



# THE UNIVERSITY of EDINBURGH

## Edinburgh Research Explorer

### Hypoglycaemia and neonatal brain injury

**Citation for published version:**

Boardman, JP 2014, 'Hypoglycaemia and neonatal brain injury' Italian Journal of Pediatrics, vol. 40, no. Suppl 2, pp. A13. DOI: 10.1186/1824-7288-40-S2-A13

**Digital Object Identifier (DOI):**

[10.1186/1824-7288-40-S2-A13](https://doi.org/10.1186/1824-7288-40-S2-A13)

**Link:**

[Link to publication record in Edinburgh Research Explorer](#)

**Document Version:**

Publisher's PDF, also known as Version of record

**Published In:**

Italian Journal of Pediatrics

**Publisher Rights Statement:**

© 2014 Boardman; licensee BioMed Central Ltd.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (<http://creativecommons.org/licenses/by/4.0/>), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. The Creative Commons Public Domain Dedication waiver (<http://creativecommons.org/publicdomain/zero/1.0/>) applies to the data made available in this article, unless otherwise stated.

**General rights**

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

**Take down policy**

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact [openaccess@ed.ac.uk](mailto:openaccess@ed.ac.uk) providing details, and we will remove access to the work immediately and investigate your claim.





MEETING ABSTRACT

Open Access

# Hypoglycaemia and neonatal brain injury

James P Boardman

From XX National Congress of the Italian Society of Neonatology  
Rome, Italy. 9-11 October 2014

The transition from fetal to neonatal life requires metabolic adaptation to ensure energy supply to vital organs and systems after separation from the placental circulation. Under normal conditions this is achieved through the mobilisation and use of alternative cerebral fuels (fatty acids, ketone bodies and lactate) when blood glucose concentration falls.

The level of blood glucose (BG) concentration that leads to cerebral injury in newborns and adverse neurodevelopmental outcome is unknown: low BG is often observed during postnatal adaptation of healthy term infants without apparent adverse consequence, and the capacity to mobilise and use alternative cerebral fuels when BG is low varies between patient groups. Severe hypoxic-ischaemic encephalopathy (HIE) is associated with impaired metabolic adaptation, and animal and human data suggest that levels of hypoglycaemia that are tolerated under normal conditions may be harmful in association with hypoxia-ischaemia.

The spectrum of cerebral injury associated with hypoglycaemia is wide and includes: white matter injury including parenchymal haemorrhage and ischaemic stroke, cortical neuronal injury, and sometimes signal change in the basal ganglia (mainly the globus pallidus) and thalamus. Vulnerability of the white matter and cortex of the posterior parietal and occipital lobes has been well reported in human imaging studies, but the site of injury is more widespread in pathological and experimental studies of neonatal hypoglycaemia. In the largest series of infants with isolated neonatal hypoglycaemia and acute neurological dysfunction, there was an association with a predominantly posterior pattern of injury in one third of the cohort, and a more extensive distribution of lesions was common.

Safe clinical management relies on the identification of infants at risk of neurological sequelae from hypoglycaemia, adequate energy provision after birth, monitoring of

blood glucose, and prompt intervention to raise the BG at specified thresholds, with the caveat that acute neurological dysfunction in association with low BG at any level should prompt urgent investigation and treatment. The optimal target blood glucose level for ensuring adequate energy provision in health and in HIE remains unknown. However, recent data support guidance to maintain blood glucose concentration  $\geq 2.5\text{ mmol/L}$  in neonates with signs of acute neurological dysfunction, which includes those with HIE, and is higher than the accepted threshold of  $\geq 2\text{ mmol/L}$  in infants without abnormal signs or hyperinsulinism.

Published: 9 October 2014

doi:10.1186/1824-7288-40-S2-A13

Cite this article as: Boardman: Hypoglycaemia and neonatal brain injury. *Italian Journal of Pediatrics* 2014 **40**(Suppl 2):A13.

Submit your next manuscript to BioMed Central  
and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your manuscript at  
[www.biomedcentral.com/submit](http://www.biomedcentral.com/submit)

