE NEUROPROTECTIVE EFFECTS OF VALPROIC ACID, AN HISTONE DEACETYLASE INHIBITOR IN A NEONATAL HYPOXIC-ISCHEMIC RAT MODEL

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Introduction Neurodegenerative diseases were associated with a decrease in histone acetylase transferase (HAT) activity, resulting in relative over-deacetylation. Histone deacetylase (HDAC) inhibitors were suggested as potentially neuroprotective agents. The aim of this study was to evaluate the neuroprotective effects of valproic acid (VPA), an histone deacetylase inhibitor, in neonatal hypoxic ischemic rat model.

Methods After being anesthetized, 7-day-old pups underwent ischemia followed by exposure to hypoxia. The pups were divided into 5 groups: sham group, vehicle group (saline group) and VPA group. VPA was administered intraperitoneally for three times; the first just after hypoxia-ischemia, the second and the third doses 24 and 48 hours after the first dose, respectively. After sacrifice, brain infarct volume, apoptosis, HDAC activity, acetylated H4 protein and caspase 3 expression, and proinflammatory cytokine concentrations were evaluated in brain tissue of rat pups.

Results Percent infarcted brain volume and number of TUNEL positive cells per unit area in hippocampus and cortex CA1 were markedly reduced with VPA treatment. HDAC activity was found to be significantly reduced in VPA group, whereas acetylated H4 protein expression was significantly increased with VPA treatment. The caspase-3 activity in VPA group was significantly lower than the control group. The proinflammatory cytokine levels also significantly decreased with VPA treatment.

Conclusion This is the first study that showed the neuroprotective effects of VPA treatment as an HDAC inhibitor by reducing percent infarcted brain volume, histone deacetylase activity, inflammation and apoptosis while increasing acetylated H4 protein levels in a neonatal hypoxic-ischemic rat model.

HYPERGLYCAEMIA AND INSULIN-INDUCED ALTERATIONS IN THE RETINA OF RAT PUPS

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Introduction Streptozotocin-induced hyperglycaemia, insulin-treatment and intravitreal injection of a potential retinoprotective agent, pituitary adenylate cyclase activating polypeptide (PACAP) on the rat pup’s retina. The aim of our study was to investigate specific effect of streptozotocin-induced hyperglycaemia, insulin-treatment and intravitreal injection of a potential retinoprotective agent, pituitary adenylate cyclase activating polypeptide (PACAP) on the rat pup’s retina.

Methods The use of 2.1% hydrogen gas mixed into synthetic air decreased activated caspase-3 protein. Gene expression for Casp-3, BDNF, MMP-2, MMP-9 and VEGFR2 was measured in tissue from prefrontal cortex and liver.

Results The use of 2.1% hydrogen gas mixed into synthetic air decreased activated caspase-3 vs. air. In liver tissue piglets resuscitated with air: 12.6 pg/mg protein SD (9.1) vs. H₂: 5.3 (4.9), p=0.031 whereas in cortex piglets resuscitated with air 26.3 pg/mg protein (14.9) vs. H₂ 15.4 (13.0), p=0.05.

There were no significant changes in gene expression in liver and cortex. Histopathology showed a tendency to less brain damage in the hydrogen group.

Conclusions Hydrogen gas used for newborn resuscitation may reduce apoptosis.

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A PIG MODEL OF THE PRETERM NEONATE: ANTHROPOMETRIC AND PHYSIOLOGICAL CHARACTERISTICS

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Background and Aims Large animal models are an essential research tool to investigate the physiology of the preterm infant, which remains poorly understood. We aim to describe the pig model of the preterm neonate in terms of growth, maturation and requirement for intensive care over a range of gestational ages and determine the effects of maternal glucocorticoid exposure and sex.

Methods Twenty-nine litters of piglets (N=305) were delivered by C-section at 91d, 94d, 97d, 100d, 104d and 113d (term 115d). Some litters received maternal betamethasone treatment (0.19mg/kg body wt; 1M) at 48h and 24h prior to delivery. At 97d piglets were resuscitated, surfactant administered, and piglets were ventilated, sedated and monitored for 6–8h post-birth using standard NICU techniques.

Results At 91d, piglets were half the weight of term animals, had fused eyelids, very thin skin, no hair, and survived a maximum of 5h due to difficulties with ventilation. At 97d piglets were able to be maintained for at least 6–8h but physiology was unstable for 1–2h. Piglets 100d and older breathed spontaneously. Only near term piglets were able to maintain body temperature. Males were heavier than females at 113d gestation (p<0.021). Exposure to maternal glucocorticoids resulted in larger females and influenced brain body wt.

Conclusions The piglet provides a useful model of preterm neonatal physiology as very preterm piglets can be survived under standard intensive care conditions. The large litters allow for parallel experiments or the use of littermates as controls.