Genetic analysis of PSORS2 markers in a UK dataset supports the association between RAPTOR SNPs and familial psoriasis

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Psoriasis [MIM 177900] is a chronic and disfiguring skin disorder, which is inherited as a multifactorial trait.1 Genome wide scans have repeatedly mapped a major disease susceptibility locus (PSORS1) to the Major Histocompatibility Complex (MHC), on chromosome 6p21 (reviewed by Capon et al2). Outside of the MHC, at least eight additional susceptibility intervals have been reported (PSORS2-9) (reviewed by Capon et al3). The PSORS2 interval [MIM 602723] was originally mapped to chromosome 17q25, in a sample of extended pedigrees presenting with disease segregation across multiple generations.3 Linkage to PSORS2 was later replicated in independently ascertained cohorts.4–6 Conversely, two distinct genome wide scans carried out by our group failed to detect any evidence for linkage to PSORS2, in United Kingdom cohorts of European descent.5,6

High density genetic analysis of the PSORS2 interval recently identified two distinct association peaks, both defined by a small number of non-coding single nucleotide polymorphisms (SNPs).7 The proximal peak spans a 20 kb genomic segment where a putative susceptibility allele, mapping between the SLC9AR1 and NAT9 genes, abolishes a RUNX1 binding site. The less characterised distal region of association lies 6 Mb away, within intron 3 of the RAPTOR gene [MIM *607130]. To define the relevance of PSORS2 genetic variation in the United Kingdom, we have analysed eight representative SNPs, selected from both association peaks. While our results do not support a pathogenic involvement of SLC9AR1/NAT9 variants, they do provide evidence for association between familial psoriasis and RAPTOR SNPs.

METHODS
Our patient cohort included a total of 233 independent parent-offspring trios of northern European origin. Of these, 116 were sampled from the family cohort previously described by Veal et al.6 The 117 remaining trios were ascertained through an affected proband, as detailed elsewhere.6 All samples were collected following approval of the Guy’s and St Thomas’ hospitals ethics Committee of King’s College, London. All subjects participating in this study gave their informed consent. Based on the reported significance of disease association the following five markers were selected from the proximal association peak: SNP8 (rs7420); SNP9 (rs734232, abolishing the RUNX1 site); SNP11 (rs895691); SNP12 (rs127977); and SNP15 (rs2305214). Based on the same criterion, three further SNPs (rs1564864, rs2019154, and rs869190) were selected from the distal association peak. Following DNA extraction from blood lymphocytes, SNP genotyping was carried out by fluorescence polarisation template directed dye incorporation as described by Speckman et al8 or with Sequenom MassArray technology (Sequenom Inc). Family based association analysis was carried out using the TRANSMIT 2.5 software9 to examine the transmission rates of marker alleles.

RESULTS AND DISCUSSION
Association analysis results are summarised in table 1. We first examined the entire dataset and found that rs2019154 (distal peak) was the only marker supporting evidence for association (p = 0.027). To reduce the heterogeneity of our cohort, we subsequently restricted the analysis to the 116 trios that had been sampled from extended pedigrees, thus defining a subset of patients with a well documented family history of psoriasis. This stratification did not modify the outcome of the proximal peak genetic analysis. However, the significance of disease association increased for all three SNPs mapping to RAPTOR SNP rs2019154 yielding a p value of 0.008. Hitherto, analysis of UK samples has failed to replicate linkage to the PSORS2 locus. The present findings emphasise the power afforded by association studies of complex trait susceptibility intervals.

Key points
• Psoriasis is a multifactorial skin disorder. The major disease susceptibility locus (PSORS1) maps to the MHC region on chromosome 6p21. Eight additional PSORS loci have been identified outside of the MHC, with the PSORS2 interval showing linkage to psoriasis in a range of independently ascertained samples.
• High density genetic analysis of the PSORS2 locus recently identified two distinct association peaks, encompassing respectively the SLC9AR1-NAT9 genes and the third intron of the RAPTOR gene. To assess the relevance of these findings to the UK population, we have analysed eight representative single nucleotide polymorphisms (SNPs), selected from both the association peaks, in a large dataset of 233 parent-offspring trios.
• The examination of the entire dataset only provided evidence for association at RAPTOR SNP rs2019154 (p = 0.027). Restricting the analysis to the 116 trios with a documented family history of psoriasis increased the significance of disease association for all three RAPTOR SNPs, with rs2019154 yielding a p value of 0.008.
• Hitherto, analysis of UK samples has failed to replicate linkage to the PSORS2 locus. The present findings emphasise the power afforded by association studies of complex trait susceptibility intervals.

Abbreviation: SNP, single nucleotide polymorphism
The failure to detect association with SNPs from the SLC9A1R1/NAT9 genomic segment is in keeping with the low relative risk that is conferred by these variants and suggests that a larger dataset might be needed to replicate the significance values reported by Helms et al. Conversely, our observation of increased RAPTOR association in the familial subset of our sample argues for strict selection criteria in patient recruitment. In this context, it is of interest that the location of the RAPTOR gene closely matches that of the original PSORS2 interval, as defined by parametric linkage analysis of multigeneration pedigrees. This suggests that RAPTOR SNPs might be specifically implicated in familial psoriasis, for example by acting as a modifier of major susceptibility loci.

The observation of significant association at the RAPTOR locus is in contrast with our previous failure to detect linkage with the PSORS2 interval. This discrepancy could be accounted for by several factors, including the informativity and spacing of the microsatellites used in our genome scans. Moreover, the psoriasis associated alleles from this region are accounted for by several factors, including the informativity of genomic intervals linked to complex traits. Mathematical modelling has clearly shown that association studies offer a greater power to detect minor susceptibility loci, compared with linkage analysis.12 Our findings demonstrate this point on experimental data, emphasising the insight afforded by association studies in the characterisation of genomic intervals linked to complex traits.

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