The gene for hereditary multiple exostoses does not map to the Langer-Giedion region (8q23-q24)

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Abstract

Hereditary multiple exostoses (HME, MIM 13370) is an autosomal dominant skeletal disorder. It is characterised by exostoses usually arising in the juxtaepiphyseal region of the long bones and sometimes the pelvis or the vertebral column. Mutations are usually inherited from a parent, often the father. They are usually dominantly inherited. Hereditary multiple exostoses is caused by mutations in the TRP II (Langer-Giedion syndrome, MIM 15025) gene. TRP II (Langer-Giedion syndrome, MIM 15025), has been previously localised (pairwise linkage \( Z = -8.96 \) at \( \theta = 0 \) with probe L48 at locus \( D8S51 \)). Multipoint linkage analysis using probes L48, L24, and L1 consistently excluded the HME gene from a large area of the distal long arm of chromosome 8, spanning the smallest region of overlap assigned to the TRP II gene. These studies support the clinical view that HME and TRP II are distinct entities.

Patients and methods

In recent years, 166 new cases of HME have been observed in the Hôpital des Enfants Malades. Of the 28 families with at least three affected subjects, six families were selected for their informativeness (47 subjects including 25 affected) (fig 1).

The criterion for diagnosis of HME syndrome was the observation of at least two exostoses of the juxtaepiphysial regions of the long bones. Exostoses associated with metaphyseal dysplasia (MIM 15625) were excluded from the study.

Owing to the existence of latent forms of HME, family studies usually included radiography of the knees. Children under 3 years were excluded because of delayed expression of the disease. A 20 ml EDTA blood sample was collected from each of the 47 subjects. DNA was extracted from lymphocyte pellets by SDS lysis, proteinase K digestion, phenol/chloroform extraction, ethanol precipitation, and Tris-EDTA resuspension. Restriction digestion of genomic DNA were performed for four hours in buffer supplied by Boehringer. DNA fragments were separated by horizontal gel electrophoresis in TEA buffer, stained with ethidium bromide for visual inspection, and denatured in situ with 0.25 mol/l HCl. The denatured DNA fragments were transferred onto a nylon membrane (Zetabind, Flo Cuno) using the technique of Southern. The DNA probe was labelled with \(^{32}P\)-dCTP by nick translation and purified on a Sephadex G-50 column to remove unincorporated nucleotides. The labelled plasmid was then competitively hybridised to sheared total human DNA to reduce background caused by common repeats in the probe. The mix was then added to 25 ml of the hybridisation solution. Membranes were hybridised in polyethylene bags overnight at 65°C. They were washed twice at room temperature in 2 x SSEP and 0.1% SDS, then washed at 65°C in 1 x SSEP and 0.1% SDS.
Results
Pairwise linkage showed negative lod score values with probe L48 (Z = -8.96 at θ = 0), excluding the HME gene from close proximity to the D8S51 locus (table). Multipoint analysis using probes L48, L24, and L1 consistently excluded the HME gene from a relatively large area of the distal long arm of chromosome 8, spanning the smallest region of overlap assigned to the TRP II gene (fig 2). The other probes tested were not informative. However, using probe MCT128, we were able to exclude the HME gene from close proximity to the D8S39 locus, which maps 50 cM distal to the TRP II gene.

Discussion
In the last few years, HME, TRP I, and TRP II have been regarded as closely related diseases and the hypothesis of either contiguous genes or allelism at a single locus has been discussed.10 Here, we exclude the HME gene from close proximity to the TRP II locus in six autosomal dominant families with multiple exostoses. The present data tend to favour the view that HME and TRP are clearly distinct entities. These results are not really surprising as a number of clinical, radiological, and laboratory features allow differentiation of the three conditions.

First, neither facial dysmorphism nor ectodermal dysplasia are present in HME, while those are consistent features in TRP syndrome. Similarly, the clinical progression of exostoses stops at the end of growth in HME while it may continue throughout adult life in TRP II, resulting in severe orthopaedic complications. Secondly, radiological features are also different in the two conditions. Metaphyseal exostoses, for instance, are pediculated in HME, while in TRP II they are more sessile. Whatever the type, the association of exostoses with shortness of the ulna, fibula, and metacarpals is more specific of HME disease, while cone shaped phalangeal epiphyses are characteristic of TRP II.11 Finally, the histopathological aspects of the cartilage are strikingly different.19 In HME, the exostoses are covered by thickened periostum that closely adheres to the cartilaginous cap. It arises in contact

Autoradiographs were obtained by exposure to Kodak X/OMAT films for one to four days at -80°C. Polymorphic DNA probes were generated by Hortshemke and Lüdecke11 from a 8q24.1 specific microdissection library of the TRP II deletion: L48 (D8S51) SacI, L24 (D8S45) XmnI, and L1 (D8S42) TaqI. Five additional probes located on both sides of the deletion on the long arm of chromosome 8 were also used, namely: PYNM3 at locus D8S17,12 LTII at locus D8S5,13 R370 at locus D8S5,14 and MCT128 at locus D8S39.16 All these DNA markers showed biallelic RFLPs.

The most likely order of markers was: cen-PYNM3-LTII-L1-L48-L24-Myc-R370-L1251-MCT128-tel. Considering the uncertainty of the exact age of onset during growth, a penetrance of 0-90 was arbitrarily assigned. Linkage analysis was performed using the MLINK and LINKMAP options of version 5.1 of the LINKAGE package.15

<table>
<thead>
<tr>
<th>Pairwise linkage analysis of HME with 8q DNA markers.</th>
<th>0</th>
<th>0.1</th>
<th>0.2</th>
<th>0.3</th>
<th>0.4</th>
</tr>
</thead>
<tbody>
<tr>
<td>PYMM3</td>
<td>-4.59</td>
<td>-0.53</td>
<td>-0.13</td>
<td>0.004</td>
<td>0.022</td>
</tr>
<tr>
<td>LTII</td>
<td>-0.39</td>
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<td>0.02</td>
<td>0.04</td>
<td>0.02</td>
</tr>
<tr>
<td>L1</td>
<td>0.40</td>
<td>0.29</td>
<td>0.19</td>
<td>0.11</td>
<td>0.05</td>
</tr>
<tr>
<td>L48</td>
<td>-8.96</td>
<td>-1.37</td>
<td>-0.49</td>
<td>0.11</td>
<td>0.02</td>
</tr>
<tr>
<td>L24</td>
<td>0.047</td>
<td>0.33</td>
<td>0.35</td>
<td>0.29</td>
<td>0.17</td>
</tr>
<tr>
<td>R370</td>
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<td>0.34</td>
<td>0.23</td>
<td>0.13</td>
<td>0.05</td>
</tr>
<tr>
<td>L1251</td>
<td>-4.35</td>
<td>-0.35</td>
<td>-0.14</td>
<td>-0.06</td>
<td>-0.03</td>
</tr>
<tr>
<td>MCT128</td>
<td>-10.3</td>
<td>-1.83</td>
<td>-0.80</td>
<td>-0.29</td>
<td>-0.06</td>
</tr>
</tbody>
</table>

Figure 1 Pedigrees of the HME families studied.

Figure 2 Multipoint linkage analysis in HME. The location score is defined as twice the natural logarithm of the odds ratio.
with the columns of growth cartilage, which are in abnormal orientation at the periphery (fig 3). In contrast, in TRP II, the exostoses seem to arise as nodules linked to the cartilage of the epiphysis by a pedicle and the cells are voluminous, rich in glycogen, and arranged in ovoid groups (fig 4). Ultrastructural studies show inclusions in chondrocytes and numerous vesicles in the cytoplasm of the osteoblasts.

In conclusion, the genetic data presented here support the clinical view that HME and TRP are probably distinct entities, especially as no deletion of chromosome 8q has been hitherto reported in cells cultured from exostoses. Nevertheless, the observation of Ogle et al suggests that this condition might be heterogeneous and future linkage studies will hopefully contribute to the mapping of the gene(s) responsible for hereditary multiple exostoses.

Addendum
Three new families with HME were tested by \textit{D8S51} (L48). One family was not informative and in the other two a recombination in each was found. Pairwise linkage analysis showed a negative lod score with probe L48 at \( Z = -18.32 \) at 0 = 0.

We thank Bernhard Horsthemke for kindly providing probes of the Langer-Giedion region. We thank Gisèle Gal and Alan Strickland for their help in preparing this manuscript. This work was supported by Association Française contre les Myopathies (AFM).