Chronic proximal spinal muscular atrophy of childhood and adolescence: problems of classification and genetic counselling

IRENA HAUSMANOWA-PETRUSEWICZ*, JACEK ZAREMBA†, AND JANINA BORKOWSKA*

From the *Department of Neurology, Medical School, and †Department of Genetics, Psychoneurological Institute, Warsaw, Poland.

SUMMARY Results obtained from a study of 354 cases of chronic proximal spinal muscular atrophy of childhood and adolescence suggest that the condition is not as homogeneous as it was previously thought. A tentative classification based on our results is proposed. Estimates of genetic risks are provided, taking into account the sex and age at clinical onset. In our opinion these factors are more reliable than the data hitherto available because they are based on a considerably larger series.

The problem of the influence of sex in the chronic proximal form of spinal muscular atrophy (CPSMA) of childhood and adolescence based upon a study of 354 cases was discussed in our previous article published in this Journal.1 The information gathered during the study mentioned above allows us to comment on two other aspects, the classification of CPSMA and genetic counselling. The material and methods were described earlier.1

Classification

Until recently CPSMA of childhood and adolescence was considered to be a single condition, because there were no grounds for dividing it into different entities.2 3 We were even doubtful about the intermediate form of Fried and Emery4 being a separate entity (see No 25 355 in McKusick5). This problem was discussed in one of our earlier publications.6 However, our recent results1 7 suggest some heterogeneity of CPSMA. In this respect the subgroup with onset at 10 to 36 months of age was of particular interest because of a sharp and significant fall of the p value of the segregation ratio (table 1, figure). It is possible that a large proportion of cases in this age range are phenocopies or new dominant mutations or both. They are distributed throughout the whole age spectrum of our material, but in the age range 10 to 36 months, in which there

Table 1 Segregation ratios and empirical risk values according to sex in different age at onset subgroups of CPSMA of childhood and adolescence.

<table>
<thead>
<tr>
<th>Sex</th>
<th>Age at onset</th>
<th>0-9 mth</th>
<th>10-36 mth</th>
<th>37 mth-8 y</th>
<th>9 y-18 y</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>p</td>
<td>r</td>
<td>p</td>
<td>r</td>
</tr>
<tr>
<td>M</td>
<td>0-25</td>
<td>1:4</td>
<td>0:036</td>
<td>1:27</td>
<td>0:22</td>
</tr>
<tr>
<td>F</td>
<td>0-20</td>
<td>1:5</td>
<td>0:046</td>
<td>1:23</td>
<td>0:16</td>
</tr>
<tr>
<td>All</td>
<td>0-22</td>
<td>1:4:5</td>
<td>0:039</td>
<td>1:25</td>
<td>0:19</td>
</tr>
</tbody>
</table>

0-24 mth 25-18 y 37 mth-18 y 0 y-18 y

| M   | 0-15        | 1:7 | 0:20| 1:5 | 0:26| 1:4 | 0-16| 1:6 |
| F   | 0-15        | 1:7 | 0:12| 1:8 | 0:09| 1:11| 0-14| 1:7 |
| All | 0-15        | 1:7 | 0:17| 1:6 | 0-17| 1:6 | 0-15| 1:7 |

*More detailed data are given in tables 2 and 3 of our previous publication.1 p=segregation ratio. r=empirical risk.

Figure. Diagram of segregation ratios on four different age at onset subgroups (compare with table 1).
may be only a few autosomal recessive cases, they become more evident. In one of our previous publications\(^8\) we considered the problem of dominant mutations and tried to explain their very infrequent occurrence (0-75\% in a large combined series of an international study on SMA\(^9\)). However, we are now reconsidering our previous opinion because there is a possibility that most cases of the dominant form of juvenile SMA (No 15 860 in McKusick\(^5\)) are new mutations, and that mild cases compatible with familial occurrence of this form of the disease are very rare, hence the apparent paucity of dominant cases of SMA. Most of them may never produce evidence of their dominance because of their greatly reduced fitness. Such cases may form an important part of the subgroup of SMA with onset at 10 to 36 months.

The basic defect of CPSMA and of other forms of SMA is not known and its clinical picture does not provide a sufficient basis for genetic classification. Therefore, we cannot be sure whether the condition we are dealing with constitutes a single entity or whether it is composed of several genetic entities. In view of the above data it is reasonable to assume that there may be three entities (table 2) as follows.

(1) The infantile chronic form characterised by age at onset usually between three and nine months with classic features of autosomal recessive inheritance with complete penetrance and the course of the disease tending to be milder in females. These cases constitute 51\% of our material (see table 2 of our previous publication\(^1\)). It should be stressed that the group of patients included here may overlap to some extent with the classic acute form of Werdnig-Hoffmann disease as defined by Pearn et al\(^10\) and Feingold et al.\(^11\) The subjects described by Pearn et al\(^12\) under the term 'subacute and chronic SMA' correspond to our infantile chronic form, but includes also (2) and (3) below.

(2) The childhood isolated form (32\% of our cases) is characterised by onset usually between 10 to 36 months with a predominance of isolated cases. Most cases described as the intermediate form or type II of the disease\(^4\)\(^14\) appear to fit well with this form, both by the range of age at onset and by the very infrequent occurrence of familial cases. The segregation ratios in this form are not compatible with autosomal recessive transmission. The assumption that this subgroup contains many dominant mutations or some phenocopies or both is more plausible.

It should be noted that Fried and Emery\(^4\) assumed that this form is recessive and deliberately excluded from their material one family in which the condition was inherited as an autosomal dominant trait. Incidentally, in their study of 14 cases there was only one other family showing features of autosomal recessive transmission. By the criteria presented here, however, this family would be included in the first form because of the age at onset under nine months.

(3) The mild childhood and adolescent form (17\% of our cases) is characterised by age of clinical onset usually after the third year of life and autosomal recessive transmission with marked sex influence, expressed by a smaller proportion of affected females and consequently by a significant reduction of penetrance.

The classification and terminology of different forms of SMA are controversial. In McKusick\(^5\) the following three forms of proximal SMA appear to be relevant to our series. (Numbers in brackets are McKusick Catalogs numbers.\(^5\)) (1) Muscular atrophy, juvenile (Kugelberg-Welander syndrome) (25 340\*). (2) Muscular atrophy, spinal, intermediate type (25 355). (3) Muscular atrophy, juvenile (Kugelberg-Welander syndrome) (15 860\*).

Initially we grouped them together because, as already stated, there were insufficient grounds for making a division. The clinical onset of CPSMA of childhood and adolescence, as our data show, may occur between birth and 18 years of life, but in our opinion there may be cases with onset of clinical expression in the early twenties. Forms 1 and 2, together with the acute form (see McKusick No 25 330\*), constitute the majority of SMA cases.

We have not included the acute form because most of our cases showed a protracted course.

*Conditions with mode of inheritance believed to be certain.

<table>
<thead>
<tr>
<th>Form of disease entity</th>
<th>Usual age at onset</th>
<th>Genetics</th>
<th>Sex influence</th>
<th>Percentage in presented material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infantile, chronic</td>
<td>3 mth-9 mth</td>
<td>R</td>
<td>Slight</td>
<td>51</td>
</tr>
<tr>
<td>Childhood, isolated</td>
<td>10 mth-36 mth</td>
<td>D mutations?</td>
<td>No evidence</td>
<td>32</td>
</tr>
<tr>
<td>Mild childhood and adolescence</td>
<td>37 mth-18 y</td>
<td>R</td>
<td>Strong</td>
<td>17</td>
</tr>
</tbody>
</table>

R=autosomal recessive.  
D=dominant.
According to Pearn et al and Feingold et al this is a distinct condition, although the available data are still ambiguous.

As we have said above, there is a possibility that some of the isolated cases in our material are new dominant mutations corresponding to form 3 (No 15 860* of McKusick). Such cases in particular may constitute an important component of the subgroup with onset at 10 to 36 months (table 1, figure).

Our data and those of other authors suggest that the following supplementary remarks and alterations in the Catalogs\(^5\) might be justifiable\(^\dag\).

1. A better delineation of the acute form (25 330*), not only by the age at clinical onset but also by the age at death according to Pearn et al and Feingold et al, and naming it 'infantile acute form of Werdnig and Hoffmann'.

2. Identifying the intermediate type of Fried and Emery\(^4\) with the 'infantile chronic' form described here (table 2) corresponding roughly to the form described by Pearn et al and adding an asterisk to the relevant No 25 355.

3. Our 'childhood isolated' form (table 2) could be considered a possible dominant mutation, identical with that designated No 15 860* in the Catalogs.\(^5\)

4. A better delineation of juvenile SMA (25 340*) corresponding to our form of 'mild childhood and adolescence' by the range of age at onset and by defining it as a sex influenced condition with reduced penetrance.

Genetic counselling

The estimation of genetic risk in CPSMA is apparently a simple matter because, in general, the disease conforms to the autosomal recessive pattern of inheritance. However, our data strongly suggest incomplete penetrance of the gene, because of the assumed 'female sparing factor' in higher age ranges, and therefore tentative values of empirical risk based on the p values can be drawn for all ages of onset subgroups, for males and females separately (table 1).

Although CPSMA of childhood and adolescence is one of the more common genetic conditions, the data on estimates of empirical risk in this disease are scanty and based on small series. The published data available are those of Bundey and Lovelace\(^15\) and Pearn et al, the latter being quoted in the book The Genetics of neurological disorders.\(^16\) In table 3 we give their risk figures together with the data from which they were drawn, as compared to our data in the same age ranges. A particularly low risk (1:15) was given by Bundey and Lovelace\(^15\) for the age at onset over 25 months. Pearn et al\(^12\) worked out the same value for 36 months onwards. We think that these figures are not reliable enough for genetic counselling in view of the very scanty series from which they were drawn. In the data of these authors, cases with age of onset over 36 months were hardly represented and a considerable proportion of the others could correspond to our subgroup of very low risk, that is, of 10 to 36 months.

Our results, although based on larger series, should also be treated with caution until more data are available. At present, the estimates of genetic risk presented in table 4 could be used for genetic counselling. If two or more sibs are affected the risk value of 1:4 should be given, independent of age at onset and sex, possibly with the exception of females in the subgroup 9 to 18 years if only male sibs were previously affected.

**Table 3. Results of segregation analysis in CPSMA with consideration of the age at onset: comparison of the present results with those of other authors.**

<table>
<thead>
<tr>
<th>Age at onset</th>
<th>Bunday and Lovelace(^13)</th>
<th>Pearn et al(^12)</th>
<th>Present series (see also ref 1)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No of probands</td>
<td>No of sibs</td>
<td>No of probands</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Affected</td>
<td>Unaffected</td>
</tr>
<tr>
<td></td>
<td>p (r)</td>
<td>p (r)</td>
<td>p (r)</td>
</tr>
<tr>
<td>Before 9 mth</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Before 24 mth</td>
<td>19</td>
<td>12</td>
<td>45</td>
</tr>
<tr>
<td></td>
<td>(0:21)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>After 24 mth</td>
<td>14</td>
<td>2</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td>(0:14)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>36 mth and later</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Overall</td>
<td>33</td>
<td>14</td>
<td>72</td>
</tr>
<tr>
<td></td>
<td>(1:6)</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

\(^{\dag}\)We take the liberty of making these comments in response to Dr McKusick's statement in the foreword to the Catalogs,\(^7\) namely that he would appreciate any suggestions aimed at increasing the usefulness of the Catalogs.
The study was supported by a grant from the Polish Academy of Sciences 10.4.05.4.1.

References


Correspondence and requests for reprints to Professor Irena Hausmanowa-Petrusewicz, Neurological Department, Medical School, ul Lindleya 4, 02005 Warsaw, Poland.
Chronic proximal spinal muscular atrophy of childhood and adolescence: problems of classification and genetic counselling.

I Hausmanowa-Petrusewicz, J Zaremba and J Borkowska

doi: 10.1136/jmg.22.5.350

Updated information and services can be found at:
http://jmg.bmj.com/content/22/5/350

**Email alerting service**

These include:

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Notes**

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/