The association between febrile convulsion in children and chronic hyperventilation in parents

M Widiastuti Samekto, MD; I Gusti Putu Ardana, MD

ABSTRACT

**Background** Febrile convulsion and chronic hyperventilation syndrome (spasmophilia) are suspected to share the same root of pathophysiology, a genetic trait abnormality related to ion channel that could cause neuronal hyperexcitability. We want to determine the prevalence ratio of parents with chronic hyperventilation syndrome between two groups of children with and without febrile convulsion.

**Methods** A cross-sectional design study was used, with a tertiary hospital setting (Kariadi Hospital). Participants were selected consecutively based on eligibility criteria. Febrile convulsion was diagnosed based on a modified Livingstone criteria. Chronic hyperventilation syndrome in parents was determined using the clinical diagnostic test of spasmophilia (88.4% sensitivity and 61.6% specificity). Statistical calculations were conducted with two by two table analysis and within the 95% confidence interval.

**Results** Sixty-two children (mean age 18 months) who met the eligibility criteria were included. The parents' mean ages were 33 years (father) and 29 years (mother). The prevalence ratio of father, mother and both parents with chronic hyperventilation between the febrile convulsion group and the non-febrile convulsion group were 2.56 (95% CI 0.53;12.31), 6.19 (95% CI 1.70;22.6) and 18.7 (95% CI 3.07;113.9), respectively.

**Conclusion** Febrile convulsion can be anticipated in children of parents who suffer from chronic hyperventilation syndrome [Paediatr Indones 2002;42:239-242].

**Keywords:** febrile convulsion, chronic hyperventilation syndrome

Chronic hyperventilation (CHV) syndrome (also known as spasmophilia or neuronal hyperexcitability syndrome) describes a set of somatic and psychological symptoms thought to result from episodic or chronic hyperventilation and usually are reproduced in whole or in part by voluntary hyperventilation. The syndrome consists of many symptoms, such as headache, fatigue, indigestion, paresthesia in extremities, cold and clammy hands and feet, anginalike chest pain, anxiety, and syncope.1-6 The incidence of this syndrome in the general population varies between 6-11% and may mimic diverse organic disorders.7 Steidle and Kalparek (1989) showed that people with spasmophilia had higher nervous excitability.8 This was also found in an unpublished data with odds ratio of 17.78 (95% CI 2.18;144.83, p=0.001).9 The presence of gated ion channels in the plasma membranes of neurons gives these cells a property of electrical excitability. A genetic aberration that results in channelopathy, especially a voltage-gated Ca channel, may underlie the increase of nervous excitability.10,11 Study on rats by Fan et al indicated that neurons in the caudal hypothalamus responsible for the modulation of respiratory output have a dominant T-type calcium current. This T-type current seems to have a role in the bursting characteristics of rhythmic firing in caudal hypothalamic neurons, thus resulting in abnormal respiratory drive.12

From the Department of Child Health, Medical School, University of Diponegoro, Indonesia.

Reprint requests to: Widiastuti Samekto, MD, Department of Child Health, Medical School, University of Diponegoro, Kariadi Hospital, Semarang 50231; Tel./Fax.: 62-24-311471
Febrile convulsion (FC), a common form of childhood seizure, occurs in young children when there is a rapid increase in their body temperature. It affects about 1 in 20 children between the age of one and four but may affect those aged between six months and five year-old. Study by Racacho et al concluded that febrile convulsion has a genetic heterogeneity. Febrile seizures are frequently linked with epilepsy although the clinical and genetic relationships are poorly understood. Berkovic indicated that idiopathic epilepsies appear to be a family of ion channelopathies, either ligand-gated or voltage-gated, suspected to be the cause of abnormal neuronal hyperexcitability. Our hypothesis was that febrile convulsions and chronic hyperventilation syndrome may share the same pathophysiological root, namely a genetic abnormality related with ion channels which might be transferred. We tried to determine whether children with febrile convulsion have parents with a history of chronic hyperventilation syndrome.

Methods

Study design and participants

A cross-sectional study was conducted in 1999 with a tertiary hospital setting. Children with fever of more than 38°C and aged between 6 months and 4 years were selected consecutively among children who were admitted to Kariadi Hospital Semarang. Patients with sepsis and altered consciousness, brain infection, hydrocephalus, intracranial space occupying lesion, brain contusion, microcephaly, cerebral palsy, and developmental delay were excluded. Parents of children eligible for study were selected based on criteria: 1) “biological” parents; 2) both parents are still alive; 3) if parents were divorced, they are still accessible for questioning; 4) they agreed to be included in the study.

Parents with hypertension, stroke or post stroke, brain space occupying lesion, post severe concussion, heart disease or heart failure, peptic ulcer were excluded. If the parents did not meet the criteria mentioned above, the child was excluded from the study group.

Determination of children with febrile convulsion

Children eligible for study were screened for febrile convulsion. The diagnosis was based on the modified Livingstone criteria: 1) the first onset of convulsion was between 6 months and 4 years; 2) the duration of convulsion was less than 15 minutes; 3) the convulsion had the characteristics of a generalized seizure; 4) the onset of convulsion was during the first 16 hours of fever; 5) neurologic examination was normal before and after febrile convulsion attack; 6) EEG was normal one week after fever abated.

Determination of parents with chronic hyperventilation syndrome

The reference standard for chronic hyperventilation syndrome in the parents was EMG result after hyperventilation provocation test. Because of motivational constrain, instead of EMG evaluation, chronic hyperventilation syndrome was determined by clinical diagnostic test. The sensitivity of the test was 88.4% and the specificity was 61.6%. Parents were regarded to have hyperventilation syndrome if they have a history of chronic syndrome consisting minimally one out of three somatic symptoms/signs (tension headache, cramps, moderate positive Chovstek’s sign), and two out of three autonomic or visceral symptoms (paresthesia, cold and clammy hands and feet, indigestion, or chest pain).

Statistical analysis

The characteristics of children with febrile convulsion and their parents were compared with the characteristics of children without febrile convulsion and their parents using t-test or chi-square test.

Two by two table analysis was used to calculate the prevalence ratio (included in the 95% confidence interval) of parents (either single or both) with chronic hyperventilation syndrome between the two groups of children.

Results

Sixty seven children were found eligible for this study; two were excluded because of developmental delay,
one because of brain infection and two because of inaccessible divorced parents. There were no significant differences in demographic characteristics among the FC group (31 subjects) and the non-FC group (also 31 subjects) (Table 1). There was also no significant difference regarding the age of their parents.

Table 1. Characteristics of Study Group

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>FC Group</th>
<th>Non-FC Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (months), mean (SD)</td>
<td>18.77 ± 9.52</td>
<td>17.10 ± 10.86</td>
</tr>
<tr>
<td>Sex (number)</td>
<td>Male 18</td>
<td>Female 19</td>
</tr>
<tr>
<td></td>
<td>Father 13</td>
<td>Mother 12</td>
</tr>
<tr>
<td>Father’s age (yrs), mean (SD)</td>
<td>32.97 ± 6.50</td>
<td>32.74 ± 6.15</td>
</tr>
<tr>
<td>Mother’s (yrs), mean (SD)</td>
<td>29.32 ± 5.42</td>
<td>28.68 ± 4.73</td>
</tr>
</tbody>
</table>

If the percentages of father, mother or both with CHV among FC and non-FC group were compared, there were significant differences (Table 2).

Table 2. Proportion of Parents with CHV in Each Study Group

<table>
<thead>
<tr>
<th>CHV</th>
<th>FC Group (n=31)</th>
<th>Non-FC Group (n=31)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Father</td>
<td>16</td>
<td>8</td>
<td>0.03</td>
</tr>
<tr>
<td>Mother</td>
<td>24</td>
<td>11</td>
<td>0.001</td>
</tr>
<tr>
<td>Both</td>
<td>11</td>
<td>5</td>
<td></td>
</tr>
</tbody>
</table>

A separate two by two table analysis was conducted to calculate the prevalence ratio of father with CHV, mother with CHV, and both parents with CHV within the FC group and the non-FC group.

To calculate the prevalence ratio of father or mother with CHV within the FC group and the non-FC group, children with both parents experiencing CHV were excluded to prevent bias (Tables 3 and 4).

Table 3. Frequency of Father within Each Study Group

<table>
<thead>
<tr>
<th></th>
<th>FC Group (n=20)</th>
<th>Non-FC Group (n=26)</th>
<th>Total n=46</th>
</tr>
</thead>
<tbody>
<tr>
<td>Father</td>
<td>5</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>Without CHV</td>
<td>15</td>
<td>23</td>
<td>38</td>
</tr>
</tbody>
</table>

Prevalence ratio: 2.56 (95% CI 0.53;12.31) p=0.211

The same way, to calculate the prevalence ratio of both parents with CHV within the FC group and the non-FC group, children with either father or mother experiencing CHV were excluded (Table 5).

Table 4. Frequency of Mother in Each Study Group

<table>
<thead>
<tr>
<th></th>
<th>FC Group (n=20)</th>
<th>Non-FC Group (n=26)</th>
<th>Total n=46</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mother With CHV</td>
<td>13</td>
<td>6</td>
<td>19</td>
</tr>
<tr>
<td>Without CHV</td>
<td>7</td>
<td>20</td>
<td>27</td>
</tr>
</tbody>
</table>

Prevalence ratio: 6.19 (95% CI 1.70;22.6) p=0.005

Table 5. Frequency of Parents in Each Study Group

<table>
<thead>
<tr>
<th></th>
<th>FC Group (n=16)</th>
<th>Non-FC Group (n=19)</th>
<th>Total n=35</th>
</tr>
</thead>
<tbody>
<tr>
<td>Both father/mother</td>
<td>11</td>
<td>2</td>
<td>13</td>
</tr>
<tr>
<td>With CHV</td>
<td>5</td>
<td>17</td>
<td>22</td>
</tr>
</tbody>
</table>

Prevalence ratio: 18.7 (95% CI 3.07;113.9) p=0.001

Discussion

Our study indicated that the prevalence of parents with CHV was significantly higher in children with febrile convulsion. If a single parent was involved, a mother with CHV was more prevalent than a father with CHV (prevalence ratio 6.19 compared with 2.56). This supports a previous study which demonstrated that women with CHV were more prevalent compared to men with CHV among outpatients of a tertiary hospital. The highest prevalence rate was found if both parents suffered from CHV (ratio 18.7). These facts support the idea that febrile convulsion and chronic hyperventilation syndrome may share the same genetic trait. This is also supported by reports describing pseudo-seizures in the absence of epilepsy caused by hyperventilation in children, suggesting that febrile convulsion and hyperventilation syndrome may have the same pathophysiology.

Study by Han et al suggested that instability of breathing in chronic hyperventilation syndrome may be caused by genetic abnormal neuronal hyperexcitability.

In conclusion, our study proved that for the majority, febrile convulsion can be anticipated in children of parents with chronic hyperventilation syndrome either from one or both parents. The results of our study might be useful for family education, guidance, and counseling. Even though successful effect of breathing therapy for chronic hyperventilation syndrome had been put forward by several researchers, would this be worth trying for children with the problem?
pensity of febrile convolution? We recommended further study concerning this issue, especially in older children since we can expect more compliance. 20,21

References