

Two Functional Lupus-Associated *BLK* Promoter Variants Control Cell-Type- and Developmental-Stage-Specific Transcription

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Efforts to identify lupus-associated causal variants in the *FAM167A/BLK* locus on 8p21 are hampered by highly associated noncausal variants. In this report, we used a trans-population mapping and sequencing strategy to identify a common variant (rs922483) in the proximal *BLK* promoter and a tri-allelic variant (rs1382568) in the upstream alternative *BLK* promoter as putative causal variants for association with systemic lupus erythematosus. The risk allele (T) at rs922483 reduced proximal promoter activity and modulated alternative promoter usage. Allelic differences at rs1382568 resulted in altered promoter activity in B progenitor cell lines. Thus, our results demonstrated that both lupus-associated functional variants contribute to the autoimmune disease association by modulating transcription of *BLK* in B cells and thus potentially altering immune responses.

Introduction

The gene structures of *BLK* (MIM 191305), a member of the *src*-family tyrosine kinases, have been described in B cells previously.¹ More recently, the *BLK*-deficiency-induced underdevelopment of IL-17-producing $\gamma\delta$ T cells has impli-

cated a critical role of expression-altering *BLK* variants in the pathogenesis of autoimmune diseases.² Studies with *Blk*-deficient mice suggest that *BLK* influences both B and T cell development and proliferation.^{2,3} This locus is associated with multiple autoimmune diseases, including systemic lupus erythematosus (SLE [MIM 152700]), systemic

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Table 1. Demographics of SLE Populations Studied

Ancestry	Affected Individuals			Control Individuals		
	Male	Female	Total	Male	Female	Total
European	344	3,617	3,980	1,181	2,365	3,546
African American	109	1,297	1,406	5,45	1,189	1,734
East Asian	101	1,171	1,272	1,158	1,112	1,270

sclerosis (MIM 181750), rheumatoid arthritis (MIM 180300) and Sjögren's syndrome (MIM 270150).^{4–11} Analyses of expression in transformed B cell lines demonstrate that risk-conferring variants within *FAM167A* (MIM 610085) and *BLK* are associated with altered mRNA expression of both *FAM167A* and *BLK*; however, the causal alleles and mechanisms remain undefined.⁷

Like other genes with TATA-less promoters, the genomic DNA upstream of exon 1 of *BLK* has two transcription start sites and promoters that drive *BLK* transcription: a ubiquitous proximal promoter (P1) and a B-lymphocyte-specific promoter (P2).¹ Recent evidence suggests that immature B cells from individuals carrying lupus risk alleles have lower amounts of *BLK* than such cells from individuals without lupus risk alleles.¹²

In this study, we leveraged the difference in linkage disequilibrium (LD) structure across populations to examine the *FAM167A/BLK* locus in a multiethnic population of SLE cases and controls and then used focused resequencing to identify additional lupus-associated variants. Functional assessment revealed the molecular mechanism impacted by the variant alleles. Using this approach, we successfully identified two functional variants that regulate transcription from the promoters in a cell-type- and developmental-stage-specific fashion.

Subjects and Methods

Study Subjects

Approval by the institutional review boards of the Oklahoma Medical Research Foundation and the collaborators' institutions was obtained prior to sample collection. All study participants provided written consent at the time of sample collection. Deidentified genomic DNA samples from individuals with SLE and control subjects were analyzed from 6,658 unrelated individuals (3,980 individuals of European ancestry [EA], 1,272 of Asian ancestry [AS], and 1,406 of African American ancestry [AA]) and 6,550 unrelated controls (3,546 EA, 1,270 of AS, and 1,734 AA) (Table 1). These samples were obtained through the Lupus Family Registry and Repository (LFRR) as part of the Oklahoma Rheumatic Disease Research and Cores Center (ORDRCC) and through collaborators from 24 additional study sites. Collaborators and the sources of all case and control individuals used in these studies are shown in Table S1 in the Supplemental Data available online.

For resequencing experiments, deidentified genomic DNA samples from individuals with SLE and controls were obtained from the Autoimmune Biomarkers Collaborative Network

(ABCOn) of the New York Cancer Project (NYCP) (191 EA SLE individuals and 96 EA controls) courtesy of Dr. Gregersen for the discovery cohort (Table S2). All individuals with SLE met classification criteria¹³ (American College of Rheumatology). All samples were independent. Only one randomly selected SLE sample was included if multiple affected individuals were available from a multiplex lupus pedigree. DNA was obtained from blood samples.

Genotyping and Quality Control

All samples were genotyped as a part of a joint effort of more than 40 investigators from around the world. These investigators contributed samples, funding, and hypotheses used for designing a custom, highly multiplexed Illumina-bead-based array method on a BeadStation system.¹⁴ Select SNPs were also assayed for genotype confirmation via TaqMan methods (Applied Biosystems). Genotyping facilities are located at the Oklahoma Medical Research Foundation, and data were sent to a central data center at Wake Forest Medical Center for quality control. These data were then distributed back to the investigators who had requested specific SNPs for final analysis and publication.

Genotype data were only used from samples with a call rate greater than 90% of the SNPs screened (98.05% of the samples). For analyses, only genotype data from SNPs with a call frequency greater than 90% in the samples tested and an Illumina GenTrain score greater than 0.7 (96.74% of all SNPs screened) were used. In addition, at least one previously genotyped sample was randomly placed on each assay plate and used for tracking samples through the genotyping process. More information on Illumina genotyping can be found at the Illumina website (Web Resources section).

Correction for Population Stratification

Following best practices in genome-wide association studies, we used all of the genotype data from all SNPs that passed quality control, including the published set of ancestry-informative makers (AIMs),¹⁵ and computed the principal components and admixture estimates. Regions of known extended LD were removed. The combination of 12,000 SNPs, including published sets of AIMs and the principal-component analysis computed across all ethnicities, generated principal components that separated ethnicities. To minimize the inflation of the test statistics, we included population-specific principal components in the logistic regression models as covariates.^{15,16} Population clustering based upon the three-dimensional plot of principal component 1 (PC1), PC2, and PC3 of the final samples used in these studies is presented (Figure S1).

Imputation-Based Association Analysis

Initially, we genotyped 372 SNPs within the *FAM167A/BLK* region (11,033,737–11,618,107 bp, hg19), and after performing quality control (HWE > 0.001 in controls and minor allele frequency [MAF] > 0.01), we had 329 SNPs in AA samples, 259 SNPs in EA samples, and 201 SNPs in AS available for imputation. To investigate the new variants in the *FAM167A/BLK* region, we used the 1000 Genomes project¹⁷ as a reference panel for imputation to estimate missing genotypes. After quality control measures (HWE > 0.001 in controls and MAF > 0.01) for the 1000 Genomes project reference panel, which contains 11,528 SNPs within the *FAM167A/BLK* region, we used 246 AA samples with 4,813 SNPs, 381 EA samples with 2,508 SNPs, and 286 AS samples with

1,847 SNPs for imputation. Imputation was carried out with MACH,^{17,18} which provided a quantitative assessment of estimate uncertainty (Rsq). All imputed SNPs were filtered with the quality controls (HWE > 0.001, MAF > 0.01, and Rsq > 0.6), and 2,137 SNPs in AA samples, 1,199 SNPs in EA samples, and 738 SNPs in AS samples were used for further analysis. At each SNP, p value, odds ratio (OR), and 95% confidence interval (CI) were calculated with gPLINK.¹⁹ We calculated allelic association results (Table 2 and Table S3) to account for imputation uncertainty with mach2 dat;²⁰ genotyped and imputed SNPs with p values ≤ 0.05 from at least one population are shown.

For each ethnic population, we used WHAP¹⁹ to calculate pairwise conditional analysis for each pair of SNPs (the most significant SNP plus each other SNP) and identify the independent effects for each SNP. We assessed whether the joint effect is explained by a single SNP. If a haplotype was significant and remained significant after we conditioned on a SNP, then that SNP did not independently account for the association. However, if the p value was no longer significant after we conditioned on a SNP, then we considered that SNP to be the source of the association.

Resequencing of *FAM167A/BLK* Exons and the Upstream Promoter Region

Resequencing was performed on 191 individuals with SLE and 96 controls from ABCoN, as detailed above (Table S2). All 13 exons and the 2.5 kb upstream promoter sequence were resequenced with whole-genome amplified genomic DNA (Cat#150045, QIAGEN). Primers for resequencing were designed to target the 13 exon regions and 2.5 kb upstream promoter region. PCR amplification was performed on genomic DNA via high-fidelity Taq polymerase according to standard protocols. PCR product purity and size were assessed on 2% agarose gels. Sanger sequencing was performed per the manufacturer's protocol. Sequence trace files were manually analyzed for variations.

Haplotype Analysis

We used the expectation-maximization algorithm in the WHAP program¹⁹ to estimate haplotype frequencies. WHAP directly calculates likelihood estimates, likelihood ratios, and p values by taking into account the information loss due to haplotype-phase uncertainty and missing genotypes. Association between inferred haplotypes and SLE was tested with an omnibus test. We used both conditional analysis and global haplotype analysis to disentangle the correlation structure in which SNPs are truly associated with phenotype. To test which of the associated SNPs were causal and which were significantly associated by LD, we performed haplotype conditional analysis on each SNP. If the global haplotype association disappeared, then the specific SNP on which we had conditioned accounted for the whole association.

Nuclear Extract Preparation

Nuclear extracts from the human Jurkat T cell line, RS4;11 pro-B cell line, Nalm-6 and Reh pre-B cell lines, Ramos immature B cell line, and Daudi mature B cell line (American Type Tissue Culture Collection) were obtained. Cells were maintained in RPMI with 10% heat-inactivated fetal bovine serum, L-glutamine (2 mM), and penicillin and streptomycin (100 units/ml). Nuclear protein extracts were prepared from cells, dialyzed against a buffer composed of 20 mM HEPES, 20% glycerol, 0.1M potassium chlo-

ride, and 0.2 mM EDTA (pH 7.9), and used in nuclear binding assays (Figures S2 and S3).²¹

Electrophoretic Mobility-Shift Assay

A forward and reverse 21 base pair synthetic oligonucleotide from the *BLK* promoter flanking the rs922483 polymorphism was purchased from Integrated DNA Technologies. All oligos were purified with polyacrylamide gel electrophoresis. Probes carrying the risk allele (T) and nonrisk allele (C) were generated, and pairs of one forward and one reverse oligonucleotide were mixed in equal molar ratios, heated, and then allowed to anneal to generate the 21 bp, double-stranded probes. T4 polynucleotide kinase (Invitrogen) was used for labeling the end of each DNA probe with (γ -³²P) adenosine triphosphate (Amersham). The nuclear extracts prepared as discussed above were incubated for 25 min at 37°C with labeled probes in binding buffer (1 μ g poly(dI-dC), 20 mM HEPES, 10% glycerol, 100 mM KCl, and 0.2 mM EDTA [pH 7.9]). DNA-protein complexes were resolved on denaturing 5% acrylamide gels. For supershift assays, varying concentrations of anti-pol II antibody (clone 8A7 and clone H-224, Santa Cruz) were added to the DNA-protein complexes; this was followed by incubation for 15 min prior to resolution on denaturing 5% acrylamide gels (Figure S3).

Luciferase Reporter Assay

We amplified the upstream sequence (−2,256 to +55 bp) of *BLK* by using genomic DNA from individuals with nonrisk haplotypes. PCR products were cloned into pCR2.1-TOPO vector (Invitrogen, Cat# K4500-01) and subcloned into pGL4 luciferase reporter vectors (Promega, Cat# E6651, Madison, WI). The construct carrying the nonrisk haplotype was used as a template for mutagenesis (Stratagene) to create other allelic haplotypes.

An internal control reporter vector, pRL-TK, containing *Renilla* luciferase driven by the thymidine kinase promoter was simultaneously transfected with our experimental vectors as a control for assay-to-assay variability. The *Renilla* luciferase activity expressed by the internal control vector was used for normalization of transfection efficiency. One to 5 μ g of each vector was transfected into the Jurkat (1 × 10⁶/sample in triplicate), RS4;11 (2 × 10⁶/sample in triplicate), Nalm-6 (3 × 10⁶/sample in triplicate), Ramos (3 × 10⁶/sample in triplicate), and Daudi (5 × 10⁶/sample in triplicate) cell lines. Cells were then incubated at 37°C for 16 hr. Luciferase activity was measured with the Dual-Luciferase Reporter Assay System (Promega, Cat# E1960). Luciferase activity was normalized through division of *BLK* risk or nonrisk construct reporter activity by the reporter activity of the pRL-TK construct. The mean and standard error of measurement were calculated on the basis of the normalized luciferase activities and used for further analysis.

Results

Trans-Population Association Testing Identified rs922483 as the Predominant SLE-Associated Causal Variant

To identify the causal variants responsible for the association of *FAM167A/BLK* with SLE, we genotyped 372 SNPs selected from the phase II HapMap in the region spanning 584.37 kb (11,033,737–11,618,107 bp, hg19) in chromosomal region 8p21 in three ethnic populations.

Table 2. Association- and Conditional-Analysis Results for Significantly Associated Peak Genotyped and Imputed SNPs

Chr.	dsSNP	BP (build37)	Allele1/Allele2	European American ^a					Asian ^b					African American ^c				
				Freq. Allele1 (Case/Control) ^d	OR (95% CI)	r ² Peak	p _{cond on} rs998683	Freq. Allele1 (Case/Control)	OR (95% CI)	r ² Peak	p _{cond on} rs1478901	Freq. Allele1 (Case/Control)	OR (95% CI)	r ² Peak	p _{cond on} rs2736345	p _{cond on} rs922483		
8	rs2409780	11,337,587	T/C	0.699/0.752	3.20 × 10 ⁻¹³ (0.77–0.82)	0.93	0.97	0.189/0.267	2.10 × 10 ⁻¹¹ (0.64–0.73)	0.85	0.29	0.822/0.865	3.913 × 10 ⁻⁰⁶ (0.73–0.83)	0.73	0.474	0.020	0.100	
8	rs1564267 ^e	11,337,887	A/G	0.154/0.167	3.54 × 10 ⁻⁰² (0.91–0.83–1)	0.08	0.91	0.166/0.237	3.11 × 10 ⁻¹⁰ (0.64–0.74)	0.72	0.38	0.429/0.462	0.01079 (0.88–0.97)	0.070	0.127	0.353		
8	rs2618444	11,338,370	A/C	0.699/0.752	3.06 × 10 ⁻¹³ (0.77–0.82)	0.93	0.97	0.189/0.267	2.30 × 10 ⁻¹¹ (0.64–0.73)	0.85	0.29	0.823/0.865	0.00000411 (0.73–0.84)	0.475	0.019	0.095		
8	rs62489069	11,338,383	A/G	0.67/0.72	1.81 × 10 ⁻¹¹ (0.79–0.85)	0.80	0.97	0.168/0.238	3.82 × 10 ⁻¹⁰ (0.64–0.74)	0.72	0.36	0.752/0.791	0.000237 (0.8–0.9)	0.238	0.061	0.166		
8	rs35393613	11,338,466	C/T	0.67/0.72	1.78 × 10 ⁻¹¹ (0.79–0.85)	0.80	0.96	0.168/0.238	5.26 × 10 ⁻¹⁰ (0.64–0.74)	0.72	0.40	0.776/0.813	0.0004116 (0.8–0.91)	0.289	0.068	0.184		
8	rs1531577	11,338,561	T/C	0.712/0.694	1.49 × 10 ⁻⁰² (1.09–1.17)	0.16	0.57	0.835/0.766	3.50 × 10 ⁻¹⁰ (1.56–1.8)	0.73	0.29	0.834/0.805	0.004446 (1.2–1.37)	0.077	0.343	0.150		
8	rs2061831	11,339,882	T/C	0.699/0.752	2.42 × 10 ⁻¹³ (0.76–0.82)	0.94	0.87	0.188/0.265	5.40 × 10 ⁻¹¹ (0.64–0.74)	0.87	0.26	0.823/0.865	4.401 × 10 ⁻⁰⁶ (0.73–0.84)	0.478	0.021	0.116		
8	rs2736332	11,339,965	C/G	0.326/0.271	1.52 × 10 ⁻¹³ (1.3–1.4)	0.82	0.31	0.813/0.735	2.93 × 10 ⁻¹¹ (1.57–1.79)	0.87	0.19	0.599/0.563	0.005471 (1.16–1.28)	0.253	0.724	0.895		
8	rs7812879 ^a	11,340,181	G/A	0.856/0.843	3.35 × 10 ⁻⁰² (1.1–1.2)	0.07	0.82	0.836/0.766	4.78 × 10 ⁻¹⁰ (1.55–1.79)	0.73	0.33	0.8/0.775	0.01928 (1.15–1.3)	0.094	0.166	0.515		
8	rs2254891 ^e	11,341,129	C/G	0.712/0.694	1.29 × 10 ⁻⁰² (1.09–1.17)	0.16	0.58	0.826/0.759	2.31 × 10 ⁻⁰⁹ (1.52–1.75)	0.76	0.68	0.848/0.828	0.02776 (1.16–1.33)	0.061	0.665	0.359		
8	rs2736336	11,341,870	G/T	0.699/0.752	2.19 × 10 ⁻¹³ (0.76–0.82)	0.94	1.00	0.197/0.272	4.19 × 10 ⁻¹⁰ (0.65–0.75)	0.90	0.87	0.794/0.838	4.237 × 10 ⁻⁰⁶ (0.74–0.84)	0.348	0.034	0.096		
8	rs2736337	11,341,880	T/C	0.699/0.752	2.24 × 10 ⁻¹³ (0.76–0.82)	0.94	0.98	0.197/0.272	3.96 × 10 ⁻¹⁰ (0.65–0.75)	0.89	0.78	0.795/0.84	2.178 × 10 ⁻⁰⁶ (0.73–0.83)	0.325	0.024	0.069		
8	rs2736338	11,341,883	A/C	0.699/0.752	2.23 × 10 ⁻¹³ (0.76–0.82)	0.94	0.98	0.197/0.272	4.00 × 10 ⁻¹⁰ (0.65–0.75)	0.90	1.00	0.795/0.84	0.00000218 (0.73–0.83)	0.325	0.024	0.069		
8	rs2254660	11,342,986	G/C	0.859/0.848	6.63 × 10 ⁻⁰² (1.09–1.19)	0.07	0.99	0.829/0.759	9.49 × 10 ⁻¹⁰ (1.54–1.77)	0.78	0.60	0.894/0.876	0.03329 (1.19–1.4)	0.030	0.411	0.217		
8	rs2254546	11,343,680	G/A	0.855/0.843	3.37 × 10 ⁻⁰² (1.1–1.2)	0.07	0.82	0.828/0.759	1.04 × 10 ⁻⁰⁹ (1.54–1.77)	0.78	0.66	0.876/0.858	0.03353 (1.17–1.36)	0.045	0.609	0.336		
8	chr11343717	11,343,717	A/G	–/–	–	–	–	–/–	–	–	–	0.979/0.97	0.03673 (1.4–1.94)	0.014	0.159	0.118		

(Continued on next page)

Table 2. Continued

Chr.	dsSNP	BP (build37)	Allele1/Allele2	European American ^a					Asian ^b					African American ^c					
				Freq. Allele1 (Case/Control) ^d	Adj. p	OR (95% CI)	r ² Peak	p _{cond on} rs998683	Freq. Allele1 (Case/Control)	Adj. p	OR (95% CI)	r ² Peak	p _{cond on} rs1478901	Freq. Allele1 (Case/Control)	Adj. p	OR (95% CI)	r ² Peak	p _{cond on} rs2736345	p _{cond on} rs922483
8	rs2736340 ^S	11,343,973	G/A	0.7/0.753	2.09 × 10 ⁻¹³	0.76 (0.71–0.82)	0.94	0.89	0.188/0.265	9.16 × 10 ⁻¹¹	0.65 (0.56–0.74)	0.87	0.31	0.824/0.866	5.323 × 10 ⁻⁰⁶	0.73 (0.64–0.84)	0.481	0.024	0.129
8	rs2618473 ^S	11,344,127	G/A	0.69/0.743	3.25 × 10 ⁻¹³	0.77 (0.71–0.83)	0.89	0.79	0.189/0.265	8.64 × 10 ⁻¹¹	0.65 (0.56–0.74)	0.87	0.32	0.552/0.582	0.01564	0.88 (0.8–0.98)	0.033	0.478	0.258
8	rs4840565 ^S	11,345,545	G/C	0.33/0.278	8.07 × 10 ⁻¹²	1.27 (1.19–1.37)	0.81	0.98	0.823/0.754	1.84 × 10 ⁻⁰⁹	1.52 (1.32–1.75)	0.81	0.88	0.36/0.312	0.00008384	1.23 (1.11–1.36)	0.529	0.192	0.431
8	rs2736342 ^S	11,347,289	A/C	0.49/0.448	3.67 × 10 ⁻⁰⁷	1.18 (1.11–1.26)	0.39	0.68	–/–	-	-	-	-	0.556/0.523	0.00846	1.14 (1.03–1.26)	0.315	0.868	0.930
8	rs1478900 ^S	11,347,660	A/G	0.854/0.844	6.64 × 10 ⁻⁰²	1.09 (0.99–1.19)	0.07	0.96	0.807/0.736	1.15 × 10 ⁻⁰⁹	1.51 (1.32–1.73)	0.89	0.30	0.874/0.857	0.04142	1.16 (1–1.35)	0.042	0.607	0.343
8	rs1478901 ^S	11,347,833	C/G	0.701/0.754	2.92 × 10 ⁻¹³	0.77 (0.71–0.82)	0.95	0.99	0.208/0.29	1.32 × 10 ⁻¹¹	0.64 (0.56–0.73)	1.00	-	0.822/0.864	0.00000525	0.73 (0.64–0.84)	0.477	0.039	0.140
8	chr11348647	11,348,647	C/A	–/–	-	-	-	-	–/–	-	-	-	-	0.982/0.987	0.02851	0.61 (0.39–0.96)	0.034	0.529	0.530
8	rs9693589	11,348,961	G/A	0.701/0.754	2.96 × 10 ⁻¹³	0.77 (0.71–0.82)	0.95	1.00	0.212/0.291	4.15 × 10 ⁻¹¹	0.65 (0.57–0.74)	0.94	collinear	0.824/0.866	5.801 × 10 ⁻⁰⁶	0.73 (0.64–0.84)	0.487	0.024	0.116
8	rs13277113 ^S	11,349,186	G/A	0.701/0.754	2.98 × 10 ⁻¹³	0.77 (0.71–0.82)	0.95	1.00	0.212/0.291	4.28 × 10 ⁻¹¹	0.65 (0.57–0.74)	0.94	collinear	0.824/0.866	5.739 × 10 ⁻⁰⁶	0.73 (0.64–0.84)	0.487	0.024	0.116
8	rs9694294 ^S	11,350,721	C/G	0.855/0.843	4.22 × 10 ⁻⁰²	1.1 (1–1.2)	0.07	0.93	0.817/0.747	8.79 × 10 ⁻¹⁰	1.52 (1.33–1.75)	0.77	0.66	0.839/0.812	0.004564	1.21 (1.06–1.38)	0.077	0.369	0.182
8	rs1478902 ^S	11,350,774	A/C	–/–	-	-	-	-	–/–	-	-	-	-	0.984/0.977	0.04526	1.44 (0.99–2.08)	0.016	0.176	0.142
8	rs4840568 ^S	11,351,019	G/A	0.675/0.73	1.46 × 10 ⁻¹³	0.77 (0.71–0.83)	0.83	0.27	0.208/0.287	5.67 × 10 ⁻¹¹	0.65 (0.57–0.74)	0.91	collinear	0.634/0.665	0.0106	0.87 (0.79–0.97)	0.162	0.246	0.597
8	rs922483 ^S	11,351,912	A/G	0.344/0.291	5.27 × 10 ⁻¹²	1.27 (1.19–1.36)	0.76	0.43	0.807/0.735	1.06 × 10 ⁻⁰⁹	1.51 (1.32–1.73)	0.83	0.98	0.308/0.252	1.151 × 10 ⁻⁰⁶	1.31 (1.17–1.47)	1.000	0.069	-
8	chr11351937	11,351,937	G/T	–/–	-	-	-	-	–/–	-	-	-	-	0.984/0.977	0.04802	1.44 (0.99–2.09)	0.016	0.196	0.158
8	rs2250788 ^S	11,352,056	G/A	0.855/0.843	3.83 × 10 ⁻⁰²	1.1 (1–1.2)	0.07	0.89	0.818/0.747	8.27 × 10 ⁻¹⁰	1.53 (1.33–1.75)	0.76	0.56	0.843/0.818	0.009211	1.19 (1.04–1.37)	0.084	0.376	0.222
8	rs13272061 ^S	11,352,261	C/A	0.5/0.459	6.15 × 10 ⁻⁰⁷	1.18 (1.1–1.26)	0.37	0.59	–/–	-	-	-	-	0.862/0.844	0.04183	1.15 (1–1.33)	0.071	0.711	0.450

(Continued on next page)

Table 2. Continued

Chr. dsSNP	BP (build37)	European American ^a				Asian ^b				African American ^c					
		Allele1/ Allele2	Allele1/ (Case/ Control) ^d	OR (95% CI)	r ² Peak	p _{cond} on rs998683	Freq Allele1 (Case/ Control)	OR (95% CI)	r ² Peak	p _{cond} on rs1478901	Freq Allele1 (Case/ Control)	OR (95% CI)	r ² Peak	p _{cond} on rs922483	
				Adj. p											
8	rs2736345 [§]	11,352,485 G/A	0.355/0.301	1.08 × 10 ⁻¹²	1.28 (1.19–1.37)	0.81	0.41	4.83 × 10 ⁻¹⁰	1.53 (1.34–1.76)	0.817/0.745	0.414/0.355	1.486 × 10 ⁻⁰⁶	1.28 (1.15–1.42)	0.626	0.152
8	rs2618476 [§]	11,352,541 A/G	0.689/0.744	6.21 × 10 ⁻¹⁴	0.76 (0.71–0.82)	1.00	collinear	1.78 × 10 ⁻¹¹	0.64 (0.56–0.73)	0.196/0.276	0.824/0.863	0.00001892	0.75 (0.65–0.86)	0.476	0.122
8	rs998683 [§]	11,353,000 G/A	0.689/0.745	5.22 × 10 ⁻¹⁴	0.76 (0.71–0.82)	1.00	-	1.18 × 10 ⁻¹⁰	0.66 (0.58–0.75)	0.208/0.286	0.824/0.864	0.00001517	0.74 (0.65–0.85)	0.472	0.111

^a3,980 case individuals and 3,546 control individuals.
^b1,271 case individuals and 1,270 control individuals.
^c1,406 case individuals and 1,734 control individuals.
^dFreq = frequency.
^eAdj. p = p value adjusted for imputation uncertainty.
^fp_{cond} = p value when conditioned on SNP.
^gGenotyped SNP.

After applying quality-control measures and adjusting for admixture within and across populations (Figure S1), we analyzed a total of 6,658 independent cases and 6,550 independent controls (Table 1 and Table S1).

To enrich the genotyped data set for nongenotyped SNPs, we imputed variants located between 11,033,737 bp and 11,618,107 bp (hg19) by using population-specific reference panels derived from the 1000 Genomes Project.²² SNP-association results for each population are shown or listed in Figures 1A–1C, Table 2, and Table S3). Considering the correlated variants that had r² > 0.6 with the peak associated SNP in each population, we observed 30 SNPs demonstrating association in the AS population (peak SNP rs1478901, p = 1.32 × 10⁻¹¹, OR = 0.64, 95% CI = 0.56–0.73) and 20 SNPs demonstrating association in the EA population (peak SNP rs998683, p = 5.22 × 10⁻¹⁴, OR = 0.76, 95% CI = 0.71–0.82) (Table 2). However, we observed only two associated SNPs (SNP rs2736345, p = 1.49 × 10⁻⁶, OR = 1.28, 95% CI = 1.15–1.42 and peak SNP rs922483, p = 1.15 × 10⁻⁶, OR = 1.31, 95% CI = 1.17–1.47) in the AA population because of the reduced LD in this region. Both variants identified in the AA population are within the subset of variants that were identified in the EA and AS samples as having r² > 0.6 relative to the peak SNPs, suggesting that the same causal variants are present in all three populations. Conditional association tests performed within each population validated rs998683, rs1478901, and rs922483 as the main SLE-associated variant for EA, AS, and AA, respectively (Table 2). Thus, rs922483 is likely to be the predominant SLE-associated variant.

We concluded that, of the common associated variants, rs922483 was the stronger functional candidate given that it is located near a putative transcript initiator (INR) site²³ (Figure S4) in a region predicted to bind RNA polymerase II (RNAPII), and its association with SLE remained significant when conditioned on rs2736345 (Table 2).

Resequencing Identified an Additional SLE-Associated triallelic SNP, rs1382568, Located within the B-Cell-Specific Promoter

To ensure identification of other uncommon and multi-allelic genetic variation in this region, we resequenced all 13 *BLK* exons and the 2.5 kb upstream promoter regions in 191 EA SLE individuals and 96 EA controls from the Autoimmune Biomarkers Collaborative Network (ABCoN) and the New York Cancer Project (NYCP), respectively. Although no additional nongenotyped or nonimputed biallelic variants were detected, an SLE-associated tri-allelic variant, rs1382568 (A/G/C), that is highly correlated with the variant (rs922483) identified in our transpopulation association study was identified (Table 3 and Table S3).

To confirm the association of these two variants, we used data obtained for these two SNPs from additional resequencing efforts on 960 subjects (710 affected individuals and 250 control individuals). Association analysis results

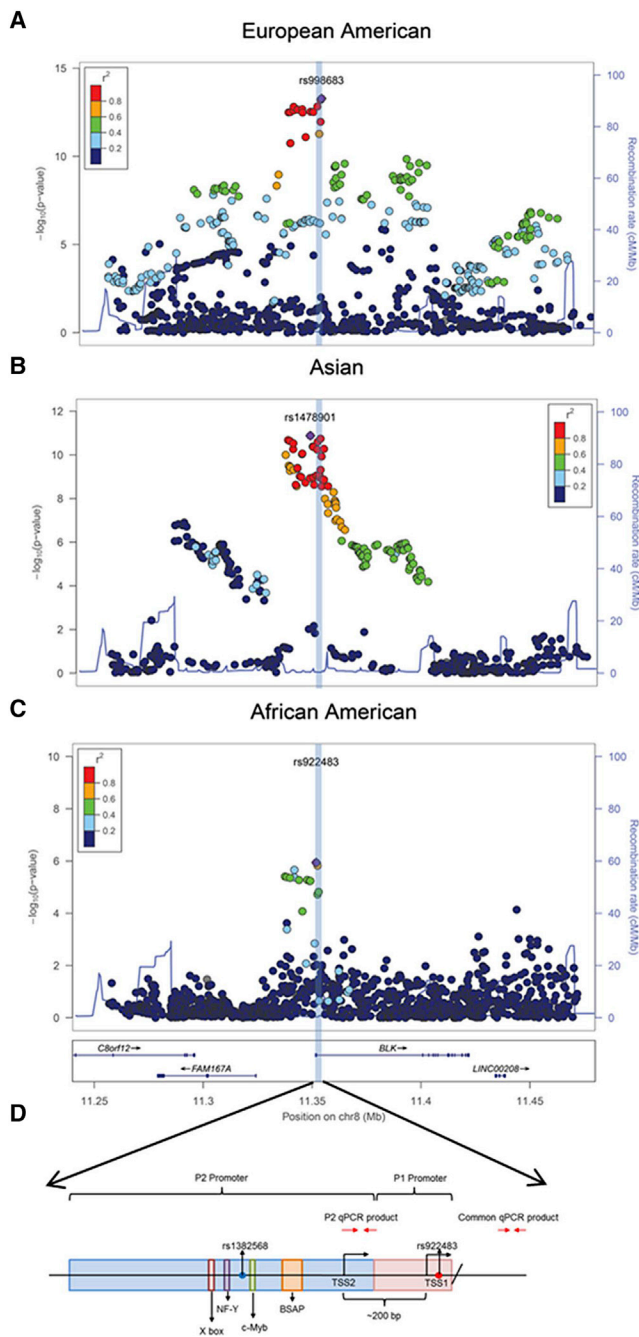


Figure 1. Genetic Association of SNPs in and around the *FAM167A/BLK* Gene Locus in SLE-Affected Individuals

SNPs in and around the *FAM167A/BLK* gene locus in individuals with SLE with (A) European ancestry, (B) Asian ancestry (C), and African American ancestry are shown. All SNPs with an $r^2 > 0.6$ (correlation with previously reported peak SNP rs13277113) are displayed. The solid blue line represents recombination rates across the region. The most significantly associated SNP in each population is colored purple, and the SNP number is indicated. (D) A schematic with key features of the *BLK* proximal promoter is shown. Probe P2 and P1 represent the 100 bp probe flanking the candidate variants, rs1382568 and rs922483. P2 and common qPCR products represent the products from luciferase gene-specific reverse transcription using product-specific primers (represented by red arrows).

from these data demonstrate that both C and A alleles at rs1382568 individually contributed to the increased SLE risk when compared to the G allele (OR 1.70, $p = 4 \times 10^{-3}$; and OR 2.53, $p = 6.66 \times 10^{-4}$, respectively). Association analysis using the combined C/A risk allele at rs1382568 had an OR = 1.90 and $p = 6.66 \times 10^{-4}$. This tri-allelic variant is located within the alternative *BLK* promoter (P2)¹ (Figure 1D). These data, and previously published results demonstrating that endogenous *BLK* expression varies with B cell developmental stage,²⁴ led us to hypothesize that the SLE-associated P2 variant might contribute to disease risk by promoting functional effects in B cells at discrete stages of development. We functionally characterized both variants (rs1382568 and rs922483) in B cell lines that phenotypically represent different stages of B cell development.

Both Risk Alleles at rs922483 (T) and rs1382568 (C) Alter *BLK* Transcription

To investigate the impact of the SLE-associated promoter variants on *BLK* transcription, we cloned the *BLK* promoter region (−2256 to +55 bp) into a firefly luciferase reporter vector and performed site-directed mutagenesis to generate all six possible haplotype combinations of the rs1382568 (P2) and rs922483 (P1) variants. B lymphoma cell lines with distinct phenotypes representing various B cell developmental stages were transfected with the reporter constructs. RS4;11 and Nalm-6 cell lines are representative of early stages of B cell development (pre- and pro-B cells), whereas Ramos and Daudi lines represent more mature B cells. The allelic effects of both *BLK* promoter variants were also tested in Jurkat cells, which are phenotypically similar to mature T cells. Endogenous *BLK* protein expression in each of these lines was confirmed to be as previously described (Figure S2).^{1,12}

Because of the small numbers of SLE-affected individuals carrying both risk alleles P1 and P2, we utilized in vitro assays to better isolate the influence of the P1 variant on *BLK* promoter activity. We assessed the average of luciferase activities of all P1-risk-allele- (T)-containing vectors, including T(P1)-C(P2), T(P1)-A(P2), and T(P1)-G(P2), as well as all P1-nonrisk-allele-containing vectors. The risk allele (T) at the P1 variant resulted in reductions of normalized luciferase expression in mature B (35%, Daudi) and mature T (32%, Jurkat) cell lines regardless of the allele at the P2 variant (p value < 0.05) (Figure 2A). The effect of the risk allele at the P1 variant on *BLK*-promoter-driven transcription was less pronounced in RS4;11 (pro-B) and Nalm-6 (pre-B) cells. Nuclear-factor binding assays demonstrated that the allelic variants at the P1 site altered nuclear-factor recruitment to the P1 promoter (Figure S3A), most likely as a result of changes in either the recruitment or the affinity of binding of the complement of nuclear factors and RNA-polymerase-complex components to this region of the *BLK* promoter, as suggested by a super-shift binding assay (Figure S3B). However, the complex nature of nuclear-factor binding to this site hampered our ability

Table 3. Analysis Results of Variations Identified in Resequencing within the Noncoding Region of BLK

Genomic Location	Nucleotide Change	dbSNP	Associated Allele	Case, Control Ratio Counts (191, 96)	OR (95% CI)	p Value	r ² (with rs13277113)
11,349,576	C>A	rs2251056	A	321:61, 150:42	1.47 (0.95–2.28)	0.0819	0.066
11,350,515	C>T	rs76610494	T	8:374, 1:191	4.10 (0.51–32.92)	0.1522	0.006
11,350,678	T>C	rs2736344	C	329:53, 149:43	1.79 (1.15–2.80)	0.0098	0.072
11,350,721	C>G	rs9694294	G	323:59, 149:43	1.58 (1.02–2.45)	0.0398	0.078
11,350,899	T>C	rs1382567	T	199:183, 93:99	1.16 (0.82–1.64)	0.4083	0.335
11,351,019	G>A	rs4840568	A	125:257, 41:151	1.79 (1.29–2.69)	0.0046	0.852
11,351,220	A>C	rs1382568 ^a	C	121:261, 38:154	1.88 (1.24–2.85)	0.0027	0.889
11,351,220	A>G	rs1382568 ^a	A	200:182, 94:98	1.15 (0.81–1.62)	0.4423	0.331
11,351,912	C>T	rs922483	T	137:245, 43:149	1.94 (1.30–2.89)	0.001	0.741
11,352,056	A>G	rs2250788	G	320:62, 148:44	1.53 (0.99–2.37)	0.0514	0.059
11,366,671	C>T	rs115856097	C	274:108, 128:64	1.27 (0.87–1.84)	0.2117	0.037
11,367,037	G>C	rs4629826	G	358:24, 176:16	1.36 (0.70–2.62)	0.3626	0.018
11,367,042	C>T	rs76154097	T	55:327, 23:169	1.24 (0.73–2.08)	0.4249	0.038
11,367,092	T>C	rs2409782	C	91:291, 45:147	1.02 (0.68–1.54)	0.9186	0.001
11,415,571	A>G	rs4841557	A	158:222, 63:129	1.46 (1.01–2.10)	0.042	0.187
11,415,596	C>T	rs4841558	C	157:223, 63:129	1.44 (1.00–2.08)	0.0484	0.181
11,418,385	G>A	rs1042695	A	145:237, 59:133	1.38 (0.95–2.00)	0.0878	0.179
11,418,765	G>A	rs368588162	A	335:45, 157:35	1.66 (1.03–2.68)	0.0375	0.019
11,418,766	G>A	rs62490888	G	340:42, 157:35	1.81 (1.11–2.94)	0.0164	0.015
11,418,772	C>T	rs4841561	T	146:234, 59:133	1.41 (0.97–2.04)	0.07	0.177
11,421,383	C>T	rs10097015	T	160:222, 68:122	1.29 (0.90–1.85)	0.1608	0.161
11,421,793	C>T	rs1042689	T	144:238, 62:130	1.27 (0.88–1.83)	0.2028	0.161
11,422,045	G>A	rs1042701	G	213:169, 98:94	1.21 (0.86–1.71)	0.2845	0.2
11,422,130	T>C	rs7843987	T	212:170, 100:92	1.15 (0.81–1.62)	0.4385	0.219

The two SNPs shown in italics (rs1382568^a and rs922483) showed association with the loci identified from the GWAS (rs13277113, OR 1.39, $p = 1 \times 10^{-10}$, ref. 1) with $r^2 > 0.5$; ^ars1382568 is a tri-allelic SNP (A/C/G); the C allele was identified as a risk allele.

to define the exact molecular interaction affected by the nucleotide variation at this site.

In order to explore the effect of P2, we compared the averaged luciferase activities from all vectors containing the P2 risk allele (C) with other vectors containing the P2 risk allele (C). We observed the most significant allelic effect at the P2 site in early B cells (RS4;11 and Nalm-6), where risk alleles A or C at the P2 site reduced luciferase expression in comparison to the nonrisk allele (G) at this variant (p value < 0.05) (Figure 2B). However, the impact of the P2 variant became insignificant when this variant was transfected into more mature B cell lines. Nuclear-factor binding assays showed that the risk allele (C) reduced the binding affinity of multiple nuclear-factor complexes to the probe containing the P2 allelic variant (Figure S3C).

The results from these assays demonstrate that the lupus-associated risk alleles at both the P1 site (rs922483) and the P2 site (rs1382568) reduce the transcriptional

activity of the *BLK* promoter in vitro. However, the effect of the risk allele at the P1 site most significantly affects *BLK* transcription in more mature B cells, whereas the effects of the risk alleles at the P2 site most significantly affect *BLK* transcription in more immature B cells.

P1 Variant Modulates Promoter Usage

Genes such as *BLK* that have multiple TSSs (transcription start sites) represent a class of genes in which changes in gene expression might be attributed to polymorphisms at multiple promoter sites. Selection of promoter use can vary on the basis of the organization of specific nuclear-factor binding sites and/or the epigenetic conformation of the genomic DNA in the promoters surrounding these TSSs. In addition, the organization of the promoters and/or TSSs and the dynamics of the transcription initiation and elongation steps of the RNA polymerase from each promoter influence which transcripts predominate within a cell. Differential promoter and TSS usage has been

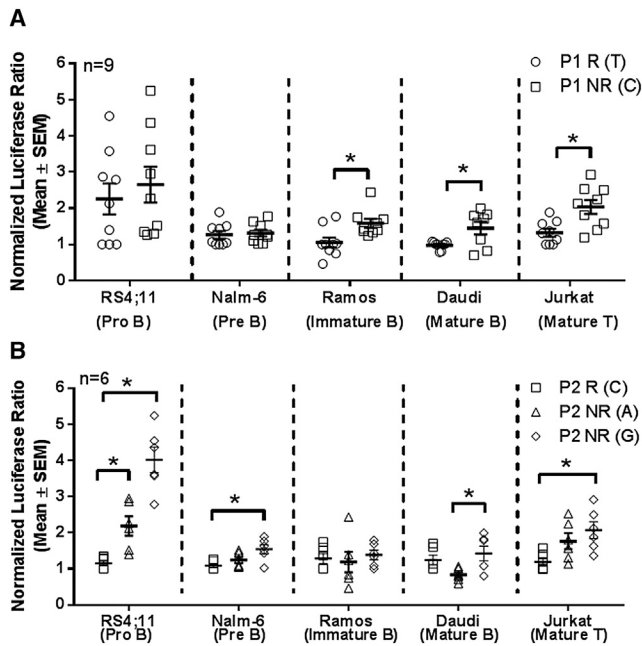


Figure 2. Both P1 and P2 Variants Affect *BLK*-Promoter-Driven Transcriptional Activity

Mean and standard error of measure (SEM) are displayed in the center, and probability density functions are represented by the sides. The effect of P1 variant with either risk or nonrisk P2 haplotype on overall luciferase expression (A) and the transcriptional activity in cell lines transfected with reporter vectors carrying one of the three SLE-associated P2 variants with a nonrisk P1 (B) is shown. Nine transfections of each vector carrying the P1 allele being compared were performed in each model cell line ($n = 9$), and triplicates were assessed for luciferase activity to give normalized means for each transfection. P1 risk [R(T)] and nonrisk [NR(C)] variants are compared (mean \pm SEM). P2 variants of each allele (G, A, or C) were assessed in six experiments. Normalized luciferase ratio = (normalized luciferase activity of the haplotype)/(normalized luciferase activity of the T allele at P1 – the luciferase activity of the C allele at P2). The normalized luciferase activity for the haplotype = luciferase activity of *BLK*:pGL4/luciferase activity of *TK*:pRL. * $p < 0.05$ in a paired t test. Means \pm SEM are shown.

elegantly demonstrated in the regulation of expression of the human *c-myc* gene (MIM 190080).²⁵ In this case, a preferred downstream promoter normally impedes (attenuates) the transcription initiated from the upstream promoter. However, inhibition of binding of the transcriptional machinery (e.g., RNA polymerase complex) prevents transcription initiation at the downstream *c-myc* promoter, removing attenuation of the upstream promoter and resulting in the upstream promoter's becoming the preferred promoter.

To determine whether such a mechanism controls *BLK* promoter selection and whether lupus-disease-associated variants in the *BLK* promoter P1 site can alter this mechanism, we used a transcript-specific luciferase reporter RT-qPCR assay to quantitate the percentage of the total *BLK* reporter transcripts in the B cell panel representing various cell stages of development. The usage of P2 and TSS2 was significantly higher in a majority of the B cell lines than

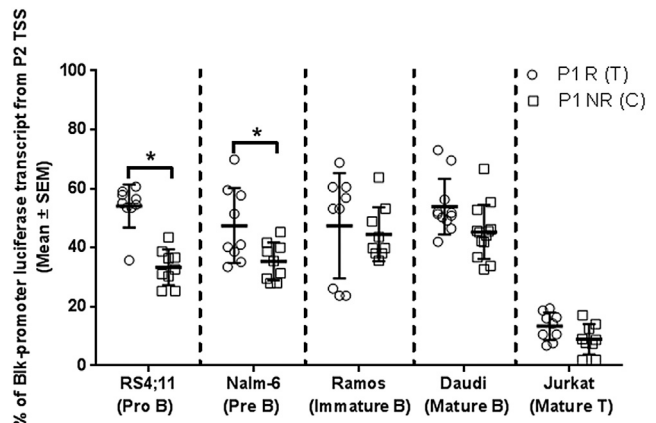


Figure 3. P1 Variant Altered Promoter Usage in RS4;11 and Nalm-6 Cell Lines

Percentages of the total *BLK* promoter-luciferase derived transcripts initiated from the P2 were determined using gene-specific RT-qPCR 16 hr post-transfection. * p -value < 0.05 using paired t test. Mean \pm SEM are shown.

in the mature T (Jurkat) cell line ($p < 0.05$) (Figure 3). This finding is consistent with the observations made by Lin et al.,¹ who showed that the P2 promoter is primarily used by B cells. The risk allele (T) at the P1 variant reduced the P1 and TSS1 contribution to the overall *BLK*-luciferase-reporter transcript levels in all cells, independent of the P2 variant ($p < 0.05$) (Figure 2A). However, the usage of P2 and TSS2 was increased by 21% and 12% in the immature B cell lines (RS4;11 and Nalm-6, respectively) in the presence of a risk-allele (T) at the P1 variant (Figure 3). These results suggest that lupus-associated risk alleles at the P1 variant decrease the effective initiation of the *BLK*-reporter transcription from P1 and TSS1. This might lower the attenuation of P2 and TSS2 in early B cells, presumably by a mechanism similar to that observed with the *c-myc* gene. These findings provide mechanistic insights as to how multiple disease-associated variants in different promoters can have a collective effect modulating expression of disease-associated genes.

Discussion

Previous studies have linked multiple genetic variants at many loci with the development of autoimmune disease.^{26–31} Genetic variants found at the *FAM167A/BLK* locus are associated with multiple autoimmune diseases, including SLE, systemic sclerosis, rheumatoid arthritis, and Sjögren's syndrome.^{4–11} Although risk-conferring variants within *FAM167A/BLK* have been shown to be associated with altered mRNA expression of both *FAM167A* and *BLK*,⁷ the causal allele or alleles remain undefined as a result of the strong association between potential causal alleles and noncausal variants. Using the trans-population mapping and sequencing strategy, we focused on two common associated variants (rs922483 and rs1382568) located

within the two promoter regions of *BLK* for additional functional analysis.

Previously published data defined the two *BLK* promoters and TSSs as a ubiquitously expressed TSS1 and a B cell-specific TSS2 located approximately 400 bp upstream of the ubiquitous promoter.¹ Because both candidate lupus-associated variants were located in functionally important loci of the *BLK* promoter, we hypothesized that they might alter unique aspects of *BLK* transcriptional regulation. The rs922483 SNP resides in the ubiquitous P1 and TSS1 site within a putative initiator of transcription (INR) site.²³ The other lupus-associated variant, rs1382568, is located in an upstream P2 region that is highly enriched for several B-cell-specific nuclear-factor binding sites. Because rs922483 and rs1382568 have a high degree of association with SLE and are located in key regions of promoters, our results confirm the possibility that these variants contribute to disease development through regulation of *BLK* promoter activity.

We used reporter assays and nuclear-factor binding in B cell lines with phenotypes representative of different developmental stages to study the effects of variants on promoter activity. We cannot exclude the possibility that fresh B cells might behave differently; it is possible that primary lymphocytes might have different expression levels and activity levels of transcription factors and that these different levels might result in altered *BLK* transcription not observed in cell lines. However, our data directly compared the effects of promoter alleles within various types of developmental stages of B cell lines characterized to represent different stages of B cell development to give a clearer picture of *BLK* transcription in early B cell development. Isolating sufficient numbers of primary progenitor B cells with all haplotypes would be prohibitive. Despite its limitations, this reporter assay allowed assessment of both the allelic and haplotype effects of these variants on *BLK* promoter activity within multiple representative cell types.

Our results demonstrated that both variants play a role in regulating *BLK* transcription. Risk alleles at these sites most likely alter the affinity and/or specificity of binding of critical nuclear factors and their interactions with RNA polymerase II subunits. Our results indicate that the degree of impact of a particular risk allele on *BLK* transcription depends both upon cell type and, in the cases of B cells, upon the developmental stage. This is consistent with observations made by Simpfordorfer et al. in primary cells, where they reported that a risk allele at rs922483 (P1 variant) led to an overall reduction in *BLK* mRNA expression in T cells from human peripheral-blood and umbilical-cord B cells.¹² Although the transcription of *BLK* was affected by the variant in early B and T cells, BLK protein level was only significantly reduced in umbilical-cord B cells.¹²

On the basis of our results and the previously published information, we propose a molecular mechanistic model depicting the cell-type- and developmental-stage-specific effect of both lupus-associated variants on the overall

BLK promoter activity (Figure 4A). In this model, the P1 promoter is the predominant promoter. When the RNA polymerase II complex binds and initiates transcription from this promoter, the P2 B-cell-specific promoter is stochastically inhibited or P2-initiated transcription is prematurely terminated by RNA polymerase complexes bound to the P1 site. Because P1 is the only active promoter in non-B cells, a switch to a risk allele at the P1 site alone will lead to a significant reduction in overall *BLK* promoter activity.

Alternatively, in B cells, production of *BLK* transcripts would be derived from both the P1 and TSS1 site and the P2 and TSS2 site. In mature B cells, P1 and TSS1 remain the preferred promoters, possibly as a result of nuclear factors and chromatin conformation at that site, which favor high-affinity RNA polymerase II binding and transcription from P1 and TSS1. When a lupus risk allele is present at the P1 site, possibly lowering the affinity of nuclear factor binding or efficiency of RNA polymerase transcription initiation, the obstruction and attenuation of P2 initiated transcription would be diminished resulting in more P2 derived transcripts. In this environment, an additional risk allele at the P2 site would result in altered nuclear-factor binding and RNA-polymerase-complex binding and initiation of transcription from this promoter. From this model, one would predict that the most dramatic decrease in *BLK* expression in immature B cells would occur when risk alleles were found at both the P1 and P2 sites and that this would result in increased risk for developing lupus.

Information accumulated from this and other studies is beginning to shape our overall understanding of how variations in *BLK* transcription expression and BLK protein levels contribute to development and/or progression of lupus.^{2,3,12,32} The emerging picture suggests that the variation of *BLK* expression is likely to result in varying functional consequences at different stages of B cell development and in different cell types (Figure 4B). Reduction in *BLK* expression by risk haplotypes could directly affect B lymphocyte development and/or impair functional responses in B cells early in development. Indeed, several previously published results indicate that the knockout of one allele of *Blk* leads to increased splenic marginal zone and peritoneal B1 B cells in older mice,³ suggesting a regulatory role for BLK. Because BLK is capable of interacting with both pre-B cell receptors and mature B cell receptors, it could play a critical role in regulating B cell selection and immune responses. Recently, BLK has also been shown to enhance BANK1 (MIM 610292) and PLC γ 1 (MIM 172420) interactions upon BCR activation to modulate B cell responses.³³ Other lupus-associated risk alleles in coding SNPs of BLK have been shown to result in reduced BLK protein stability.¹⁰ In addition, *BLK* deficiency can impair early T cell development as well as the development of IL-17-producing $\gamma\delta$ T cells.² Although there has been a suggestion that BLK is also an important signal transduction molecule in plasmacytoid dendritic

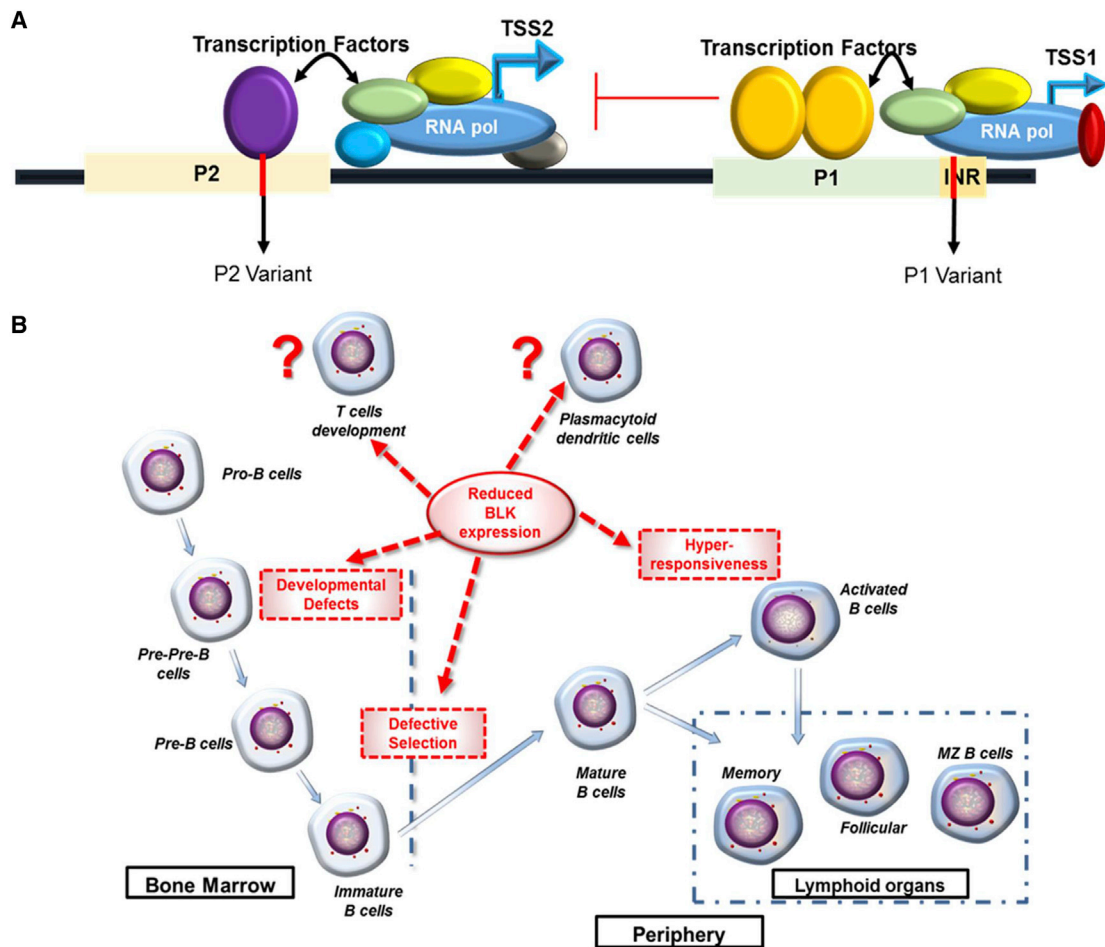


Figure 4. Proposed Molecular Model of Transcriptional Control of *BLK* Expression

A proposed model depicting how the lupus-associated risk alleles at the P1 and P2 sites alter *BLK* transcription (A) and a proposed model representing ways in which *BLK* might affect B cell development and control of autoimmune responses (B) are shown.

cells (pDCs), further investigations are necessary for the evaluation of the association between *BLK*-related alterations in pDCs and autoimmune diseases.³⁴ Our results indicate that *BLK* risk alleles alter *BLK* promoter activity and might thus contribute to autoreactive or regulatory cell responses.

Supplemental Data

Supplemental data include four figures, three tables, and Supplemental Acknowledgments and can be found with this article online at <http://www.cell.com/ajhg/>.

Consortia

Members of the BIOLUPUS Network are as follows: Sandra D'Alfonso (Italy), Rafaella Scorza (Italy), Gian Domenico Sebastiani (Italy), Mario Galeazzi (Italy), Johan Frostegård (Sweden), Bernard Lawerys and Fredric Houssieu (Belgium), Peter Junker and Helle Lastrup (Denmark), Marc Bijl and Cees Kallenberg (Holland), Eموke Endreffy (Hungary), Carlos Vasconcelos and Berta Martins da Silva (Portugal), Ana Suarez (Spain), Iñigo Rúa-Figueroa (Spain), and Norberto Ortego-Centeno (Spain) and Enrique de Ramon Garrido (Spain).

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Web Resources

The URLs for data presented herein are as follows:

Applied Biosystems, <http://www.appliedbiosystems.com>

gPLINK, <http://pngu.mgh.harvard.edu/~purcell/plink/>

Illumina, <http://www.illumina.com/>

Lupus Family Registry and Repository (LFRR), <http://omrf.ouhsc.edu/lupus>

Online Mendelian Inheritance in Man (OMIM), <http://www.omim.org>

WHAP, <http://pngu.mgh.harvard.edu/~purcell/whap/>

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Supplemental Data

Two Functional Lupus-Associated *BLK* Promoter Variants Control Cell-Type- and Developmental-Stage-Specific Transcription

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Figure S1.

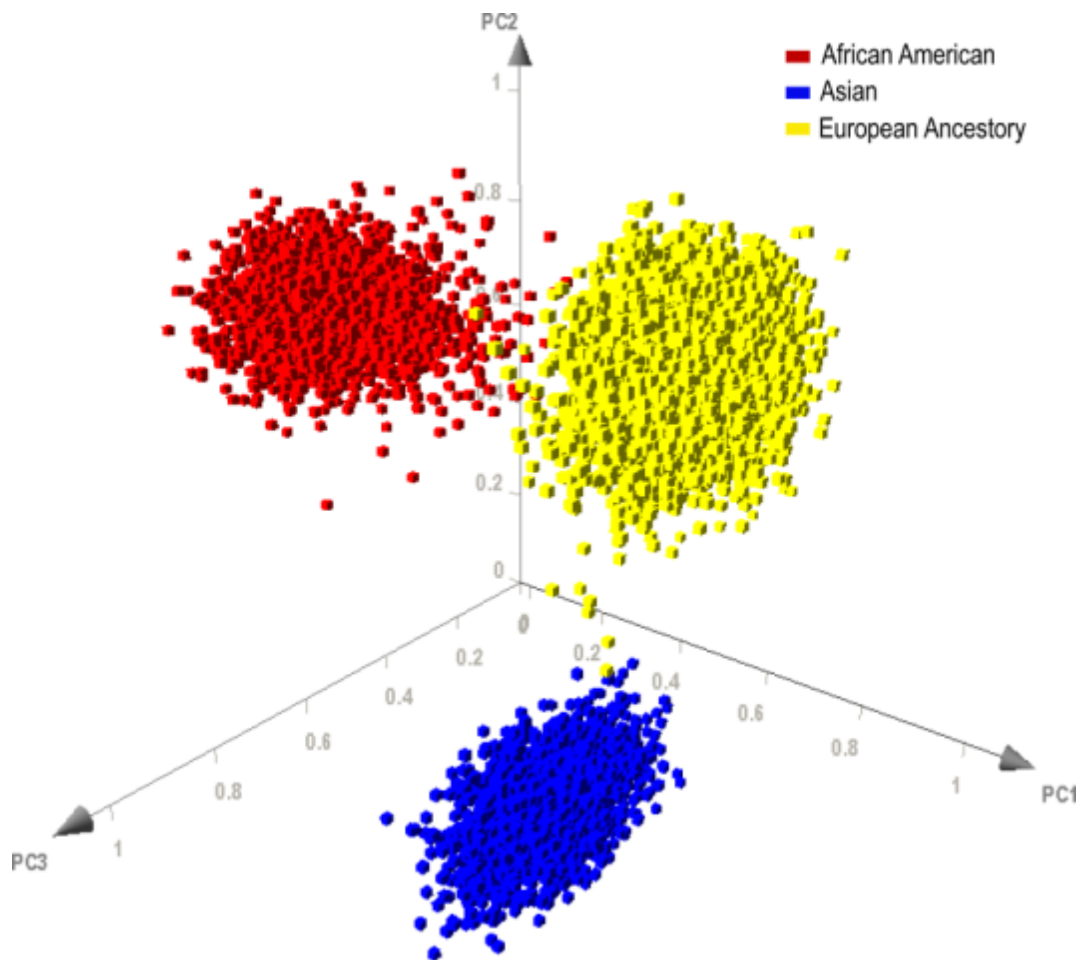


Figure S1. Population substructure of study participants used in genetic association analysis.

The population segregation based on the first three principle components calculated for this dataset are shown. Red = African-American; Blue = Asian; Yellow = European-American

Figure S2.

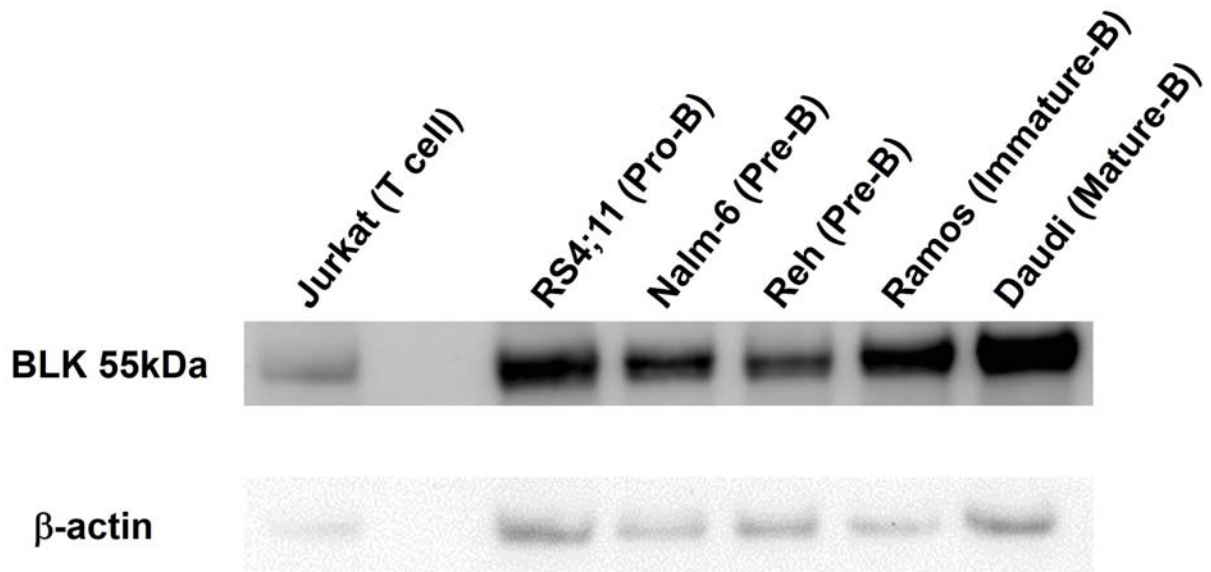


Figure S2: BLK protein expression in the cell lines used in reporter assays. The 55 kDa BLK band was detected in all cell lines used in the study using the anti-human BLK antibody (clone 7A12, Novus Biologicals). The antibody has been shown to specifically bind to BLK native and recombinant proteins.

Figure S3.

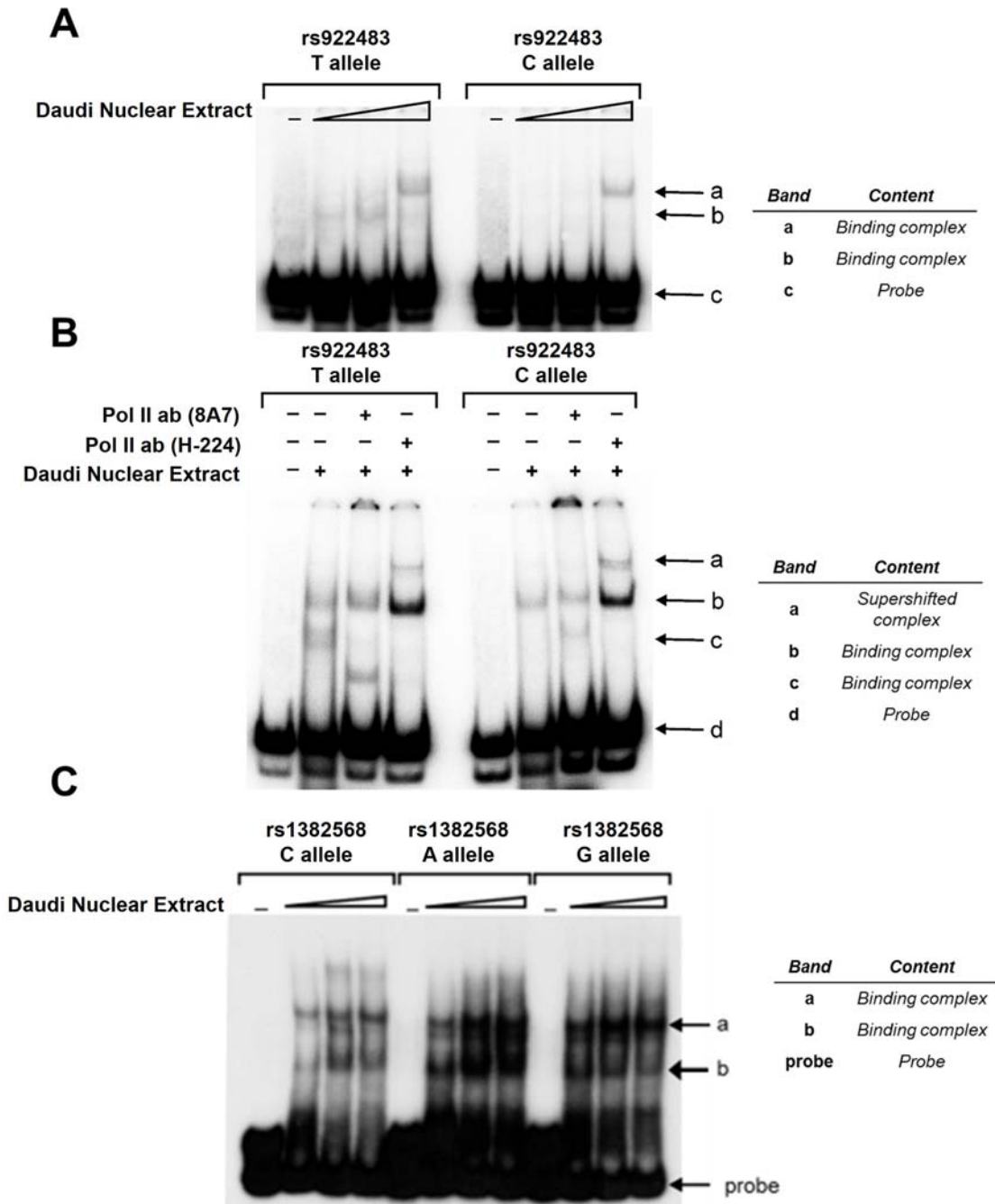


Figure S3: Risk alleles at rs922483 and rs1382568 influence binding of specific transcriptional factors (TFs) complexes. Electrophoretic mobility shift assay confirming differences in nuclear

factor binding of Probe P1 (100 bp) flanking the P1 SNP variant (rs922483) is shown in panel (A). “-” represents labeled DNA probe only control. Subsequent lanes represent the binding of increasing amounts of nuclear factors extracted from Daudi nuclei. Supershift assay of P1 probe is shown in panel (B). Two ug of anti-RNAPII antibodies were co-incubated with nuclear factor prior to the addition of labeled probes. The electrophoretic mobility shift of probe P2 flanking P2 SNP (rs1382568) is shown in panel (C). Increasing amount of nuclear factors extracted from Daudi nuclei were used to titrate the binding affinity. All results shown are of a representative experiment.

Figure S4.

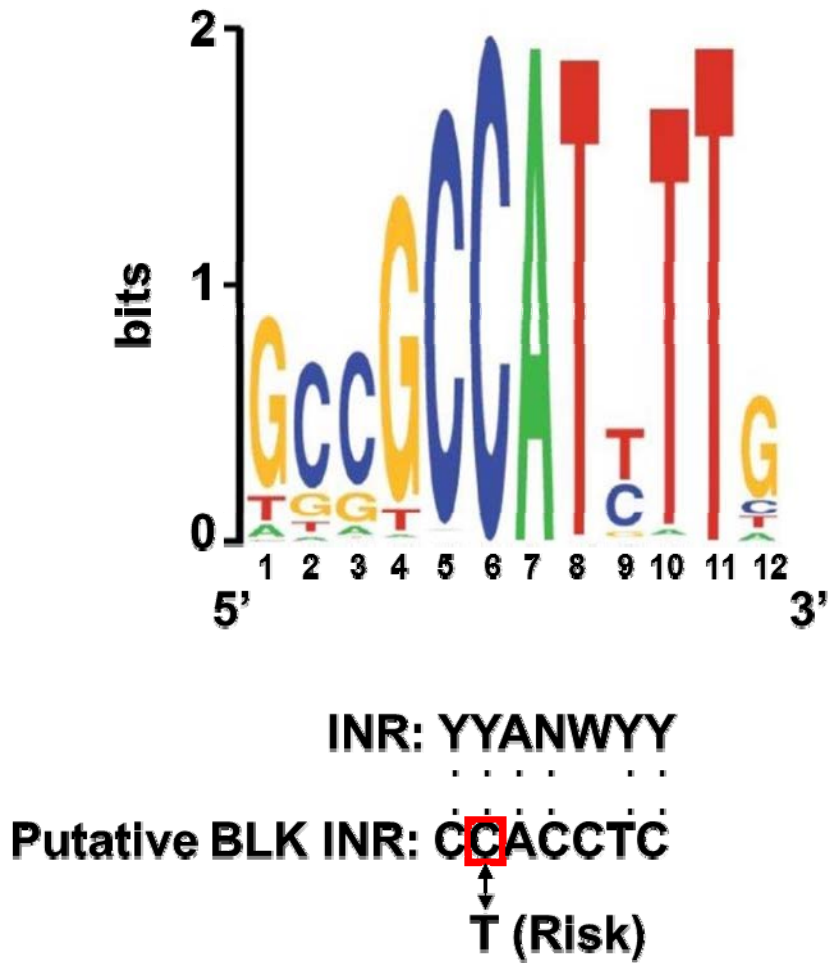


Figure S4. The variant rs922483 resides within a human Initiation (INR) motif. A weblogo diagram (<http://weblogo.berkeley.edu/logo.cgi>) based on alignments of INR motifs done by Yarden et.al.¹ and the corresponding *BLK* putative INR sequence with the location of variant rs922483 is shown (red box). *BLK* sequence location is shown in **Figure 1D**.

Table S1: Source and demographics of samples used in the SNP genotyping association analyses presented.

Contributor/Group	<i>European-Ancestry</i>		<i>Asian</i>		<i>African-American</i>		<i>Total</i>	
	Case	Control	Case	Control	Case	Control	Case	Control
Stevens: United States (Washington)	19	0	10	0	2	0	31	0
Tsao: United States (California), China	30	0	534	483	70	6	634	489
Jacob: United States (California)	108	0	82	0	0	0	190	0
^a PROFILE: Brown: United States (Alabama)	731	1083	0	0	575	637	1306	1720
CLU & SLEIGH: Gilkeson, Kamen: United States (South Carolina)	73	175	0	0	112	65	185	240
^b LFRR/ORDRCC: Harley, James: United States (Oklahoma)	667	553	0	0	428	209	1095	762
^c LuMNS: Gaffney, Sivils : United States (Minnesota)	445	0	0	0	0	0	445	0
ORDRCC: James: United States (Oklahoma)	122	137	2	46	52	55	176	238
Merrill: United States (Oklahoma)	78	2	0	0	28	5	106	7
Vyse: England	248	528	0	0	0	0	248	528
Scofield: United States (Oklahoma)	30	66	0	0	0	0	30	66
^d TRIDOM: Niewold: United States (Illinois, Minnesota)	79	0	0	0	139	0	218	0
Criswell: United States (California)	79	0	0	0	0	0	79	0
Boackle: United States (Colorado)	14	0	2	0	0	0	16	0
^e Gregersen: United States (New York)	0	0	0	0	0	299	0	299
Freedman: United States (North Carolina)	0	0	0	0	0	458	0	458
Bae: South Korea	0	0	642	741	0	0	642	741
BIOLUPUS: Sweden	78	0	0	0	0	0	78	0
BIOLUPUS: Belgium	39	53	0	0	0	0	39	53
Pons-Estel: Argentina (Europeans)	44	0	0	0	0	0	44	0
BIOLUPUS: Spain	464	419	0	0	0	0	464	419
BIOLUPUS: Portugal	186	174	0	0	0	0	186	174
BIOLUPUS: Hungary	25	47	0	0	0	0	25	47
BIOLUPUS: Italy	339	309	0	0	0	0	339	309
BIOLUPUS: Holland	65	0	0	0	0	0	65	0
BIOLUPUS: Danish	17	0	0	0	0	0	17	0
Totals:	3980	3546	1272	1270	1406	1734		

^aPROFILE (US multi-ethnic, multi-regional cohort of lupus patients), ^bLFRR (Lupus Family Registry and Repository), ^cLuMNAS (Univ. of Minnesota Lupus Association Cohort), ^dTRIDOM (University of Chicago Translational Research in the Department of Medicine registry), ^eFeinstein Institute of Medical Research (laboratory of Dr. Gregersen).

Table S2: Demographics of the ABCoN/NYCP re-sequencing samples

Ancestry	Case			Control		
	<i>Male</i>	<i>Female</i>	Total	<i>Male</i>	<i>Female</i>	Total
European-Ancestry	0	191	191	0	96	96

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Supplemental References

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