Iron supplementation reduces the frequency and severity of breath-holding attacks in non-anaemic children

Anthony Zehetner
Department of Paediatrics, Teaching and Research Unit, The University of Newcastle, Gosford Hospital, Gosford, Australia

Abstract

Iron supplementation reduces the frequency and severity of breath-holding attacks (BHA}s), particularly in children with iron deficiency. The issue of iron supplementation is less clear for Westernized children with BHAs who present to an outpatient community clinic and are not iron-deficient. This is the first reported case series of iron-replete children with frequent and disabling breath-holding attacks who have responded to a course of oral iron supplementation. This intervention is safe, improves quality of life for both child and carer, and is significantly cost-effective in terms of health resource utilization.

Introduction

For several years, the medical literature has associated the incidence of breath-holding attacks (BHAs), particularly in children with iron deficiency.1,3 A recent Cochrane Systematic Review suggests that iron supplementation reduces the frequency and severity of breath-holding attacks, particularly in children with iron deficiency (OR 76.48; 95% CI 15.65 to 373.72; P< 0.00001).4 Due to study design, trial limitations and the populations studied (high rate of consanguinity and an up to 10% prevalence of haemoglobinopathy), the issue of iron supplementation is less clear for Westernized children with breath-holding attacks (BHAs) who present to an outpatient community clinic and are not iron-deficient.

I wish to report a small case series of iron-replete children with frequent and disabling breath-holding attacks who have responded to a course of oral iron supplementation in my practice.

Case Reports

Case #1
A 2-year-9-month-old girl presented predominantly cyanotic BHAs since the age of 3 months. He had 2-8 episodes per day, with transient loss of consciousness, triggered at the onset of tantrums or if his father leaves the room. He had no thalassaemia, his ECG and EEG were both normal, and there was no family history of BHAs. Reduction in BHAs to one every 2-3 days was noted at 8 weeks, falling to one every 3 weeks at 12 weeks of treatment. Thereafter, no loss of consciousness was noted with further BHAs. The child is now able to leave his father’s side and attend day care.

Case #2
A 15-month-old boy presented with mixed cyanotic and pallid BHAs since the age of 8 months. Her electrocardiogram (ECG) and electroencephalogram (EEG) were both normal. Her paternal aunt had BHAs during childhood. Haemoglobin increased from 106 g/L to 145 g/L at the end of the treatment period. Subsequent BHAs were not associated with loss of consciousness.

Discussion

All children were administered elemental iron at 5 mg/kg/day for a period of three months (12-16 weeks) in divided doses, as per the Cochrane protocol.4 Compliance was monitored from medication usage (prescription refills and the presence of black stools) and patients reviewed at 4-6 weekly visits. A BHA diary was kept from baseline until the end of the treatment period. All children tolerated the oral iron supplementation for the three month course and no adverse events were reported (such as gastro-intestinal intolerance).

From Table 1, there was a 67-99% (mean 86%) reduction in the frequency of BHAs from baseline. In all cases, the severity (progression to loss of consciousness) was reduced. From follow-up parental reports, this effect has been maintained after the iron supplementation was ceased at 12-16 weeks. Parents and child care workers also reported improvements in the child’s behaviour.

While children with BHAs are expected to outgrow this condition by the age of seven to eight years;5 some will also do this from the age of three-to-four years. However none of the children in this series fall into this age bracket, and the dramatic reduction in episodes over three months appears swift, so the effect is considered attributable to the iron intervention rather than ageing.

While the number of cases is small and the intervention is not blinded or controlled for with a placebo, the magnitude of the results and clinical outcomes warrant further study. This finding is significant as it is the first documented case in the medical literature of non-iron deficient children with BHAs responding to a trial of iron supplementation. It is also the first published case reports of BHAs in an Australian population. To study this effect further, an application for an NHMRC-funded multicentre double-blinded placebo-controlled randomized control trial in an Australian setting has been submitted.

The cost-effectiveness (and tolerability) of the intervention cannot be ignored. From 1 July 2011 in NSW, the cost of an ambulance call-out is A$320 + $2.89 per kilometre to the nearest hospital (up to a total cost of A$5248).6 For case C, whose parents called an ambulance out on three occasions within one month, this cost is considerable (at least $1000) in comparison with the price of a bottle of Ferro-liquid at A$15. Freeing up health resources ever under high demands has benefits to our society beyond monetary costs. Similarly, the quality of life gained for the caregiver and child is considerable as the BHAs resolve.

Although investigations, such as an EEG, ECG and haemoglobin electrophoresis, were carried out in these cases, the diagnosis of...
breath-holding attacks is a clinical one made on a typical history, which all of these children possessed. Frequently these other investigations are performed for differential diagnostic purposes and prior to presentation at the initial clinic visit. Iron deficiency is implicated in conditions other than anaemia (a late manifestation) and BHAs. It is increasingly recognized to be a cause of restless legs, febrile seizures, thrombosis, impaired immunity and poor behaviour. It is known that children who have experienced BHAs may become adolescents with syncopal episodes. A recent retrospective cohort study of largely non-iron deficient children with BHAs showed a 29.4% incidence of concentration problems.

### Table 1. Patient demographics and effect of iron supplementation on breath-holding attacks.

| Case | Gender | Age at clinic presentation (months) | Ethnicity | Family history of BHAs | Baseline Hb (g/L) | Av BHA frequency (No. per week) | % reduction in BHA frequency |
|------|--------|----------------------------------|-----------|------------------------|------------------|------------------------------|-----------------------------|
| A    | F      | 31                               | Anglo-Saxon | Y                      | 106              | 2.3                          | 0.17                         | 93%                         |
| B    | M      | 15                               | Asian     | N                      | 112              | 35                          | 0.33                         | 99%                         |
| C    | M      | 14                               | Arabic    | N                      | 120              | 0.5                          | 0.17                         | 67%                         |

BHAs, breath-holding attacks

### References

1. Colina KF, Abelson HT. Resolution of breath-holding spells with treatment of concomitant anaemia. J Pediatr 1995;126:395-7.
2. Daoud AS, Batieha A, Al-Sheyyab M, et al. Effectiveness of iron therapy on breath-holding spells. J Pediatr 1997;130:547-50.
3. Mocan H, Yildiran A, Orhan F, et al. Breath holding spells in 91 children and response to treatment with iron. Arch Dis Child 1999;81:261-2.
4. Zehetner AA, Orr N, Buckmaster A, et al. Iron supplementation for breath-holding attacks in children [Systematic Review]. Cochrane Database Syst Rev 2010;5:CD008132.
5. Tam DA, Rash FC. Breath-holding spells in a patient with transient erythroblastopenia of childhood. J Pediatr 1997;130:651-3.
6. Ambulance Service of NSW. Provision of Ambulance Services: New Fees (2011). http://www.ambulance.nsw.gov.au/Media/docs/fees%2020110701-640bc2b5-0a82-40ed-a92f-65e68cd34e11-0.pdf Accessed: 21 July 2011.
7. Yadav D, Chandra J. Iron deficiency: beyond anaemia. Indian J Pediatr 2011;78:65-72.
8. Olsen AL, Mathiasen R, Rasmussen NH, Knudsen FU. Long-term prognosis for children with breath-holding spells. Dan Med Bull 2010;57:A4217.