# Large-Scale Exome-wide Association Analysis Identifies Loci for White Blood Cell Traits and Pleiotropy with Immune-Mediated Diseases

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White blood cells play diverse roles in innate and adaptive immunity. Genetic association analyses of phenotypic variation in circulating white blood cell (WBC) counts from large samples of otherwise healthy individuals can provide insights into genes and biologic pathways involved in production, differentiation, or clearance of particular WBC lineages (myeloid, lymphoid) and also potentially inform the genetic basis of autoimmune, allergic, and blood diseases. We performed an exome array-based meta-analysis of total WBC and sub-type counts (neutrophils, monocytes, lymphocytes, basophils, and eosinophils) in a multi-ancestry discovery and replication sample of ~157,622 individuals from 25 studies. We identified 16 common variants (8 of which were coding variants) associated with one or more WBC traits, the majority of which are pleiotropically associated with autoimmune diseases. Based on functional annotation, these loci included genes encoding surface markers of myeloid, lymphoid, or hematopoietic stem cell differentiation (*CD69, CD33, CD87*), transcription factors regulating lineage specification during hematopoiesis (*ASXL1, IRF8, IKZF1, JMJD1C, ETS2-PSMG1*), and molecules involved in neutrophil clearance/apoptosis (*C100rf54, LTA*), adhesion (*TNXB*), or centrosome and microtubule structure/function (*KIF9, TUBD1*). Together with recent reports of somatic *ASXL1* mutations among individuals with idiopathic cytopenias or clonal hematopoiesis of undetermined significance, the identification of a common regulatory 3' UTR variant of *ASXL1* suggests that both germline and somatic *ASXL1* mutations contribute to lower blood counts in otherwise asymptomatic individuals. These association results shed light on genetic mechanisms that regulate circulating WBC counts and suggest a prominent shared genetic architecture with inflammatory and autoimmune diseases.

## Introduction

White blood cells (WBCs) are major constituents of the blood and lymphatic system. They are classified into two

lineages: myeloid (neutrophils, basophils, eosinophils, and monocytes) and lymphoid (lymphocytes). Lineage commitment of hematopoietic stem cells involves precise transcriptional and epigenetic regulation, creating the

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specific bone marrow microenvironment to produce each distinct mature blood cell type.<sup>1</sup> Mature WBCs play diverse, choreographed roles in innate and adaptive immunity including detection, neutralization, and elimination of invading pathogens, response to tissue injury, and wound healing. In addition, WBCs are associated with the development of chronic inflammatory, allergic, and autoimmune diseases.<sup>2</sup> Therefore, total and differential WBC counts are important clinical measures of susceptibility to infection and used to monitor disease activity and tolerability to therapeutic regimens for oncologic and rheumatologic diseases.

Total and differential WBC counts are complex, polygenic traits with estimated heritability of 50%-60%.<sup>3</sup> Previous genome-wide association studies (GWASs) have characterized common and lower frequency variation contributing to WBC counts in European, African, and Asian ancestry populations (N.P., U.M.S., J.B.-J., and M.-H.C., unpublished data).<sup>3–12</sup> More than 30 distinct genetic loci have been discovered; in some instances, these genetic studies have provided important biologic insights into the development, maturation, or regulation of WBC types. Nonetheless, these studies have explained only a small proportion (<10%) of the estimated heritability of

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Population	Total WBC	Neutrophils	Monocytes	Lymphocytes	Basophils	Eosinophils
Discovery						
European ancestry	108,596	60,851	44,325	47,105	44,138	32,517
African ancestry	23,250	10,119	9,790	9,808	9,509	8,282
Hispanic American	5,536	4,825	3,452	3,450	3,453	3,450
East Asian	968	965	-	_	_	-
South Asian	464	463	-	_	_	-
Replication						
European ancestry	18,808	17,066	17,066	17,109	16,189	15,327
Total	157,622	94,289	74,633	77,472	73,289	59,576

WBC traits in European ancestry populations<sup>6</sup> and less than 25% in African ancestry (AA) populations (in AA, a substantial proportion of the variation in WBC counts is attributed to a single variant—rs2814778—in *DARC* [Duffy Antigen Receptor for Chemokines (MIM: 613665)]).<sup>3,13</sup> In an effort to augment the discoveries from GWASs and to identify additional functional loci contributing to variation in WBC counts, we performed exome array-based meta-analysis of total and differential counts in a multiancestry samples from 25 studies.

## Material and Methods

## **Study Subjects**

The Blood-Cell Consortium (BCX) is an international collaboration with the goal of identifying common and rare variants associated with blood cell traits through exome genotyping arrays (Table S1). The consortium, which is comprised of multi-ancestry cohorts including European ancestry (EA), African ancestry (AA), Hispanic ancestry (HA), East Asian ancestry (EAS), and South Asian ancestry (SA), is divided into three main working groups: red blood cell (RBC), platelet, and WBC. For exome-wide association analysis of WBC traits, the discovery and replication phases included a total of 157,622 participants from 25 cohorts (Tables 1, S2, and S3). The discovery sample consisted of up to 138,814 individuals from 21 studies. The replication sample included 18,808 independent individuals from 4 additional studies. The division of discovery and replication samples was dictated by timing; we collected all available studies for initial discovery and then identified others who could participate only at a later point in time and hence were used for replication. A summary of descriptive statistics for total WBC, neutrophils, monocytes, lymphocytes, basophils, and eosinophils is shown in Table S4. All participants provided informed consent and the study was approved by the Institutional Review Board of each participating study.

#### Genotyping and Quality Control

Each participating study used one of the following exome content genotyping arrays: Illumina ExomeChip v.1.0, Illumina ExomeChip v.1.1\_A, Illumina ExomeChip-12 v.1.1, Affymetrix Axiom Biobank Plus GSKBB1, or Illumina HumanOmniExpressExome Chip. Genotypes were called either using a combination of the Illumina GenomeStudio and zCall software or using the Exomechip joint calling plan developed by the Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Consortium<sup>14</sup> (Table S1). Standard quality-control criteria were applied by each study. Exclusion criteria included sample call rates of less than 98%, excess heterozygosity rates, Hardy-Weinberg equilibrium p values  $< 1 \times 10^{-6}$ , and sex mismatch. Additionally, ancestry was confirmed through principal components or multi-dimensional scaling analyses using linkage disequilibrium (LD) pruned markers ( $r^2 < 0.2$ ) with minor allele frequency greater than 1%. Scatterplots anchored using the 1000 Genomes Project populations were visually inspected and ancestry outliers were excluded. Insertion and deletion variants and variants mapping to the Y chromosome, the pseudo-autosomal region, or mitochondrial sequence were removed, leaving only those on the autosomal and X chromosomes. All remaining variants (including monomorphic variants) were aligned to the forward strand and alleles were checked to ensure that the correct reference allele was specified. After all quality-control procedures, each study generated an indexed variant call file (VCF) for subsequent analyses. The VCF files were checked for allele alignment with the checkVCF package.

We performed study-specific quality control on each trait association result using the EasyQC protocol.<sup>15</sup> Variant allele frequencies from each study were plotted against ethnicity-specific reference population allele frequencies to identify allele frequency deviations and the presence of flipped alleles. In order to assess proper trait transformation in each cohort, a scatterplot of the median standard error versus study-specific sample size was visually inspected for deviations.

#### **Statistical Analysis**

To assess the association between WBC-related traits and Exomechip variants, white blood cell and differential counts (total WBC, neutrophils, monocytes, lymphocytes, eosinophils, and basophils) were obtained from complete blood cell count. Each of the WBC-related traits was log<sub>10</sub> transformed to normalize the distribution of the traits. In each participating study, residuals for each WBC trait were calculated from linear regression models adjusted for age, age-squared, sex, study center (where applicable), and principal components. Residuals from this model were then transformed using the rank-based inverse normal transformation to control type I error.<sup>16</sup> Autosomal and X chromosome variants were then tested for association with each WBC trait using either Rvtests or RAREMETALWORKER software packages. Both packages generate single variant association score summary statistics, variance-covariance matrices containing LD relationships between variants within a 1 MB window, and variant-specific parameters including minor allele frequency, chromosome position, strand, genotype call rate, and Hardy-Weinberg equilibrium p values.

## **Discovery Association Meta-analysis**

For each WBC trait, we performed three distinct discovery metaanalyses: in EA only, AA only, and combined across all five ancestry groups. Ancestry-stratified (EA and AA) and combined all ancestry (EA, AA, HA, EAS, and SA) meta-analyses of single variant association results were carried out using the Cochran-Mantel-Haenszel approach implemented in RareMETALS.<sup>17</sup> We included variants in the meta-analysis if the genotype call rate was  $\geq$  95%. For palindromic variants (i.e., A/T and C/G variants), we compared allele frequencies taken across the entire consortium in order to detect flipped alleles. We kept variants with an allele frequency difference <0.3 or <0.6 for ancestry-specific (EA, AA) or combined all ancestry analyses, respectively.<sup>15</sup> Using singlevariant score statistics and variance-covariance matrices of LD estimates, we performed two types of gene-based tests across the contributing studies: (1) a burden test that assumes all qualifying rare variants in a gene are associated with a trait with the same direction of effect (variable threshold test), and (2) the sequence kernel association test (SKAT) that accounts for rare variants in a gene having opposing direction of effects.<sup>17</sup> For all gene-based tests performed, we considered single-nucleotide variants (SNVs) with an allele frequency of  $\leq 1\%$  and annotated as missense, nonsense, and splice site variants; the latter two categories include loss-of-function variants. Similar to the single-variant analyses, results were generated for EA, AA, and for the combined all ancestry samples. For the discovery single variant and gene-based association analyses, the statistical significance threshold was set as p value  $< 2 \times 10^{-7}$  and  $< 3 \times 10^{-6}$ , respectively.

### **Conditional Analysis**

To identify multiple independent associations within a region, using the RareMETALS software we performed stepwise conditional analyses adjusting for the most significant single variant in a 1 MB window, across the entire Exomechip array. This step was repeated until there was no new association signals identified in each region, defined as a p value  $< 2 \times 10^{-7}$ . Further, to assess whether SNVs identified by the present study were independent of any previously reported WBC-associated variants, we conditioned our regression models on known GWAS sentinel variants or their proxies (LD  $r^2 \ge 0.80$ ). For regions of the genome where there is extended LD structure spanning more than 1 MB, we performed a stepwise conditional analysis in GCTA software<sup>18</sup> conditioning on the most significant variant in the region first (or the GWAS sentinel variant or LD proxy).

#### **Replication Meta-analysis**

We sought replication of association results using four independent European ancestry cohorts (Tables 1 and S3). The singlevariant association results from each replication cohort were combined using the Cochran-Mantel-Haenszel method in RareMETALS. Contributing replication cohorts adhered to the quality control and association analysis procedures described previously for the discovery analysis. Replication of association findings were considered significant if the variants demonstrated the same direction of effect as the discovery association metaanalyses with a replication p value < 0.05. A meta-analysis of discovery and replication results was performed using an inverse-variance weighting method as implemented in METAL.<sup>19</sup> We also performed replication of gene-based associations in independent ~2,900 EA samples.

## Phenome-wide Association Study Analysis

In 29,722 EA samples from the BioVU study,<sup>20</sup> we performed phenome-wide association study (PheWAS) analysis<sup>21</sup> to assess the association between our WBC-related loci and 1,502 International Classification of Disease, Ninth Revision (ICD-9) code curated clinical phenotypes.<sup>21</sup> Variants were included in the analysis if there were ten cases with at least one copy of the minor allele. Associations between SNVs and phenotypes were assessed using a logistic regression model adjusted for sex and five principal components. Empirical significance was estimated by permutation test. The permutation test was performed by assigning each vector of clinical phenotypes to a random subject 50,000 times, and then scanning all SNV-phenotype combinations with association tests. We then created a ranked distribution of the maximum test statistics over all SNV-phenotype combinations in each of the 50,000 permutations. The 95th percentile of the distribution of maximum test statistics across the 1,502 clinical phenotypes and 95 SNVs equates to a threshold that controls the family-wise error rate at 0.05. This threshold accounts for multiple testing across SNVs and phenotypes. Our observed test statistics greater than this 95<sup>th</sup> percentile were considered statistically significant.

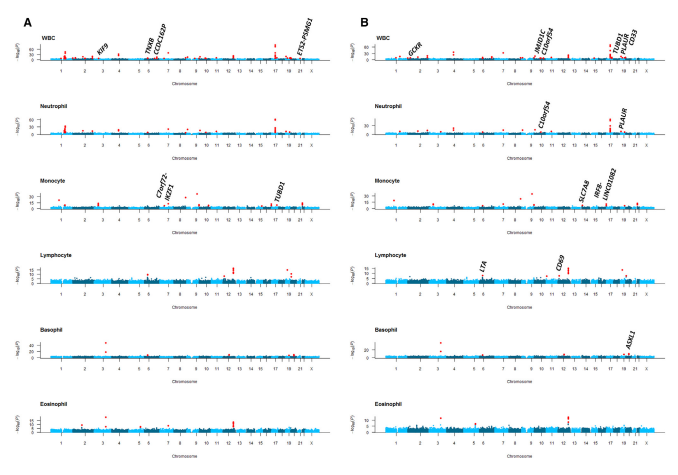
To further assess pleiotropy between WBC-associated variants and inflammatory diseases, we performed lookups in published GWASs of various autoimmune diseases (celiac disease [MIM: 212750], inflammatory bowel disease [IBD; MIM: 266600], multiple sclerosis [MS; MIM: 126200], primary biliary cirrhosis [PBC; MIM: 109720], psoriasis [MIM: 177900], rheumatoid arthritis [RA; MIM: 180300], systemic lupus erythematosus [SLE; MIM: 152700], type 1 diabetes mellitus [T1D; MIM: 222100]) and coronary artery disease (MIM: 608901).<sup>22–30</sup> We supplemented the full GWAS summary statistics lookups with the GRASP database<sup>31</sup> to include other immunologically relevant clinical phenotypes and quantitative traits. Similarly, to assess whether the WBC variants were associated with other blood cell traits, we obtained effect sizes and p values for these variants from RBC- and platelet-related traits exome array analyses within the BCX consortium.<sup>32,33</sup>

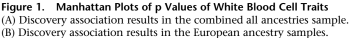
### **Functional Annotation of Variants**

To assess the functional consequences of coding and non-coding variants associated with WBC traits, we utilized a variety of existing variant annotation resources. Using a curated collection of more than 100 separate expression quantitative trait loci (eQTL) datasets, we queried whether our list of WBC-trait loci were also associated with transcript expression in blood-cell-specific eQTL datasets. A general overview of a subset of >50 eQTL studies has been published,<sup>34</sup> with specific citations for the blood-cell-specific eQTL datasets shown in Table S5. Additional in silico functional annotations were performed with ANNOVAR.<sup>35</sup> The deleteriousness of each variant was estimated with the Combined Annotation-Dependent Depletion (CADD) score where each variant is assigned a scaled C score; a score of greater than 10 is suggested to indicate deleteriousness.<sup>36</sup>

### Results

We conducted an exome-wide association analyses of total WBC and differential counts (neutrophils, monocytes,





The combined all ancestry sample include European, African, Hispanic, East Asian, and South Asian ancestries. Genetic variants that passed the array-wide significance threshold (p value  $< 2.0 \times 10^{-7}$ ) are highlighted in red. Discovery genetic loci that replicated in independent samples are shown.

lymphocytes, basophils, and eosinophils) in a discovery sample of ~138,814 individuals of European, African, Hispanic, East Asian, and South Asian ancestries across 21 cohorts (Tables 1 and S3). Quantile-quantile plots with genomic inflation factors and their respective Manhattan plots for each discovery meta-analysis are presented in Figures 1, S1, and S2. The discovery effort yielded 144 arraywide significant SNV associations (p value  $< 2.0 \times 10^{-7}$ ) (Table S6). After stepwise conditional analyses, we refined this list to 28 independent SNV associations with WBC counts that were not previously reported (Table S7). Of these 28 variant associations, 16 were replicated (p value < 0.05 and consistent direction of effect) in 17,897 independent EA individuals (Figure 1, Table 2). Fourteen of the replicated loci are located in genomic regions not previously associated with WBC traits. The remaining two loci (TNXB rs185819 and IRF8 rs11642873) represent secondary, independent signals located within a 1 MB window of a previously reported WBC locus. Of the 16 replicated loci, 10 were significantly associated with total WBC count, 2 with neutrophil count, 4 with monocyte count, 2 with lymphocyte count, and 1 with

basophil count. As described further below, several loci were associated with more than one WBC trait (Table 2); the WBC-subtype-specific association results for each of the 16 replicated variants are shown in Table S8. For each locus, the allele frequencies stratified by ancestry are shown in Table S9. The full summary Exomechip association results for all traits are publicly available online (see Web Resources).

## Total WBC

We found missense variants in a number of genes that were associated with total WBC. In *GCKR* (MIM: 600842), rs126032 (p.Leu446Pro [c.1337T>C]) was associated with lower total WBC in the EA meta-analysis (p value =  $8.13 \times 10^{-13}$ ). This variant was also nominally associated with lower neutrophil, lymphocyte, and basophil counts in EAs, consistent with its association with total WBC. The rs126032 variant was also associated with lower total WBC in AAs (p value = 0.014). In *KIF9* (MIM: 607910), rs2276853 (p.Arg573Trp [c.1717C>T]) was associated with increased total WBC in the multi-ancestry meta-analysis (p value =  $3.29 \times 10^{-9}$ ). The signal was largely driven

Trait				Alt/				AA	Discove	ry		Replica	ntion			Combin	ed M	eta-analysi	is	
(Population)	dbSNPID	Chr	Pos	-		Gene	Annotation	AA Substitution	N	Beta (SE)	р	N	EAF	Beta (SE)	р	N	EAF	Beta (SE)	р	p <sub>het</sub>
WBC (EA)	rs1260326	2	27,730,940	C/T	0.58	GCKR	missense, splice site	p.Leu446Pro	108,596	-0.030 (0.005)	$4.01 \times 10^{-10}$	17,897	0.6	-0.044 (0.012)	$\frac{2.66 \times 10^{-4}}{10^{-4}}$	126,493	0.58	-0.032 (0.004)	$8.13 \times 10^{-13}$	0.28
WBC (All)	rs2276853	3	47,282,303	A/G	0.58	KIF9	missense	p.Arg573Trp	132,764	0.023 (0.004)	$3.65 \times 10^{-8}$	17,897	0.6	0.025 (0.012)	$3.00 \times 10^{-2}$	150,661	0.58	0.023 (0.004)	$3.29 \times 10^{-9}$	0.86
WBC (All)	rs185819 <sup>a</sup>	6	32,050,067	C/T	0.51	TNXB	missense	p.His1161Arg	132,764	0.031 (0.005)	$4.02 \times 10^{-10}$	17,897	0.47	0.034 (0.015)	$2.24 \times 10^{-2}$	150,661	0.51	0.031 (0.005)	$2.85 \times 10^{-11}$	0.83
WBC (All)	rs9374080	6	109,616,420	C/T	0.43	CCDC162P	intronic regulatory		132,764	0.023 (0.004)	$^{4.01}_{10^{-8}} \times$	17,897	0.46	0.025 (0.011)	$2.55 \times 10^{-2}$	150,661	0.43	0.023 (0.004)	$3.15 \times 10^{-9}$	0.84
WBC (EA)	rs3747869	10	73,520,632	C/A	0.9	C10orf54 (DD1a)	missense	p.Asp187Glu	108,596	0.040 (0.007)	$4.26 \times 10^{-8}$	17,897	0.9	0.083 (0.018)	$6.40 \times 10^{-6}$	126,493	0.9	0.046 (0.007)	$1.42 \times 10^{-11}$	0.03
WBC (EA)	rs1935	10	64,927,823	G/C	0.49	JMJD1C	missense	p.Glu2353Asp	108,596	-0.026 (0.005)	$2.46 \times 10^{-8}$	17,897	0.46	-0.027 (0.012)	$2.06 \times 10^{-2}$	126,493	0.49	-0.026 (0.004)	$\frac{1.57}{10^{-9}}$ ×	0.93
WBC (EA)	rs1292053	17	57,963,537	G/A	0.45	TUBD1	missense	p.Met76Thr	108,596	-0.03 (0.004)	$1.28 \times 10^{-11}$	17,897	0.44	-0.027 (0.011)	$1.51 \times 10^{-2}$	126,493	0.45	-0.030 (0.004)	$6.55 \times 10^{-13}$	0.78
WBC (EA)	rs4760	19	44,153,100	G/A	0.16	CD87 (PLAUR)	missense	p.Leu272Pro	85,685	-0.043 (0.007)	$2.51 \times 10^{-10}$	17,897	0.15	-0.052 (0.015)	$7.13 \times 10^{-4}$	103,582	0.16	-0.044 (0.006)	$8.34 \times 10^{-13}$	0.6
WBC (EA)	rs3865444	19	51,727,962	A/C	0.31	CD33	upstream		86,936	-0.037 (0.005)	$3.51 \times 10^{-12}$	17,897	0.32	-0.033 (0.012)	$5.14 \times 10^{-3}$	104,833	0.31	-0.036 (0.005)	$6.81 \times 10^{-14}$	0.77
WBC (All)	rs2836878	21	40,465,534	A/G	0.26	ETS2- PSMG1	intergenic		132,764	-0.025 (0.005)	$8.36 \times 10^{-8}$	17,897	0.26	-0.026 (0.012)	$3.44 \times 10^{-2}$	150,661	0.26	-0.025 (0.004)	$8.41 \times 10^{-9}$	0.89
NEU (EA)	rs3747869	10	73,520,632	C/A	0.9	C10orf54 (DD1a)	missense	p.Asp187Glu	60,851	0.053 (0.010)	$2.11 \times 10^{-8}$	16,669	0.9	0.073 (0.019)	$1.17 \times 10^{-4}$	77,520	0.9	0.057 (0.009)	$1.65 \times 10^{-11}$	0.34
NEU (EA)	rs4760	19	44,153,100	G/A	0.16	CD87 (PLAUR)	missense	p.Leu272Pro	56,112	-0.047 (0.008)	$\frac{1.54}{10^{-8}}$ ×	16,669	0.15	-0.044 (0.016)	$5.55 \times 10^{-3}$	72,781	0.16	-0.046 (0.007)	$3.01 \times 10^{-10}$	0.87
MON (All)	rs4917014	7	50,305,863	G/T	0.28	C7orf72- IKZF1	intergenic		57,183	-0.038 (0.007)	$1.97 \times 10^{-8}$	16,669	0.32	-0.048 (0.012)	$\frac{8.92 \times 10^{-5}}{10^{-5}}$	73,852	0.29	-0.040 (0.006)	9.75 × 10 <sup>-12</sup>	0.48
MON (EA)	rs11625112	14	23,596,740	G/A	0.46	SLC7A8	intronic		44,325	-0.038 (0.007)	$3.82 \times 10^{-8}$	16,669	0.45	-0.031 (0.012)	$7.04 \times 10^{-3}$	60,994	0.46	-0.036 (0.006)	$1.03 \times 10^{-9}$	0.62
MON (EA)	rs11642873 <sup>a</sup>	16	85,991,705	C/A	0.2	IRF8- LINC01082	intergenic		44,325	0.057 (0.008)	$1.41 \times 10^{-11}$	16,669	0.2	0.113 (0.014)	$6.17 \times 10^{-15}$	60,994	0.2	0.072 (0.007)	$1.40 \times 10^{-22}$	0.00
MON (All)	rs1292053	17	57,963,537	G/A	0.45	TUBD1	missense	p.Met76Thr	57,183	-0.036 (0.006)	$2.55 \times 10^{-9}$	16,669	0.44	-0.040 (0.012)	$6.08 \times 10^{-4}$	73,852	0.45	-0.037 (0.005)	$6.53 \times 10^{-12}$	0.76
LYM (EA)	rs2229094	6	31,540,556	C/T	0.26	LTA	missense	p.Cys13Arg	47,105	0.046 (0.008)	$\frac{1.89}{10^{-8}}$ ×	16,711	0.25	0.078 (0.018)	$8.54 \times 10^{-6}$	63,816	0.26	0.051 (0.007)	$3.14 \times 10^{-12}$	0.09

			414				Discovery	ď	Replication	uo		<b>Combined Meta-analysis</b>	ysis	
(Population) dbSNPID Chr Pos	dignetid	Chr		F EAI	Ally Ref EAF Gene	Annotation Substitution	z	Beta (SE) p N	<b>–</b>	EAF Beta (SE) p	Эр	N EAF Beta (SE) p	E) p	Phet
YM (EA) r	s4763879	12	LYM (EA) rs4763879 12 9,910,164 A/G 0.36 CD69	3 0.3	6 CD69	intronic regulatory	47,105 -0.038 (0.007)	$(3 3.08 \times 1)^{-8}$	16,711 0	.36 -0.038 (0.012)	$1.36 \times 10^{-3}$	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	$1.59 \times 10^{-10}$	0.99
BAS (EA) rs	s2295764	20	rs2295764 20 31,025,163 G/A 0.36 ASXL1	4 0.3	6 ASXL1	3' UTR	44,138 -0.042 (0.007)	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	15,770 0	.36 -0.031 (0.012)	$9.78 \times 10^{-3}$	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	$1.46 \times 10^{-10}$	0.43

by the association in EAs (p value =  $1.39 \times 10^{-6}$ ) and was apparent for both neutrophil and lymphocyte counts in EAs and in multi-ancestry meta-analyses. In TNXB (MIM: 600985), rs185819 (p.His1161Arg [c.3428A>G]) was associated with increased total WBCs in the multi-ancestry meta-analysis (p value =  $2.85 \times 10^{-11}$ ). The association was consistently significant across EA and AA populations and for all WBC sub-types. The effect allele frequency was comparable between EAs and AAs but varied in the other ancestry groups. In C10orf54 (MIM: 615608), rs3747869 (p.Asp187Glu [c.561T>G]) was associated with increased total WBC in the EA meta-analysis (p value =  $1.42 \times$  $10^{-11}$ ). Although rs3747869 was also associated with neutrophil, monocyte, and eosinophil counts, the signal was not consistent across ancestry groups. The effect allele frequencies were markedly different between EA, AA, HA, SA, and EAS ancestry groups. In JMJD1C (MIM: 604503), rs1935 (p.Glu2353Asp [c.7059G>C]) was associated with lower total WBC (p value =  $1.57 \times 10^{-9}$ ) in the EA meta-analysis. Although the rs1935 variant was not consistently associated with total WBC across all the major ethnic groups, it was significant in the HAs (p value =  $5.58 \times 10^{-3}$ ). Significantly low neutrophil, lymphocyte, and eosinophil counts were also observed for rs1935. In TUBD1 (MIM: 607344), rs1292053 (p.Met76Thr [c.227T>C]) was associated with lower total WBC in the EA meta-analysis (p value =  $6.55 \times 10^{-13}$ ). This association was similar in EAs and AAs and for neutrophil, monocyte, and lymphocyte counts. Finally, in PLAUR (MIM: 173391) the rs4760 (p.Leu272Pro [c.815T>C]) variant was associated with lower total WBC (p value =  $8.34 \times 10^{-13}$ ) in the EA meta-analysis. The effect allele frequencies were highly discrepant across ancestries, perhaps explaining why the association was observed only in EAs. The rs4760 association with total WBC was almost entirely due to its strong association with neutrophil counts. Outside of coding regions, an intronic variant (rs9374080) in CCDC162P was associated with increased total WBC in the multi-ancestry meta-analysis (p value =  $3.15 \times$ 10<sup>-9</sup>). The association was consistent across EAs and AAs

and was observed for neutrophil and monocyte counts and was especially strong for basophil counts. The rs3865444 variant, just upstream of CD33 (MIM: 159590), was associated with lower total WBC in the EA metaanalysis (p value =  $6.81 \times 10^{-14}$ ). The allele frequencies were highly discrepant across ancestry groups and rs3865444 was not significantly associated with total WBC outside of the EAs. However, the association was consistent across neutrophil, monocyte, and eosinophil counts. Finally, an intergenic variant (rs2836878) near ETS2 (MIM: 164740) and PSMG1 (MIM: 605296) was associated with lower total WBC in the multi-ancestry meta-analysis (p value =  $8.41 \times 10^{-9}$ ). The association was driven by the EA signal, and the variant had different allele frequencies across ancestry groups. The association with total WBC was consistent across neutrophil, monocyte, and basophil counts.

We identified a rare, missense variant in *OR4C6* (rs144349650, p.Leu112Val [c.334C>G], EAF = 0.00042) that was significantly associated with lower total WBC in the EA discovery analysis (p value =  $1.87 \times 10^{-11}$ ; Table S7). The allele frequency was rare in all ancestry groups and did not replicate in additional samples of >17,000 EAs, perhaps due to low statistical power. Likewise, we identified a burden of rare, missense variants in *TAF3* (MIM: 606576) that was significantly associated with increased total WBC in the EA discovery set ( $p_{VT} = 1.58 \times 10^{-6}$ ; Table S10). However, the signal did not replicate in an additional independent 2,898 samples.

## Neutrophil Count

In addition to the associations with total WBC, we identified two missense variants that were associated with neutrophil count at exome-wide significance levels. The effect estimate of the rs3747869 variant in *C100rf54* for total WBC appeared to be a combination of effects from neutrophil, monocyte, and eosinophil counts, though the effect was strongest for neutrophils, largely explaining the overall association with total WBC. The association between rs4760 in *PLAUR* and total WBC also appeared to be explained by the association with neutrophil counts.

The association between the rare, missense rs144349650 variant in *OR4C6* was observed for neutrophil counts as well as total WBC in the EA and multi-ancestry discovery sets. In gene-based test, *OR4C6* was associated with neutrophil count ( $p_{SKAT} = 2.56 \times 10^{-8}$ ; Table S10). Likewise, a burden of rare, missense variants in *ZNF439* was associated for neutrophil counts in the AA set ( $p_{VT} = 9.57 \times 10^{-7}$ ; Table S10). Neither the *ZNF439* nor the *OR4C6* gene-based association signals replicated.

## Monocyte Count

We found mostly non-coding variants associated with monocyte counts at the exome-wide level. One exception was the rs1292053 (p.Met76Thr) missense variant in TUBD1, for the multi-ancestry meta-analysis (p value = 6.53 ×  $10^{-12}$ ). Although the association was consistent across neutrophil and lymphocyte counts, the association with total WBC was almost entirely driven by the strong association with monocyte counts. An intergenic variant (rs4917014) near C7orf72-IKZF1 (MIM: 603023) was associated with lower monocyte count in the multi-ancestry meta-analysis (p value =  $9.75 \times 10^{-12}$ ). It was not associated with any other WBC sub-type. An intronic variant (rs11625112) in SLC7A8 (MIM: 600749) was associated with lower monocyte counts in the EA meta-analysis (p value =  $1.03 \times 10^{-9}$ ). We also found a secondary signal, rs11642873 near IRF8 (MIM: 601565),<sup>37</sup> that was associated with higher monocyte count in the EA meta-analysis (discovery beta [p value] =  $0.072 [1.40 \times 10^{-22}]$ , conditional beta [p value] =  $0.054 [1.41 \times 10^{-11}]$ ). Similar to their association with monocyte count, both rs11625112 in SLC7A8 and rs11642873 near IRF8 had consistent associations with basophil and eosinophil counts, but were not seen in AAs and HAs.

## Lymphocyte Count

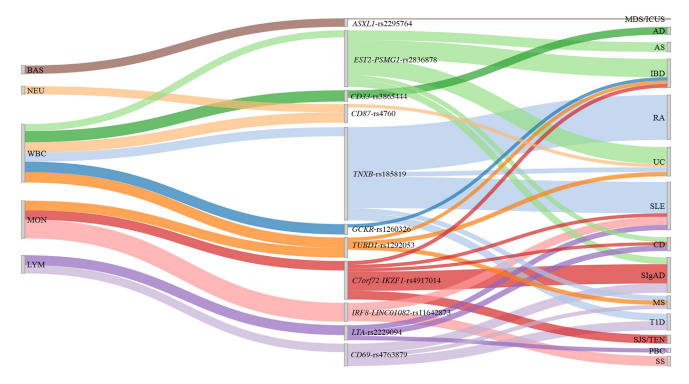
An intronic variant (rs4763879) in CD69 (MIM: 107273) was associated with decreased lymphocyte count in the EA meta-analysis (p value =  $1.59 \times 10^{-10}$ ). None of the other sub-types showed an association with rs4763879. The signal was not observed in AAs or HAs. A secondary missense variant (rs2229094, p.Cys13Arg [c.37T>C]) in LTA (MIM: 153440) was associated with higher lymphocyte count in the EA meta-analysis (p value =  $3.14 \times$  $10^{-12}$ ). The association was consistent across EAs and AAs, as well as for neutrophil counts, basophil counts, and for total WBC. LTA-rs2229094 is located near a previously reported WBC-associated SNP rs2524079 in LOC101929772,<sup>6</sup> though the LD between these variants is quite low ( $r^2 = 0.04$ ). Finally, although we observed a rare, missense variant in TRIM6 (MIM: 607564) (rs199694284, p.Val258Ala [c.773T>C], EAF in EAs =  $5.25 \times 10^{-5}$ , discovery p value =  $7.56 \times 10^{-8}$ ) associated with lymphocyte counts in EAs (Table S7), the association did not replicate.

## **Basophil Count**

In the EA meta-analysis, we identified a 3' UTR variant (rs2295764) in *ASXL1* (MIM: 612990) associated with lower basophil count (p value =  $1.46 \times 10^{-10}$ ). This variant was also associated with lower eosinophil and monocyte counts. The allele frequencies differed across ethnic groups and the association was not observed in AAs or HAs.

# Shared Associations of WBC Loci with Disease Phenotypes

To assess the shared association between these WBC loci and immune-mediated diseases and other relevant clinical phenotypes, we performed a PheWAS in 29,722 individuals and queried published GWAS databases of autoimmune diseases including IBD, MS, RA, SLE, and T1D. The majority of WBC variants discovered by the present study were associated with multiple autoimmune diseases. PheWAS identified TNXB (rs185819, p.His1161Arg) associated with risk of MS and SLE (Figure 2, Table 3). In lookups of GWAS databases, after correcting for multiple testing of 16 variants and 15 inflammatory diseases (p value  $< 2.08 \times$  $10^{-4}$ ), disease-variant associations were additionally detected for MS (CD69, TUBD1), IBD (GCKR, LTA, TNXB, IKZF1, TUBD1, ETS2-PSMG1), SLE (LTA, IRF8, TNXB, IKZF1), RA (TNXB), PBC (LTA), and T1D (CD69, TNXB). Additional associations between immunologically relevant clinical phenotypes and WBC trait variants included selective immunoglobulin A deficiency (MIM: 137100) with CD69 and IKZF1 (p value  $< 1.90 \times 10^{-11}$ ) and between IRF8 and systemic sclerosis (MIM: 181750) (p value = 2.30  $\times$  10<sup>-12</sup>). The inflammatory marker C-reactive protein (CRP) was strongly associated with GCKR and ETS2-*PSMG1* (p value <  $4.00 \times 10^{-8}$ ) (Tables 3 and S11).



#### Figure 2. Pleiotropy Plot Showing Shared Genetic Loci between WBC Traits and Autoimmune Inflammatory and Other Immune-Mediated Diseases

The thickness of each line connecting genes with WBC subtypes and immune-mediated diseases corresponds to the observed strength of association in p values. p values for gene-disease associations were derived from published genome-wide association studies (see Material and Methods section for references). Abbreviations are as follows: AD, Alzheimer disease; AS, ankylosing spondylitis; BAS, basophils; CD, Crohn disease; ICUS, idiopathic cytopenia of undetermined significance; IBD, inflammatory bowel diseases; LYM, lymphocytes; MON, monocytes; MDS, myelodysplastic syndrome; MS, multiple sclerosis; NEU, neutrophils; PBC, primary biliary cirrhosis; RA, rheumatoid arthritis; SIgAD, selective immunoglobulin A deficiency; SJS/TEN, Stevens-Johnson syndrome/toxic epidermal necrolysis; SLE, systemic lupus erythematosus; SS, systemic sclerosis; T1D, type 1 diabetes mellitus; UC, ulcerative colitis, WBC, white blood cells.

## Discussion

In this large-scale exome-wide association meta-analysis of WBC related traits in ~157,622 discovery and replication samples from five ancestries, we discovered 14 primary and 2 secondary SNV associations with total WBC and differential counts in EAs and the combined multi-ancestry samples, substantially increasing the number of loci associated with these hematologic traits. We observed shared genetic mechanisms influencing variations in WBC counts and susceptibility to chronic inflammatory and autoimmune diseases. These include genes and pathways involved in hematopoietic stem cell differentiation, apoptosis, cell adhesion, centrosome, and microtubule function.

Our statistical thresholds to declare significance at the discovery stage (p < 2 ×  $10^{-7}$  in the single-variant analyses) was adjusted for the approximate number of variants genotyped on the ExomeChip. Although we did not explicitly correct for testing multiple traits, the p values of our reported variants (Table 2) all pass the 5.0 ×  $10^{-8}$  standard of evidence for genomewide association studies of correlated traits.<sup>38</sup> Furthermore, we relied on independent replication to confirm our observed associations. Despite the limited size of

our replication set, it is noteworthy that we robustly replicated both known and novel WBC variants, suggesting a very low probability of reporting false-positive associations.

To quantitatively assess the contribution of loci identified by our Exomechip analysis, we have performed a comparative analysis of the proportion of total WBC phenotypic variance explained in a random sub-sample of 17,306 EAs from our largest discovery cohort, the WHI study. The proportion of variance in total WBC explained by the 28 previously known GWAS loci is 0.0137. The proportion of variance explained by the combination of known GWAS loci plus the ten additional Exomechipidentified loci we report is 0.0183. Thus, our Exomechip analysis has resulted in a 34% increase in the proportion of variance explained for total WBC in whites. These results suggest the possibility that exonic variants and/or variants not well-captured by traditional GWAS arrays may make an important contribution to the genetic architecture of WBC traits.

## Loci Involving Hematopoietic Lineage Differentiation and Activation of Cell Surface Receptors

Consistent with the pattern of association of the *CD33* index SNP rs3865444 with lower total WBC count involving

Trait (Population)	dbSNP ID	Chr	Pos	Alt/Ref	Gene	Phenotype	Sample Size	p Value <sup>a</sup>
WBC (EA)	rs1260328	2	27,730,940	C/T	GCKR	inflammatory bowel disease	96,486	$1.27 \times 10^{-4}$
LYM (EA)	rs2229094 <sup>b</sup>	6	31,540,556	C/T	LTA	Crohn disease	69,268	$7.81 \times 10^{-7}$
LYM (EA)	rs2229094 <sup>b</sup>	6	31,540,556	C/T	LTA	systemic lupus erythematosus	23,209	$3.09 \times 10^{-7}$
LYM (EA)	rs2229094 <sup>b</sup>	6	31,540,556	C/T	LTA	primary biliary cirrhosis	21,216	$1.31 \times 10^{-5}$
WBC (All)	rs185819	6	32,050,067	C/T	TNXB	multiple sclerosis <sup>c</sup>	22,850	$2.16 \times 10^{-6}$
WBC (All)	rs185819	6	32,050,067	C/T	TNXB	type 1 diabetes	33,394	$3.29 \times 10^{-9}$
WBC (All)	rs185819	6	32,050,067	C/T	TNXB	ulcerative colitis	72,647	$2.91 \times 10^{-6}$
WBC (All)	rs185819	6	32,050,067	C/T	TNXB	systemic lupus erythematosus <sup>c</sup>	23,209	$2.32 \times 10^{-3}$
WBC (All)	rs185819	6	32,050,067	C/T	TNXB	rheumatoid arthritis	103,558	$3.90 \times 10^{-53}$
MON (All)	rs4917014	7	50,305,863	G/T	C7orf72-IKZF1	systemic lupus erythematosus	32,444	$8.10 \times 10^{-5}$
MON (All)	rs4917014	7	50,305,863	G/T	C7orf72-IKZF1	inflammatory bowel disease	96,486	$4.59 \times 10^{-5}$
MON (All)	rs4917014	7	50,305,863	G/T	C7orf72-IKZF1	Crohn disease	69,268	$1.49 \times 10^{-4}$
MON (All)	rs4917014	7	50,305,863	G/T	C7orf72-IKZF1	Stevens-Johnson syndrome/ toxic epidermal necrolysis	1,129	$8.00 \times 10^{-1}$
MON (All)	rs4917014	7	50,305,863	G/T	C7orf72-IKZF1	selective immunoglobulin a deficiency	2,748	$2.80 \times 10^{-2}$
LYM (EA)	rs4763879	12	9,910,164	A/G	CD69	type 1 diabetes	38,522	$1.90 \times 10^{-1}$
LYM (EA)	rs4763879	12	9,910,164	A/G	CD69	multiple sclerosis	38,135	$2.18 \times 10^{-5}$
LYM (EA)	rs4763879	12	9,910,164	A/G	CD69	selective immunoglobulin A deficiency	2,748	$1.90 \times 10^{-1}$
MON (EA)	rs11642873	16	85,991,705	C/A	IRF8-LINC01082	systemic lupus erythematosus	23,209	$3.56 \times 10^{-10}$
MON (EA)	rs11642873	16	85,991,705	C/A	IRF8-LINC01082	systemic sclerosis	14,853	$2.30 \times 10^{-1}$
WBC (EA); MON (All)	rs1292053	17	57,963,537	G/A	TUBD1	multiple sclerosis	38,135	$7.47 \times 10^{-6}$
WBC (EA); MON (All)	rs1292053	17	57,963,537	G/A	TUBD1	Crohn disease	96,486	$8.53 \times 10^{-6}$
WBC (EA); MON (All)	rs1292053	17	57,963,537	G/A	TUBD1	inflammatory bowel disease	96,486	$9.61 \times 10^{-5}$
WBC (EA); NEU (EA)	rs4760	19	44,153,100	G/A	CD87 (PLAUR)	ulcerative colitis	72,647	$1.51 \times 10^{-4}$
WBC (EA)	rs3865444	19	51,727,962	A/C	CD33	Alzheimer disease	59,716	$1.60 \times 10^{-9}$
BAS (EA)	rs2295764	20	31,025,163	G/A	ASXL1	somatic mutations in MDS, CML, and ICUS	_	_
WBC (All)	rs2836878	21	40,465,534	A/G	ETS2-PSMG1	ankylosing spondylitis	9,609	$4.90 \times 10^{-1}$
WBC (All)	rs2836878	21	40,465,534	A/G	ETS2-PSMG1	Crohn disease	69,268	$2.43 \times 10^{-6}$
WBC (All)	rs2836878	21	40,465,534	A/G	ETS2-PSMG1	ulcerative colitis	72,647	$2.05 \times 10^{-2}$
WBC (All)	rs2836878	21	40,465,534	A/G	ETS2-PSMG1	inflammatory bowel disease	96,486	$3.70 \times 10^{-2}$
WBC (All)	rs2836878	21	40,465,534	A/G	ETS2-PSMG1	selective immunoglobulin A deficiency	2,748	$1.40 \times 10^{-8}$

Abbreviations: Chr, chromosome; Pos, basepair position; Alt, effect allele; Ref, reference allele; CML, chronic myelogenous leukemia; ICUS, Idiopathic cytopenia of undetermined significance; MDS, myelodysplastic syndrome, WBC, white blood cell; NEU, neutrophil; MON, monocyte; LYM, lymphocyte; BAS, basophil. <sup>a</sup>Significant results are shown after correcting for multiple testing of 16 variants and 15 diseases ( $p < 2.08 \times 10^{-4}$ ). When multiple studies report the same variant-trait associations, results from the largest sample size are presented here.

<sup>b</sup>LD r<sup>2</sup> between rs2229094 and rs1799964 is 0.75.

<sup>c</sup>Phenome-wide association results. Permutation p value for association with multiple sclerosis was 0.0122.

all myeloid lineages (and lower platelet count) (Table S12), CD33 is an early myeloid differentiation antigen and cell surface receptor that binds sialic acid-containing ligands and mediates diverse inhibitory functions of WBC in the innate immune system.<sup>39</sup> *CD33* is also highly

expressed on the surface of acute myeloid leukemia (AML) cells. *CD33* rs3865444 is in complete LD with *CD33* rs12459419 (p.Ala14Val), the presumed functional variant that results in lower full-length *CD33* expression due to skipping of exon 2.<sup>40</sup>

*PLAUR* encodes for the glycosyl-phosphatidylinositolanchored urokinase plasminogen activator receptor (*UPAR*). UPAR, also known as CD87, is a differentiation antigen on cells of the myelomonocytic lineage and also an activation antigen on monocytes and T lymphocytes.<sup>41,42</sup> The deleterious coding variant of *CD87* rs4760A>G (p.Leu272Pro)<sup>36</sup> is also a strong eQTL for *CD87* expression in monocytes and whole blood (Table S13). In addition to its role in plasminogen activation and fibrinolysis, UPAR is involved in cell adhesion, migration, and chemotaxis and is a regulator of the uptake by macrophages of apoptotic neutrophils.<sup>43</sup> It is possible that the latter mechanism might explain the selective association of rs4760 with lower neutrophil count.

The *CD69* intronic allele rs4763879G>A was associated with lower lymphocyte count but not with other WBC types. Accordingly, *CD69* encodes a calcium-dependent lectin superfamily of type II transmembrane cell surface receptor involved in regulation of lymphocyte proliferation.<sup>44</sup> As an early activation marker of lymphocytes, CD69 inhibits egress of lymphocytes into the circulation by downregulating sphingosine-1-phosphate receptor type 1 (MIM: 601974).<sup>44</sup> Notably, *CD69* rs4763879 correlates with the expression of *CD69* in monocytes and with the expression of C-type lectin domain family member genes *CLECL1* and *CLEC2D* in lymphoid cells (Table S13).

The intronic variant rs9374080 of non-coding RNA/ pseudogene CCDC162P has been previously associated with red blood cell traits-lower mean corpuscular volume, mean corpuscular hemoglobin<sup>45</sup>—and with platelet traits (Tables S11 and S12). In this study, we extend the association of rs9374080 to higher total WBC and myeloidderived cell counts, including basophil count (Table S8). The index SNP is located ~70 kb 3' of CD164 (endolyn) (MIM: 603356), which encodes a small transmembrane sialomucin protein on the surface of early hematopoietic progenitors, maturing erythroid cells, and activated basophils.<sup>46</sup> CD164 regulates CXCR4/CXCL12 signaling in hematopoietic precursor cells.<sup>47</sup> The region of association is located within a putative regulatory region enriched in epigenomic marks and ChIP-seq sites for various hematopoietic transcription factors (GATA1, TAL1) in K562 ervthroleukemia and lymphoblastoid cell lines.<sup>48</sup> These observations fit with the broad pattern of association of this variant with multiple blood cell lineages.

## Loci Involving Hematopoietic Transcription Factors and Epigenetic Modifiers

We identified variants in or near multiple genes encoding hematopoietic transcription factors that are associated with WBC traits. These loci include *IRF8-LINC01082*, *C7orf72-IKZF1*, *SLC7A8-CEBPE*, *JMJD1C*, *ASXL1*, and *ETS2-PSMG1*.

The 3' UTR variant rs2295764 of *ASXL1*, which was significantly associated with *ASXL1* transcript expression, was associated with lower basophil count and to a lesser

degree with lower monocyte and eosinophil counts and also to some extent with higher red cell distribution width (Tables S12 and S13). ASXL1 is a chromatin binding transcriptional regulator of the polycomb group and hematopoietic tumor suppressor gene.<sup>49</sup> *JMJD1C* is also an epigenetic regulator of gene expression, probably through histone demethylation.<sup>50</sup> The association between *JMJD1C* and lower WBC counts (this study), platelet count, mean platelet volume, and platelet reactivity<sup>51</sup> indicate multilineage effects on hematopoiesis. JMJD1C was originally identified as a ligand-dependent interacting partner of thyroid hormone and androgen receptors.<sup>52</sup> In human myeloid leukemia cells, *JMJD1C* functions as a coactivator for the leukemogenic transcriptional complex RUNX1-RUNX1T1 to increase AML cell proliferation and survival.<sup>53</sup> An intergenic variant rs2836878 located between ETS2 and PSMG1 showed evidence of multi-lineage association with lower total WBC count across all myeloid cell types and to a lesser extent with lower platelet count and higher hemoglobin (Table S12); rs2836878 is a wholeblood eQTL for ETS2 (Table S13). ETS2 is another protooncogene that encodes for a transcription factor involved in stem cell development, cell senescence, and death, whereas the product of PSMG1 is involved in maturation of proteasomes. ETS2, which is highly expressed in monocytes but not in granulocytes, has been shown to be involved in macrophage differentiation, regulation of megakaryocytic gene expression, T cell development, and phenotypic switch from erythroid to megakaryocytic development in hematopoietic cells.<sup>54</sup>

We identified several variants associated with monocyte count in loci that involve hematopoietic transcription factor genes (IRF8, SLC7A8, and IKZF1), further supporting their role in regulation of myelopoiesis and granulocyte/ monocyte lineage fate. The minor C allele of rs11642873, located 35 kb 3' of IRF8, was associated with higher monocyte count (and to a lesser degree with higher eosinophil and basophil counts) (Table S8). In eQTL analysis, an IRF8 variant rs17445836 is in moderate LD with the IRF8 rs11642873 variant ( $r^2 = 0.48$ ) that has a *cis*-regulatory effect on IRF8 expression in CD14<sup>+</sup> monocytes (Table S13). IRF8 encodes a transcription factor critical for myeloid lineage commitment by promoting differentiation of monocytes/dendritic cells and suppressing granulpoiesis.<sup>55</sup>  $Irf8^{-/-}$  mice have a myeloproliferative disorder with markedly increased number of macrophages and granulocytes in bone marrow, spleen, and lymph nodes as well as increased number of granulocytes in peripheral blood, suggesting a tumor-suppressive role of IRF8.<sup>56</sup>

Another non-coding variant associated with lower monocyte count, and to a lesser extent with lower basophil and eosinophil counts, was the intronic variant rs11625112 of *SLC7A8*, which encodes an amino acid transporter highly expressed in absorptive epithelia of the kidney and small intestine and also in the brain.<sup>57</sup> The index SNP is located within a blood cell DNase hypersensitivity site ~8 kb upstream of *CEBPE*, which encodes a

hematopoietic transcription factor essential for terminal differentiation and functional maturation of granulocytes.<sup>58</sup> Recent data also suggest a role of *CEBPE* isoforms in differential regulation of eosinophil production as well as in the monocyte-granulocyte lineage decision.<sup>59</sup>

The transcription factor encoded by IKZF1 or Ikaros was initially described as a regulator of lymphoid lineage differentiation and hematopoietic progenitor cell self-renewal.<sup>60</sup> An Ikaros isoform selectively expressed in myeloid precursor cells was subsequently found to regulate myeloid differentiation.<sup>60</sup> The minor allele of intergenic variant rs4917014 in C7orf72-IKZF1 associated selectively with lower monocyte count is located ~50 kb upstream of IKZF1 within an LD block enriched in hematopoietic cell DNase hypersensitivity sites and enhancer histone markers, several of which are also located within ChIPseq binding sites for the myeloid transcription factor PU.1.<sup>48</sup> The index SNP is also a monocyte and whole blood trans-eQTL for several immune response genes (Table S13). Further studies are required to assess whether the upstream IKZF1 or CEBPE regulatory elements harboring the index SNP are important for isoform- or lineage-specific monocyte development.

# Loci Involved in Regulation of Cell Death and Apoptosis

Apoptosis regulates hematopoietic stem cells and maintains the balance between cell proliferation and cell death.<sup>61</sup> Altered apoptotic processes contribute to the development of autoimmune and other inflammatory diseases.<sup>62</sup> We identified associations between WBC traits and coding variants in two additional genes involved in apoptosis. C10orf54 rs3747869 (p.Asp187Glu) was associated with higher total WBC and neutrophil counts. The product of C10orf54 (also known as Death Domain 1-alpha, DD1a), a direct transcriptional target of p53, regulates apoptosis and clearance of apoptotic cells, processes that are critical for resolution of inflammation, immune tolerance, and regulation of autoimmune responses.<sup>63</sup> DD1a is exclusively expressed within the hematopoietic compartment (monocytes, mature T cells, and macrophages) and functions as a negative immune checkpoint regulator for T cell activation and response.<sup>64</sup>

*LTA* rs2229094 (p.Cys13Arg) was associated with higher lymphocyte count. *LTA* encodes a member of the tumor necrosis factor family involved in lymphoid organ development and apoptosis.<sup>65</sup> Loss of *LTA* was associated with a 4-fold increase in B lymphocytes in peripheral blood count in mice.<sup>65</sup> The index missense SNP is also a *cis*-eQTL for *LTA* and *NFKBIL1* (MIM: 601022) (Table S13).

# Loci Involved in Other Cellular and Inflammatory Processes

We identified several missense variants (*TNXB* rs185819 [p.His1161Arg], *TUBD1* rs1292053 [p.Met76Thr], and *KIF9* rs2276853 [p.Arg573Trp]) in genes involved in other cellular processes that might be relevant to WBC produc-

tion or immune function. *TNXB* encodes a member of the tenascin family of extracellular matrix glycoproteins and inhibits cell adhesion and migration.<sup>66</sup> The index SNP localizes to the major histocompatibility complex class III region on chromosome 6 and overlaps *ATF6B* and *CYP21A2* at its 5' and 3' ends, respectively. The missense SNP is also an eQTL in blood or lymphoblastoid cell lines for several class II HLA genes (Table S13). The pattern of association of *TNXB* rs185819 suggests an effect at an early stage of myeloid and lymphoid differentiation. *ATF6B*, a member of the ATF6-related family of transcription factors that operate in the unfolded protein response,<sup>67</sup> is also a key virulence factor for *Toxoplasma gondii*.<sup>68</sup>

Although the role of *TUBD1* and *KIF9* on hematopoiesis is not known, both genes are involved in the structure and function of microtubules and centrosomes that are important for cell division and proliferation.<sup>69</sup> TUBD1 encodes for delta-tubulin microtubule protein that is associated with centrosome structure and function. The TUBD1 rs1292053 (p.Met76Thr), which was associated with both total WBC and monocyte counts, and to some extent with red cell and platelet parameters, is in LD with a number of SNPs in neighboring genes RPS6KB1 and RNFT1 and is a blood eQTL for RNFT1 (Tables S12 and S13). RPS6KB1 encodes a member of the ribosomal S6 kinase family of serine/threonine kinases and is part of the PI3K/AKT/ mTOR signaling pathway that plays a central role in a wide spectrum of cellular activities, including cell proliferation, survival, and differentiation.<sup>70</sup> The PI3K pathway is also involved in Toll-like receptor (TLR) signaling and release of cytokines from macrophages,<sup>71</sup> and a proxy SNP of TUBD1 rs1292053 has been associated with CRP.<sup>72</sup> KIF9 is a member of the kinesin family of genes related to microtubule binding and microtubule motor activity. The KIF9 rs2276853 variant is in LD with about 50 other variants spanning two other genes, SETD2 and KLHL18, several of which are within epigenomic blood cell marks and eQTLs for KIF9, KLHL18, and NBEAL2.<sup>48</sup>

The *GCKR* rs1260326 variant is an eQTL for *SNX17*, which has been associated with T cell activation and is a binding protein for human papillomavirus L2 capsid protein and for *NRBP1*, which binds a Dengue virus protein.<sup>48,73</sup>

## Relationship of WBC Loci to Autoimmune and Chronic Inflammatory Diseases

Abnormal immune response by lymphocytes and other white blood cells directed against self-antigens can lead to tissue injury and development of autoimmune disease.<sup>74</sup> Our results add to recent evidence that genetic factors controlling WBC and immune cell counts contribute to autoimmune disease risk.<sup>75</sup> Several loci involve regulation of cellular mechanisms critical in the development of autoimmune diseases such as modulation of autoimmune reactivity  $(CD69)^{76}$  and apoptosis (*LTA*, *DD1a*, *CD87*).<sup>43,63</sup>

The majority of our WBC-associated loci that showed substantial overlap were also associated with risk of various autoimmune and inflammatory diseases including IBD, RA, SLE, T1D, PBC, systemic sclerosis, Alzheimer disease, and Stevens-Johnson syndrome (Figure 2, Table 3). Although many of these genetic susceptibility loci are shared between different autoimmune diseases, other loci appear to be more restricted to particular cellular contexts. For example, there is an over-representation of SLE loci expressed selectively in B cells; RA-associated loci are preferentially expressed in CD4<sup>+</sup> effector T memory cells; epithelial-associated stimulated dendritic cell genes in Crohn disease; and monocyte-specific eQTLs among neurodegenerative disease variants.<sup>77,78</sup>

Abnormal inflammatory response and activation of microglial cells are linked with the development of AD and other neurodegenerative diseases. The WBC-associated gene CD33 is among the inflammation-related AD risk loci identified by GWASs.<sup>79</sup> A variant in this gene was shown to modulate CD33 exon 2 splicing efficiency, leading to abnormal activation of microglial cells that are tissue-resident macrophages of the brain derived from monocyte lineage cells.<sup>79</sup> In eQTL analysis of neuropathologically normal human brain tissues, CD33 rs3865444 is a cis-eQTL of C-type lectin domain family 11 member A (CLEC11A) that functions as growth factor for hematopoietic progenitor cells.<sup>80</sup> Several of the same loci are involved in susceptibility to infectious diseases (IRF8 and mendelian susceptibility to mycobacterial disease [MIM: 209950],<sup>81</sup> TNXB associated with T. gonadii and climatic adaptation,68,82 malaria with ABO [MIM: 110300] and DARC,<sup>3,83</sup> CD87 with clearance of bacteria<sup>84</sup>), highlighting the evolutionary trade-offs between protection against pathogens and risk of chronic disease later in life.

# Relationship of WBC Loci to Hematologic Disease and Therapy

Hematopoiesis is controlled by the differential expression of key transcription factors that act cooperatively to maintain a well-orchestrated balance of hematopoietic stem cell self-renewal and differentiation.<sup>85</sup> These functions of transcription factors are frequently dysregulated in leukemia by chromosomal translocations, mutations, or aberrant expression and lead to abnormal self-renewal. Several of the WBC loci have additional relationships to hematologic disease and therapeutics. CD33 is expressed in the brain and on AML blasts and leukemic stem cells and has therefore been exploited therapeutically as a target for anti-leukemic therapy.40 The CD33 rs3865444 and rs12459419 variants associated with lower WBC count and alternative splicing of exon 2, respectively, have been associated with both Alzheimer disease risk and AML treatment efficacy.<sup>40</sup> The exon 2 region of CD33 is important for sialic acid binding, microglial cell phagocytosis of beta-amyloid, and an epitope recognized by the antibody-targeted chemotherapy agent gemtuzumab ozogamicin.<sup>40,86</sup> CD87 is expressed on various immune cells

including neutrophils, monocytes, macrophages, T cells, and basophils, as well as endothelial cells and hepatocytes.<sup>41,87</sup> The cleaved soluble form of CD87 might have a role in hematopoietic stem/progenitor cell mobilization.<sup>88</sup>

Somatic mutations in ASXL1 are associated with risk of myelodysplastic syndrome (MDS [MIM: 614286]), chronic myelomonocytic leukemia (CMML [MIM: 607785]), and idiopathic cytopenia of undetermined significance (ICUS).<sup>49,89,90</sup> Knockdown of Asxl1 in mouse results in impaired lymphoid and myeloid differentiation and multi-lineage cytopenias.<sup>91</sup> Collectively, these results suggest that both germline and somatic mutations in ASXL1 cause lower blood cell counts. The transcription factor ETS2 has been shown to regulate phenotypic switch from erythroid to megakaryocyte in acute megakaryocytic leukemia (AMKL), and overexpression of ETS2 results in altered sensitivity to chemotherapy drugs.<sup>54</sup> Recent studies have shown that IKZF1 deletions and mutations that caused reduction of Ikaros activity are highly associated with development of acute lymphoblastic leukemia.<sup>92,93</sup> On the other hand, depletion of *JMJD1C* leads to growth impairment of a variety of leukemic cell types without noticeable effects on normal hematopoietic cells.<sup>52</sup> Therefore, JMJD1C is a potentially relevant drug target for leukemia.

Besides the single-variant association results, we confirmed previously reported gene-based association results for WBC count (CXCR2)<sup>12</sup> and monocytes (IL17RA) (N.P., U.M.S., J.B.-J., and M.-H.C., unpublished data). We also identified an additional gene putatively associated with WBC count (TAF3). IL17RA is widely expressed in myelomonocytic cells, lymphocytes, and bone marrow stromal cells and is part of the IL-17 cytokine signaling pathway that plays role in hematopoiesis, promotes inflammation, and is implicated in autoimmune diseases such as psoriasis, RA, and IBD.<sup>94</sup> TAF3, which encodes for a TATA-box binding protein, is located near GATA3, a transcription factor important for T lymphocyte differentiation. Variants in TAF3 are associated with mean corpuscular hemoglobin concentration<sup>95</sup> whereas GATA3 variants have been associated with susceptibility to hematologic malignancies.<sup>96</sup> Despite our large sample size, power to detect rare variants of more modest effect, either individually or aggregated into gene-based tests, may be limited. Future studies will require enormous sample sizes, probably considerably larger than in the current study, in order to detect additional rare variants (both individually and in aggregate) of moderate effect sizes associated with complex traits.

Our study has both strengths and limitations. By combining data from 25 studies world-wide, we were able to investigate the effect sizes and allele frequencies of variants in multiple ancestry groups. Variants with consistent effects across ancestries serve as strong candidates for causal variants. In addition to our ability to investigate how genetic variants influence WBC sub-types, our discovery analyses were well powered to detect moderate effect sizes. Indeed, although we did not correct for testing seven different phenotypes in three different meta-analyses, the combined p values of our reported variants (Table 2) all pass the  $5.0 \times 10^{-8}$  standard of evidence for genome-wide association studies of correlated traits.<sup>38</sup> We note that some cohorts did not measure a differential WBC in addition to total WBC, which limited our ability to assess associations with specific WBC subtypes in some instances.

In conclusion, by combining WBC exome-array analysis with PheWAS and functional annotation of variants, we identified likely causal variants associated with total and differential WBC counts as well as risk of autoimmune and inflammatory diseases. These results shed light on genetic mechanisms that regulate WBC counts and suggest a shared genetic architecture with predisposition to autoimmune and chronic inflammatory diseases. Future studies in model organisms are required to elucidate the underlying molecular mechanisms of how these genes result in variations in WBC count and development of autoimmune diseases.

## Supplemental Data

Supplemental Data include 2 figures, 13 tables, and additional funding information and can be found with this article online at http://dx.doi.org/10.1016/j.ajhg.2016.05.003.

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## Web Resources

1000 Genomes, http://www.1000genomes.org

- BCX ExomeChip association results, http://www.mhihumangenetics.org/en/resources
- CheckVCF, https://github.com/zhanxw/checkVCF

GRASP, http://grasp.nhlbi.nih.gov/Overview.aspx

HPC @ NIH, http://hpc.nih.gov

OMIM, http://www.omim.org/

R statistical software, http://www.r-project.org/

RareMETALS, http://genome.sph.umich.edu/wiki/RareMETALS

- RareMetalWorker, http://genome.sph.umich.edu/wiki/ RAREMETALWORKER
- RvTests, http://genome.sph.umich.edu/wiki/RvTests

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# **Supplemental Data**

# Large-Scale Exome-wide Association Analysis

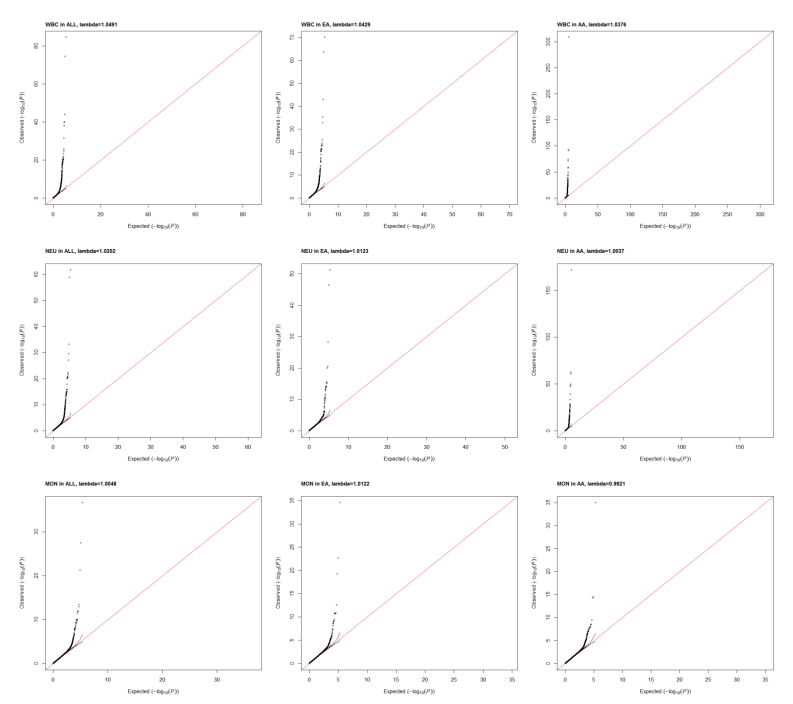
# Identifies Loci for White Blood Cell Traits

# and Pleiotropy with Immune-Mediated Diseases

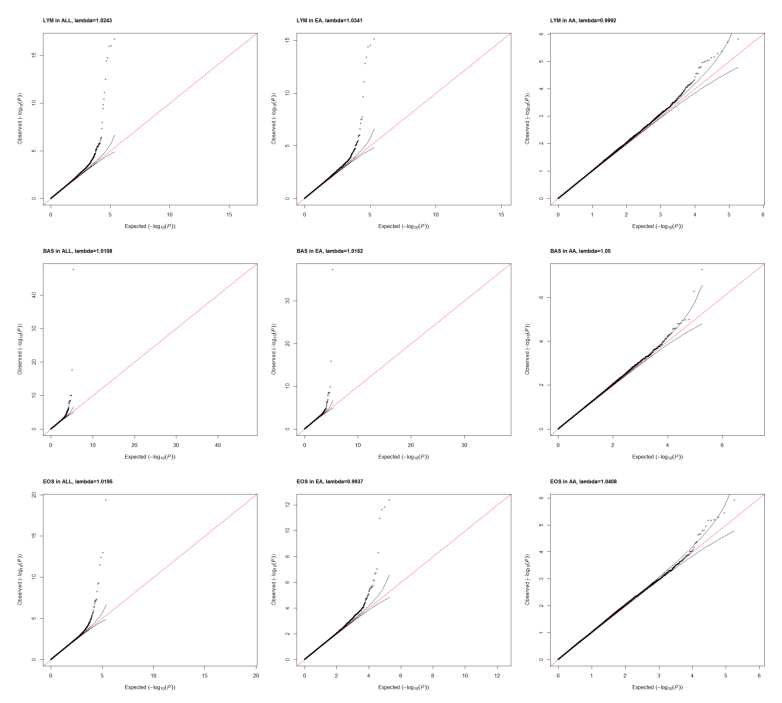
Salman M. Tajuddin, Ursula M. Schick, John D. Eicher, Nathalie Chami, Ayush Giri, Jennifer A. Brody, W. David Hill, Tim Kacprowski, Jin Li, Leo-Pekka Lyytikäinen, Ani Manichaikul, Evelin Mihailov, Michelle L. O'Donoghue, Nathan Pankratz, Raha Pazoki, Linda M. Polfus, Albert Vernon Smith, Claudia Schurmann, Caterina Vacchi-Suzzi, Dawn M. Waterworth, Evangelos Evangelou, Lisa R. Yanek, Amber Burt, Ming-Huei Chen, Frank J.A. van Rooij, James S. Floyd, Andreas Greinacher, Tamara B. Harris, Heather M. Highland, Leslie A. Lange, Yongmei Liu, Reedik Mägi, Mike A. Nalls, Rasika A. Mathias, Deborah A. Nickerson, Kjell John Starr. Jean-Claude Tardif. Ioanna Tzoulaki. Nikus. М. Digna R. Velez Edwards, Lars Wallentin, Traci M. Bartz, Lewis C. Becker, Joshua C. Denny, Laura M. Raffield, John D. Rioux, Nele Friedrich, Myriam Fornage, He Gao, Joel N. Hirschhorn, David C.M. Liewald, Stephen S. Rich, Andre Uitterlinden, Lisa Bastarache, Diane M. Becker, Eric Boerwinkle, Simon de Denus, Erwin P. Bottinger, Caroline Hayward, Albert Hofman, Georg Homuth, Ethan Lange, Lenore J. Lehtimäki, Yingchang Launer. Terho Lu, Andres Metspalu. Chris J. O'Donnell, Rakale C. Quarells, Melissa Richard, Eric S. Torstenson, Kent D. Taylor, Anne-Claire Vergnaud, Alan B. Zonderman, David R. Crosslin, Ian J. Deary, Marcus Dörr, Paul Elliott, Michele K. Evans, Vilmundur Gudnason, Mika Kähönen, Bruce M. Psaty, Jerome I. Rotter, Andrew J. Slater, Abbas Dehghan, Harvey D. White, Santhi K. Ganesh, Ruth J.F. Loos, Tõnu Esko, Nauder Faraday, James G. Wilson, Mary Cushman, Andrew D. Johnson, Todd L. Edwards, Neil A. Zakai, Guillaume Lettre, Alex P. Reiner, and Paul L. Auer

# **Supplemental Figure**

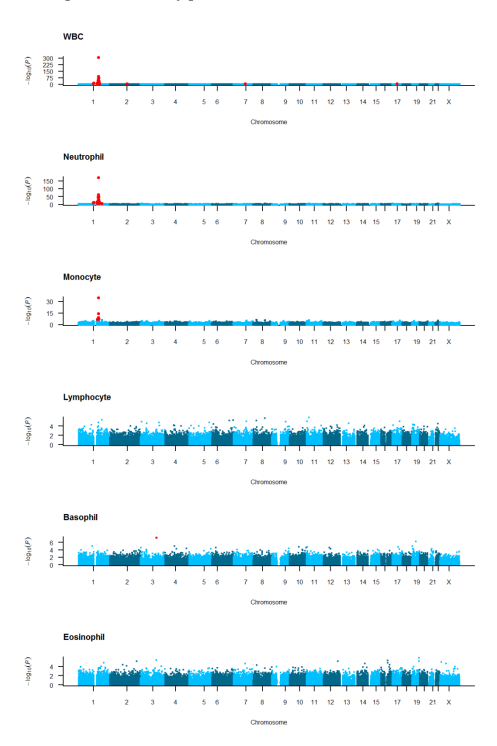
**Figure S1. Quantile-quantile plots of p-values of white blood cell traits.** Results of the three sets of discovery meta-analyses in combined all ancestries (ALL), European ancestry (EA), and African ancestry (AA) individuals are shown here. The combined all ancestry samples include Hispanic Americans, East Asians, and South Asians in addition to EAs and AAs.



# Figure S1. Continued







# **Supplemental Tables**

See Table S1 and Table S2 in the accompanying Excel file.

# Table S3. Sample size across ethnicities for white blood cell exome-wide association analysis

-		То	tal WBC					Neutrophil				Monocyte	9	L	ymphocy	te		Basophi			Eosinoph	ul
Study		E	thnicity					Ethnicity				Ethnicity			Ethnicity			Ethnicity	,		Ethnicity	Y
· -	EA	AA	HA	EAS	SA	EA	AA	HA	EAS	SA	EA	AA	HA	EA	AA	HA	EA	AA	HA/HL	EA	AA	HA
Discovery																						
AGES	2,955					2,954					2,954	368		2,954						2,954		
ARIC	10,345	2,821				6,792	368				6,792			6,792	368		6,792	368		6,792	368	
BIOME	1,043	2,328	3,319			636	1,981	2,688			10,209	1,981	2,688	636	1,981	2,688	636	1,981	2,688	636	1,981	2,688
BIOVU - I	21,276	1,921				10,208	909				636	910		10,218	911		14,903	1,357				
BIOVU - II		1,020					385					384			385			393				
CARDIA	2,110	1,946				2,110	1,942				2,095	1,928		2,110	1,942		1,637	1,350		1,955	1,762	
CHS	4,197	776																				
ECGUT	974					973					973			973			962			968		
FHS	5,610					2,314					2,314			2,314			2,314			2,314		
SOLID TIMI-52	8,025	198	874	254	116	8,026	198	874	251	116	_,			_,			_,			_,		
STABILITY	8,490	118	500	714	348	8,487	118	500	714	347												
HABC	1,251	110	200		510	1,251		200	/11	517	1,251			1,251			1,251			1,251		
HANDLS	1,201	834				1,201	834				1,201	834		1,201	834		1,201	813		1,201	832	
JHS		2,039					2,027					2,026			2,028			1,898			1,989	
LBC1921	475	2,057				473	2,027				473	2,020		473	2,020			1,070			1,707	
LBC1921	963					963					963			963								
MESA	1.266	750	843			1,122	673	763			1,123	675	764	1,121	675	762	1,123	673	765	1,121	671	762
MHIBB	7,911	750	045			6,978	075	105			6,978	075	704	6,978	075	762	6,978	075	705	6,978	0/1	762
REGARDS	7,911	5,030				0,778					0,778			0,778			0,778			0,778		
RS	2,794	5,050												2,758								
SHIP	7,251					4,076					4,076			4,076			4,075			4,077		
WHI	21,660	3,469				3,488	684				3,488	684		3,488	684		3,467	676		3,471	679	
Total per ancestry	108,596	23,250	5,536	968	464	60,851	10,119	4,825	965	463	44,325	9,790	3,452	47,105	9.808	3,450	44,138	9,509	3,453	32,517	8,282	3,450
Total per trait	108,570	25,250	5,550	708	138,814	00,001	10,117	4,025	705	77,223	44,525	),1)0	57,567	47,105	7,000	60,363	44,150	),50)	57,100	52,517	0,202	44,249
Replication					150,014					11,225			57,507			00,505			57,100			,2-1>
AIRWAVE	14,887					14,887					14,887			14,887			14,887			14,887		
BIOVU-Replication																				11,007		
(EA)	1,987					888					888			888			905					
FINCAVAS	911					397					397			397			397			440		
GeneSTAR	1.023					894					894			937								
Total per ancestry	18,808					17,066					17,066			17,109			16,189			15,327		
Total per trait	1				18,808					17,066	1		17,066			17,109	.,		16,189	/-		15,327
Grand Total					157,622					94,289			74,633			77,472			73,289			59,576

Abbreviations: EA, European Ancestry; AA, African Ancestry; HA, Hispanic American; EAS, East Asian; SA, South Asian.

Table S4. Mean and standard deviations of white blood cell traits in participating cohorts of the Blood-Cell Consortium

Study	Age	% Female	Total WBC	Neutrophil	Monocyte	Lymphocyte	Basophil	Eosinophil
AIRWAVE	40.87 (9.03)	37.3	6.52 (1.71)	3.94 (1.34)	0.4 (0.18)	1.8 (0.53)	0.06 (0.04)	0.2 (0.12)
n	14887	5550	14887	14886	14887	14886	14865	14866
AGES	76.4 (5.5)	57.9	6.05 (1.81)	3.55 (1.30)	0.541 (0.0178)	1.74 (0.972)	0.0286 (0.0256)	0.207 (0.145)
n	2953	1711	2955	2954	2954	2954	2954	2954
ARIC-AA	56.5 (5.8)	62.8	5.6 (1.9)	3.1 (1.4)	0.3 (0.1)	2.0 (0.6)	0.1 (0.0)	0.1 (0.1)
n	2832	1779	2832	369	369	369	160	355
ARIC-EA	57.3 (5.7)	53	6.2 (2.0)	3.8 (1.4)	0.4 (0.2)	1.8 (0.7)	0.1 (0.0)	0.2 (0.1)
n	10347	5489	10346	6792	6792	6792	2351	3893
BioME-AA	57 (14)	38.6	6.6 (2.2)	3.8 (1.8)	0.5 (0.2)	2.0 (0.8)	0.0 (0.0)	0.2 (0.1)
n	2653	1024	2329	1966	1966	1966	1966	1966
BioME-EA	68 (11)	38.9	7.2 (2.9)	4.7 (2.4)	0.6 (0.2)	1.7 (1.6)	0.0 (0.0)	0.2 (0.1)
n	1090	424	1046	634	634	634	634	634
BioME-HA	61 (15)	62	7.4 (2.3)	4.6 (1.9)	0.6 (0.2)	2.0 (0.8)	0.0 (0.0)	0.2 (0.2)
n	3640	2258	3319	2660	2660	2660	2660	2660
BioVU-I-EA	56.42 (16.82)	54.8	7.87 (5.34)	5.03 (3.16)	0.64 (0.54)	1.88 (2.76)	0.0456 (0.0433)	NA
N	21276	11654	21276	10208	10209	10218	14903	NA
BioVU-I-AA	49.58 (18.37)	61.1	7.30 (3.45)	4.56 (0.04)	0.59 (0.32)	2.06 (3.44)	0.0337 (0.0341)	NA
n	1921	1174	1921	909	910	911	1357	NA
BioVU-II-EA	50.93 (15.22)	11/4	7.75 (2.58)	4.93 (2.58)	0.58 (0.27)	1.9 (0.81)	0.036 (0.026)	NA
n	2298	2298	1987	888	888	888	928	NA
n BioVU-II-AA	39.40 (13.60)	100	7.63 (3.17)	4.58 (3.14)	0.56 (0.33)	2.53 (5.91)	928 0.0346 (0.0497)	NA
			. ,		0.36 (0.33) 384			
	1375	1375	1020	385		385	393	NA 0.17 (0.14)
CARDIA-EA	25.41 (3.36)	52.7 1113	6.25 (1.78) 2110	3.50 (1.40) 2110	0.32 (0.18) 2095	2.13 (0.65) 2110	0.04 (0.05) 1637	0.17 (0.14) 1955
n CADDIA AA	2110		2110					
CARDIA-AA	24.23 (3.77)	56.3	5.90 (1.95)	3.12 (1.49)	0.31 (0.17)	2.24 (0.83)	0.05 (0.04)	0.16 (0.13)
n Grug F.	1946	1095	1946	1942	1928	1942	1348	1760
CHS-EA	72.8 (5.6)	56.3	6.4 (2.0)	NA	NA	NA	NA	NA
n	4197	2361	4197	NA	NA	NA	NA	NA
CHS-AA	72.8 (5.6)	62.2	5.9 (1.9)	NA	NA	NA	NA	NA
n	776	483	776	NA	NA	NA	NA	NA
EGCUT	47.08 (16.72)	46.9	6.36 (1.7842)	3.70 (1.4080)	0.51 (0.1773)	1.96 (0.6362)	0.0299 (0.0194)	0.154 (0.1217
n	974	457	974	973	973	973	974	973
FHS	42.6 (13.9)	53.1	6.1 (1.6)	3.6 (1.2)	0.5 (0.1)	1.7 (0.5)	0.0 (0.0)	0.2 (0.1)
n	5790	3077	5610	2315	2315	2315	2315	2315
FINCAVAS	53.3 (13.9)	41.9	7.3 (2.5)	4.1 (2.2)	0.5 (0.2)	2.2 (1.1)	0.04 (0.03)	0.23 (0.2)
n	913	383	912	396	396	396	396	436
GeneSTAR-EA	42.47 (12.67)	52.20	6.64 (2.00)	4.21 (1.55)	0.38 (0.18)	2.09 (0.63)	NA	NA
n	1023	534	1023	894	894	937	NA	NA
GeneSTAR-AA	42.40 (12.34)	61.99	6.01 (1.87)	3.47 (1.47)	0.3 (0.16)	2.23 (0.7)	NA	NA
n	613	380	613	543	543	562	NA	NA
HABC-EA	75.72 (2.83)	52.5	6.593 (3.69)	3.837 (1.44)	0.564 (0.20)	1.93 (3.20)	0.0636 (0.04)	0.19 (0.17)
n	1251	657	1251	1251	1251	1251	1251	1251
HANDLS	48.13 (8.90)	54.7	6.23 (2.06)	3.55 (1.66)	0.40 (0.16)	2.08 (0.72)	0.03 (0.02)	0.16 (0.13)
n	870	476	834	834	834	834	813	832
JHS	53.0 (13.0)	37.5	5.59 (1.87)	3.09 (1.48)	0.39 (0.14)	1.93 (0.66)	0.03 (0.02)	0.14 (0.13)
n	2258	846	2044	2032	2030	2032	1903	2000
LBC1936	69.6 (0.8)	49.6	7.0 (1.9)	4.4 (1.5)	0.5 (0.2)	1.8 (0.8)	NA	NA
n	964	49.0 478	963	963	963	963	NA	NA
LBC1921	964 79.1 (0.6)	478 58.3	965 7.1 (1.9)	903 4.6 (1.4)	905 0.5 (0.2)	903 1.7 (1.1)	NA	NA
	475	58.5 277	7.1 (1.9) 475	4.6 (1.4) 473	0.5 (0.2) 473	473	NA	NA NA
n MESA EA								
MESA-EA	69.49 (9.55) 1267	50.1	6.139 (1.706) 1266	3.79 (1.299)	0.483 (0.1842)	1.666 (0.5854)	0.028 (0.0445)	0.171 (0.1275
n MESA AA	1267	635 52.0	1266	1193	1194	1193	1179	1186
MESA-AA	69.54 (9.10)	53.9	5.779 (1.93)	3.205 (1.4436)	0.428 (0.166)	1.883 (0.6357)	0.018 (0.0341)	0.162 (0.1284
n NEGA HA	750	404	750	669	670	670	653	667
MESA-HA	68.93 (9.43)	52	6.185 (1.6204)	3.68647 (1.3285)	0.433 (0.1618)	1.789 (0.5582)	0.020 (0.0385)	0.1721(0.132
n	845	439	843	623	622	622	609	613
MESA-EAS	65.12 (10.01)	47.1	5.314 (1.3629)	3.0834 (1.0671)	0.290 (0.108)	1.640 (0.5039)	0.030 (0.0311)	0.122 (0.1224
n	119	56	119	94	94	94	82	86
MHIBB	63.9 (10.9)	36.6	7.3 (2.5)	4.6 (2.2)	0.6 (0.2)	1.8 (1.0)	0.0 (0.1)	0.2 (0.1)
n	7911	2899	7911	7911	7911	7911	7911	7911
REGARDS	63.5 (9.2)	68.0	5.6 (2.0)	NA	NA	NA	NA	NA
	5039	3427	5030	NA	NA	NA	NA	NA
n	5039	5727	5050	11/1	11/1	111/1	19/1	11/1

n	3016	1618	2794	NA	NA	2758	NA	NA
SHIP	49.19 (16.09)	51	6.74 (2.02)	NA	NA	NA	NA	NA
n	3164	1613	3159	NA	NA	NA	NA	NA
SHIP-Trend	52.00 (15.40)	51.3	6.15 (1.94)	3.64 (1.41)	0.53 (0.18)	1.75 (0.59)	0.03 (0.02)	0.16 (0.12)
n	4245	2179	4238	4222	4222	4222	4221	4223
SOLID-TIMI 52-AA	60.50 (10.13)	33.8	6.909 (2.137)	4.3402 (1.773)	NA	NA	NA	NA
n	201	68	200	200	NA	NA	NA	NA
SOLID-TIMI 52-EA	64.55 (9.25)	25.8	7.5738 (2.22)	5.0038 (1.81)	NA	NA	NA	NA
n	8111	2092	8035	8036	NA	NA	NA	NA
SOLID-TIMI 52-EAS	63.59 (10.4)	19.6	6.8109 (2.005)	4.2585 (1.65)	NA	NA	NA	NA
n	255	50	255	252	NA	NA	NA	NA
SOLID-TIMI 52-SA	58.28 (9.67)	15.4	8.1573 (2.396)	5.6544 (2.074)	NA	NA	NA	NA
n	117	18	116	116	NA	NA	NA	NA
SOLID-TIMI 52-HA	63.29 (9.74)	28.3	7.4106 (2.171)	4.8 (1.792)	NA	NA	NA	NA
n	893	253	875	875	NA	NA	NA	NA
STABILITY-AA	62.86 (10.22)	25.4	5.6873 (1.67)	3.3362 (1.379)	NA	NA	NA	NA
n	118	30	118	118	NA	NA	NA	NA
STABILITY-EA	65.91 (8.98)	17.6	6.8245 (1.839)	4.3007 (1.427)	NA	NA	NA	NA
n	8499	1499	8490	8487	NA	NA	NA	NA
STABILITY-EAS	64.25 (8.72)	22.3	6.7628 (1.87)	4.1576 (1.466)	NA	NA	NA	NA
n	714	159	714	714	NA	NA	NA	NA
STABILITY-SA	58.35 (10.84)	13	7.5481 (1.977)	4.8583 (1.697)	NA	NA	NA	NA
n	354	46	348	347	NA	NA	NA	NA
STABILITY-HL	66.34 (8.41)	17.2	6.6986 (1.768)	4.2348 (1.444)	NA	NA	NA	NA
n	500	86	500	500	NA	NA	NA	NA
WHI-EA	66.2 (6.7)	100	6.1 (1.7)	3.9 (1.4)	0.6 (0.2)	1.8 (1.0)	0.0 (0.0)	0.2 (0.1)
n	21867	21867	21841	3669	3669	3669	3669	3669
WHI-AA	65.0 (6.6)	100	5.6 (1.7)	3.2 (1.5)	0.5 (0.2)	1.9 (1.0)	0.0 (0.0)	0.2 (0.2)
n	3501	3501	3497	712	712	712	713	712
Total	134,770		133,273	78,123	58,450	61,248	56,263	42,615

Abbreviations: EA, European Ancestry; AA, African Ancestry; HA, Hispanic American; EAS, East Asian; SA, South Asian.

Table S5. Blood cell specific expression quantitative trait loci (eQTL) datasets used for lookups of white blood cell trait variants identified in the present study

Whole blod samples18344981, 21829388, 22692066, 23818875, 23359819, 23880221, 24013639, 23157493, 23715323, 24092820, 24314549, 24956270, 24592274, 24728292, 24740359, 25609184, 22563384, 25578447Lymphoblastoid cell lines derived from asthmatic children17873877, 23345460Lymphoblastoid cell lines derived from 3 HapMap populations17873874Lymphoblastoid cell lines derived from HapMap CEU populations18193047Lymphoblastoid cell lines from population samples19644074, 22286170, 22941192, 23755361, 23995691, 25010687, 25951796Neutrophils26151758, 26259071CD19+ B cells22446964Primary phytohaemagglutinin (PHA)-stimulated T cells19644074, 23755361CD4+ T cells2083654Peripheral blood monocytes19222302,20502693,22446964, 23300628, 25951796, 26019233CD14+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interferon- gamma24604202CD14+ dendritic cells before and after stimulation with lipopolysaccharide (LPS) or interferon- interferon-beta24604203Micro-RNA QTLs21691150, 26020509DNase-I QTLs2307276	Datasets	Reference [PMID]
Lakoxyte sind kokoyte samples in individuals with Celia disease         192478           Whole blood samples         1834988, 12809388, 2209006, 23818875, 23359819, 2380721, 2401359, 23015493, 23715323, 24002820, 24314509, 24056270, 24972202, 24710359, 25060184, 2256334, 25474530, 2581634, 25578447           Lymphoblascid cell lines derived from athmatic children         1787387, 23345460           Lymphoblascid cell lines derived from 3Hapdap opulations         1787387           Lymphoblascid cell lines derived from Hapdap CEU populations         1819307           Lymphoblascid cell lines form population samples         0644074, 2236170, 22941192, 23755361, 23995691, 25010687, 25951796           Natrophils         0541738, 26259071           CD1+ 4 cells         2446964           Prinary phytohaenagglutin (PHA)-stimulated Teells         19644074, 23755361           Prinary phytohaenagglutin (PHA)-stimulated Teells         19622302,2024692446964, 23300628, 25951796, 26019233           CD1+ readi         1922302,202069,22446964, 23300628, 25951796, 26019233           CD1+ readi for a dafter Mytobacteriant taberculosis infection         22238106           CD1+ dericit cells before and after Mytobacteriant taberculosis infection         2230102           CD1+ dericit cells before and after Mytobacteriant taberculosis infection         230076           Dandrit cells before and after Mytobacteriant taberculosis infection         23017           Dandrit cells before and after Myt	Blood cell related eQTL studies included fresh lymphocytes	17873875
Whole blood samples         18340981, 1382988 229/02602, 32818875, 23359819, 23880221, 24013639, 25578407           Lymphoblastoid cell lines derived from ashmatic children         17873877, 23345460           Lymphoblastoid cell lines derived from 3 hapMap populations         17873874           Lymphoblastoid cell lines derived from HapMap CRU populations         18193047           Lymphoblastoid cell lines derived from HapMap CRU populations         18193047           Lymphoblastoid cell lines derived from HapMap CRU populations         18193047           Lymphoblastoid cell lines derived from HapMap CRU populations         18193047           Lymphoblastoid cell lines derived from HapMap CRU populations         18193047           CD19 I B cells         26151758, 26259071           CD19 I B cells         19644074, 23755361, 23995691, 25010687, 25951796           Primary phytobacmagglutinin (PHA)-stimulated T cells         19624074, 23755361           Primary phytobacmagglutinin (PHA)-stimulated T cells         19624074, 23755361           CD1 r cells         0833654           CD1 r deribric cells before and after Mycobacterium tuberculosis infection         2233810           CD1 r deribritic cells before and after Mycobacterium tuberculosis infection         2233810           CD1 r deribritic cells before and after Mycobacterium tuberculosis infection         2233810           Demotriti cells before and after Mycobacterium tuberculosis in	Fresh leukocytes	19966804
Whole mode samples24962274, 24728292, 24740359, 25609184, 22563384, 25474530, 2581634, 25578447Lymphoblastic cell lines drived from asthmatic children1787877, 2334540Lymphoblastic cell lines drived from Tampha populations1787877, 2334540Lymphoblastic cell lines drived from Tampha populations1890477Lymphoblastic cell lines drived from Tampha populations1964074, 22286170, 22941192, 23755361, 23995691, 25010687, 25951796Neutrophil26151785, 2629071CD19-B cells264064CD19-B cells1964074, 23236170, 22941192, 23755361, 23995691, 25010687, 25951796CD14-T cells2083054CD4-T cells2083054CD14-T cells2083054CD14-T cells relation with lipoplysaccharide (LPS) or inform aman2233100288, 25951796, 26019233CD14-t noncytes before and fler Mycobacterium tuberculosis infection2233810CD14-t cells before and after Mycobacterium tuberculosis infection223810CD14-t cells before and after Mycobacterium tuberculosis infection2030764CD14-t cells before and after Mycobacterium tuberculosis infection2030764CD14-t cells before and after Mycobacterium tuberculosis infection2030764Nace-QTLS2030764Nace-QTLS2030764Stoping QTLS2050789Splicing QTLS2568889Micro-RNA QTLS2568889Micro-RNA QTLS259142Splicing QTLS250423Splicing QTLS2504324Splicing QTLS2504324Splicing QTLS2505889Gontye-Tuse Expression (GTEx) whole Mod QTLS	Leukocyte samples in individuals with Celiac disease	19128478
Jumphoblastoid cell lines derived from 3 HapMap populations17873874Lymphoblastoid cell lines derived from HapMap CEU populations18193047Lymphoblastoid cell lines form population samples19644074, 22286170, 22941192, 23755361, 23995691, 25010687, 25991796Neurophils26151758, 26250710, 22941192, 23755361, 23995691, 25010687, 25991796CD19+ B cells26446964DPinary phytohemagglutinin (PHA)-stimulated T cells19644074, 23755361CD1+ T cells20833654CD4+ T cells20833052, 20502693, 22446964, 23300c28, 25991796, 26019233CD1+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interfere guman2464402CD1+ dedritic cells before or after stimulation with lipopolysaccharide (LPS) or interfere interferon-beta2404202CD1+ dedritic cells before or after stimulation with lipopolysaccharide (LPS) influenza or interferon-beta2406402CD1+ dedritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or 	Whole blood samples	
Lymphoblastoid cell lines derived from HpMpCEU populations         18193047           Lymphoblastoid cell lines from population samples         19644074, 22286170, 22941192, 2375361, 23995691, 25010687, 25951796           Neurophils         26151758, 26259071           CD19+ B cells         22446964           Primary phytohaemagglutinin (PHA)-stimulated T cells         19644074, 2375361           CD4+ T cells         20833654           Peripheral blood monocytes         1922202, 20202693, 22446964, 23300628, 25951796, 26019233           CD14+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interform gumma         24604202           CD14+ dordhitic cells before and after stimulation with lipopolysaccharide (LPS) or interform gumma         24604203           Netro-RNA QTLs         21691150, 26002069           Dask-1 QTLs         21691150, 26002069           Netro-RNA QTLs         2307276           Riboonal occupancy QTLs         25657249           Splicing QTLs         25685889           Seeding QTLs         25685889           SanDB eQTL data*         http://www.scandb.org/newinterface/about.html           Gentype-Tissue Expression (GTEx) whole blood cQTL <sup>b</sup> 2371523, www.gtexportal.org           QTL databases at University of Chicago         http://eqt.uchicago.edu/Home.html	Lymphoblastoid cell lines derived from asthmatic children	17873877, 23345460
Lymphoblastoid cell lines from population samples19644074, 22286170, 22941192, 23755361, 23995691, 25010687, 25951796Neutrophils26151758, 26259071CD19+ B cells22446964Primary phytohaemagglutinin (PHA)-stimulated T cells19644074, 23755361CD1+ T cells20833654Peripheral blood monocytes19222302,20502693, 22446964, 23300628, 25951796, 26019233CD1+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interform gumma24604202CD1+ dendritic cells before or after Stimulation with lipopolysaccharide (LPS), influenza or interferon-beta24604203Dendritic cells before and after Mycobacterium tuberculosis infection2233810Dendritic cells before or after Stimulation with lipopolysaccharide (LPS), influenza or interferon-beta26002509Norse-NA QTLs26002509DNase-1 QTLs2507276Ribosomal occupancy QTLs25052749Splicing QTLs25655889Splicing QTLs25655889Splicing QTLs25655889Splicing QTLs1pt//www.scandb.org/newinterface/about.htmlGenotype-Tissae Expression (GTEs), whole blood eQTL <sup>5</sup> 23715323, www.gtexportal.orgeQTL databases at University of Chicagohtp://eqtl.uchicago.edu/Home.html	Lymphoblastoid cell lines derived from 3 HapMap populations	17873874
Neurophils26151758, 26259071CD19+ B cells22446964Primary phytohaemagglutinin (PHA)-stimulated T cells19644074, 23755361CD4+ T cells20833654Peripheral blood monocytes19222302,20502633,22446964, 23300628, 25951796, 26019233CD14+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interferon- gamma24604202CD14+ dendritic cells before and after Mycobacterium tuberculosis infection2223810Dendritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or infereron-beta24604203Dendritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or infereron-beta2237810Dendritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or infereron-beta2207276Nicro-RNA QTLs2307276Disonal occupancy QTLs25657249Splicing QTLs25657249Splicing QTLs25657249Splicing QTLs25657249Splicing QTLs25685889SanDB eqTL data <sup>a</sup> http://www.scandb.org/newinterface/about.htmlGenotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 2715323, www.gtexportal.orgeqTL databases at University of Chicagohttp://www.scandb.org/newinterface/about.html	Lymphoblastoid cell lines derived from HapMap CEU populations	18193047
CD194 B cells22446964Primary phytohaemagglutinin (PHA)-stimulated T cells19644074, 23755361CD4+ T cells0833654Peripheral blood monocytes19222302,2050693,22446964, 23300628, 25951796, 26019233CD14+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interferor gamma2404202CD11+ dendritic cells before and after Mycobacterium tuberculosis infection2233810Dendritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or interferor-beta24604203Micro-RNA QTLs21691150, 26020509DNas-I QTLs21691150, 26020509DNas-I QTLs2307276Ribonal occupancy QTLs2567249Splicing QTLs2567249Splicing QTLs2567249Splicing QTLs2568889ScanDB eQTL data*http://www.scandh.org/newinterface/about.htmlGentype-Tissue Expression (GTEx) whole blood eQTL*23715323, www.gtexportal.orgeqTL databases at University of Chicagohttp://eqtl.uchicago.edu/Home.html	Lymphoblastoid cell lines from population samples	19644074, 22286170, 22941192, 23755361, 23995691, 25010687, 25951796
Primary phytohaemagglutinin (PHA)-stimulated T. cells1644074. 2375561CD4+ T. cells2083654Peripheral blood monocytes1922302.0502693.22446664. 23300628. 25951796, 26019233CD14+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interfero gamma2404020CD11+ dendritic cells before and after stimulation with lipopolysaccharide (LPS) or interfero interferon-beta2203810Dendritic cells before and after Mycobacterium tuberculosis infection22030200209.2014064.Dirditic cells before and after Mycobacterium tuberculosis infection2404020Nicro-RNA QTLs1691150, 26020509Dnsa-f QTLs2007276Histone acetylation QTLs2007276Fibrion QTLs2505249Splicing QTLs2505249Splicing QTLs2665889SanDB eQTL data*1691/92/92/92/92/92/92/92/92/92Splicing QTLs2665889Gorotype-Tissue Expression (GTEx) whole blood eQTLb*2017523, www.gtexportal.orge_QTL databases at University of Chicagobtp://qtuchicago.edu/Home.html	Neutrophils	26151758, 26259071
CD4+ T cells2083654Peripheral blood monocytes1922302.0502693,22466964, 23300628, 25951796, 26019233CD14+ monocytes before and after stimulation with lippolysaccharide (LPS) or interform gamma2203810CD11+ dendritic cells before and after Mycobacterium tuberculosis infection2223810Dendritic cells before or after stimulation with lippolysaccharide (LPS), influenza or interferon-beta24604203Micro-RNA QTLs1691150, 26020509DNase-I QTLs2307276Histone acetylation QTLs25079442Ribosonal occupancy QTLs2567249Splicing QTLs2567249Splicing QTLs2568889Airo-RNA QTLs2568889Splicing QTLs2568889SanDB eQTL data1tp://www.scandb.org/newinterface/about.htmlGentype-Tissue Expression (GTEx) whole blood eQTLb2315323, www.gtexportal.orgeqTL databases at University of Chicagohttp://eqtLuchicago.edu/Home.html	CD19+ B cells	22446964
Peripheral blood monocytes1922302,02502693,22446964,23300628,25951796,26019233CD14+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interfore gamma2404202CD11+ dendritic cells before and after Mycobacterium tuberculosis infection2233810Dendritic cells before and after stimulation with lipopolysaccharide (LPS), influenza or interforo-beta2406403Micro-RNA QTLs21091150, 26020509DNase-1 QTLs2307276Ristone accupator QTLs2507249Ristone accupator QTLs2507249Splicing QTLs25052499Splicing QTLs25052499Splicing QTLs25052499Splicing QTLs2505889SanDB eQTL data*1tp://www.scandb.org/newinterface/about.htmlGentype-Tissue Expression (GTEx) whole blood eQTL*215323, www.gtexportal.orgeQTL databases at University of Chicagottp://eqtLuchicago.edu/Home.html	Primary phytohaemagglutinin (PHA)-stimulated T cells	19644074, 23755361
CD14+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interfero24604202CD11+ dendritic cells before and after Mycobacterium tuberculosis infection2233810Dendritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or inferferon-beha24604203Micro-RNA QTLs21691150, 26020509DNase-I QTLs2307276Fistone acetylation QTLs2567249Ribosmal occupancy QTLs2567249Splicing QTLs25685889Splicing QTLs25685889SonDB eQTL data <sup>®</sup> 2568589Gentype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 271532, www.steandb.org/newinterface/about.htmlGentype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 2715323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqtLuchicago.edu/Home.html	CD4+ T cells	20833654
gamma2404202CD11+ dendritic cells before and after Mycobacterium tuberculosis infection2233810Dendritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or interferon-beta24604203Micro-RNA QTLs21691150, 26020509DNase-I QTLs2307276Histone acetylation QTLs25657249Splicing QTLs25657249Splicing QTLs2565789Splicing QTLs25658889SanDB eQTL data <sup>4</sup> http://www.scandb.org/newinterface/about.htmlGenotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 2315323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqtLuchicago.edu/Home.html	Peripheral blood monocytes	19222302,20502693,22446964, 23300628, 25951796, 26019233
Dendritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or interferon-beta24604203Micro-RNA QTLs1691150, 26020509DNase-I QTLs2307276Histone acetylation QTLs2579442Ribosomal occupancy QTLs2657249Splicing QTLs25685889Micro-RNA QTLs25791433Splicing QTLs25685899Splicing QTLs25685899SoaDB eQTL data <sup>a</sup> http://www.scandb.org/newinterface/about.htmlGenotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqtLuchicago.edu/Home.html	CD14+ monocytes before and after stimulation with lipopolysaccharide (LPS) or interferon- gamma	24604202
interferon-beta24004203Micro-RNA QTLs21691150, 26020509DNase-I QTLs2307276Histone acetylation QTLs25799442Splicing QTLs25657249Splicing QTLs25685889Micro-RNA QTLs25695889Splicing QTLs25685889Splicing QTLs25685889Splicing QTLs25685889Splicing QTLs2569523Splicing QTLs2569523Splicing QTLs2569523Splicing QTLs2569523Square Splicing QTLs2569523Square Splicing QTLs2569523Square Splicing QTL data <sup>a</sup> http://www.scandb.org/newinterface/about.htmlGenotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqt.luchicago.edu/Home.html	CD11+ dendritic cells before and after Mycobacterium tuberculosis infection	22233810
DNase-I QTLs22307276Histone acetylation QTLs2579442Ribosomal occupancy QTLs25657249Splicing QTLs25685889Micro-RNA QTLs25791433Splicing QTLs25685889Splicing QTLs25685889ScanDB eQTL data <sup>a</sup> http://www.scandb.org/newinterface/about.htmlGenotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqtLuchicago.edu/Home.html	Dendritic cells before or after stimulation with lipopolysaccharide (LPS), influenza or interferon-beta	24604203
Histone acetylation QTLs25799442Ribosomal occupancy QTLs25657249Splicing QTLs25685889Micro-RNA QTLs25791433Splicing QTLs25685889Splicing QTLs25685889ScanDB eQTL data <sup>a</sup> http://www.scandb.org/newinterface/about.htmlGenotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqtl.uchicago.edu/Home.html	Micro-RNA QTLs	21691150, 26020509
Ribosomal occupancy QTLs25657249Splicing QTLs25685889Micro-RNA QTLs25791433Splicing QTLs25685889ScanDB eQTL data <sup>a</sup> http://www.scandb.org/newinterface/about.htmlGenotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqtl.uchicago.edu/Home.html	DNase-I QTLs	22307276
Splicing QTLs25685889Micro-RNA QTLs25791433Splicing QTLs2568589ScanDB eQTL data <sup>a</sup> <u>http://www.scandb.org/newinterface/about.html</u> Genotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqtl.uchicago.edu/Home.html	Histone acetylation QTLs	25799442
Micro-RNA QTLs25791433Splicing QTLs25685889ScanDB eQTL data <sup>a</sup> http://www.scandb.org/newinterface/about.htmlGenotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.orgeQTL databases at University of Chicagohttp://eqtl.uchicago.edu/Home.html	Ribosomal occupancy QTLs	25657249
Splicing QTLs       25685889         ScanDB eQTL data <sup>a</sup> <u>http://www.scandb.org/newinterface/about.html</u> Genotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.org         eQTL databases at University of Chicago       http://eqtl.uchicago.edu/Home.html	Splicing QTLs	25685889
ScanDB eQTL data <sup>a</sup> http://www.scandb.org/newinterface/about.html         Genotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.org         eQTL databases at University of Chicago       http://eqtl.uchicago.edu/Home.html	Micro-RNA QTLs	25791433
Genotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup> 23715323, www.gtexportal.org         eQTL databases at University of Chicago       http://eqtl.uchicago.edu/Home.html	Splicing QTLs	25685889
eQTL databases at University of Chicago http://eqtl.uchicago.edu/Home.html	ScanDB eQTL data <sup>a</sup>	http://www.scandb.org/newinterface/about.html
	Genotype-Tissue Expression (GTEx) whole blood eQTL <sup>b</sup>	23715323, www.gtexportal.org
Long non-coding RNAs in monocytes 25025429	eQTL databases at University of Chicago	http://eqtl.uchicago.edu/Home.html
	Long non-coding RNAs in monocytes	25025429

<sup>a</sup>ScanDB cis-eQTLs were limited to those with P<1.0E-6 and trans-eQTLs with P<5.0E-8.

<sup>b</sup>GTEx Analysis V4 for 13 tissues were downloaded from the GTEx Portal (www.gtexportal.org) and results were filtered using the following criteria: Splicing QTL (sQTL) results generated with sQTLseeker with false discovery rate  $P \le 0.05$  were retained. For all gene-level eQTLs, if at least 1 SNP passed the tissue-specific empirical threshold in GTEx, the best SNP for that eQTL was always retained. All gene-level eQTL SNPs with P < 1.67E-11 were also retained, reflecting a global threshold correction of P = 0.05/(30,000 genes X 1,000,000 tests).

See Table S6 in the accombanying Excel file.

											Discovery				Replicati	on			
rait population)	dbSNPID	Chr	Pos	Alt	Ref	EAF	Gene	Annotation	AA Substitution	CADD Score	N	Beta	SE	Р	Ν	EAF	Beta	SE	Р
BC (EA)	rs4925663	1	247,614,617	Т	С	0.39	OR2B11	Missense	p.Gly223Asp	24.10	107,345	0.0251	0.0046	4.55E-08	17,897	0.4	0.01693	0.0112	0.132
BC (All)	rs4925663	1	247,614,617	Т	С	0.37	OR2B11	Missense	p.Gly223Asp	24.10	131,513	0.026	0.0042	5.85E-10	17,897	0.4	0.01693	0.0112	0.132
/BC (All)	rs1260326	2	27,730,940	С	Т	0.62	GCKR	Missense	p.Leu446Pro	0.11	132,764	- 0.0299	0.0044	1.44E-11	17,897	0.6	-0.0437	0.012	0.00027
/BC (EA)	rs1260326	2	27,730,940	С	Т	0.58	GCKR	Missense	p.Leu446Pro	0.11	108,596	- 0.0298	0.0048	4.01E-10	17,897	0.6	-0.0437	0.012	0.00027
/BC (All)	rs2276853	3	47,282,303	А	G	0.58	KIF9	Missense	p.Arg573Trp	32.00	132,764	0.023	0.0042	3.65E-08	17,897	0.6	0.02516	0.0116	0.03
/BC (All)	rs9374080	6	109,616,420	С	Т	0.43	CCDC162P	Intronic		3.79	132,764	0.0227	0.0041	4.01E-08	17,897	0.46	0.02512	0.0113	0.0255
YM (All)	rs2229094	6	31,540,556	С	Т	0.26	LTA	Missense	p.Cys13Arg	0.03	59,978	0.0451	0.0071	1.55E-10	16,711	0.25	0.07796	0.0175	8.54E-0
YM (EA)	rs2229094	6	31,540,556	С	Т	0.26	LTA	Missense	p.Cys13Arg	0.03	47,105	0.0456	0.0081	1.89E-08	16,711	0.25	0.07796	0.0175	8.54E-0
EU (All)	rs185819ª	6	32,050,067	С	Т	0.52	TNXB	Missense	p.His1161Ar g	0.00	76,838	0.0308	0.0058	9.65E-08	16,669	0.47	0.02635	0.0158	0.0948
/BC (All)	rs185819	6	32,050,067	С	Т	0.51	TNXB	Missense	p.His1161Ar g	0.00	132,764	0.0307	0.0049	4.02E-10	17,897	0.47	0.03395	0.0149	0.0224
/BC (All)	rs1050331	7	44,808,091	G	Т	0.48	ZMIZ2	3'UTR	5	5.30	123,297	0.0235	0.0041	1.25E-08	15,910	0.48	0.01149	0.0117	0.325
ION (All)	rs4917014	7	50,305,863	G	Т	0.28	C7orf72-IKZF1	Intergenic		1.91	57,183	-0.038	0.0068	1.97E-08	16,669	0.32	-0.0478	0.0122	8.92E-0
ION (EA)	rs10107630	8	130,603,635	Т	С	0.57	CCDC26	Intronic		5.39	44,325	- 0.0683	0.0069	2.45E-23	15,781	0.57	-0.1069	0.0118	1.06E-1
ION (All)	rs10107630	8	130,603,635	Т	С	0.56	CCDC26	Intronic		5.39	57,183	- 0.0663	0.006	3.11E-28	15,781	0.57	-0.1069	0.0118	1.06E-
BC (All)	rs1982151	9	86,617,265	G	А	0.71	RMII	Missense	p.Asn455Ser	0.00	132,764	0.0255	0.0044	6.97E-09	17,897	0.74	0.02095	0.0123	0.0896
BC (EA)	rs4409764	10	101,284,237	G	Т	0.51	GOT1- LINC01475	Intergenic		7.60	108,596	- 0.0255	0.0046	3.34E-08	17,897	0.53	-4.00E- 04	0.0115	0.974
ION (All)	rs6584283	10	101,291,593	С	Т	0.55	LINC01475	Intronic ncRN	A	-	57,183	-0.035	0.0061	9.55E-09	16,669	0.53	-0.0121	0.0121	0.319
/BC (EA)	rs1935	10	64,927,823	G	С	0.49	JMJD1C	Missense	p.Glu2353As p	16.64	108,596	- 0.0261	0.0047	2.46E-08	17,897	0.46	-0.0271	0.0117	0.0206
EU (All)	rs3747869	10	73,520,632	С	А	0.9	C10orf54 (DD1α)	Missense	p.Asp187Glu	10.88	76,838	0.0494	0.0087	1.58E-08	16,669	0.9	0.07326	0.019	0.0001
/BC (EA)	rs3747869	10	73,520,632	С	А	0.9	C10orf54 (DD1α)	Missense	p.Asp187Glu	10.88	108,596	0.0398	0.0073	4.26E-08	17,897	0.9	0.08289	0.0184	6.40E-0
EU (EA)	rs3747869	10	73,520,632	С	А	0.9	(DD1a) (DD1a)	Missense	p.Asp187Glu	10.88	60,851	0.0533	0.0095	2.11E-08	16,669	0.9	0.07326	0.019	0.00012
/BC (All)	rs3747869	10	73,520,632	С	А	0.91	(DD1a) (DD1a)	Missense	p.Asp187Glu	10.88	132,764	0.0381	0.0068	2.31E-08	17,897	0.9	0.08289	0.0184	6.40E-0
EU (EA)	rs144349650	11	55,432,976	G	С	0.0006 9	OR4C6	Missense	p.Leu112Val	13.09	60,851	0.7176	0.1096	5.95E-11	16,669	Monomorph	NA	NA	NA
EU (All)	rs144349650	11	55,432,976	G	С	0.0007	OR4C6	Missense	p.Leu112Val	13.09	76,838	0.6648	0.0962	4.92E-12	16,669	Monomorph ic	NA	NA	NA
/BC (EA)	rs144349650	11	55,432,976	G	С	0.0004 2	OR4C6	Missense	p.Leu112Val	13.09	108,596	0.7207	0.1073	1.87E-11	17,897	2.81E-05	-0.6178	0.9965	0.535
/BC (All)	rs144349650	11	55,432,976	G	С	0.0004 4	OR4C6	Missense	p.Leu112Val	13.09	132,764	0.6379	0.0942	1.26E-11	17,897	2.81E-05	-0.6178	0.9965	0.535
YM (EA)	rs199694284	11	5,632,403	С	Т	5.3E- 05	TRIM6	Missense	p.Val258Ala	25.20	40,313	- 2.3971	0.4458	7.56E-08	16,711	5.98E-05	-0.9067	0.6995	0.195
AS (EA)	rs4430553	12	66,698,895	С	Т	0.54	HELB	Missense	p.Leu191Pro	5.73	44,138	0.0404	0.0068	3.02E-09	15,770	0.53	0.02111	0.0116	0.0681
AS (All)	rs4430553	12	66,698,895	С	Т	0.56	HELB	Missense	p.Leu191Pro	5.73	56,707	0.0391	0.006	9.73E-11	15,770	0.53	0.02111	0.0116	0.0681
YM (All)	rs4763879	12	9,910,164	А	G	0.32	CD69	Intronic		3.03	59,978	- 0.0366	0.0064	1.08E-08	16,711	0.36	-0.0381	0.0119	0.0013
YM (EA)	rs4763879	12	9,910,164	А	G	0.36	CD69	Intronic		3.03	47,105	- 0.0379	0.0069	3.08E-08	16,711	0.36	-0.0381	0.0119	0.0013
ION (EA)	rs11625112	14	23,596,740	G	А	0.46	SLC7A8	Intronic		6.34	44,325	- 0.0376	0.0068	3.82E-08	16,669	0.45	-0.031	0.0115	0.0070
/BC (EA)	rs2306331	15	51,217,361	С	Т	0.46	AP4E1	Missense	p.Cys88Arg	6.49	100,571	- 0.0249	0.0046	6.15E-08	15,910	0.47	-0.0174	0.0116	0.134
ION (EA)	rs11642873ª	16	85,991,705	С	А	0.2	IRF8- LINC01082	Intergenic	-	0.75	44,325	0.0249	0.0085	1.41E-11	16,669	0.2	0.11251	0.0144	6.17E-

Table S7. Independent single nucleotide variants (N=28) associated with white blood cell traits in European only and combined All ancestries meta-analysis

MON (All)	rs11642873	16	85,991,705	С	А	0.17	IRF8- LINC01082	Intergenic		0.75	57,183	0.0561	0.0081	3.24E-12	16,669	0.2	0.11251	0.0144	6.17E-15
MON (All)	rs1292053	17	57,963,537	G	А	0.45	TUBD1	Missense	p.Met76Thr	5.53	57,183	- 0.0359	0.006	2.55E-09	16,669	0.44	-0.0398	0.0116	0.00061
WBC (All)	rs1292053	17	57,963,537	G	А	0.45	TUBD1	Missense	p.Met76Thr	5.53	132,764	- 0.0287	0.004	1.06E-12	17,897	0.44	-0.027	0.0111	0.0151
WBC (EA)	rs1292053	17	57,963,537	G	А	0.45	TUBD1	Missense	p.Met76Thr	5.53	108,596	- 0.0304	0.0045	1.28E-11	17,897	0.44	-0.027	0.0111	0.0151
BAS (All)	rs736289	19	33,757,062	С	Т	0.43	SLC7A10- CEBPA	Intergenic		0.41	56,707	0.0343	0.0061	2.34E-08	15,770	0.4	0.01229	0.0116	0.291
BAS (EA)	rs736289	19	33,757,062	С	Т	0.39	SLC7A10- CEBPA	Intergenic		0.41	44,138	0.0402	0.0069	7.02E-09	15,770	0.4	0.01229	0.0116	0.291
NEU (All)	rs4760	19	44,153,100	G	А	0.14	CD87 (PLAUR)	Missense	p.Lue272Pro	24.20	71,415	- 0.0471	0.0079	2.33E-09	16,669	0.15	-0.044	0.0159	0.00555
WBC (EA)	rs4760	19	44,153,100	G	А	0.16	CD87 (PLAUR)	Missense	p.Lue272Pro	24.20	85,685	-0.043	0.0068	2.51E-10	17,897	0.15	-0.0518	0.0153	0.00071
WBC (All)	rs4760	19	44,153,100	G	А	0.14	CD87 (PLAUR)	Missense	p.Lue272Pro	24.20	106,384	- 0.0424	0.0065	8.63E-11	17,897	0.15	-0.0518	0.0153	0.00071
NEU (EA)	rs4760	19	44,153,100	G	А	0.16	CD87 (PLAUR)	Missense	p.Lue272Pro	24.20	56,112	- 0.0467	0.0083	1.54E-08	16,669	0.15	-0.044	0.0159	0.00555
WBC (All)	rs3865444	19	51,727,962	А	С	0.28	CD33	Upstream		3.79	107,635	-0.034	0.005	6.25E-12	17,897	0.32	-0.0329	0.0118	0.00514
WBC (EA)	rs3865444	19	51,727,962	А	С	0.31	CD33	Upstream		3.79	86,936	- 0.0367	0.0053	3.51E-12	17,897	0.32	-0.0329	0.0118	0.00514
LYM (All)	rs6136489	20	1,923,734	G	Т	0.4	SIRPA-PDYN	Intergenic		2.52	59,978	- 0.0406	0.0061	3.99E-11	16,711	0.33	-0.0096	0.0121	0.428
WBC (All)	rs6136489	20	1,923,734	G	Т	0.39	SIRPA-PDYN	Intergenic		2.52	132,764	- 0.0246	0.0042	4.03E-09	17,897	0.33	-0.0179	0.0116	0.122
LYM (EA)	rs6136489	20	1,923,734	G	Т	0.34	SIRPA-PDYN	Intergenic		2.52	47,105	- 0.0384	0.007	3.33E-08	16,711	0.33	-0.0096	0.0121	0.428
BAS (EA)	rs2295764	20	31,025,163	G	А	0.36	ASXL1	3'UTR		0.10	44,138	- 0.0419	0.0071	3.28E-09	15,770	0.36	-0.031	0.012	0.00978
BAS (All)	rs2295764	20	31,025,163	G	А	0.33	ASXL1	3'UTR		0.10	56,707	- 0.0367	0.0064 1	1.07E-08	15,770	0.36	-0.031	0.012	0.00978
WBC (All)	rs2836878	21	40,465,534	А	G	0.26	ETS2-PSMG1	Intergenic		1.82	132,764	- 0.0246	0.0046	8.36E-08	17,897	0.26	-0.0264	0.0125	0.0344

Abbreviations: Chr, chromosome; Pos, basepair position; Alt, effect allele; Ref, reference allele; EAF, effect allele frequency; AA, amino acid; CADD, Combined annotation dependent depletion; EA, European ancestry, All, combined European, African, Hispanic American, East Asian and South Asian ancestries; WBC, white blood cell; NEU, neutrophil; MON, monocyte; LYM, lymphocyte; BAS, basophil.

<sup>a</sup>Secondary signal identified through conditional analysis.

Table S8. Association between 16 replicated loci and white blood cell subtypes in the discovery meta-analysis. Primary associations are shown in	l
bold.	

	2 3	27,730,940 27,730,940 47,282,303	GCKR KIF9	C/T	(population) WBC (EA) WBC (All) WBC (AA) NEU (EA) NEU (All) LYM (All) BAS (EA) BAS (All)	<b>108,596</b> 132,764 17,200 60,851 76,838 47,105 59,978 44,138	0.58 0.62 0.85 0.58 0.62 0.59 0.64	-0.030 -0.030 -0.039 -0.020 -0.019 -0.016 -0.013	0.005 0.004 0.016 0.006 0.006 0.007	<b>4.01E-10</b> 1.44E-11 1.44E-02 9.81E-04 6.46E-04 2.15E-02
rs2276853	3	47,282,303	KIF9		WBC (AA) NEU (EA) NEU (All) LYM (EA) LYM (All) BAS (EA)	17,200 60,851 76,838 47,105 59,978	0.85 0.58 0.62 0.59	-0.039 -0.020 -0.019 -0.016	0.016 0.006 0.006 0.007	1.44E-02 9.81E-04 6.46E-04
rs2276853	3	47,282,303	KIF9		NEU (EA) NEU (All) LYM (EA) LYM (All) BAS (EA)	60,851 76,838 47,105 59,978	0.58 0.62 0.59	-0.020 -0.019 -0.016	0.006 0.006 0.007	9.81E-04 6.46E-04
rs2276853	3	47,282,303	KIF9	10	NEU (All) LYM (EA) LYM (All) BAS (EA)	76,838 47,105 59,978	0.62 0.59	-0.019 -0.016	0.006 0.007	6.46E-04
rs2276853	3	47,282,303	KIF9		LYM (EA) LYM (All) BAS (EA)	47,105 59,978	0.59	-0.016	0.007	
rs2276853	3	47,282,303	KIF9		LYM (All) BAS (EA)	59,978				2.15E-02
rs2276853	3	47,282,303	KIF9		BAS (EA)		0.64	0.013		
rs2276853	3	47,282,303	KIF9	N/0	. ,	44,138		-0.015	0.006	3.91E-02
rs2276853	3	47,282,303	KIF9		BAS (All)		0.59	-0.022	0.007	2.17E-03
rs2276853	3	47,282,303	KIF9		Di lo (i lii)	56,707	0.64	-0.021	0.006	1.04E-03
				A/G	WBC (All)	132,764	0.58	0.023	0.004	3.65E-08
					WBC (EA)	108,596	0.60	0.022	0.005	1.39E-06
					NEU (EA)	60,851	0.60	0.017	0.006	4.56E-03
					NEU (All)	76,838	0.59	0.018	0.005	8.24E-04
					LYM (EA)	47,105	0.60	0.015	0.007	2.70E-02
					LYM (All)	59,978	0.58	0.018	0.006	2.52E-03
					LYM (AA)	9,423	0.53	0.039	0.015	7.86E-03
rs2229094	6	31,540,556	LTA	C/T	LYM (EA)	47,105	0.26	0.046	0.008	1.89E-08
					LYM (All)	59,978	0.26	0.045	0.007	1.55E-10
					LYM (AA)	9,423	0.28	0.044	0.017	7.30E-03
					WBC (EA)	108,596	0.26	0.035	0.006	2.73E-09
					WBC (All)	132,764	0.26	0.031	0.005	5.40E-09
					NEU (All)	76,838	0.26	0.015	0.006	1.74E-02
					BAS (EA)	44,138	0.26	0.026	0.008	1.37E-03
					BAS (All)	56,707	0.27	0.020	0.007	5.59E-03
rs185819	6	32,050,067	TNXB	C/T	WBC (All)	132,764	0.51	0.031	0.005	4.02E-10
		- ,,			WBC (EA)	108,596	0.52	0.029	0.006	2.62E-07
					WBC (AA)	17,200	0.47	0.035	0.012	4.38E-03
					NEU (EA)	60,851	0.52	0.025	0.007	1.33E-04
					NEU (All)	76,838	0.52	0.031	0.006	9.65E-08
					NEU (AA)	9,734	0.47	0.058	0.016	1.80E-04
					MON (EA)	44,325	0.51	0.020	0.008	1.14E-02
					MON (All)	57,183	0.51	0.021	0.007	1.89E-03
					LYM (EA)	47,105	0.51	0.015	0.007	3.93E-02
					LYM (All)	59,978	0.50	0.019	0.006	3.23E-03
					LYM (AA)	9,423	0.47	0.041	0.015	5.79E-03
					EOS (EA)	32,517	0.48	0.019	0.009	4.39E-02
					EOS (All)	44,249	0.48	0.016	0.008	3.58E-02
					BAS (EA)	44,138	0.51	0.020	0.007	6.87E-03
					BAS (All)	56,707	0.51	0.020	0.006	1.90E-03
rs9374080	6	109,616,420	CCDC162P	C/T	WBC (All)	132,764	0.43	0.023	0.004	4.01E-08
	U	107,010,420	00001021	U/ 1	WBC (AII) WBC (EA)	108,596	0.43	0.023	0.004	2.67E-06
					WBC (EA)	17,200	0.44	0.021	0.003	2.07E-00 3.74E-02
					NEU (EA)	60,851	0.30	0.024	0.006	1.08E-02
					NEU (EA) NEU (All)	76,831 76,838	0.44	0.013	0.005	5.69E-04
								0.018	0.005	
					MON (All) BAS (EA)	57,183 44 138	0.43			2.71E-02
					BAS (EA) BAS (All)	44,138 56,707	0.45 0.43	0.026 0.024	0.007 0.006	1.34E-04 9.85E-05

rs4917014	7	50,305,863	C7orf72-IKZF1	G/T	MON (All)	57,183	0.28458	-0.038	0.00677	1.97E-08
					MON (EA)	44,325	0.32243	-0.0336	0.00726	3.84E-06
					MON (AA)	9,406	0.08266	-0.0606	0.02683	2.40E-02
rs3747869	10	73,520,632	C10orf54 (DD1a)	C/A	WBC (EA)	108,596	0.90	0.040	0.007	4.26E-08
					WBC (All)	132,764	0.91	0.038	0.007	2.31E-08
					NEU (EA)	60,851	0.90	0.053	0.010	2.11E-08
					NEU (All)	76,838	0.90	0.049	0.009	1.58E-08
					MON (EA)	44,325	0.90	0.041	0.011	2.44E-04
					MON (All)	57,183	0.91	0.037	0.010	3.77E-04
					EOS (EA)	32,517	0.82	0.033	0.014	1.58E-02
					EOS (All)	44,249	0.85	0.025	0.012	4.23E-02
rs1935	10	64,927,823	JMJD1C	G/C	WBC (EA)	108,596	0.49	-0.026	0.005	2.46E-08
					WBC (All)	132,764	0.46	-0.025	0.004	6.57E-09
					NEU (EA)	60,851	0.49	-0.024	0.006	4.00E-05
					NEU (All)	76,838	0.46	-0.022	0.005	4.99E-05
					LYM (All)	59,978	0.45	-0.013	0.006	2.86E-02
					EOS (All)	44,249	0.41	-0.020	0.007	6.33E-03
rs4763879	12	9,910,164	CD69	A/G	LYM (EA)	47,105	0.36	-0.038	0.007	3.08E-08
					LYM (All)	59,978	0.32	-0.037	0.006	1.08E-08
					WBC (EA)	108,596	0.36	-0.014	0.005	2.63E-03
					WBC (All)	132,764	0.33	-0.013	0.004	2.94E-03
rs11625112	14	23,596,740	SLC7A8	G/A	MON (EA)	44,325	0.46	-0.038	0.007	3.82E-08
1311023112	11	23,370,710	SECTIO	0,11	MON (All)	57,183	0.46	-0.030	0.006	4.20E-07
					EOS (EA)	32,517	0.42	-0.021	0.008	1.17E-02
					EOS (All)	44,249	0.43	-0.021	0.007	2.77E-03
					BAS (EA)	44,138	0.46	-0.021	0.007	7.76E-06
					BAS (EA) BAS (All)	56,707	0.46	-0.026	0.007	1.07E-05
	16	85 001 705	IDEQ LINCOLOGO	C/A	MON (EA)				0.000	
rs11642873	10	85,991,705	IRF8-LINC01082	C/A		<b>44,325</b>	0.20	0.057		1.41E-11
					MON (All)	57,183	0.17	0.056	0.008	3.24E-12
					EOS (EA)	32,517	0.18	0.036	0.010	5.94E-04
					EOS (All)	44,249	0.15	0.033	0.010	8.18E-04
					BAS (EA)	44,138	0.20	0.021	0.009	1.53E-02
				~	BAS (All)	56,707	0.17	0.016	0.008	4.73E-02
rs1292053	17	57,963,537	TUBD1	G/A	WBC (EA)	108,596	0.45	-0.030	0.004	1.28E-11
					WBC (All)	132,764	0.45	-0.029	0.004	1.06E-12
					WBC (AA)	17,200	0.48	-0.029	0.011	8.31E-03
					NEU (EA)	60,851	0.45	-0.023	0.006	1.04E-04
					NEU (All)	76,838	0.45	-0.020	0.005	1.28E-04
					MON (EA)	44,325	0.45	-0.032	0.007	2.25E-06
					MON (All)	57,183	0.45	-0.036	0.006	2.55E-09
					MON (AA)	9,406	0.48	-0.035	0.015	1.80E-02
					LYM (EA)	47,105	0.44	-0.016	0.007	1.46E-02
					LYM (All)	59,978	0.45	-0.013	0.006	3.05E-02
rs4760	19	44,153,100	CD87 (PLAUR)	G/A	NEU (EA)	56,112	0.16	-0.047	0.008	1.54E-08
					NEU (All)	71,415	0.14	-0.047	0.008	2.33E-09
					WBC (EA)	85,685	0.16	-0.043	0.007	2.51E-10
							0.1.4	0.040		
					WBC (All)	106,384	0.14	-0.042	0.007	8.63E-11
rs3865444	19	51,727,962	CD33	A/C	WBC (All) WBC (EA)	106,384 <b>86,936</b>	0.14 0.31	-0.042 -0.037	0.007 0.005	8.63E-11 3.51E-12

					NEU (EA)	57,363	0.31	-0.025	0.006	1.15E-04
					NEU (All)	72,666	0.28	-0.021	0.006	3.26E-04
					MON (EA)	40,837	0.31	-0.020	0.008	9.46E-03
					MON (All)	53,011	0.28	-0.016	0.007	2.51E-02
					EOS (EA)	29,046	0.28	-0.029	0.010	2.16E-03
					EOS (All)	40,099	0.25	-0.025	0.009	3.09E-03
rs2295764	20	31,025,163	ASXL1	G/A	BAS (EA)	44,138	0.36	-0.042	0.007	3.28E-09
					BAS (All)	56,707	0.33	-0.037	0.006	1.07E-08
					WBC (AA)	17,200	0.20	-0.029	0.013	3.23E-02
					MON (EA)	44,325	0.36	-0.032	0.007	5.64E-06
					MON (All)	57,183	0.33	-0.026	0.006	3.95E-05
					EOS (EA)	32,517	0.32	-0.027	0.009	2.14E-03
					EOS (All)	44,249	0.30	-0.025	0.008	9.81E-04
rs2836878	21	40,465,534	ETS2-PSMG1	A/G	WBC (All)	132,764	0.26	-0.025	0.005	8.36E-08
					WBC (EA)	108,596	0.27	-0.024	0.005	7.49E-07
					NEU (EA)	60,851	0.28	-0.025	0.006	1.08E-04
					NEU (All)	76,838	0.26	-0.025	0.006	3.38E-05
					MON (EA)	44,325	0.28	-0.026	0.008	7.28E-04
					MON (All)	57,183	0.25	-0.022	0.007	1.69E-03
					BAS (EA)	44,138	0.28	-0.019	0.008	1.29E-02
					BAS (All)	56,707	0.25	-0.018	0.007	9.80E-03

Abbreviations: Chr, chromosome; Pos, basepair position; Alt, effect allele; Ref, reference allele; EAF, effect allele frequency; EA, European ancestry; AA, African ancestry; All, combined European, African, Hispanic American, East Asian and South Asian ancestries; WBC, white blood cell; NEU, neutrophil; MON, monocyte; LYM, lymphocyte; EOS, eosinophils; BAS, basophils.

See Table S9 in the accompanying Excel file

					Number	Discovery						Replicat	ion			
Trait	Population	Gene	Test	Chr	of variants	Ν	Р	Beta	S.D.	MAF cutoff	cMAC	Test	Ν	Р	Beta	S.D.
WBC	EA	CXCR2	SKAT	2	9	108,596	1.241E-14	-0.223	0.029	1	3,725	SKAT	2,898	0.0173	-0.406	0.169
WBC	All	CXCR2	SKAT	2	8	138,814	9.48E-15	-0.193	0.027	1	10,438					
WBC	EA	CXCR2	VT	2	9	108,596	7.239E-14	-0.223	0.029	0.012	3,725	VT	2,898	0.053	-0.406	0.169
WBC	All	CXCR2	VT	2	5	138,814	3.56E-13	-0.21	0.028	0.0023	10,438					
WBC	All	JAK2	VT	9	6	138,814	2.682E-06	0.334	0.066	0.00036	8,045	VT	2,898	0.0091	-0.505	0.163
WBC	EA	TAF3	VT	10	6	108,596	1.583E-06	0.328	0.064	0.00037	6,848	VT	2,898	NA	NA	NA
NEU	EA	CXCR2	SKAT	2	8	60,851	6.853E-09	-0.221	0.038	1	2,950	SKAT	1,285	0.190	-0.017	0.236
NEU	All	CXCR2	SKAT	2	7	77,223	5.062E-09	-0.203	0.035	1	9,252					
NEU	EA	CXCR2	VT	2	8	60,851	2.495E-08	-0.221	0.038	0.0185	2,950	VT	1,285	0.580	-0.392	0.335
NEU	All	CXCR2	VT	2	4	77,223	2.128E-08	-0.218	0.037	0.0023	9,252					
NEU	All	JAK2	VT	9	6	77,223	1.553E-07	0.431	0.077	0.00054	6,388	VT	1,285	NA	NA	NA
NEU	EA	OR4C6	SKAT	11	15	60,851	2.561E-08	0.109	0.044	1	512	SKAT	1,285	0.4844	-0.1455	0.198
NEU	EA	IL17RA	SKAT	22	18	44,325	1.004E-16	-0.128	0.026	1	10,016	SKAT	1,285	0.949	-0.018	0.176
NEU	AA	ZNF439	VT	19	4	10,119	9.566E-07	-1.099	0.221	0.083	1,688	VT	1,285	NA	NA	NA
MON	All	IL17RA	SKAT	22	19	57,567	1.174E-18	-0.104	0.023	1	23,595	SKAT	1,285	0.972	-0.004	0.176
LYM	All	TBX3	VT	12	4	60,363	1.959E-06	0.419	0.083	0.00060	144	VT	1,285	NA	NA	NA

Table S10. Genes associated with white blood cell and differential counts identified using gene-based association meta-analysis

Potentially novel genes are shown in bold.

Abbreviations: Chr, Chromosome; MAF, minor allele frequency; cMAC, cumulative minor allele count; WBC, white blood cell; NEU, neutrophil; MON, monocytes; LYM, lymphocytes; SKAT, sequence kernel association test; VT, variable-threshold test; ; EA, European ancestry; AA, African ancestry; All, combined European, African, Hispanic American, East Asian and South Asian ancestries.

Table S11. Association between white blood cell trait associated variants and immunologically relevant quantitative traits in previous genome-wide association studies (P < 1.64E-04).

Trait (population)	dbSNPID	Chr	Pos	Alt/Ref	Gene	Quantitative traits	Sample size	P <sup>a</sup>	PMID
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	C-reactive protein	70,410	3.80E-43	23263486, 21300955, 23505291, 22939635, 18439548
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Plasma protein-C levels	9,424	2.00E-17	20802025
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Serum urate	50,337	5.90E-17	20884846, 23263486, 21768215, 19503597, 21943158
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Platelet count	37,438	9.10E-10	22139419
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	2-hour glucose tolerance test	133,010	9.00E-15	22885924, 20081857, 23263486
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Fasting blood glucose	133,010	2.20E-41	22885924, 20081857, 20081858, 23263486
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Fasting insulin	133,010	2.70E-22	22885924, 20081857, 20081858, 23263486
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Factor VII activity in plasma	31,212	6.20E-24	21676895, 20231535, 21676895
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Total cholesterol	140,059	4.40E-28	20686565, 23063622, 20339536, 20339536, 19060906
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Triglycerides	140,059	1.30E-139	20686565, 22629316, 23063622, 19936222, 18193043, 20657596, 19060911, 19802338, 20339536, 23505323, 20139978, 19060910, 23236364, 21943158, 18454146, 19913121, 21862451, 19060906, 22171074
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	HDL	26,768	6.30E-36	19936222
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	Height	183,727	9.40E-05	20881960
WBC (EA)	rs1260326	2	27,730,940	C/T	GCKR	HOMA-IR	122,744	9.20E-07	20081858, 20081857, 23263486
WBC (All)	rs185819	6	32,050,067	C/T	TNXB	Height	183,727	1.70E-06	20881960, 18391951
WBC (All)	rs9374080	6	109,616,420	C/T	CCDC162P	Mean corpuscular hemoglobin concentration	135,367	3.00E-21	23222517
WBC (All)	rs9374080	6	109,616,420	C/T	CCDC162P	Mean corpuscular volume	135,367	2.30E-18	23222517, 19862010, 20139978
WBC (All)	rs9374080	6	109,616,420	C/T	CCDC162P	Red blood cell count	135,367	1.60E-15	23222517
MON (All)	rs4917014	7	50,305,863	G/T	C7orf72-IKZF1	HDL	140,059	2.40E-05	20686565
WBC (EA)	rs1935	10	64,927,823	G/C	JMJD1C	Platelet count	16,388	2.70E-08	22423221
WBC (EA)	rs1935	10	64,927,823	G/C	JMJD1C	Triglycerides	140,059	1.50E-07	20686565, 19060906
MON (EA)	rs11625112 <sup>b</sup>	14	23,596,740	G/A	SLC7A8	Triglycerides	140,059	4.90E-05	20686565, 19060906
WBC (All)	rs2836878	21	40,465,534	A/G	ETS2-PSMG1	C-reactive protein	82,725	4.00E-08	21300955, 22939635

<sup>a</sup>Lookup of 16 single nucleotide variants was performed and results with multiple testing corrected p-value < 1.64E-04 (16 variants and 19 quantitative traits) are shown here. When multiple studies report the same variant-trait associations, results from the largest sample size are presented here.

<sup>b</sup> LD r<sup>2</sup> between rs11625112 and rs2239633 in *SLC7A8* is 0.63.

Abbreviations: Chr, chromosome; Pos, basepair position; Alt, effect allele; Ref, reference allele; HDL, high density lipoprotein; HOMA-IR, homeostatic model assessment-insulin resistance; WBC, white blood cell; NEU, neutrophil; MON, monocyte; LYM, lymphocyte; BAS, basophil.

dbSNPID	Chr	Pos	Gene	Alt/Ref	EAF	Trait	Beta	SE	Р
European ances	stry								
rs1260326	2	27,730,940	GCKR	C/T	0.58	WBC	-0.03	0.005	4.01E-10
						PLT	-0.037	0.005	7.15E-14
						RBC	0.022	0.006	5.73E-04
						HCT	0.023	0.005	7.53E-06
						HGB	0.022	0.005	3.74E-06
rs2229094	6	31,540,556	LTA	C/T	0.26	LYM	0.046	0.008	1.89E-08
						PLT	0.019	0.006	2.04E-03
rs1935	10	64,927,823	JMJD1C	G/C	0.49	WBC	-0.026	0.005	2.46E-08
151700	10	01,727,020	0	0,0	0115	PLT	0.05	0.005	3.11E-25
						MPV	-0.112	0.008	7.64E-41
						RDW	-0.017	0.007	1.75E-02
						MCH	-0.017	0.007	1.18E-02
						MCT	0.017	0.007	1.16E-02
rs4763879	12	9,910,164	CD69	A/G	0.36	LYM	-0.038	0.007	3.08E-08
						HGB	-0.01	0.005	3.44E-02
						MCHC	-0.012	0.006	3.38E-02
				~					
rs11625112	14	23,596,740	SLC7A8	G/A	0.46	MON	-0.038	0.007	3.82E-08
						HCT	0.012	0.005	1.76E-02
rs1292053	17	57,963,537	TUBD1	G/A	0.45	WBC	-0.03	0.004	1.28E-11
						PLT	-0.017	0.005	1.94E-04
						RDW	0.024	0.007	3.85E-04
						HCT	-0.015	0.005	2.05E-03
						HGB	-0.016	0.005	5.11E-04
						MCV	-0.012	0.006	4.34E-02
						MCH	-0.016	0.006	1.45E-02
rs3865444	19	51,727,962	CD33	A/C	0.31	WBC	-0.037	0.005	3.51E-12
						PLT	-0.026	0.005	1.11E-06
						MCHC	0.016	0.006	8.70E-03
rs2205761	20	21 025 162	ASVI 1	G/A	0.26	DVC	-0.042	0.007	3.28E-09
rs2295764	20	31,025,163	ASXL1	G/A	0.36	BAS RDW	-0.042 0.034	0.007 0.007	3.28E-09 2.35E-06
All combined a	ncestries						0.001	0.007	2.552.00
rs2276853	3	47,282,303	KIF9	A/G	0.58	WBC	0.023	0.004	3.65E-08
		, ,				HGB	-0.011	0.004	1.14E-02
								-	
rs9374080	6	109,616,420	CCDC162P	C/T	0.43	WBC	0.023	0.004	4.01E-08
						PLT	0.015	0.004	3.07E-04
						MPV	-0.024	0.007	9.28E-04
						RBC	-0.043	0.005	3.13E-15
						RDW	-0.041	0.006	6.19E-11

Table S12. White blood cell trait variants associated with platelet and red blood cell related traits in the European and combined All ancestries of the Blood-Cell Consortium.

						MCV	0.06	0.006	8.84E-28
						MCH	0.062	0.006	3.10E-26
						MCHC	0.023	0.005	5.13E-06
rs185819	6	32,050,067	TNXB	C/T	0.51	WBC	0.031	0.005	4.02E-10
						PLT	0.017	0.005	6.16E-04
						MPV	-0.023	0.009	8.78E-03
						MCH	-0.015	0.007	2.94E-02
rs1292053	17	57,963,537	TUBD1	G/A	0.45	MON	-0.036	0.006	2.55E-09
						PLT	-0.013	0.004	1.40E-03
						RBC	-0.012	0.005	2.51E-02
						RDW	0.023	0.006	2.30E-04
						HCT	-0.013	0.004	2.97E-03
						HGB	-0.013	0.004	1.02E-03
rs2836878	21	40,465,534	ETS2-PSMG1	A/G	0.26	WBC	-0.025	0.005	8.36E-08
						PLT	-0.014	0.005	3.44E-03
						RDW	-0.025	0.007	3.28E-04
						HCT	0.016	0.005	1.37E-03
						HGB	0.022	0.005	2.72E-06
						MCV	0.014	0.006	2.14E-02
						MCH	0.024	0.007	2.78E-04
						MCHC	0.019	0.006	9.14E-04

Abbreviations: Chr, chromosome; Pos, basepair position; Alt, effect allele; Ref, reference allele; EAF, effect allele frequency; EA, European ancestry, All, combined European, African, Hispanic American, East Asian and South Asian ancestries; BAS, basophil; HCT, hematocrit; HGB, hemoglobin; LYM, lymphocyte; MCH, mean corpuscular hemoglobin; MCHC, mean corpuscular hemoglobin concentration; MCV, mean corpuscular volume; MON, monocytes; MPV, mean platelet volume; PLT, platelet; RBC, red blood cell; RDW, red cell distribution width; WBC, white blood cell.

See Table S13 in the accompanying Excel file.

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## BioMe

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