Confirmation of genetic heterogeneity in familial psoriasis

Deborah Matthews, Lionel Fry, Anne Powles, Jean Weissenbach, Robert Williamson

Abstract
Psoriasis affects approximately 2% of the European population and is often familial. Linkage of a subset of psoriasis families to loci on chromosome 17q has recently been reported. We have studied members of a large multiply affected family from the north east of England and analysed genotypes for markers from 17q, including the polymorphic microsatellite markers AFM210x5, AFM163y1, AFM044x3, AFM355y1, and AFM217yd19. Two point and multipoint analysis clearly show exclusion of linkage between the telomeric region of 17q and psoriasis in this family. This confirms the genetic heterogeneity of psoriasis and the existence of at least one other major psoriasis locus.


Psoriasis presents with red scaly patches on the skin, characterised by epidermal hyperplasia and an inflammatory cell infiltration into both the dermis and epidermis. There is also a large increase in cytokine production. Maturation of keratinocytes is incomplete and cell division occurs at an earlier point in the cell cycle, resulting in skin up to 16 times the thickness of normal skin.

Psoriasis affects about 2% of the European population. Evidence that psoriasis has a genetic component has come from studies reporting that approximately 90% of those affected have a family history of the disease, with 8% of first degree relatives and 3% of second degree relatives affected. Twin studies have shown a concordance rate of 70% in monozygotic twins and 23% in dizygotic twins.

The phenotype of psoriasis is variable. There are several different clinical types; the most common form, affecting 90% of patients, is plaque forming psoriasis. However, less frequent guttate psoriasis may occur in the same family as plaque forming psoriasis. Five percent of psoriatics go on to develop arthritis.

The usual age of onset of psoriasis is between 15 and 25; it may, however, occur at any age, but once present it rarely regresses. The onset may be preceded by one of several environmental triggers, including stress, change of climate, trauma to the skin and, infections, particularly streptococcal infection.

Various HLA associations have been reported with psoriasis. There is increased frequency of the occurrence of class I antigens B13, B17, Cw6, and Cw7, and class II antigens DR7 and DR4. B13, B57, and Cw6 show a particularly strong association with age of onset and family history. There is no evidence for linkage between the disease and HLA loci. However, the HLA associations and the fact that several treatments for psoriasis are immunosuppressive support the hypothesis that psoriasis is an autoimmune disease.

Using a dominant model of inheritance with full penetrance, Tomofahrde et al. showed a linkage of familial psoriasis to the distal end of chromosome 17q in a proportion of white American families. A maximum two point lod score of 5.33 was obtained in one family with marker D17S784 at a recombination value of 0.04; other families appeared to show exclusion, but only at full penetrance. The aim of this study was to determine if the gene predisposing to psoriasis is located on chromosome 17q in a large English family, or if heterogeneity of the locus for psoriasis can be confirmed in our family.

Patients
Twenty seven people from one family participated in this study. Consent was obtained from all people and each was given a full medical examination. Medical histories were taken, including age of onset, treatments, and other medical conditions. On the basis of this information members of the family were assigned the following status: 12 affected, 11 unaffected, and four of unknown affected status.

Persons were designated “affected” if they had clear psoriasis lesions. Unaffected subjects have no history of the disease and no dermatological signs on physical examination. The four people in generation IV designated “unaffected status” are all over the age of 25 and have absolutely no clinical signs of psoriasis. However, the four people in the same generation designated “unknown status” have signs which indicate they may develop the disease, but which are not definitive, such as pitted nails and dry skin on the elbows and scalp.

The affected status of II-2 is not known as he died before this study started and no clinical data are available.

Methods
The DNA was extracted from 10 ml of blood using standard methods. The polymorphic markers AFM210x5, AFM163y1, AFM044x3, and AFM217yd10 were selected from a chromosomal map of 17q. The se-
Confirmation of genetic heterogeneity in familial psoriasis

Figure 1  Chromosome 17q haplotypes. Marker order from top to bottom: AFM210xa5, AFM163yg1, AFM044xg3, AFMa353yg1, and AFM217yd10. Solid symbols: affected persons; open symbols: unknown affected status; N: unaffected persons. Inferred haplotypes are given in brackets.

...quence of AFMa353yg1 was obtained directly from Genethon, but is now available on public databases. The markers were used to genotype the family according to standard conditions. CEPH DNA 1347-02 was included on each electrophoresis gel as a control of allele size. The marker allele frequencies were based on the genotypes of a subset of the CEPH panel.

Familial psoriasis is assumed to be an autosomal dominant trait, although the penetrance value is unknown; the penetrance was therefore varied between 0·5 and 0·99. The MLINK program (version 5·2) was used to calculate two point lod scores between psoriasis and each individual microsatellite marker. Multipoint linkage analysis was performed using the LINKMAP program. The genetic map used for this was telomere-AFM217yd10(0·11)-AFMa353yg1(0·01)-AFM044xg3(0·04)-AFM163yg1(0·06)-AFM210xa5. The recombination fractions between adjacent loci are in parentheses.

Results

Using markers AFM210xa5, AFM163yg1, AFM044g3, AFMa353yg1, and AFM217yd10 the patients were genotyped and haplotypes were derived. The segregation of the haplotypes within the family can be seen in Fig 1.

Pairwise lod scores were generated between each marker and psoriasis. The population prevalence of psoriasis was assumed to be 2% and the penetrance was varied between 0·5 and 0·99. The pairwise lod scores at penetrance values of 60%, 80%, and 99% are shown in the table. The maximum lod score obtained by Tomfohrde et al was 5·33 with locus D17S784 (AFM044xg3) at 4 cM and 99% penetrance. With this family the corresponding lod score (D17S784 at 4 cM and 99% penetrance) is −3·146.

Multilocus linkage analysis on the same data produced further evidence that the psoriasis locus and the distal region of 17q are not linked.

<table>
<thead>
<tr>
<th>Penetrance</th>
<th>Marker</th>
<th>Recombination fraction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0.00</td>
</tr>
<tr>
<td>0.99</td>
<td>AFM210xa5</td>
<td>-6.481</td>
</tr>
<tr>
<td></td>
<td>AFM163yg1</td>
<td>-2.673</td>
</tr>
<tr>
<td></td>
<td>AFM044xg3</td>
<td>-4.463</td>
</tr>
<tr>
<td></td>
<td>AFMa353yg1</td>
<td>-5.345</td>
</tr>
<tr>
<td></td>
<td>AFM217yd10</td>
<td>0.085</td>
</tr>
<tr>
<td>0.80</td>
<td>AFM210xa5</td>
<td>-2.615</td>
</tr>
<tr>
<td></td>
<td>AFM163yg1</td>
<td>-1.074</td>
</tr>
<tr>
<td></td>
<td>AFM044xg3</td>
<td>-2.047</td>
</tr>
<tr>
<td></td>
<td>AFMa353yg1</td>
<td>-1.735</td>
</tr>
<tr>
<td></td>
<td>AFM217yd10</td>
<td>0.089</td>
</tr>
<tr>
<td>0.60</td>
<td>AFM210xa5</td>
<td>-1.786</td>
</tr>
<tr>
<td></td>
<td>AFM163yg1</td>
<td>-0.543</td>
</tr>
<tr>
<td></td>
<td>AFM044xg3</td>
<td>-1.476</td>
</tr>
<tr>
<td></td>
<td>AFMa353yg1</td>
<td>-1.089</td>
</tr>
<tr>
<td></td>
<td>AFM217yd10</td>
<td>0.100</td>
</tr>
</tbody>
</table>
penetrance.

Figure 2: Multipoint exclusion maps of the distal region of 17q at 60%, 80%, and 90% penetrance.

in this family. The program Linkmap was used to perform likelihood calculations on fitting the psoriasis locus into the intervals on the 17q map. Fig 2 is the exclusion map produced. The three maps were calculated for 60%, 80%, and 90% penetrance.

Discussion

Our data show that psoriasis is not linked to the 17q region in this family. This confirms that familial psoriasis is a genetically heterogeneous disease with at least two loci involved, one at 17q and one other. The result is not surprising as it is in agreement with population studies which concluded that the inheritance of psoriasis could not be explained as a simple dominant disorder resulting from mutations at one locus. 28-30

The large clinical variation observed with psoriasis can also be more easily explained if psoriasis is caused by mutations at more than one gene. Whether the two or more genes involved act independently or whether epistasis is involved is yet to be determined.

We thank all the patients who participated in this study, particularly Thomas Cockburn for his help in contacting and organizing the family members, and Dr. Jean Weissenbach, for marker sequence data in advance of publication. This study was supported by Eurogen.

Confirmation of genetic heterogeneity in familial psoriasis.

D Matthews, L Fry, A Powles, J Weissenbach and R Williamson

doi: 10.1136/jmg.32.7.546

Updated information and services can be found at:
http://jmg.bmj.com/content/32/7/546

These include:

**Email alerting service**

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/