Identification of fifteen new psoriasis susceptibility loci highlights the role of innate immunity

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Study	Cases (N)	Controls (N)	Total	λGC
Kiel ¹	474	1,146	1,620	1.09
CASP ²	1,359	1,400	2,759	1.06
WTCCC2 ³	2,178	5,175	7,353	1.04
PAGE	3,580	5,902	9,482	0.992
GAPC	2,997	9,183	12,180	0.963
TOTAL	10,588	22,806	33,394	1.11

Supplementary Table 1. Description of the 5 datasets used in the meta-analysis

Supplementary Table 2. Description of individual Immunochip samples

'Group' refers to the group that provided the samples and corresponds to the membership of consortia.

Dataset	Population Sample	Group (cases)	Group (controls)	Cases	Controls
	USA	UMich/NPF/HFH	UMich/FIMR/NPF	1,351	2,694
	Canada	UToronto/MU	UToronto	362	20
PAGE Immunochip	Estonia	UTartu/EGCUT	EGCUT	1,295	898
	Germany	CAU Kiel	CAU Kiel/KORA/HNR	572	2,290
	Total			3,580	5,902
	UK	KCL/Glasgow/Sheffield	WTCCC2	207	4822
	Finland	Helsinki	DILGOM	240	490
	Spain	Barcelona	Barcelona	269	202
	The Netherlands	Nijmegan	Groningen	152	1,107
GAPC Immunochip	Austria	Graz	-	310	0
	Sweden	Gothenburg/Stockholm	-	859	0
	Italy	Rome	Groningen	73	509
	Germany	Erlangen	PopGen	826	1,984
	Ireland	Dublin	Dublin	61	69
	Total			2,997	9,183

Supplementary Table 3. Association results for each of the 5 studies for the most strongly associated SNPs

Showing the 21 known and 15 newly identified loci. The overall OR was calculated using the effective sample size-weighted approach.

Known Loci	i				K	(iel				CAS	P			ωтс	CC2			PA	GE			GAPC			
SNP	Chr.	Position	Risk/ Non-risk allele	Р		RAF) (ctrl)	OR (95% CI)	Р	RAF (case)		OR (95% CI)	Р		RAF) (ctrl)	OR (95% CI)	Р		RAF (ctrl)	OR (95% CI)	Р		RAF) (ctrl)	OR (95% CI)	Combined P-value	
rs7552167	1	24,518,643	G/A	8.0x10 ⁻¹	0.86	0.86	1.03	1.1x10 ⁻²	0.88	0.86	1.29	1.6x10 ⁻⁴	0.89	0.86	1.24	4.5x10 ⁻⁵	0.87	0.85	1.22	4.6x10 ⁻⁴	0.88	0.86	1.20	8.5×10 ⁻¹²	1.21
rs9988642	1	67,726,104	T/C	9.5x10 ⁻⁴	0.95	0.92	(0.83-1.28) 1.79 (1.24-2.57)	2.4x10 ⁻⁷	0.95	0.91	(1.06-1.57) 1.82 (1.44-2.29)	2.6x10 ⁻⁶	0.95	0.93	(1.11-1.38) 1.45 (1.24-1.69)	3.8x10 ⁻⁸	0.95	0.93	(1.12-1.33) 1.43 (1.25-1.64)	1.8x10 ⁻⁸	0.96	0.93	(1.09-1.31) 1.53 (1.33-1.76)	1.1×10 ⁻²⁶	1.52
rs6677595	1	152,590,187	T/C	2.4x10 ⁻⁴	0.68		1.36 (1.15-1.61)	7.6x10 ⁻⁴	0.68		1.22 (1.09-1.37)	2.7x10 ⁻¹⁰	0.71		1.29 (1.19-1.40)	2.7x10 ⁻⁷	0.67		1.18 (1.10-1.26)	2.8x10 ⁻¹⁵	0.70		1.31 (1.22-1.40)	2.1×10 ⁻³³	1.26
rs62149416	2	61,083,506	T/C	2.5x10 ⁻¹	0.67	0.65	1.10 (0.93-1.29)	1.8x10 ⁻⁴	0.68		1.25	1.4x10 ⁻⁷	0.69		1.24 (1.14-1.34)	7.2x10 ⁻⁴	0.66		1.12 (1.05-1.19)	6.3x10 ⁻⁷	0.67		1.16	1.8×10 ⁻¹⁷	1.17
rs17716942	2	163,260,691	T/C	8.6x10 ⁻²	0.90	0.88	,	3.0x10 ⁻²	0.88	0.86	1.19 (1.02-1.40)	3.4x10⁻ ⁸	0.90	0.86	1.38 (1.23-1.54)	6.7x10 ⁻⁹	0.90	0.86	1.31 (1.20-1.45)	8.2x10 ⁻⁴	0.88	0.86	```	3.3×10 ⁻¹⁸	1.27
rs27432	5	96,119,273	A/G	9.8x10 ⁻²	0.31		1.15 (0.97-1.35)	1.3x10 ⁻¹	0.30		1.10 (0.97-1.23)	5.2x10 ⁻⁸	0.31		1.26 (1.16-1.37)	1.1x10 ⁻⁵	0.30		1.16 (1.09-1.24)	6.4x10 ⁻¹⁰	0.32	0.28	1.23 (1.15-1.32)	1.9×10 ⁻²⁰	1.20
rs1295685	5	131,996,445	G/A	4.2x10 ⁻¹	0.80	0.78	1.08 (0.89-1.31)	1.5x10 ⁻⁶	0.83		1.41 (1.22-1.61)	2.1x10 ⁻²	0.84		1.12 (1.02-1.24)	8.6x10 ⁻³	0.78		1.15 (1.06-1.23)	1.9x10 ⁻⁴	0.81		1.20 (1.11-1.30)	3.4×10 ⁻¹⁰	1.18
rs2233278	5	150,467,189	C/G	5.3x10 ⁻²	0.07		1.39 (1.00-1.92)	5.5x10 ⁻⁸	0.09		1.86 (1.48-2.34)	5.5x10 ⁻¹¹	0.09		1.68 (1.44-1.96)	2.7x10 ⁻¹⁴	0.10		1.54 (1.37-1.72)	2.8x10 ⁻¹⁴	0.08		1.54 (1.37-1.74)	2.2×10 ⁻⁴²	1.59
rs12188300	5	158,829,527	T/A	8.1x10 ⁻⁶	0.15	0.11	1.80 (1.39-2.33)	1.9x10 ⁻⁷	0.14		1.68 (1.38-2.05)	8.6x10 ⁻¹⁴	0.15		1.64 (1.44-1.86)	6.9x10 ⁻²¹	0.13		1.58 (1.43-1.75)	2.1x10 ⁻¹³	0.12		1.48 (1.34-1.63)	3.2×10 ⁻⁵³	1.58
rs4406273	6	31,266,090	A/G	4.2x10 ⁻⁴²	0.28		5.18 (4.04-6.63)	3.1x10 ⁻⁵³	0.23		4.13 (3.41-4.99)	1.3x10 ⁻²²⁹	0.29		6.33 (5.66-7.08)	1.8x10 ⁻¹⁶⁹	0.25		3.35 (3.06-3.67)	6.9x10 ⁻²⁶⁵	0.27		4.24 (3.88-4.64)	4.5×10 ⁻⁷²³	4.32
rs33980500	6	111,913,262	T/C	5.8x10 ⁻⁵	0.12		1.67 (1.31-2.14)	3.3x10 ⁻⁴	0.10		1.42 (1.17-1.72)	3.5x10 ⁻¹⁵	0.11		1.72 (1.50-1.97)	5.2x10 ⁻¹⁰	0.11		1.38 (1.25-1.53)	3.6x10 ⁻¹⁹	0.11	0.07	1.54 (1.39-1.71)	4.2×10 ⁻⁴⁵	1.52
rs582757	6	138,197,824	C/T	5.9x10 ⁻⁴	0.33	0.27	1.34 (1.14-1.59)	6.5x10 ⁻⁶	0.32		1.31 (1.16-1.47)	3.5x10 ⁻⁸	0.31		1.26 (1.16-1.37)	1.7x10 ⁻⁶	0.31		1.17 (1.09-1.25)	4.0x10 ⁻⁸	0.32	0.28	1.22 (1.14-1.31)	2.2×10 ⁻²⁵	1.23
rs1250546	10	81,032,532	A/G	3.7x10 ⁻²	0.65		1.19 (1.01-1.39)	4.7x10 ⁻⁴	0.61		1.21 (1.09-1.35)	1.9x10 ⁻¹	0.59		1.05 (0.97-1.14)	1.1x10 ⁻²	0.61		1.09 (1.03-1.16)	4.6x10 ⁻³	0.60		1.09 (1.03-1.16)	6.8x10 ⁻⁷	1.10
rs645078	11	64,135,298	A/C	4.2x10 ⁻¹	0.63		1.07 (0.91-1.25)	4.8x10 ⁻³	0.64		1.17 (1.05-1.31)	1.4x10 ⁻¹	0.62		1.06 (0.98-1.14)	1.7x10 ⁻³	0.63		1.12 (1.05-1.19)	6.4x10 ⁻²	0.62	0.61	1.06 (0.98-1.13)	2.2x10 ⁻⁶	1.09
rs2066819	12	56,750,204	C/T	6.4x10 ⁻²	0.95		1.41 (0.97-2.03)	2.0x10 ⁻⁵	0.96		1.84 (1.38-2.44)	9.3x10 ⁻⁸	0.95		1.57 (1.33-1.85)	4.1x10 ⁻⁴	0.94		1.27 (1.12-1.44)	6.4x10 ⁻⁵	0.95		1.28 (1.12-1.47)	5.4×10 ⁻¹⁷	1.39
rs8016947	14	35,832,666	G/T	6.8x10 ⁻⁴	0.64		1.31 (1.12-1.53)	5.5x10 ⁻³	0.59		1.16 (1.05-1.30)	4.6x10 ⁻⁶	0.61		1.19 (1.11-1.29)	2.6x10 ⁻⁴	0.59		1.12 (1.05-1.19)	1.1x10 ⁻⁶	0.60	0.56	1.17 (1.10-1.25)	2.5×10 ⁻¹⁷	1.16
rs12445568	16	31,004,812	C/T	1.6x10 ⁻³	0.43	0.37	1.28 (1.10-1.49)	1.2x10 ⁻³	0.40		1.21 (1.08-1.36)	5.2x10 ⁻³	0.40		1.12 (1.03-1.20)	1.4x10 ⁻⁶	0.41		1.16 (1.09-1.23)	2.9x10 ⁻⁶	0.40	0.37	1.16 (1.09-1.24)	1.2×10 ⁻¹⁶	1.16
rs28998802	17	26,124,908	A/G	4.0x10 ⁻³	0.20	0.16	1.36 (1.11-1.68)	3.2x10 ⁻²	0.18		1.18 (1.01-1.38)	7.7x10 ⁻⁴	0.18		1.20 (1.08-1.34)	2.0x10 ⁻⁸	0.16		1.27 (1.17-1.38)	9.1x10 ⁻⁵	0.16		1.18 (1.08-1.28)	3.3×10 ⁻¹⁶	1.22
rs34536443	19	10,463,118	G/C	1.3x10 ⁻²	0.97	0.95	1.77 (1.11-2.81)	NA	NA	NA	NA	4.7x10 ⁻⁹	0.97		1.81 (1.48-2.21)	3.4x10 ⁻¹⁰	0.98		1.76 (1.46-2.10)	9.0x10 ⁻¹⁴	0.98		2.09 (1.72-2.54)	9.1×10 ⁻³¹	1.88
rs1056198	20	48,556,229	C/T	3.5x10 ⁻¹	0.59		1.07 (0.92-1.25)	2.4x10 ⁻⁴	0.63		1.22 (1.10-1.36)	1.6x10 ⁻⁶	0.63		1.20 (1.11-1.30)		0.59		1.17 (1.10-1.25)	4.9x10 ⁻³	0.58		1.11 (1.05-1.18)	1.5×10 ⁻¹⁴	1.16
rs4821124	22	21,979,289	C/T	1.2x10 ⁻¹	0.22		1.17 (0.96-1.41)	8.9x10 ⁻²	0.22	0.20		5.2x10 ⁻⁴	0.20	0.18		2.0x10 ⁻⁴	0.20	0.18		8.4x10 ⁻²	0.22	0.20		3.8×10⁻ ⁸	1.13

Newly I	dentifie	ed Loci			Ki	el			CAS	SP			wтс	CC2			PAG	GE			GAP	с			
SNP	Chr.	Position	Risk/ Non-risk allele	Р	RAF (case)	RAF (ctrl)	OR (95% CI)	Р		RAF) (ctrl)	OR (95% CI)	Р		RAF) (ctrl)	OR (95% CI)	Р		RAF (ctrl)	OR (95% CI)	Р	RAF (case)		OR (95% CI)	Combined P-value	
rs11121129	1	8,268,095	A/G	9.2x10 ⁻³	0.31	0.27	1.25 (1.06-1.47)	3.2x10 ⁻²	0.30		1.14 (1.01-1.29)		0.33		1.11 (1.03-1.20)	-	0.29		1.06 (0.99-1.13)	2.1x10 ⁻⁵	0.31		1.20 (1.12-1.28)	1.7×10 ⁻⁸	1.13
rs7536201	1	25,293,084	C/T	1.5x10 ⁻¹	0.51	0.48	()	3.3x10 ⁻²	0.52	0.49	1.12 (1.01-1.25)	1.8x10 ⁻³	0.54	0.50	· ,	1.4x10 ⁻⁷	0.53	0.49	` '	3.2x10 ⁻³	0.52	0.50	- /	2.3×10 ⁻¹²	1.13
rs10865331	2	62,551,472	A/G	4.8x10 ⁻¹	0.38	0.37	1.06 (0.91-1.23)	2.0x10 ⁻²	0.41		1.14 (1.02-1.27)		0.40	0.37	1.11 (1.03-1.20)	6.0x10 ⁻⁸	0.41	0.38	. ,	6.0x10 ⁻²	0.39		1.08 (1.01-1.15)	4.7×10 ⁻¹⁰	1.12
rs9504361	6	577,820	A/G	1.5x10 ⁻²	0.58		1.21 (1.04-1.42)		0.57		1.12 (1.01-1.25)		0.60		1.17 (1.09-1.27)		0.57		1.06 (1.00-1.13)	1.7x10 ⁻⁵	0.57		1.14 (1.07-1.21)	2.1×10 ⁻¹¹	1.12
rs2451258	6	159,506,600	C/T	9.4x10 ⁻¹	0.33		1.01 (0.86-1.18)		0.38		1.15 (1.03-1.29)		0.40		1.12 (1.04-1.21)		0.34		1.14 (1.07-1.22)	1.3x10 ⁻²	0.36		1.11 (1.04-1.18)	3.4×10 ⁻⁸	1.12
rs2700987	7	37,386,237	A/C	7.8x10 ⁻²	0.61	0.57	1.16 (0.98-1.36)	1.8x10 ⁻²	0.59		1.14 (1.02-1.27)		0.61		1.18 (1.10-1.27)		0.58		1.09 (1.02-1.16)	1.5x10 ⁻²	0.59		1.07 (1.01-1.14)	4.3×10 ⁻⁹	1.11
rs11795343	9	32,523,737	T/C	2.3x10 ⁻³	0.65	0.59	1.28 (1.09-1.49)		0.63		1.17 (1.05-1.30)		0.64		1.15 (1.06-1.24)		0.63		1.12 (1.05-1.19)	3.9x10 ⁻²	0.61		1.05 (0.99-1.12)	8.4×10 ⁻¹¹	1.11
rs10979182	9	110,817,020	A/G	6.9x10 ⁻³	0.64		1.24 (1.06-1.45)		0.61		1.11 (1.00-1.24)		0.63		1.13 (1.04-1.22)		0.61		1.08 (1.02-1.15)	3.2x10⁻³	0.61		1.12 (1.06-1.20)	2.3×10 ⁻⁸	1.12
rs4561177	11	109,962,432	A/G	9.0x10 ⁻¹	0.59		1.01 (0.87-1.18)		0.61		1.12 (1.01-1.25)		0.61		1.15 (1.07-1.24)		0.61		1.13 (1.07-1.21)	1.1x10⁻⁵	0.63		1.16 (1.09-1.24)	7.7×10 ⁻¹³	1.14
rs3802826	11	128,406,438	A/G	2.3x10 ⁻¹	0.51		1.10 (0.94-1.28)	4.2x10 ⁻²	0.51		1.12 (1.00-1.24)		0.52		1.10 (1.02-1.18)		0.49		1.12 (1.06-1.19)	1.3x10 ⁻⁴	0.51		1.15 (1.08-1.22)	9.5×10 ⁻¹⁰	1.12
rs367569	16	11,365,500	C/T	2.5x10 ⁻¹	0.73	0.71	1.11 (0.93-1.31)	6.8x10 ⁻²	0.73		1.12 (0.99-1.27)		0.73		1.14 (1.05-1.24)		0.72		1.10 (1.03-1.18)	3.6x10 ⁻³	0.74		1.15 (1.07-1.23)	4.9×10 ⁻⁸	1.13
rs963986	17	40,561,579	C/G	9.6x10 ⁻⁴	0.19		1.43 (1.16-1.76)	2.7x10 ⁻²	0.17		1.19 (1.02-1.38)		0.17		1.12 (1.01-1.24)		0.16		1.07 (0.98-1.16)	1.3x10⁻⁵	0.18		1.22 (1.13-1.33)	5.3×10-9	1.15
rs11652075	17	78,178,893	C/T	5.8x10 ⁻¹	0.52	0.51	1.04 (0.90-1.21)		0.53		1.18 (1.00-1.40)		0.52		1.11 (1.03-1.19)		0.52		1.02 (0.96-1.08)	2.2x10 ⁻⁸	0.55		1.19 (1.11-1.26)	3.4×10 ⁻⁸	1.11
rs545979	18	51,819,750	T/C	3.8x10 ⁻²	0.34		1.19 (1.01-1.40)		0.33		1.17 (1.04-1.31)		0.32		1.16 (1.07-1.26)		0.32		1.09 (1.02-1.17)	_	0.31		1.11 (1.04-1.19)	3.5×10 ⁻¹⁰	1.12
rs892085	19	10,818,092	A/G	3.2x10 ⁻²	0.62		1.19 (1.01-1.39)	1.5x10 ⁻³	0.58		1.27 (1.10-1.48)		0.57		1.16 (1.08-1.26)		0.60		1.10 (1.03-1.17)		0.60		1.20 (1.13-1.28)	3.0×10 ⁻¹⁷	1.17

Supplementary Table 4. Disease overlap and SNP functional annotation for the known and newly identified SNPs.

The 'disease overlap' is defined as an associated SNP identified in the same region (within 500kb) as the top psoriasis SNP (using NHGRI GWAS catalog and Immunochip results for Celiac disease⁴). Underlined entries have SNPs that are in LD ($r^2>0.7$) with the identified SNP. *denotes association of the same SNP in the same direction. AD: Atopic dermatitis, AS: Ankylosing spondylitis, BD: Behcet's disease, CD: Crohn's disease, CeD: Celiac disease, IgE: Serum IgE, IgA: Selective Immunoglobulin A deficiency, LE: Leprosy, MS: Multiple Sclerosis, PBC: Primary biliary cirrhosis, RA: Rheumatoid arthritis, SI: Soluble ICAM-1, SLE: Systemic lupus erythematosus, SS: Systemic sclerosis, T1D: Type I Diabetes, UC: Ulcerative colitis. Supplementary Figure 5 shows graphical view of the disease overlap.

SNP	Chr.	Position	Risk/ Non-risk allele	Variant Annotation	Disease Overlap (based on the GWAS catalog)	All genes in locus (+/- 500kb)
Kr	nown	loci				
rs7552167	1	24,518,643	G/A	4878 bp upstream of IL28RA		LOC100132287;SRRM1;LOC100133331;CNR2;C1orf130;PNRC2;OR4F3;OR4F29; SFRS13A;RPL11;MYOM3;LOC284632;TCEB3;NIPAL3;C1orf128;C1orf201;HMGCL; GALE;IL22RA1;OR4F16;FUCA1;LOC100132062;IL28RA;LYPLA2;GRHL3;RCAN3
rs9988642	1	67,726,104	T/C	454 bp downstream of IL23R	<u>AS, UC, CD</u> , BD, LE, PBC	LOC100132287;LOC100133331;IL23R;GNG12;IL12RB2;MIER1;SLC35D1;C1orf141 ;INSL5;TCTEX1D1;WDR78;OR4F3;OR4F29;SERBP1;GADD45A; OR4F16; LOC100132062
rs6677595	1	152,590,187	T/C	3613 bp downstream of LCE3B		LOC100132287;LOC100133331;FLG;IVL;SPRR4;SPRR3;OR4F3;OR4F29;SPRR1A ;SPRR1B;RPTN;LCE2C;LCE2B;LCE2A;LCE2D;CRCT1;LCE4A;LCE6A;KPRP;FLG2; SMCP;C1orf68;CRNN;OR4F16;SPRR2E;SPRR2D;SPRR2F;SPRR2A;SPRR2B;LOC1 00132062;LCE1F;LCE1D;LCE1E;LCE1B;LCE1C;HRNR;LCE1A; LCE3D; LCE3E;LCE3A;LCE3B;LCE3C;LCE5A
rs62149416	2	61,083,506	T/C	FLJ16341 intron	<u>RA</u> , UC, CD, CeD	REL;USP34;PEX13;KIAA1841;AHSA2;PAPOLG;C2orf74;PUS10;BCL11A
rs17716942	2	163,260,691	T/C	KCNH7 intron	T1D, IgA	SLC4A10;IFIH1;FAP;KCNH7;GCG;GCA;DPP4
rs27432	5	96,119,273	A/G	ERAP1 intron	<u>AS,</u> CD	LNPEP;CAST;RIOK2;ERAP2;ERAP1;LIX1;PCSK1
rs1295685	5	131,996,445	G/A	IL13 3'UTR	IgE, CD, AD, platelet counts, C-reactive protein, eosinophil counts and fibrinogen	IL13; C5orf56; IRF1; ANKRD43; IL4; IL5; AFF4 ;HSPA4; UQCRQ; ZCCHC10; SLC22A4;SLC22A5;CCNI2;GDF9;P4HA2;KIF3A;PDLIM4;SHROOM1; LEAP2; SEPT8;RAD50
rs2233278	5	150,467,189	C/G	TNIP1 5'UTR	SLE, SS, CD	SLC36A3;SLC36A1;SLC36A2;MYOZ3;RBM22;TNIP1;GPX3;ZNF300;IRGM;LOC134 466;ANXA6;CCDC69;SYNPO;FAT2;DCTN4;C5orf62;GM2A
rs12188300	5	158,829,527	T/A	Intergenic	MS, CD, AS, UC	LOC285627;IL12B;UBLCP1;EBF1;RNF145
rs4406273	6	31,266,090	A/G	Intergenic	UC, AS, SS, SLE, Vitiligo, AIDS progression, Grave's Disease, Hepatitis B vaccine response, Follicular lymphoma, CD4:CD8 lymphocyte ratio	MUC21;LY6G6C;LY6G6D;HLA-B; SFTA2; APOM; DPCR1; LY6G5C; PSORS1C2; CLIC1; AIF1; LY6G6E; LY6G6F; LSM2; PSORS1C1; PSORS1C3; LST1; C6orf26; DDAH2 ; C6orf27 ; C6orf25; TNF ; BAT5; BAT2; BAT1; C6orf47; LY6G5B; DDR1; GTF2H4; MSH5; HLA-C; LTA; MCCD1; ATP6V1G2; VARS2; POU5F1; TCF19; NFKBIL1; CCHCR1; SNORA38; VARS; HCP5; CSNK2B; HCG27; HCG22; LTB; MICB;SNORD117;CDSN;C6orf15;HCG26;BAT4;BAT3;NCR3;MICA;SNORD84
rs33980500	6	111,913,262		missense mutation inTRAF3IP2		TUBE1; REV3L;WISP3;SLC16A10;FYN;TRAF3IP2;C6orf225;KIAA1919
rs582757	6	138,197,824	C/T	TNFAIP3 intron	CeD, RA, UC, SLE	PERP;OLIG3;TNFAIP3;PBOV1;KIAA1244

rs1250546	10	81,032,532	A/G	ZMIZ1 intron	MS, CD, CeD, Vitiligo	LOC283050;ZCCHC24;EIF5AL1;PPIF;MIR1256;ZMIZ1;LOC650623;SFTPA2; SFTPA1
rs645078	11	64,135,298	A/C	RPS6KA4 intron		NRXN2;MACROD1;BAD;SLC22A12;SLC22A11;TRMT112;MARK2;MEN1;RPS6KA ;STIP1;MIR1237;KCNK4;FERMT3;TRPT1;SF1;PLCB3;DNAJC4;ESRRA;C11orf20; PPP1R14B;GPR137;COX8A;RASGRP2;CDC42BPG;PYGM;NUDT22;MAP4K2;EHD1 PRDX5;OTUB1;RCOR2;CCDC88B;VEGFB;NAA40;FKBP2;FLRT1
rs2066819	12	56,750,204	C/T	STAT2 intron		ZC3H10;CNPY2;OBFC2B;APOF;SPRYD4;ATP5B;NACA;MYL6B;WIBG;PA2G4; SMARCC2;RNF41;ESYT1;CDK2;SILV;BAZ2A;SLC39A5;MIP;ANKRD52;TIMELESS; CS;ERBB3;RBMS2;PRIM1;SNORD59B;SNORD59A;RAB5B;RPS26;HSD17B6;PAN2 IL23A;SUOX;COQ10A;DGKA;GLS2;RPL41;MYL6;STAT2;PTGES3;IKZF4
rs8016947	14	35,832,666	G/T	Intergenic		KIAA0391;PPP2R3C;BRMS1L;INSM2;NFKBIA;RALGAPA1;PSMA6; C14orf19; FAM177A1;SRP54;BAZ1A
rs12445568	16	31,004,812	C/T	STX1B intron		FBXL19;ITGAX;ITGAL;ITGAM;ITGAD;CTF1;C16orf58;PYCARD;PRSS36;FUS; PYDC1;RNF40;BCKDK;TRIM72;NCRNA00095;PRSS8;MYST1;PRR14;PHKG2; ORAI3; BCL7C; C16orf93; ZNF843; HSD3B7; PRSS53; TGFB111; SNORA30; ZNF688; ZNF689; SETD1A; COX6A2; ZNF646; ZNF668; FBRS; ARMC5; MIR762; SRCAP;ZNF629; VKORC1; STX1B;ZNF785;ZNF768;ZNF764;SLC5A2;STX4;ZNF747
rs28998802	17	26,124,908	A/G	NOS2 intron		KSR1;LGALS9;FLJ40504;PYY2;WSB1;NOS2;C17orf108;NLK;PPY2
rs34536443	19	10,463,118	G/C	Missense mutation in <i>TYK2</i>	CD, T1D, SI	ATG4D;S1PR2;P2RY11;SLC44A2;ILF3;CDC37;PPAN-P2RY11 ; C19orf38; MRPL4 MIR638; MIR1238; SNORD105; TYK2; ZGLP1; COL5A3; OLFM2; AP1M2; EIF3G; CDKN2D; FDX1L; LOC147727; ANGPTL6; C3P1; KRI1 ;PPAN; MIR199A1; ICAM5; ICAM4 ;ICAM3 ;ICAM1 ;TMED1
rs1056198	20	48,556,229	C/T	RNF114 intron		TMEM189-UBE2V1; KCNB1; B4GALT5; TMEM189; SLC9A8; CEBPB; UBE2V1; RNF114; SPATA2; SNAI1; PTGIS
rs4821124	22	21,979,289	C/T	966 bp downstream of <i>UBE2L3</i>	SLE, <u>CeD</u> *, <u>RA</u> , <u>CD</u> , MS	CCDC116;YDJC;PPIL2;POM121L8P;PI4KAP2;TOP3B;PPM1F;YPEL1;MIR130B; RIMBP3B;MIR301B;SDF2L1;MAPK1;RIMBP3C;HIC2;UBE2L3
Newly lo	denti	fied Loci				
rs11121129	1	8,268,095	A/G	Intergenic	UC, CeD	LOC100132287;LOC100133331;PARK7;PER3;OR4F3;OR4F29;TNFRSF9;RERE; VAMP3;UTS2;ERRFI1;SLC45A1;CAMTA1;OR4F16;LOC100132062
rs7536201	1	25,293,084	C/T	1583 bp upstream of <i>RUNX3</i>	<u>AS, CeD</u>	LOC100132287;CLIC4;SRRM1;RHD;LOC100133331;C1orf130;TMEM57;OR4F3; OR4F29;RUNX3;NIPAL3;C1orf63;TMEM50A;OR4F16;SYF2;LOC100132062;RHCE; RCAN3
rs10865331	2	62,551,472	A/G	Intergenic	<u>AS</u> *	B3GNT2;TMEM17;CCT4;EHBP1;COMMD1;FAM161A
rs9504361	6	577,820	A/G	EXOC2 intron	CeD, BCC, PSP	IRF4;DUSP22;LOC285768;HUS1B;EXOC2
rs2451258	6	159,506,600	C/T	Intergenic	MS, CeD, CD, <u>RA</u>	RSPH3;EZR;TMEM181;DYNLT1;FNDC1;OSTCL;SYTL3;TAGAP
rs2700987	7	37,386,237	A/C	ELMO1 intron	PBC, CeD, RA	GPR141;ELMO1;MIR1200
rs11795343	9	32,523,737	T/C	DDX58 intron		APTX;TOPORS;NDUFB6;DDX58;TAF1L;ACO1;TMEM215
rs10979182	9	110,817,020	A/G	Intergenic		KLF4

rs4561177	11	109,962,432	A/G	1655 bp upstream of ZC3H12C		ARHGAP20;RDX;ZC3H12C;FDX1
rs3802826	11	128,406,438	A/G	ETS1 intron	SLE, CeD	TP53AIP1;ETS1;FLI1;C11orf45;ARHGAP32;KCNJ1;KCNJ5
rs367569	16	11,365,500	C/T	1664 bp downstream of <i>PRM3</i>	CeD, T1D, PBC	C16orf75;DEXI;LITAF;ZC3H7A;SNN;SOCS1;FAM18A;CIITA;TNP2;PRM1;PRM3; PRM2;TXNDC11;CLEC16A
rs963986	17	40,561,579		PTRF intron	CD, MS	GHDC;KCNH4;FAM134C;LOC100190938;ATP6V0A1;TTC25;CNTNAP1;TUBG1; STAT3;HSPB9;G6PC;ACLY;NKIRAS2;LOC388387;WNK4;DNAJC7;CCDC56;PSME3; PSMC3IP;TUBG2;PLEKHH3;CNTD1;RAB5C;COASY;LOC90586;AOC3;AOC2; NAGLU;STAT5B;STAT5A;BECN1;KAT2A;DHX58;CCR10;HCRT;VPS25;MLX;CNP; RAMP2;EZH1;HSD17B1;PTRF
rs11652075	17	78,178,893		Missense mutation in CARD14		CBX2;NPTX1;FLJ35220;CBX8;SLC26A11;LOC100294362;CCDC40;SGSH;RPTOR ;EIF4A3;CBX4;GAA;TBC1D16;CARD14;RNF213;ENPP7
rs545979	18	51,819,750		POLI intron		POLI;MBD2;C18orf26;STARD6;SNORA37;C18orf54
rs892085	19	10,818,092		QTRT1 intron		ATG4D;S1PR2;LDLR;SLC44A2;ILF3;CDC37;C19orf52;C19orf38;MRPL4;KANK2; MIR638;MIR1238;TYK2;DOCK6;ZGLP1;AP1M2;CDKN2D;FDX1L;LOC147727;KRI1; SMARCA4;SPC24;MIR199A1;CARM1;ICAM5;ICAM4;ICAM3;ICAM1;TMED1;QTRT1;k EAP1;RAVER1;YIPF2;MIR1181;S1PR5;PDE4A;DNM2

Supplementary Table 5. Association results for rs892085

The association of rs892085 before and after conditioning on either of two signals in *TYK2*: i) rs12720356, the most significantly associated SNP in this region from a previous study³; ii) rs34536443, the most significant SNP in this region

			F	P value		
Analysis	Kiel	CASP	WTCCC2	PAGE	GAPC	Meta
No conditioning	3.21x10 ⁻²	1.48x10 ⁻³	1.02x10 ⁻⁴	2.22x10 ⁻³	4.18x10 ⁻¹⁰	2.95x10 ⁻¹⁷
Conditioning on rs12720356	8.72x10 ⁻²	2.88x10 ⁻³	2.40x10 ⁻³	1.26x10 ⁻²	3.68x10 ⁻⁸	7.36x10 ⁻¹³
Conditioning on rs34536443	5.40x10 ⁻²	NA	7.70x10 ⁻⁴	9.69x10 ⁻³	1.84x10 ⁻⁸	4.22x10 ⁻¹²

Supplementary Table 6. Significant results for the meta-conditional analysis

Signals achieving genome-wide significance when conditioning on the most strongly associated SNPs of the 19 known and 15 new loci that achieve genome-wide significance in this study. Because the strongest SNP in the *TYK2* region was poorly imputed in CASP GWAS, the second strongest SNP (rs2304256) was used in the conditional analysis for this dataset; the CASP GWAS was not included in the conditional meta-analysis for the *TYK2* locus. Underlined shared diseases indicate the indentified SNPs are in high LD (r2>0.7). RAF: Risk Allele Frequency.

SNP	Chr.	Position(bp)	GWAS P-value (meta)	Immunochip P-value (meta)	Combined P-value	Risk/Non- risk alleles	RAF (Case)	RAF (ctrl)	OR (meta)	Notable genes	Disease overlap ^c
rs2111485	2	163,110,536	7.9×10 ⁻⁴	9.5×10 ⁻⁶	2.7×10 ⁻⁸	G/A	0.647	0.610	1.14	IFIH1	T1D, IgA
rs2910686	5	96,252,589	2.3×10 ⁻⁵	1.3×10 ⁻⁴	2.0×10 ⁻⁸	C/T	0.442	0.437	1.12	ERAP2	<u>CD</u> , AS
rs4379175	5	158,804,928	4.8×10 ⁻²⁰	6.9×10 ⁻²²	9.0×10 ⁻⁴⁰	G/T	0.737	0.678	1.31	IL12B	MS, CD, AS, UC
rs13437088	6	31,355,119	2.8×10 ⁻¹⁷	1.1×10 ⁻²⁴	3.1×10 ⁻⁴⁰	T/C	0.342	0.251	1.32	MICA	AS, GD [¥]
rs12720356	19	10,469,975	9.7×10 ⁻⁶	1.1×10 ⁻⁵	3.2×10 ⁻¹⁰	A/C	0.929	0.911	1.25	TYK2	<u>CD[*],</u> T1D, SI

^c AS: Ankylosing spondylitis, CD: Crohn's Disease; GD: Graves' disease, IgA: Selective Immunoglobulin A deficiency, MS: Multiple Sclerosis, SI: Soluble ICAM-1, T1D: Type 1 Diabetes; UC: Ulcerative Colitis. ^{*}denotes association with the same SNP. ^{*} locus also associated with Systemic sclerosis, CD4:CD8 ratio, Vitiligo, AIDS progression, white blood cell types, Dengue shock syndrome, and Nevirapine-induced rash.

Supplementary Table 7. Association results for the most significant SNPs of the 5 significant loci in the conditional analysis for each of the datasets.

					I	Kiel			CA	SP			WTC	CC2			PA	GE			GAPC		
SNP	Chr.	Position	Risk/ Non- risk allele	Р		RAF) (ctrl)	OR (95% CI)	Р	RAF (case)		OR (95% CI)	Р	RAF (case)		OR (95% CI)	Р	RAF (case)		OR (95% CI)	Р	RAF RAF (case)(ctrl	- OR) (95% CI)	Combined P-value
rs2111485	2	163,110,536	G/A	8.5x10 ⁻	² 0.67		1.19 (0.98-1.44)		0.63		1.13 (0.98-1.30)		0.66		1.11 (1.03-1.21)		² 0.64		1.10 (1.02-1.18)		0.65 0.61	1.20 (1.11-1.29)	2.7×10 ⁻⁸
rs2910686	5	96,252,589	C/T	4.9x10 [°]	³ 0.46		1.32 (1.09-1.59)		0.44		1.25 (1.09-1.42)		0.46		1.09 (1.01-1.18)		³ 0.44		1.09 (1.02-1.18)		0.43 0.44	1.12 (1.04-1.20)	2.0×10 ⁻⁸
rs4379175	5	158,804,928	G/T	4.3x10 [`]	4 0.76		1.42 (1.17-1.72)		0.75		1.40 (1.23-1.60)		² 0.73		1.35 (1.24-1.46)		² 0.75		1.30 (1.20-1.39)		0.73 0.68	3 1.25 (1.16-1.35)	9.0×10 ⁻⁴⁰
rs13437088	36	31,355,119	T/C	4.0x10 ⁻	² 0.34		1.23 (1.01-1.51)		0.35		1.28 (1.12-1.45)		⁵ 0.36		1.44 (1.31-1.58)		¹ 0.33		1.26 (1.18-1.36)		0.34 0.25	5 1.33 (1.23-1.43)	3.1×10 ⁻⁴⁰
rs12720356	6 19	10,469,975	A/C	1.7x10 ⁻	² 0.95		1.62 (1.08-2.42)	NA	NA	NA	NA	1.7x10 ⁻⁵	0.92		1.34 (1.17-1.53)		² 0.93		1.18 (1.04-0.74)		0.94 0.91	1.21 (1.06-1.38)	3.2×10 ⁻¹⁰

Supplementary Table 8. Epistasis results

Pairwise combinations of psoriasis loci having the strongest evidence for interaction. The p-values and Z-scores are for the interaction terms from the meta-analysis of the epistasis results. The p-values for *LCE-HLA-C* and *ERAP1-HLA-C* remain statistically significant after Bonferroni correction.

Gene1	SNP	Gene2	SNP 2	P-value	Z-score
LCE	rs6677595	HLA-C	rs4406273	1.8×10 ⁻⁶	4.78
ERAP1	rs27432	HLA-C	rs4406273	2.8×10 ⁻¹⁰	6.31

Supplementary Table 9. Differential expression analysis results for genes in each of the 39 psoriasis loci identified by the primary or conditional association analysis.

For each locus, a genomic region in strong LD ($r^2>0.7$) with the most significantly associated SNP was defined using the tag SNP function of PLINK; the tagged regions were then extended by 50 kb on each side. Microarray results⁵ for all genes overlapping these extended regions are shown. Results for two-sample tests of differential expression are shown, including the false discovery rate (FDR), fold-change in expression levels (FC), and whether expression is up- or down-regulated in lesional skin (using FDR ≤ 0.05 and FC ≤ 0.67 or \geq 1.50 as criteria for differential expression).

	Gene	FDR	FC	Differentially expressed
Known loci	IL28RA	2.43E-01	0.92	
	LOC284632	9.43E-01	1.02	
	IL22RA1	8.85E-08	1.32	
	IL23R	1.77E-01	1.05	
	IL12RB2	5.99E-49	1.62	Up
	LCE3D	0.00E+00	24.42	Up
	PAPOLG	1.27E-04	0.84	
	REL	2.73E-19	1.74	Up
	PUS10	1.52E-40	1.75	Up
	GCA	1.00E+00	1.02	
	KCNH7	1.00E+00	1.02	
	ERAP1	1.14E-45	0.69	
	ERAP2	5.33E-03	1.21	
	CAST	1.14E-45	0.69	
	KIF3A	1.88E-13	0.73	
	IL13	1.62E-04	1.12	
	IL4	2.53E-01	1.04	
	RAD50	4.24E-06	0.85	
	TNIP1	1.22E-15	1.32	
	ANXA6	1.73E-04	1.11	
	LOC285627	6.54E-01	1.03	
	PSORS1C3	0.34E-01 1.84E-01	1.05	
	HCG27	1.54E-01	0.95	
	C6orf15		1.03	
		8.90E-01	1.03	
	POU5F1	1.00E+00		
	HCG22	4.47E-03	1.09	
	HLA-B	2.26E-15	1.26	
	PSORS1C1	1.00E+00	0.99	
	PSORS1C2	6.72E-04	1.39	
	HLA-C	1.70E-14	1.25	
	CDSN	2.80E-16	2.00	Up
	CCHCR1	7.78E-05	0.90	
	MICA	1.22E-17	0.60	Down
	TCF19	6.08E-30	1.41	
	TRAF3IP2	1.56E-03	1.10	
	TNFAIP3	5.18E-01	1.05	
	STAT2	8.77E-21	1.37	
	SLC39A5	5.39E-03	1.11	
	CS	7.01E-02	0.93	
	IL23A	4.32E-22	1.34	
	RNF41	1.17E-01	0.93	
	OBFC2B	8.28E-05	1.16	
	ANKRD52	1.52E-01	1.05	
	APOF	1.00E+00	1.01	
	COQ10A	1.00E+00	0.99	
	SMARCC2	1.47E-26	0.69	
	PAN2	4.93E-21	0.73	
	CNPY2	9.14E-25	1.30	
	NFKBIA	1.01E-02	1.09	
	PSMAG	1.01E-02	1.05	

4.99E-32

1.35

PSMA6

	NCRNA00095	1.41E-01	1.05	
	PRSS53	4.92E-120	2.66	Up
	BCKDK	7.01E-25	1.32	
	VKORC1	1.36E-20	1.30	
	MIR762	4.28E-03	0.92	
	STX4	5.95E-07	1.14	
	BCL7C	4.28E-03	0.92	
	FBXL19	6.32E-38	1.43	
	ORAI3	2.59E-19	0.73	
	ZNF646	4.29E-08	1.22	
	SETD1A	1.01E-03	1.11	
	HSD3B7	4.15E-02	0.89	
	MYST1	7.35E-21	0.78	
	PRSS8	4.78E-19	1.80	Up
	ZNF668	2.02E-06	1.15	
	STX1B	3.67E-02	1.07	
	PRSS36	1.12E-04	1.11	
	CTF1	1.00E+00	1.00	
	NOS2	4.78E-25	1.54	Up
	CDC37	3.03E-19	1.26	
	RAVER1	1.00E+00	1.00	
	ICAM4	2.30E-03	1.10	
	MIR1181	3.03E-19	1.26	
	PDE4A	8.49E-25	0.70	
	ICAM3	2.61E-09	1.24	
	ZGLP1	2.38E-02	1.09	
	TYK2	8.31E-02	0.94	
	ICAM1	1.12E-10	1.21	
	FDX1L	3.55E-06	1.17	
	ICAM5	1.28E-03	1.10	
	SLC9A8	5.56E-05	0.90	
	SPATA2 SNAI1	8.65E-21	0.77	
	RNF114	1.30E-01	1.06	Douro
	CCDC116	2.28E-21 2.29E-07	0.66 0.87	Down
	PPIL2	2.29E-07 2.17E-08	1.18	
	YDJC	2.17E-08 2.92E-19	1.26	
	PI4KAP2	2.92E-19 2.29E-07	0.87	
	MIR301B	2.29E-07	0.87	
	RIMBP3B	8.97E-33	0.68	
	UBE2L3	6.67E-38	1.39	
	RIMBP3C	8.97E-33	0.68	
	MIR130B	2.29E-07	0.87	
	SDF2L1	1.22E-12	1.34	
New loci	RUNX3	1.35E-08	1.20	
	EXOC2	8.20E-11	1.25	
	TAGAP	5.59E-04	1.10	
	ELMO1	1.59E-02	0.87	
	TOPORS	5.18E-05	0.86	
	NDUFB6	3.21E-12	1.20	
	DDX58	7.68E-63	3.30	Up
	ZC3H12C	4.50E-31	1.61	Up
	ETS1	8.93E-11	1.31	
	TNP2	8.77E-01	1.02	
	SOCS1	6.56E-21	1.53	Up
	PRM3	2.78E-04	1.11	
	PRM1	1.00E+00	1.02	
	PRM2	4.80E-03	1.09	
	PTRF	3.73E-36	0.67	Down
	STAT3	1.08E-63	2.13	Up
	ATP6V0A1	1.73E-02	1.14	
	SGSH	9.47E-02	0.94	
	SLC26A11	5.95E-13	0.83	

	CARD14	1.29E-68	2.01	Up
	MBD2	2.50E-08	1.16	_
	POLI	6.56E-21	0.54	Down
	STARD6	1.00E+00	1.02	
	SNORA37	7.12E-01	1.03	
	C18orf54	3.55E-02	1.09	
	<i>MIR199A1</i>	3.64E-08	1.21	
	TMED1	1.00E+00	0.99	
	DNM2	2.13E-10	1.18	
	ILF3	1.35E-26	1.37	
	LOC147727	1.00E-08	0.76	
	MIR638	3.64E-08	1.21	
	QTRT1	9.08E-02	1.07	
Conditional Analysi	s IFIH1	1.14E-50	2.79	Up
	FAP	8.25E-01	1.09	
	LNPEP	9.47E-35	0.73	
	ERAP2	5.33E-03	1.21	
	IL12B	2.70E-14	1.22	
	HLA-B	2.26E-15	1.26	
	MICA	1.22E-17	0.60	Down
	HCG26	7.96E-03	0.91	
	CDC37	3.03E-19	1.26	
	RAVER1	1.00E+00	1.00	
	ICAM4	2.30E-03	1.10	
	KEAP1	7.00E-10	1.18	
	MRPL4	2.44E-44	1.44	
	MIR1181	3.03E-19	1.26	
	PDE4A	8.49E-25	0.70	
	ICAM3	2.61E-09	1.24	
	ZGLP1	2.38E-02	1.09	
	TYK2	8.31E-02	0.94	
	ICAM1	1.12E-10	1.21	
	FDX1L	3.55E-06	1.17	
	ICAM5	1.28E-03	1.10	

Supplementary Table 10. Variance in liability

The variance in liability explained by each locus, as determined under a liability model⁶. Prevalence of psoriasis was set as 2% when estimating the variance. Assuming the multiple loci have an additive effect on the risk of psoriasis, the variance in liability explained by the 19 known loci, 15 new loci, and the 5 secondary signals identified by conditional analysis are 11.35%, 1.60%, and 1.36%, respectively.

	Notable	Risk allele Freq	Risk allele Freq		Variance in	
SNP	nearby genes	(case)	(control)	OR	RR	liability
Known loci						
rs7552167	IL28RA	0.878	0.858	1.21	1.21	0.14%
rs9988642	IL23R	0.952	0.929	1.52		0.14%
rs6677595	LCE3D	0.689	0.64	1.52		0.30%
rs62149416	REL	0.671	0.635	1.17		0.40%
rs17716942	IFIH1	0.891	0.863	1.17		0.19%
rs27432	ERAP1	0.309	0.863	1.27	1.27	0.22%
	IL13/IL4					
rs1295685		0.807	0.798	1.18		0.14%
rs2233278	TNIP1	0.09	0.058	1.59		0.41%
rs12188300	IL12B	0.132	0.095	1.58		0.62%
rs4406273	HLA-C	0.259	0.092	4.32		6.44%
rs33980500	TRAF3IP2	0.108	0.074	1.52		0.41%
rs582757	TNFAIP3	0.315	0.273	1.23		0.28%
rs2066819	IL23A/STAT2	0.948	0.934	1.39		0.21%
rs8016947	NFKBIA	0.6	0.564	1.16		0.18%
rs12445568	FBXL19	0.403	0.368	1.16		0.17%
rs28998802	NOS2	0.17	0.145	1.22		0.16%
rs34536443	TYK2	0.974	0.953	1.88		0.54%
rs1056198	RNF114	0.6	0.573	1.16		0.18%
rs4821124	UBE2L3	0.208	0.189	1.13	1.13	0.08%
<u>New loci</u>						
rs11121129	SLC45A1	0.308	0.287	1.13		0.10%
rs7536201	RUNX3	0.528	0.494	1.13		0.12%
rs10865331	B3GNT2	0.404	0.374	1.12		0.10%
rs9504361	EXOC2/IRF4	0.574	0.546	1.12		0.10%
rs2451258	TAGAP	0.362	0.348	1.12		0.10%
rs2700987	ELMO1	0.591	0.564	1.11		0.09%
rs11795343	DDX58	0.628	0.597	1.11		0.09%
rs10979182	KLF4	0.617	0.591	1.12	1.12	0.10%
rs4561177	ZC3H12C	0.617	0.581	1.14	1.14	0.14%
rs3802826	ETS1	0.505	0.484	1.12	1.12	0.11%
rs367569	SOCS1	0.729	0.709	1.13	1.13	0.10%
rs963986	STAT3,	0.169	0.154	1.15	1.15	0.08%
rs11652075	CARD14	0.53	0.502	1.11	1.11	0.09%
rs545979	STARD6,POLI,	0.317	0.291	1.12	1.12	0.09%
rs892085	ILF3,CARM1	0.593	0.558	1.17	1.17	0.20%
Conditional analysis loci						
rs2111485	IFIH1	0.647	0.61	1.14	1.14	0.13%
rs2910686	ERAP2	0.442	0.437	1.12		0.10%
rs4379175	IL12B	0.737	0.678	1.31		0.51%
rs13437088	MICA	0.342	0.251	1.32		0.48%
rs12720356	TYK2	0.929	0.911	1.25		0.13%

Supplementary Figures

Supplementary Figure 1: Manhattan plot for meta-analysis

The 34 susceptibility loci that achieve genome-wide significance (above the green line) in the meta-analysis. The 19 known loci are colored blue, and the 15 new loci are colored red. Only SNPs with P-values $\ge 1 \times 10^{-50}$ are shown.



Supplementary Figure 2. Regional association plots

Regional association plots using LocusZoom⁷ to show the combined p-values in each of the 15 new loci. The most significant SNP from each locus was used as the index SNP to compute the linkage





11

11.2

11.4

Position on chr16 (Mb)

11.6

11.8









Supplementary Figure 3. Manhattan plot for the results of the conditional analysis. Green dots shown SNPs in the five loci that achieve genome-wide significance (P=5x10⁻⁸, denoted by red line).





Supplementary Figure. 4. Regional association plots for the significant loci in the conditional meta-analysis.



Supplementary Figure 5. Plots illustrating the phenotype overlap for the susceptibility variants listed in Table 1.

Only disease associations that are listed in the NHGRI GWAS catalog

Α.

(http://www.genome.gov/gwastudies) and Immunochip results for Celiac disease⁴ were included. Genes are shown in red and phenotypes in black. A) Shows the overlap with phenotypes where the identified variant is in LD (r2>0.7) with the identified psoriasis variant. B) Shows all loci shared with phenotypes that have identified variants within 500kb of the identified psoriasis variant for that locus. Phenotypes sharing the most loci with psoriasis are generally more central to the plot, while those sharing only one or two loci are situated on the outside (Plots were produced using Gephi available at http://gephi.org/)⁸. Owing to the number of connections with other phenotypes, the HLA locus was removed from the plot.

AD: Atopic dermatitis, AS: Ankylosing spondylitis, BD: Behcet's disease, CD: Crohn's disease, CeD: Celiac disease, IgE: Serum IgE, IgA: Selective Immunoglobulin A deficiency, LE: Leprosy, MS: Multiple Sclerosis, PBC: primary biliary cirrhosis, PS: Psoriasis, , RA: rheumatoid arthritis, SI: Soluble ICAM-1, SLE: Systemic lupus erythematosus, SS: Systemic sclerosis T1D:Type I Diabetes, UC: Ulcerative colitis.





Supplementary note

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Additional Methods and results

Association Analysis

Logistic regression was used to perform association analysis for the imputed dosages for the GWAS datasets: Kiel (mach2dat^{7,9}), CASP (mach2dat), and the WTCCC2 (SNPTESTv2¹⁰). Linear mixed modeling as implemented in EMMAX¹¹ was used to perform association analyses for the two Immunochip datasets separately due to its ability to control for population stratification and cryptic relatedness. The Balding-Nicholls algorithm¹² was used to generate the kinship matrix.

Meta-analysis

Sample size weighting was employed to combine P-values across the 5 studies using METAL, employing effective sample sizes¹³. To calculate the meta-odds ratios (ORs) for each SNP, we performed logistic regression analysis for the Immunochip data, using the top ten PCs to adjust for population stratification. The meta-ORs were then calculated as follows:

$$\beta_{weighted} = \frac{\sum_{i=1}^{D} N_i^{eff} \beta_i}{\sum_{i=1}^{D} N_i^{eff}}$$
$$OR_{meta} = e^{\beta_{weighted}}$$

where β_i is the coefficient estimated from the logistic regression analysis, and D is the number of datasets.

Conditional analysis of the newly identified signal at 10.82 Mb on chr19

Conditional analysis was performed to determine if the signal tagged by rs892085 at 10.82 Mb on chromosome 19 is new signal associated with psoriasis. We conditioned on two SNPs from the known locus near *TYK2*: i) the previously identified SNP rs12720356³; ii) the current best SNP from our meta-analysis rs34536443. Each conditional analysis was first performed individually for each study (logistic regression framework for the three GWAS; linear mixed model for the Immunochip datasets). Meta-analysis based on the conditional results was then performed using METAL¹³. Since rs34536443 has poor imputation quality (r2=0.17) in the CASP dataset, we excluded this dataset when conditioning on rs34536443. As shown in **Supplementary Table 5**, rs892085 achieves genome-wide significance after conditioning on either of the two *TYK2* SNPs, indicating that it is an independent signal for psoriasis susceptibility.

Conditional analysis

We performed conditional analysis for the 3 GWAS datasets using logistic regression, whereas we used the linear mixed modeling implemented in EMMAX for the two Immunochip datasets. The most strongly associated SNPs from the 19 known loci achieving genome-wide significance

in this study and the 15 new loci (**Table 1**) were used as covariates for all five studies. Because the CASP GWAS did not have good imputation quality ($r^2>0.3$) for SNP rs34536443; the second best SNP in the 19q13.2 region (rs2304256: $P_{comb} = 1.20 \times 10^{-20}$) was used in the conditional analysis of the CASP dataset. We next used METAL to combine the conditional analysis results; for the 19q13.2 region (ie ±500 kb surrounding SNP rs34536443) the meta-analysis excluded the CASP dataset, whereas we used all five datasets for the other regions.

Conditional analysis on the ERAP2 signal

For the follow-up conditional analysis of the *ERAP2* signal, we used only the best ERAP1 SNP (rs27432) as a covariate. Conditioning only on rs27432, the most significant *ERAP1* SNP in the unconditional analysis, continued to support the *ERAP2* signal ($P = 3.6 \times 10^{-7}$). When considered together with the LD results between the 2 SNPs (r2=0.17 and D'=0.75 for PAGE; r2=0.18 and D'=0.76 for GAPC), these suggest the two loci might have opposite effects arising from the same haplotype. To test this, we performed a haplotype association test using the PAGE phased genotypes data for rs27432 and rs2910686. The risk/non-risk alleles for these SNPs are A/G and C/T, respectively, and their haplotype frequencies are: 0.03 (AC), 0.25 (AT), 0.41 (GC), and 0.31 (GT). The haplotype association analysis for ERAP1-ERAP2 was performed using logistic regression, with haplotype counts of AT (ERAP1 risk / ERAP2 nonrisk), GC (ERAP1 nonrisk / ERAP2 risk), and AC (ERAP1 risk / ERAP2 risk) as 3 independent variables, and the top 10 PCs as covariates. We found that the AT (ERAP1 risk / ERAP2 risk) haplotype is strongly associated ($P = 3.1 \times 10^{-6}$), the GC (ERAP1 nonrisk / ERAP2 risk) haplotype is not associated (P = 0.25), suggesting that the genetic effect of *ERAP2* is masked by *ERAP1*.

Causal SNP lookup

We identified SNPs in strong LD ≹0.9) with the most significant SNP from each of the known and new loci listed in Table 1, including the secondary signals identified by the conditional analysis. LD among SNPs was computed from 379 European-ancestry samples in the 1000 Genomes project (May 21st 2011 version). We then used ANNOVAR¹⁴ to annotate each of these SNPs. All identified SNPs affecting the predicted protein sequence were missense variants; none were nonsense or splicing mutations. SIFT¹⁵ and PolyPhen¹⁶ were used to predict the impact of the mutations on the function of the protein.

Epistasis

We performed an analysis of epistasis using the most significantly associated SNP from each of the 34 loci in **Tables 1** reaching genome-wide significance in this study. Logistic regression was used to model epistasis in the five datasets using a risk allele dosage model; for the PAGE and GAPC datasets the top 10 PCs were included as covariates. For each pair of SNPs, the likelihood ratio test was employed to compute the p-value of the interaction term for each dataset. Epistasis results were combined using METAL, again omitting the CASP dataset for the 19q13.2 region.

Gene expression

We retrieved the SNPs that reside within 500 kb (3 Mb for MHC) of each of the most strongly associated SNPs identified from the known or new loci, including secondary signals from the conditional analysis. Using European-ancestry samples from the 1000 Genomes Project, we then used the tag SNP function from PLINK to identify genomic regions in strong LD ($r^2>0.7$) with the most significant SNP; we then extended the tagged regions by 50kb on each side. We identified genes overlapping any of the extended regions, and we used false discovery rate (FDR) ≤ 0.05 and fold-change (FC) ≤ 0.67 or ≥ 1.5 to declare genes as differentially expressed in psoriatic skin lesions from a previous microarray experiment^{17,18}.

eQTL lookup

To check whether any of the 34 genome-wide significant SNPs that were known or new and the 5 secondary signals identified by conditional analysis were eQTLs, we queried the cis-psoriasis eQTL database (in normal and psoriatic skin) compiled by Ding *et al.*¹⁹, and we used $P<1x10^{-7}$ as criteria to look for eQTLs using expression of microarray probesets corresponding to Entrez genes. (available at http://www.sph.umich.edu/csg/junding/eQTL/TableDownload/).

Since the imputation was performed using Hapmap reference panel in Ding *et al.*, we performed eQTL analysis using our imputed CASP GWAS data (using 1000 genomes for reference panel) and their corresponding gene expression microarray data (data as described in Ding *et al.*¹⁹). The significant results are consistent with the cis-psoriasis eQTL database.

Heritability explained by psoriasis-associated SNPs

We estimated the variance in liability (locus-specific heritability)²⁰ that can be explained by the known, new, and the secondary signals of psoriasis-associated SNPs using the approach described by So *et al.*⁶. We set the prevalence of the disease as 0.02, and calculated risk ratios (RR) from our estimated ORs using an iterative approach²¹.

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