Genome-wide association study identifies a psoriasis susceptibility locus at TRAF3IP2


Psoriasis is a multifactorial skin disease characterized by epidermal hyperproliferation and chronic inflammation, the most common form of which is psoriasis vulgaris (PsV). We present a genome-wide association study of 2,339,118 SNPs in 472 PsV cases and 1,146 controls from Germany, with follow-up of the 147 most significant SNPs in 2,746 PsV cases and 4,140 controls from three independent replication panels. We identified an association at TRAF3IP2 on 6q21 and genotyped two SNPs at this locus in two additional replication panels (the combined discovery and replication panels consisted of 6,487 cases and 8,037 controls; combined P = 2.36 × 10^{-10} for rs13210247 and combined P = 1.24 × 10^{-16} for rs33980500). About 15% of psoriasis cases develop psoriatic arthritis (PsA). A stratified analysis of our datasets including only PsA cases (1,922 cases compared to 8,037 controls, P = 4.57 × 10^{-12} for rs33980500) suggested that TRAF3IP2 represents a shared susceptibility for PsV and PsA. TRAF3IP2 encodes a protein involved in IL-17 signaling and which interacts with members of the Rel/NF-κB transcription factor family.

Psoriasis is a chronic immune-mediated and hyperproliferative disorder of the skin that affects up to 3% of individuals in populations of European ancestry. The most common form, PsV, is characterized by red, raised, scaly plaques that commonly occur on the elbows, knees, scalp and lower back. Disease concordance in monozygotic twin pairs amounts to at most 70% (ref. 5), and the sibling recurrence risk, s, of PsV has been estimated to range between 4 and 11 (refs. 5, 6). Part of the genetic susceptibility can be explained by the established susceptibility locus at PSORS1 in the human leukocyte antigen (HLA) complex on chromosome 6p21.3 (HLA-C) as well as polymorphisms at the IL12B (ref. 7), IL23R (refs. 8–11), IL4-IL13 (ref. 12), IL23A, TNIP1 and TNFAIP3 (ref. 13) loci. Because these susceptibility loci only account for a λc ≤ 1.35 (ref. 13), a large fraction of the heritability for PsV remains unexplained. In addition, a deletion of two LCE cluster genes has been proposed to be a risk factor for the development of psoriasis, implying that structural variants could also contribute to overall disease susceptibility.

To identify additional PsV susceptibility loci, we performed genome-wide SNP genotyping of 487 German PsV cases and 1,161 controls (screening panel A; Supplementary Table 1) using the Illumina HumanHap 550K array. For genotype imputation with phased HapMap data as a reference and for subsequent statistical analysis, we used a dataset that passed stringent quality control filters. This cleaned dataset consisted of 504,742 autosomal SNPs genotyped in 472 cases and 1,146 controls. Imputation served to considerably increase the genomic coverage of our study to a total number of 2,339,118 SNPs. Conservatively accounting for multiple testing by Bonferroni correction, the threshold for genome-wide significance was P ≤ 2.14 × 10^{-8} in the imputed dataset. A moderate genomic control value of λGC = 1.065 indicated a minimal overall inflation of the test statistics due to population stratification. Furthermore, the multidimensional scaling analysis showed genuine European ancestry of panel A samples, and identity-by-state analysis revealed neither non-European outliers nor cryptically related individuals after quality control screening (Supplementary Fig. 1). Our screening panel A included 177 cases and 2,034 controls from Germany and 122 cases and 2,220 controls from Canada.

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had 80% power to detect a variant with an odds ratio of 1.38 or higher at the 5% significance level, assuming a frequency of the disease-associated allele of at least 30% in controls.

The initial comparison of case-control frequencies confirmed the association of PsV with the established susceptibility loci at HLA-C (rs12191877, \( P = 4.21 \times 10^{-32} \), odds ratio (OR) = 2.79, 95% CI 2.35–3.33) and IL12B (rs2546890, \( P = 8.83 \times 10^{-8} \), OR = 0.65, 95% CI 0.56–0.76). Suggestive evidence for association was found for IL23R (rs1004819, \( P = 6.17 \times 10^{-4} \), OR = 1.33, 95% CI 1.13–1.57) and on 1q21 22 kb upstream of LCE3C (rs4112788, \( P = 6.36 \times 10^{-4} \), OR = 0.76, 95% CI 0.64–0.89) (Supplementary Fig. 2). The lead SNPs rs2546890 and rs1004819 were in moderate linkage disequilibrium (LD) (\( r^2 > 0.50 \)) with the previously described associated variants at the IL12B (rs7709212)\(^9,\)\(^12\) and IL23R (rs2201841)\(^13\) loci, respectively.

To identify additional susceptibility loci, we excluded all SNPs within the extended HLA complex (chromosome 6p21 at 25–34 Mb) and visually inspected the cluster plots of all genotyped index SNPs with a \( P \) value less than 0.001 after the clumping procedure. The 180 strongly associated SNPs were subsequently selected for replication analysis and genotyped in the German replication panel B (681 cases and 1,824 controls). After quality control, we excluded 33 follow-up SNPs.

In addition, an in silico replication was performed using available genome-wide association study (GWAS) datasets for panel C from the United States (1,303 cases and 1,322 controls; Collaborative Association Study of Psoriasis (CASP))\(^13\) and for panel D from Canada (762 cases and 994 controls; S.D., J.V.R., M.B., H.F. and C.R., unpublished data). We imputed both datasets, that is, the CASP dataset, which was genotyped by Perlegen Sciences, and the Canadian dataset, which was generated using the Illumina Human 1M BeadChip. Of the 147 SNPs genotyped in panel B, we obtained genotypes for 126 SNPs in panel C and for 144 SNPs in panel D. Detailed association results including genotype counts of the 147 SNPs are given in Supplementary Table 2.

Two SNPs upstream of IL12B, included as a positive control for our experiment, achieved genome-wide significance in the combined replication panels B through D (rs2546890: replication stage \( P = 9.07 \times 10^{-15} \), rs953861: replication stage \( P = 6.50 \times 10^{-14} \)). A previously unidentified association in the combined analysis of the three replication panels was obtained for rs13210247 (replication stage \( P = 6.25 \times 10^{-6} \)), risk allele frequency for cases = 0.09, risk allele frequency for controls = 0.07 with association robust to Bonferroni correction (corrected \( P \) value for panels A through F = 2.36 \times 10^{-10})). This intronic SNP is located in TRAF3IP2, which encodes the TRAF3 interacting protein 2 and which is also known as ACT1, encoding the NF-kB activator 1 protein. In the combined panels A through D, consisting of a total of 3,218 PsV cases and 5,286 controls, this SNP achieved genome-wide significance (combined \( P = 7.31 \times 10^{-9} \) (Fig. 1). We observed suggestive evidence for association for four other SNPs; however, these associations did not remain significant after Bonferroni correcting for multiple testing. These four SNPs are located in RYR2 (rs2485558, replication stage \( P = 0.01 \), combined \( P = 1.53 \times 10^{-5} \)), DPP6 (rs916514, replication stage \( P = 0.0496 \), combined \( P = 6.03 \times 10^{-4} \)), MMP27 (rs1939015, replication stage \( P = 0.026 \), combined \( P = 7.09 \times 10^{-4} \)) and 31 kb downstream of NFkBIA (rs2145623, replication stage \( P = 2.29 \times 10^{-3} \), combined \( P = 5.01 \times 10^{-4} \)), respectively (Table 1).

We also confirmed the previously reported association of PsV and the LCE3C-LCE3B deletion\(^14\) at \( P = 3.80 \times 10^{-3} \) (for details on genotyping and copy number analyses, see Online Methods).

To further validate the newly associated PsV susceptibility gene TRAF3IP2, we genotyped the lead SNP, rs13210247, in the two additional case-control panels E (1,987 cases and 1,661 controls from Michigan) and F (1,282 cases and 1,090 controls from Canada). The combined replication panels B through F comprised 6,015 PsV cases and 6,891 healthy controls, yielding a replication \( P \) value of \( 1.00 \times 10^{-7} \) with the same direction of effect for all five panels (combined \( P \) value for panels A through F = 2.36 \times 10^{-10})).

We additionally selected all validated coding SNPs within TRAF3IP2 from the NCBI dbSNP\(^16\) build 130 (rs1043730, rs13190932 and rs33980500) for genotyping in replication panel B. Of the existing three missense SNPs, only rs33980500, located in exon 2, showed a significant association (\( P = 0.0265 \), OR = 1.28, 95% CI 1.03–1.58). Therefore, we also genotyped this SNP in the replication panels C through F. Association analysis of the obtained replication cohort (panel B through F) yielded a highly significant \( P \) value of \( 8.00 \times 10^{-14} \) (combined \( P \) value for panels A through F = 1.24 \times 10^{-16} (Table 2). Because the missense SNP was not available from the HapMap II CEU imputation reference, we performed imputation based on CEU haplotypes generated by the 1000 Genomes Project for panels A, C and D in order to fine map a region of 700 kb around TRAF3IP2 containing 1,935 SNPs (Fig. 1).

The intronic SNP rs13210247 in replication panel B and the missense SNP rs33980500 were genotyped in the replication panels, and the combined \( P \) values of panels A through F are indicated for both SNPs (Table 2). The magnitude of linkage disequilibrium (LD) with the central SNP rs13210247, measured by \( r^2 \), is reflected by the color of each SNP symbol (for color coding, see upper right corner of the plot). Recombination activity (cM/Mb) is depicted by a blue line.

Figure 1 Regional plot of TRAF3IP2. Regional plot of the negative decadic logarithm of the combined \( P \) values from the imputed panels A, C and D in an ~700-kb window around the intronic lead SNP rs13210247 (blue filled diamond) and the missense SNP rs33980500 (\( r^2 = 0.63 \)). Panels A, C and D were imputed with CEU haplotypes generated by the 1000 Genomes Project (August 2009 release) as a reference. The strongest signal of association was confined to TRAF3IP2. The intronic SNP rs13210247 and the missense SNP rs33980500 were genotyped in the replication panels, and the combined \( P \) values of panels A through F are indicated for both SNPs (Table 2). The magnitude of linkage disequilibrium (LD) with the central SNP rs13210247, measured by \( r^2 \), is reflected by the color of each SNP symbol (for color coding, see upper right corner of the plot). Recombination activity (cM/Mb) is depicted by a blue line.

The missense SNP rs33980500 (Asp19Asn) causes a mutation from aspartic acid to asparagine in the protein sequence and the resulting
change in charge (a negative electric charge to nonpolar) might have an impact on the protein structure and, hence, its function. Additionally, the mutation is located in a region that is more than 90% conserved among different species (Supplementary Fig. 3), which further implies a possible functional consequence of this change.

Furthermore, we performed in silico analyses and found a putative TRAF6-binding peptide (amino acids 13 to 21) containing the SNP Asp19Asn. This peptide is similar to a known TRAF6-binding motif of CD40, a member of the tumor necrosis factor receptor superfamily. Notably, the mutation of the accordant seventh peptide residue (asparagine to aspartate) has previously been observed to change the binding affinity of TRAF6 to CD40 significantly. Therefore, rs33980500 may also affect the interaction of TRAF3IP2 to TRAFs and, thus, the involved inflammatory pathways.

About 15% of individuals with psoriasis develop psoriatic arthritis (PsA), which is an inflammatory, disabling arthritis. This suggests that PsV and PsA share common susceptibility factors.

To determine the association of the TRAF3IP2 missense SNP with PsA, we performed a PsA stratified analysis of panels A through F. Association analysis of all individuals with PsA and controls within our combined sample (1,919 cases and 8,037 controls) yielded a P value of 4.57 × 10⁻¹² for rs33980500 (OR = 1.57, 95% CI 1.38–1.78). In an equally sized subgroup of randomly extracted PsV cases without PsA, we obtained a P value of 2.04 × 10⁻⁶ (OR = 1.37, 95% CI 1.20–1.57). The overlap in confidence intervals for the two associations, along with the fact that no significant difference was obtained when comparing the 1,919 PsA cases versus the randomly selected 1,919 PsV cases suggests that the TRAF3IP2 gene locus represents a shared susceptibility locus for PsA and PsV.

Gene expression of psoriatic and unaffected skin differs significantly for hundreds of genes that are involved in immune response or in the regulation of cellular differentiation and proliferation. To check if altered gene activity might trigger disease progression, we examined expression levels of several loci in biopsies from 57

Table 2 Summary of association results for the two TRAF3IP2 SNPs genotyped in additional replication panels

<table>
<thead>
<tr>
<th>Chr. position (bp)</th>
<th>dbSNP ID</th>
<th>Nearby genes (relative position)</th>
<th>A1</th>
<th>A2</th>
<th>A1</th>
<th>A2</th>
<th>A1</th>
<th>A2</th>
<th>A1</th>
<th>A2</th>
<th>A1</th>
<th>A2</th>
<th>OR (95% CI)</th>
<th>P repl.</th>
<th>P comb.</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>112,029,413</td>
<td>rs13210247 (intron)</td>
<td>G</td>
<td>0.10</td>
<td>0.06</td>
<td>0.07</td>
<td>0.09</td>
<td>0.07</td>
<td>0.09</td>
<td>0.07</td>
<td>0.09</td>
<td>0.07</td>
<td>1.00 × 10⁻⁷</td>
<td>2.36 × 10⁻¹⁰</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>112,019,955</td>
<td>rs33980500 (missense)</td>
<td>C</td>
<td>0.10</td>
<td>0.06</td>
<td>0.11</td>
<td>0.07</td>
<td>0.09</td>
<td>0.06</td>
<td>0.11</td>
<td>0.08</td>
<td>0.11</td>
<td>0.07</td>
<td>8.00 × 10⁻¹⁴</td>
<td>1.24 × 10⁻¹⁶</td>
</tr>
</tbody>
</table>

The lead SNP rs13210247 and the missense SNP rs33980500 were genotyped in five independent PsV case-control panels (B through F; Supplementary Table 1). The number of cases and controls of each panel is shown in the top column. Nucleotide positions refer to NCBI build 36. Chr. denotes the rare allele and A2 is the common allele. Allele frequencies of A1 are shown (AF). The combined P value of the meta-analysis is shown for the independent replication panels B through D (P_repl.) as well as for the GWAS panel together with the independent replication panels (panels A through F; P_comb.).
healthy controls and compared these to expression levels of biopsies of involved and uninvolved skin from 53 PsV cases (subgroup of panel C). We analyzed the TRAF3IP2 locus and some genes that potentially act downstream of TRAF3IP2. For the TRAF3IP2 locus, the expression level was slightly altered between involved and uninvolved skin (P = 2.2 × 10⁻⁵) (Supplementary Fig. 4). However, the dosage of risk alleles at the according SNPs did not correlate with transcript levels for the genes in involved, uninvolved or normal skin (Supplementary Table 3).

Genetic variants in the TRAF3IP2 locus are implicated in PsV and PsA susceptibility for the first time, to our knowledge by our study. The gene product of TRAF3IP2 is a positive signaling adaptor required for IL-17-mediated T-cell immune responses. TRAF3IP2 interacts with tumor necrosis factor receptor-associated factor (TRAF) proteins and either IκB kinase or mitogen-activated protein kinase to activate either NF-κB or JNK kinase. Upon recruitment to CD40 and the BAFF receptor in B cells, TRAF3IP2 also negatively regulates B-cell survival through its interaction with TRAF3 (ref. 22). The negative regulation of CD40-BAFF–mediated B-cell functions results in ‘hyper-T-cell-dependent’ and T-cell–dependent immune responses in TRAF3IP2-deficient mice. Researchers in a previous study also found that TRAF3IP2 is a key component of IL-17–mediated signaling and that IL-17–mediated gene expression is completely abolished in TRAF3IP2-deficient mouse embryonic fibroblasts. In addition, epithelial cells have been identified as the critical cell type in which TRAF3IP2 mediates IL-17–dependent inflammatory disease. In PsV, a subset of T cells expressing IL-17 plays a major role. The IL-17–expressing T cells that are abundant in the epidermis of psoriatic lesions but which are absent in healthy donor epidermis, mainly IL-17–expressing CD4+ (but also CD8+) T cells are strongly increased relative to healthy skin. Epidermal hyperplasia and the production of innate immune peptides such as human β-defensin 2 (hBD-2) are associated with these IL-17+ T cells. The crucial role of IL-17 is also supported by the observation that in psoriasis, treatment response to TNF inhibitors correlates with suppressed IL-17 signaling. Moreover, we showed that in immortalized N-TERT keratinocytes (N-TERT KCs) and in normal human keratinocytes (NHKs), IL-17A and TNF-α together induce the expression of DEF6b4 (encoding hBD-2), which is absent in N-TERT KCs and NHKs with silenced TRAF3IP2 (Supplementary Fig. 5). In epithelial cells, the binding of IL-17A and/or IL-17F to the heterodimeric IL-17R leads to the recruitment of TRAF3IP2 through homotypic interactions between conserved (SEFIR) domains. This allows the incorporation of TRAF6 into the signaling complex followed by downstream activation of the NF-κB and mitogen-activated protein kinase pathways. We thus speculate that a dysregulation of TRAF3IP2 might have a major impact on IL-17 signaling and, hence, on the activation of NFκB-pathways, leading to the upregulation of pro-inflammatory factors.

For the TRAF3IP2 locus newly identified here, fine mapping and resequencing efforts together with extensive functional studies are required to detect all potential causal variants and thus to specify the contribution of the locus to overall disease susceptibility.

METHODS

Methods and any associated references are available in the online version of the paper at http://www.nature.com/naturegenetics/.

Accession codes. The CASP GWAS dataset (panel C) is deposited in dbGaP under the accession code phs000019.v1.p1.

Note: Supplementary information is available on the Nature Genetics website.

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AUTHOR CONTRIBUTIONS


COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Recruitment of cases and controls. Samples were organized in panels that corresponded to the successive steps of the present study. All individual panels (A–F) were independent from each other.

All German cases in panels A and B were recruited either at the Department of Dermatology of the Christian-Albrechts-University Kiel or the Department of Dermatology and Allergy of the Technical University Munich through local outpatient services. Individuals were considered to be affected if chronic plaque or guttate psoriasis lesions covered more than 1% of their total body surface area or if at least two skin, scalp, nail or joint lesions were clinically diagnosed as being caused by psoriasis. Psoriatic arthritis (PsA) was diagnosed by a clinical finding of joint complaints and radiologic and rheumatologic confirmation by criteria according to Moll and Wright27, or more recently, to the Classification Criteria for Psoriatic Arthritis (CASPAR)28.

2,510 German healthy control individuals in panels A and B were obtained from the PopGen biobank29. 483 German healthy controls were selected from the KORA S4 survey, an independent population-based sample from the general population living in the region of Augsburg, southern Germany30.

The American study population (panel C) consisted of 1,303 psoriasis cases and 1,322 controls after quality control measures. The datasets used for the analyses described in this manuscript were obtained from the database of Genotype and Phenotype (dbGaP). The genotyping of samples was provided through the Genetic Association Information Network (GAIN). Samples and associated phenotype data were provided by the Collaborative Association Study of Psoriasis (CASP). Funding support for CASP was provided by the US National Institutes of Health, the Foundation for NICH's Genetic Information Network and the National Psoriasis Foundation.

The Canadian population sample used for panel D consisted of 762 psoriasis cases and 994 controls sampled from the Quebec founder population (QFP). Membership in the QFP was defined as having four grandparents with French-Canadian family names who were born in the Province of Quebec, Canada or in adjacent areas in the provinces of New Brunswick and Ontario or in New England or New York State. This criterion assured that all subjects were descendants of French-Canadians living before the 1960s, after which time admixture with non-French-Canadians became more common.

For replication of the two TRAF3IP2 SNPs (rs13210247 and rs33980500), panel E was tested for an association. This panel comprised 1,987 psoriasis cases and 1,661 controls of European ancestry. The sample was collected at the University of Michigan, and selection criteria for cases included a requirement of at least two psoriatic plaques or a single plaque occupying >1% of an individual’s total body surface area outside the scalp. Individuals that presented only palmoplantar psoriasis, inverse psoriasis or sebopsoriasis were excluded. Individuals used as controls were older than 18 years and had no history of psoriasis and no family history of psoriasis.

Panel E also used only for replication of the two TRAF3IP2 SNPs, consisted of 1,282 psoriasis cases and 1,090 controls from Toronto and Newfoundland, Canada. Psoriasis was diagnosed by a dermatologist. Control individuals showed no evidence of psoriasis, rheumatoid arthritis or other autoimmune disorders.

Written, informed consent was obtained from all study participants, and all protocols were approved by the respective institutional ethical review committees of the participating centers.

SNP genotyping for the genome-wide screen. The genotyping for the GWAS, which was part of the German GWAS initiative funded by the National Genome Research Network (NGFN), was performed by Illumina’s service facility using the Illumina HumanHap 550K v1 with 561,466 SNP markers. All experimental steps were carried out according to standard protocols.

We excluded five samples with more than 10% missing genotypes (that is, having a call rate <90%). Individuals who showed statistically relevant genetic dissimilarity to the other subjects (population outliers) or who showed evidence for cryptic relatedness to other study participants (unexpected duplicates, first- or second-degree relatives) were removed (Supplementary Fig. 1). These quality control measures left 472 psoriasis samples and 1,146 control samples for inclusion in screening panel A. All gender assignments could be verified by reference to the proportion of heterozygous SNPs on the X chromosome. Before analysis, we excluded 56,724 markers (10% of the total number of SNPs) that had a low genotype call rate (<95% in cases or controls; n = 4,254), were monomorphic or rare (minor allele frequency <2% in cases or controls; n = 30,862), deviated from Hardy-Weinberg equilibrium (HWE) in the control sample (HWE P < 0.01; n = 7,600) or that were nonautosomal SNPs (n = 14,008).

Imputation. Genotype imputation was performed using a hidden Markov model algorithm implemented in the software program MACH v.1.0.1633 to infer missing genotypes in silico. As a reference, HapMap II CEU phased haplotypes22 were used.

As input for the imputation, only genotyped SNPs that passed quality control were used. Of the imputed SNPs, we analyzed only those SNPs that could be imputed with a relatively high confidence (estimated r2 between imputed SNP and true genotypes >0.3), had a minor allele frequency >2% in cases or controls and a HWE P value < 0.01 in the control sample. To take imputation uncertainty into account, we used allelic dosage association as implemented in the program MACH2DAT31. The allelic dosage is the weighted sum of the genotype class probabilities.

SNP selection for replication. SNPs in the genome-wide scan that passed quality control were analyzed using gPLINK v2.049 in combination with PLINK v1.0532. The SNP list (including imputed and genotyped SNPs) was pruned for redundancy due to linkage disequilibrium by using the −clump command in PLINK. For all genotyped index SNPs with P < 10−8, a visual inspection of the cluster plots was performed. SNPs that did not pass the visual inspection were excluded from further analyses. The 180 most strongly associated index SNPs of the clumps (P < 2.6 × 10−4) were ordered as genotyping assays. If the genotyping assay design was not possible for a SNP, the next best SNP from the clump was chosen for the assay design.

SNPlex and TaqMan genotyping. The ligation-based SNPlex genotyping system and functionally tested TaqMan SNP Genotyping Assays (Applied Biosystems) were used to genotype variants in replication panel B. For technical replication of the results of the two TRAF3IP2 SNPs, TaqMan SNP Genotyping Assays were used in panels A–E and the Sequenom Platform was used for panel F.

Of the 180 selected SNPs, 147 SNPs passed quality control measures. These SNPs had a high call rate (>90% in cases or controls; 25 SNPs were removed), were not nonmonomorphic (minor allele frequency >1% in cases or controls; 2 SNPs were removed) and did not deviate from Hardy–Weinberg equilibrium in the control population (HWE P > 0.0001; 5 SNPs were removed). One SNP failed genotyping and was excluded from the analysis.

Copy number variation genotyping and quality control. The functionally tested TaqMan Copy Number Assays Hs 02550639_cn (for LCE3C) and Hs 02878369_cn (for LCE3B) (Applied Biosystems) were used to genotype copy number variation within the LCE gene cluster in samples of panel A and B (754 cases and 1,052 controls). Genotyping was carried out according to Applied Biosystems’ standard protocols with 5 ng of dried DNA per well.

All cases were genotyped three times and all control samples were genotyped four times. The generated data was analyzed with the analysis software CopyCaller v1.0. The analysis settings were selected as recommended by the CopyCaller software user guide. The chosen confidence threshold of the associated predicted copy number was 295%. Control samples were removed when more than one of the four measurements did not pass quality control measures, whereas cases were removed when one of the three measurements did not pass quality control. Altogether, 736 cases and 932 controls remained for association analysis (94% of all samples).

Statistical analyses. Power calculations were carried out using PS Power and Sample Size v3.1231. GWAS data were analyzed using R statistical environment version 2.10.0 and gPLINK v2.049 in combination with PLINK v1.0533. The −clump command was used to reduce the number of SNPs for follow-up by removing correlated hit SNPs. The meta-analysis of the different panels was performed with METAL35.

TRAF3IP2 silencing in keratinocytes. Small hairpin RNA (shRNA) targeting enhanced green fluorescent protein (EGFP) was cloned into the pLenti4/Block-it-DEST (Invitrogen) as previously described36. shRNA lentiviral
constructs directed against TRAF3IP2 were obtained from Sigma. Lentiviral particles were produced in 293FT cells and used to infect immortalized N-TERT keratinocytes (kindly provided by J. Rheinwald) and normal human keratinocytes (NHK) at 10–20% confluence as previously described. After 72 h of infection, the cells were incubated in basal medium and stimulated for 30 h with IL17 (10 ng/ml), IL22 (10 ng/ml) and/or TNF-α (20 ng/ml). Total RNA was isolated using RNasy Mini Kits with on-column DNase digestion according to the manufacturer’s instructions (Qiagen) and were reverse transcribed using the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems). Real-time PCR analysis for DEFB4, TRAF3IP2 and the control gene RPLP0 (encoding ribosomal protein P0) was performed using prevalidated TaqMan gene expression assays from Applied Biosystems according to the manufacturer’s instructions. Target gene expression was normalized to the control gene RPLP0, and mRNA levels were expressed as a percent of RPLP0.


