Hydroxykynurenine/hydroxyanthranilic acid ratios and febrile convulsions

J MCKIERNAN, D MELLOR, S COURT, J EDSON, AND K LACEY

Department of Child Health and University Hospital, Queen's Medical Centre, Nottingham

SUMMARY Hydroxykynurenine/hydroxyanthranilic acid ratios were measured in children with febrile convulsions, afebrile fits, and fever, as well as in healthy controls. Increased ratios were found not only in the children who had fits but also in the children who were febrile and did not have fits. It is suggested that a raised hydroxykynurenine/hydroxyanthranilic acid ratio does not necessarily indicate vitamin B6 deficiency but may represent a nonspecific response of tryptophan metabolism to stress.

Febrile convulsions are a common problem in children, affecting between 2 and 5% in the 6-month to 5-year age range. The pathogenesis of these seizures is imperfectly understood. The elucidation of the syndromes of vitamin B6 deficiency and dependency in which fits are a major clinical feature, raised the possibility that a specific biochemical basis might also be found for the more common convulsive disorders of childhood. The tryptophan load test has been traditionally regarded as a test of vitamin B6 status although it is now appreciated that other influences—such as oestrogenic and corticosteroid hormones, stress, and the general state of protein metabolism—may affect the result. Tryptophan load tests were carried out on 20 children with febrile convulsions by Carredu et al. and abnormalities in metabolite excretion were found. It was therefore suggested that a relationship exists between febrile convulsions and pyridoxine deficiency. Lennox-Buchthal suggested that the association between vitamin B6 and febrile convulsions be further explored, noting that in the syndrome of vitamin B6 dependency, fever increases the need for pyridoxine and may precipitate convulsions.

Tryptophan load tests in the past have mainly been carried out using timed collections of urine after the tryptophan load, and then by measuring a number of urinary metabolites. Such collections are impractical in many children. In addition, great variability in metabolite excretion is found. We used a modification of the tryptophan load test, and measured only two urinary metabolites—hydroxykynurenine (HK) and hydroxyanthranilic acid (HA), which were then expressed as a ratio—KH/HA. This ratio is thought to reflect the activity of the liver enzyme kynureninase, which requires vitamin B6 as a cofactor (Fig. 1). It is considered to be a reliable index of pyridoxine deficiency and, because it is a ratio, it has the advantage of not requiring a carefully timed urine collection. We estimated HK/HA ratios after a tryptophan load in normal children, children who were febrile, children who had afebrile fits, and children with febrile convulsions, and report our results.

Patients and methods

Each of the 69 children with febrile convulsions had been admitted to hospital after a first or second febrile fit. Ages ranged between 6 months and 5 years. The diagnosis was based on the usual criteria—that
The HK/HA results with height of relationship between HK/HA ratio and age, convulsions had to ratio tryptophanisions. The higher (P < 0.05) significantly cortisol. To measured using urine samples. The eluted methods of column recoveries of measurement hydrolysed % were supplied samples. Tryptophan load after minutes urine single child was well. ‘Fever only’ group. Six children, aged between 2 and 5 years who had a first nonfebrile fit, who were otherwise normal and not on anticonvulsant treatment, comprised the ‘afebrile fit’ group. Seven healthy children, who were not in hospital, acted as controls.

**Tryptophan load test.** Tryptophan powder 100 mg/kg body weight was given as a sweetened suspension. A single urine specimen was collected at least 20 minutes after the ingestion of the dose and frozen until assay.

**Measurement of HK and HA.** Fractionation of tryptophan metabolites from urine was carried out by an ion exchange resin technique based on the methods of Brown and Price. Urine samples were hydrolysed using the method described by Heeley. The eluted HK and HA were measured spectrophotometrically using Brown’s method. A blank and a duplicate test were performed on each sample. Column recoveries were carried out with each batch of tests and the recovery of metabolites added to urine samples was also determined; recovery was 89% ± 7 (mean ± SD) for both HK and HA.

**Plasma cortisol.** Blood samples were collected from 15 children with febrile convulsions at the time of admission to hospital and plasma cortisol was measured using a solid phase radioimmunoassay supplied by the Tenovus Institute, Cardiff.

All results were analysed by Student’s t test for unpaired samples.

**Results**

The HK/HA ratio in the healthy control children was 0.80 ± 0.16 (mean ± SE) (Fig. 2). Ratios were significantly higher (P < 0.05) in the groups of children with fever, afebrile fits, and febrile convulsions. The majority of children with febrile convulsions had raised HK/HA ratios, but no relationship between HK/HA ratio and age, sex, height of fever, or complexity of fit could be found.

In 30 cases it was possible to relate the HK/HA ratio to the time interval between the convulsion and the tryptophan load test (Fig. 3). Increased ratios were found up to 4 days after the fit. In 6 cases the tryptophan load test was repeated one month or more after the convulsion, when the child was well. Ratios in the normal range were found in 4 children and in 2 values were still raised. We found no relationship between the HK/HA ratios and plasma cortisol in 15 children with febrile convulsions.
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Discussion

This study shows that in a small group of healthy children, HK/HA ratios after a tryptophan load were within the range found in healthy women, and we could not find any results for healthy children. From our results it is also clear that febrile illnesses, not accompanied by convulsions, may induce a rise in the HK/HA ratio. Even in a child who had had a single nonfebrile fit, the ratio was higher than in the control.

We have confirmed, therefore, that tryptophan metabolism is disordered in children with febrile convulsions as was suggested by Carredu et al. Interestingly, Rabe and Plonko found abnormal xanthurenic acid excretion after a tryptophan load in a small number of children with infections, some of whom had febrile fits. None showed evidence of pyridoxine depletion on the basis of the 4-pyridoxic acid excretion test.

Despite the refinements in the tryptophan load test, we feel that an abnormal HK/HA ratio is not a specific test of pyridoxine deficiency. Coon and Nagler stressed that the test might be influenced by other factors—such as stress, adrenal glucocorticoids, sex hormones, and the general state of protein metabolism. Increased hormone activity, as happens in stress, and increased protein catabolism may increase the flow of metabolites down the kynurenine pathway and result in raised HK/HA ratios even in the presence of normal amounts of pyridoxine (Fig. 1). We feel that it is likely that some such mechanism may underlie the increased HK/HA ratios that we have found in children with fever, fits, and febrile convulsions although we were unable to demonstrate any relationship between plasma cortisol levels and the HK/HA ratios.

Despite these considerations, our results do not exclude the possibility that fever or infection can induce a transient state in which vitamin B6 demand may exceed supply. Such a state of relative B6 deficiency could conceivably be the underlying mechanism in the pathogenesis of febrile convulsions. If so, this would have clear implications for prophylaxis in children after an initial febrile convolution.

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References


Correspondence to Dr J McKiernan, Paediatric Unit, Limerick Regional Hospital, Dooradoyle, Limerick, Eire.

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