

ORIGINAL ARTICLE

Genome-wide association study identifies seven novel loci associating with circulating cytokines and cell adhesion molecules in Finns

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ABSTRACT

Background Inflammatory processes contribute to the pathophysiology of multiple chronic conditions. Genetic factors play a crucial role in modulating the inflammatory load, but the exact mechanisms are incompletely understood.

Objective To assess genetic determinants of 16 circulating cytokines and cell adhesion molecules (inflammatory phenotypes) in Finns.

Methods Genome-wide associations of the inflammatory phenotypes were studied in Northern Finland Birth Cohort 1966 (N=5284). A subsequent meta-analysis was completed for 10 phenotypes available in a previous genome-wide association study, adding up to 13 577 individuals in the study. Complementary association tests were performed to study the effect of the ABO blood types on soluble adhesion molecule levels.

Results We identified seven novel and six previously reported genetic associations ($p<3.1\times10^{-9}$). Three loci were associated with soluble vascular cell adhesion molecule-1 (sVCAM-1) level, one of which was the *ABO* locus that has been previously associated with soluble E-selectin (sE-selectin) and intercellular adhesion molecule-1 (sICAM-1) levels. Our findings suggest that the blood type B associates primarily with sVCAM-1 level, while the A1 subtype shows a robust effect on sE-selectin and sICAM-1 levels. The genotypes in the *ABO* locus associating with higher soluble adhesion molecule levels tend to associate with lower circulating cholesterol levels and lower cardiovascular disease risk.

Conclusion The present results extend the knowledge about genetic factors contributing to the inflammatory load. Our findings suggest that two distinct mechanisms contribute to the soluble adhesion molecule levels in the *ABO* locus and that elevated soluble adhesion molecule levels per se may not increase risk for cardiovascular disease.

INTRODUCTION

It is currently established that inflammatory load may play a role in the aetiology of autoimmune and infectious diseases, but also in a broad range of other diseases, such as chronic cardiometabolic disorders,¹ neurodegenerative diseases² and cancer.³ The risk for these diseases increases with age,⁴ and due to the world's ageing population⁵ their prevalence is likely to expand. Moreover, these diseases often co-occur, which is likely due to shared inflammation-related pathophysiology.⁶

Inflammation is the body's physiological response to harmful stimuli involving multiple molecular and cellular interactions attempting to restore disturbances in tissue or systemic homeostasis. Circulating cytokines, growth factors, chemokines and cell adhesion molecules (CAMs) (hereafter inflammatory phenotypes) are fundamental mediators of inflammatory responses. Genes encoding these molecules and their receptors play a crucial role in mediating the related functions. Previous studies have identified loci associating with levels of inflammatory phenotypes,^{7–9} but the understanding of the exact regulatory mechanisms is still incomplete.

To add insights into the genetic mechanisms contributing to the inflammatory load, we performed a genome-wide association study (GWAS) of 16 circulating inflammatory phenotypes in 5284 individuals from Northern Finland Birth Cohort 1966 (NFBC1966) and a subsequent meta-analysis of 10 phenotypes in three other Finnish population cohorts,⁷ adding up to a total of 13 577 individuals in the study. We report identification of seven novel and replication of six loci associating with levels of the circulating inflammatory markers.

METHODS

Study populations, genotyping and inflammatory phenotype quantification

Northern Finland Birth Cohort 1966

The NFBC1966 comprises 96% of all births during 1966 in the two northernmost provinces in Finland; altogether 12 058 children were live-born into the cohort, and the follow-ups occurred at the ages of 1, 14, 31 and 46 years.¹⁰ The data analysed in the present study are from the 31 years' follow-up when clinical examinations and blood sampling were completed for altogether 6033 individuals, 5284 of whom had body mass index (BMI), inflammatory phenotypes and genotype data available (a maximum number of individuals per inflammatory marker of 5100). Genotyping of the samples was completed

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Complex traits

using 370 k Illumina HumanHap arrays (Illumina, California, USA), and subsequent imputation was performed based on the 1000 Genomes reference panel. A total of 16 inflammatory phenotypes were quantified from overnight fasting plasma samples using Bio-Rad's Bio-Plex 200 system (Bio-Rad Laboratories, California, USA) with Milliplex Human Chemokine/Cytokine and CVD/Cytokine kits (Cat# HCYTOMAG-60K-12 and Cat# SPR349; Millipore, St Charles, Missouri, USA) and Bio-Plex Manager Software V.4.3 as previously described.¹¹ The 16 inflammatory phenotypes studied in the NFBC1966 were interleukin (IL) 1-alpha, IL1-beta (IL1β), IL4, IL6, IL8, IL17, IL1 receptor antagonist (IL1ra), interferon gamma-induced protein 10 (IP10), monocyte chemoattractant protein 1 (MCP1), tumour necrosis factor alpha (TNFα), vascular endothelial growth factor (VEGF), plasminogen activator inhibitor 1, soluble CD40 ligand, soluble E-selectin (sE-selectin), soluble intercellular adhesion molecule-1 (sICAM-1) and soluble vascular cell adhesion molecule 1 (sVCAM-1).

GWAS summary statistics from three Finnish population cohorts

Meta-analyses were conducted for 10 phenotypes available in a previous GWAS.⁷ The study included up to 8293 Finnish individuals from the Cardiovascular Risk in Young Finns Study (YFS)¹² and FINRISK (www.thl.fi/finriski),¹³ adding up to 13 577 individuals studied in the present meta-analyses. Shortly, YFS is a population-based follow-up study started in 1980 comprising randomly chosen individuals from Finnish cities Helsinki, Kuopio, Tampere, Oulu and Turku. The YFS data included in the previous GWAS are from 2019 individuals who participated in the follow-up in 2007 and who had both inflammatory phenotype and genotype data available. FINRISK is a Finnish population survey conducted every 5 years to monitor chronic diseases and their risk factors. The surveys use independent, random and representative samples from different geographical areas of Finland. The data included in the present meta-analyses were from participants of the 1997 and 2002 surveys. Genotypes were obtained using 670 k Illumina HumanHap arrays (Illumina) and imputed based on 1000 Genomes reference panel. Inflammatory markers were quantified using Bio-Rad's premixed Bio-Plex Pro Human Cytokine 27-plex Assay and 21-plex Assay, and Bio-Plex 200 reader with Bio-Plex V.6.0 software (Bio-Rad Laboratories) as previously described.¹⁴ Samples were serum in YFS, EDTA plasma in FINRISK1997 and heparin plasma in FINRISK2002.

Statistical analyses

GWAS and meta-analysis

To allow meta-analysis between the present results and the previous GWAS, the data processing and analysis model were done according to Ahola-Olli *et al.*⁷ First, rank-based inverse transformation was applied to normalise the phenotypes. Preceding the analyses, linear regression models were fitted to adjust the transformed inflammatory phenotypes for age, sex, BMI and the 10 first genetic principal components to control for population stratification. The resulting residuals were again normalised with inverse transformation, and the adjusted and transformed residuals were used as phenotypes in the analyses.

Genome-wide association tests were performed using snptest V.2.5.1 software.¹⁵ Allele effects were estimated using an additive model (-frequentist 1), and the option to centre and scale the phenotypes was disabled (-use_raw_phenotypes). The GWAS results were filtered by including markers with model fit info >0.8 and minor allele count >10. Filtered data were used to perform meta-analyses by METAL software (V.2011-03-25)¹⁶ for the 10 phenotypes (IL1 β , IL1ra, IL4, IL6, IL8, IL17, IP10, MCP1, TNF α

and VEGF) available in the previous GWAS.⁷ Genomic control correction was enabled (GENOMICCONTROL ON) to account for population stratification and cryptic relatedness. To estimate the heterogeneity of effect sizes between NFBC1966 and the previous GWAS, calculation of heterogeneity statistics based on Cochrane's Q-test was enabled (ANALYZE HETEROGENEITY).

Supplemental genome-wide tests in NFBC1966

Individuals showing symptoms of an acute infection were omitted from the supplemental genome-wide tests performed in the NFBC1966 population. Here, individuals reported having fever at the time of the blood sampling and individuals having C-reactive protein (CRP) level > 10 mg/L were excluded. Otherwise the analysis models were as above.

Conditional analyses and variance explained

To assess whether the identified loci harbour multiple independent association signals, we conducted conditional analyses by further adjusting the models with the locus-specific lead variants. The association tests were repeated within a 2 Mb window around the lead SNP for the phenotypes studied in the NFBC1966 population only. For the meta-analysed phenotypes, we applied a method proposed by Yang *et al*¹⁷ that enables conditional analyses of GWAS summary statistics. NFBC1966 was used as a reference sample to estimate linkage disequilibrium (LD) corrections in these analyses. The proportion of variance explained was calculated using all independent variants using the following formula:

Variance explained =
$$\left(\beta x \sqrt{2 \ x \ MAF \left(1 - MAF\right)}\right)^2$$

Here β is the variant's effect estimate on the inflammatory phenotype and MAF denotes minor allele frequency.

Complementary association tests on soluble adhesion molecule levels

Complementary association tests were conducted to better evaluate the molecular mechanism explaining the two potentially independent association signals with soluble CAM levels in the *ABO* locus. Here, linear models were repeated within a 2 Mb window and further adjusted for the ABO blood type or rs507666 genotype tagging the A1 subtype.¹⁸ In addition, we determined the effect estimates of ABO blood types and ABO blood types stratified by rs507666 genotype on sE-selectin, sICAM-1 and sVCAM-1 levels: the adjusted and transformed CAM concentrations were as outcomes in the linear models and ABO blood types as categorical variables (individuals with blood type A vs non-A, and so on). Corresponding models were fitted for the rs507666-stratified blood types (individuals with blood type A and rs507666 G/G vs others, and so on).

Shared genetic influences on inflammatory and cardiovascular phenotypes

As previous evidence suggests that elevated concentrations of circulating markers of inflammation increase the risk of cardio-vascular diseases (CVD),^{19 20} we further evaluated how variants in the loci associating with inflammatory phenotypes may relate to other cardiovascular traits. We used the gwas-pw method developed by Pickrell *et al*²¹ that estimates whether a locus harbours a genetic variant influencing one of the two phenotypes compared (models 1 and 2), if the same variant influences both phenotypes (model 3), or if separate variants within a locus influence the two phenotypes (model 4). Using the gwas-pw and open-access data provided by CARDIOGRAM,²² MEGASTROKE consortium²³ and

Table 1	Basic characteristics of the Northern Finland Birth Cohort
1966 stud	dy population

Characteristics	
Total number of individuals	5284
Number of men (%)	2543 (48.1)
Age, years	31.1±0.4
Body mass index, kg/m ²	24.4±4.0
Glucose, mmol/L	5.1±0.7
Low-density lipoprotein-cholesterol, mmol/L	3.0±0.9
High-density lipoprotein-cholesterol, mmol/L	1.6±0.4
Systolic blood pressure, mm Hg	124.2±13.6
Diastolic blood pressure, mm Hg	76.8±11.7

Values are mean±SD.

Global Lipids Genetics Consortium,²⁴ we evaluated the shared genetic determinants of circulating levels of low-density lipoprotein cholesterol (LDL-C), high-density lipoprotein cholesterol (HDL-C), total cholesterol (TotC) and total triglycerides (TG), as well as risk of coronary artery disease (CAD), ischaemic stroke and the inflammatory phenotypes showing significant genetic associations in the present study. The genome breakpoint data set for individuals with European ancestry provided at https://bitbucket.org/nygcresearch/ldetect-data was used in the gwas-pw analyses to split the genome into approximately independent blocks.²⁵

RESULTS

The basic characteristics of the NFBC1966 study population are provided in table 1. Inflammatory phenotype distributions are tabulated in online supplementary table S1, and their correlation structure is shown in online supplementary figure S1. Using a threshold of $p < 3.1 \times 10^{-9}$ for statistical significance (standard genome-wide significance level $p < 5 \times 10^{-8}$ corrected for 16 phenotypes tested), we identified seven novel and six previously reported loci associating with one or more of the inflammatory phenotypes. The results are summarised in table 2 and combined Manhattan plots are shown in figure 1. Manhattan plots and Q-Q plots for each inflammatory phenotype are provided in online supplementary figure S2 A-Z. Genomic inflation factor values range between 0.99 and 1.02, suggesting no inflation in the test statistics (online supplementary table S2). Online supplementary table S3 lists the traits associated previously with the loci showing novel associations with inflammatory phenotypes in the present study.

Cell adhesion molecules

The *ABO* locus shows large effects on sE-selectin, sICAM-1 and sVCAM-1 levels

We observed a novel effect on sVCAM-1 concentration in 9q34.2 near *ABO* (ABO, alpha 1–3 n-acetylgalactosaminyltransferase and alpha 1–3-galactosyltransferase) in the NFBC1966 population. This locus showed a robust association also with sE-selectin and sICAM-1 concentrations as previously reported.¹⁸ ²⁶ ²⁷ Noteworthy, the lead variant for sE-selectin and sICAM-1 associations (rs2519093) was different from the lead variant for sVCAM-1 association (rs8176746). The former variant is in LD ($r^2=1$ in NFBC1966) with rs507666 tagging the ABO blood type A subtype A1, whereas the latter variant tags the blood type B.¹⁸

As the GWAS results suggested two potentially separate association signals with the soluble CAM levels in the *ABO* locus, we conducted complementary association tests to better evaluate the molecular mechanisms explaining the associations. The association of the rs8176746 with sVCAM-1 concentration was significant when adjusted for the rs507666 indicative of the A1 subtype ($p=4.98 \times 10^{-15}$). On the contrary, the associations of the rs2519093 with concentrations of sE-selectin and sVCAM-1 were highly significant when adjusted for the ABO blood type ($p=3.40 \times 10^{-123}$ and $p=3.43 \times 10^{-17}$, respectively). Overall, these results suggest that the association of the rs8176746 with sVCAM-1 level is independent of the A1 subtype, while the association of the rs2519093 with sE-selectin and sICAM-1 levels is independent of the ABO blood type. Statistical significances were abolished when the rs8176746 association with sVCAM-1 was adjusted for ABO blood type and rs2519093 association with sE-selectin or sICAM-1 was adjusted for rs507666.

To further evaluate the related molecular mechanisms, we determined the effect estimates of the ABO blood types and ABO blood types stratified by rs507666 genotype on soluble CAM levels. The blood type A showed negative associations with the levels of all the three CAMs and the effect was the most robust on the sE-selectin level (figure 2, left panel). However, major discrepancies in the effect directions were seen when the analyses were stratified by the rs507666 genotype (figure 2, right panel). Congruent with previous reports,^{18 26} the present results suggest that the A1 subtype/rs507666 influences sE-selectin or sICAM-1 levels. In contrast, the blood type B seems to attribute predominantly to sVCAM-1 level, while the A1 subtype/rs507666 shows only a modest effect on sVCAM-1.

HSP90B1 and ABCA8 loci associate with sVCAM-1 levels

We identified two other novel loci for sVCAM-1 (12q23.3 and 17q24.2) in the NFBC1966 population. In chr12 the lead variant rs117238625 is in LD ($r^2=1$ in NFBC1966) with rs117468318 that locates in the 5' untranslated region (UTR) region of *HSP90B1* (heat shock protein 90 kDa beta member 1) and, according to RegulomeDB,²⁸ is likely to affect transcription factor binding. The association signal in chr17 locates near *ABCA8* (ATP binding cassette subfamily A member 8) encoding one of the ATP binding cassette transporters.

Variations in sialyltransferase encoding genes show an effect on sE-selectin level

For sE-selectin level, we identified a novel association in 11q24.2 in the region of *ST3GAL4* (ST3 beta-galactoside alpha-2,3-si-alyltransferase 4). We identified a suggestive signal with sE-selectin level also in 3q12.1 near *ST3GAL6* (ST3 beta-galactoside alpha-2,3-sialyltransferase 6), but the association was not significant after multiple correction ($p=1.75 \times 10^{-08}$). Both of the sialyltransferase genes have been implicated in the production of functional E-selectin, P-selectin and L-selectin ligands in mice.²⁹

Two independent association signals on sICAM-1 level near ICAM1

We replicated the previously reported association for sICAM-1 level in 19p13.2 near *ICAM1* (intracellular adhesion molecule 1).^{18 30} When the primary association test was conditioned for the lead variant rs117960796, another significant association was detected (rs74428614, p= 1.14×10^{-16}) indicative of more than one independent variant contributing to sICAM-1 level in this locus.

Vascular endothelial growth factor

In the meta-analyses, we identified a novel locus 4p16.2 with a large effect on VEGF (β =-2.38 SD). This locus harbours genes *EVC* (EvC ciliary complex subunit 1), *EVC2* (EvC ciliary complex subunit 2) and *STK32B* (serine/threonine kinase 32B). In addition, we replicated two previously reported loci associating with

Table 2 Si	gnificant lo	ci associati	ng with the circu	Table 2 Significant loci associating with the circulating inflammatory phenotypes	ry phenotypes									
C truck	no du che	-	Chundacittion.	Candidate acus	Macurat monola		dh CNDfouran		141	040 0			honiclaus scariol	Total variance
Suud	INIALKEL	rocus		canulate gene	(c)allagi lealest	AIIIOLALIOII	canulate gene livearest gene(s) Annotation apping reference		CA CAL	Deld	r value	пецгиа	или в кранат	expidined
NFBC1966	sE-selectin	9q34.2	9:136 141 870	ABO	ABO	Intronic	rs2519093	0.994	T 0.188	-0.903	4.48e-305	NA	0.249	0.258
		11q24.2	11:126 266 665	ST3GAL4	ST3GAL4	Intronic	rs11220471	0.967	G 0.212	-0.162	7.72e-12	NA	0.009	
	sICAM-1	9q34.2	9:136 141 870	ABO	ABO	Intronic	rs2519093	0.994	T 0.188	-0.352	7.43e-48	NA	0.038	0.118
		19p13.2	19:10 383 403	ICAM1	ICAM1	Intronic	rs117960796	0.802	A 0.012	-1.669	8.03e-40	NA	0.066	
		19p13.2	19:10 497 360	ICAM1	CDC37	Intergenic	rs74428614	0.992	A 0.163	0.226	1.14e-16*	NA	0.014	
	sVCAM-1	9q34.2	9 :136 131 322	ABO	ABO	Missense	rs8176746	1.000	T 0.129	0.256	5.06e-19	NA	0.015	0.038
		12q23.3	12 :104 448 391	HSP90B1	GLT8D2	Intronic	rs117238625	0.981	A 0.023	0.510	2.90e-14	NA	0.012	
		17q24.2	17:66 823 805	ABCA8	ABCA8	Intergenic	rs112001035	0.883	A 0.060	-0.324	1.04e-13	NA	0.012	
Meta-analyses	IL1B	6p22.1	6 :30 017 071		HLA locus	Intronic	rs6917603	1.000	C 0.251	-0.163	1.76e-12	1.00	0.010	0.015
		6p22.1	6 :30 013 887		HLA locus	Intronic	rs9261224	1.000	T 0.035	0.261	1.31e-09 *	1.00	0.005	
	IP10	4q21.1	4:76 899 176	CXCL10	SAD1	Intronic	rs192716315	0.851	C 0.003	1.513	2.71e-13	1.00	0.014	0.014
	MCP1	1q23.2	1:159 175 354	ACKR1	ACKR1	Missense	rs12075	1.000	A 0.469	0.148	1.43e-33	1.51e-13	0.011	0.011
	TNFac	13q14.3	13: 51 141 997	DLEU1	DLEU1	Intronic	rs17074575	0.803	G 0.002	2.131	2.71e-09	1.00	0.018	0.018
	VEGF	4p16.2	4:5 636 073	STK32B	EVC2	Intronic	rs186725382	0.875	A 0.001	-2.380	4.53e-10	1.00	0.011	0.052
		6p21.1	6:43 927 050	VEGFA	C6orf223	Intergenic	rs7767396	1.000	A 0.523	0.284	8.35e-105	1.22e-69	0.040	0.056
		9p24.2	9:2 686 273	VLDLR	VLDLR, KCNV2	Intergenic	rs7030781	0.959	T 0.373	-0.099	1.57e-13	5.34e-04	0.005	
Statistical sign	ificance is con	sidered at p<	:3.1×10 ⁻⁹ . Novel fin	Statistical significance is considered at $p<3.1\times10^{-9}$. Novel findings are highlighted with bold font. All positions correspond to human genome build 37	with bold font. All p	ositions corres	spond to human gene	ome build	37.					
*Indicates asso	ciations that.	are significar	nt after conditioning	*Indicates associations that are significant after conditioning the analyses on the locus-specific lead variant on the preceding row	ocus-specific lead va	riant on the pi	receding row.							
EA, effect allel	e; EAF, effect a	Ilele frequen.	cy; HLA, human leuk	cocyte antigen; HetPV	al, p value of heterog	Jeneity as esti	mated by Cochrane's	Q-test; IL	1β, interleu	kin 1-beta;	INFO, imputa	tion score in	EA, effect allele; EAF, effect allele frequency; HLA, human leukocyte antigen; HetPVal, p value of heterogeneity as estimated by Cochrane's Q-test; IL1B, interleukin 1-beta ; INFO, imputation score in NFBC1966; IP10, interferon gamma-induced	eron gamma-induced

protein 10; MCP1, monocyte chemoattractant protein 1; NA, not available; NFBC1966, Northern Finland Birth Cohort 1966; TNFo; tumour necrosis factor alpha; VEGF, vascular endothelial growth factor; dbSNP, single nucleotide polymorphism database; SE-selectin; slCAM-1, soluble intercellular adhesion molecule -1; sVCAM-1, soluble vascular cell adhesion molecule-1.

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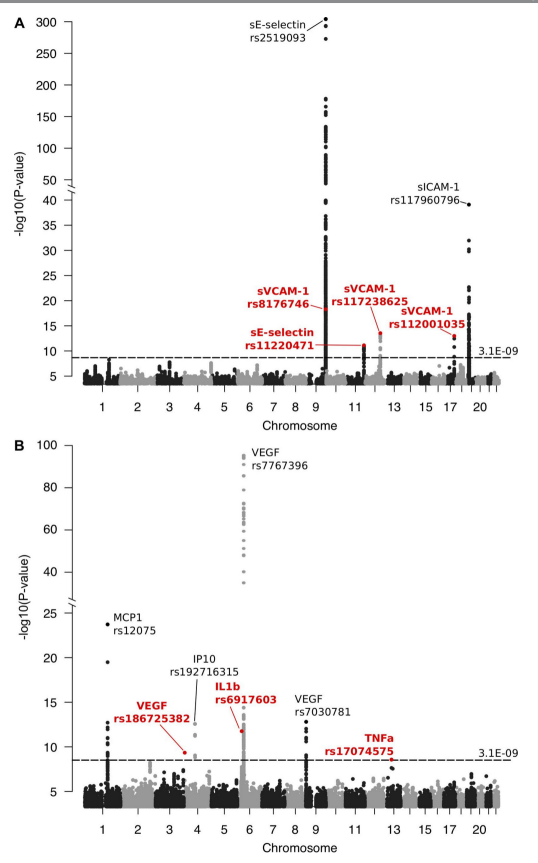


Figure 1 The combined Manhattan plots for significant associations with inflammatory markers studied in (A) Northern Finland Birth Cohort 1966 and in (B) meta-analyses with three other Finnish population cohorts. Significance threshold $p < 3.1 \times 10^{-9}$ derives from the standard p value limit for genome-wide significance $p < 5 \times 10^{-8}$ corrected for 16 markers examined in the present study. Novel association signals are highlighted with red font and replicated loci are marked with black font. sE-selectin, soluble E-selectin; IL1b, interleukin 1-beta; IP10, interferon gamma-induced protein 10; MCP1, monocyte chemoattractant protein 1; sICAM-1, soluble intercellular adhesion molecule-1; sVCAM-1, soluble vascular cell adhesion molecule-1; TNFa, tumour necrosis factor alpha; VEGF, vascular endothelial growth factor.

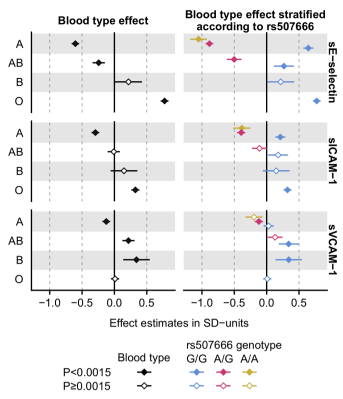


Figure 2 The effects of the ABO blood types and the A1 subtype on soluble adhesion molecule levels. The effects of the ABO blood types on sE-selectin, sICAM-1 and sVCAM-1 levels were evaluated in linear models, where adjusted (sex, age, body mass index and the 10 first genetic principal components) and transformed soluble adhesion molecule concentrations were used as outcomes and the ABO blood type served as categorical variable (A vs non-A, and so on). Corresponding models were fitted for the ABO blood types stratified by the rs507666-A allele count (0, 1 or 2), where the A allele tags the ABO subtype A1 having enhanced glycosyltransferase activity.¹⁸ No individuals were found to have B or O blood type and one or more copies of the rs507666-A allele, and thus it was not possible to perform stratification within these blood types. sE-selectin, soluble E-selectin; sICAM-1, soluble intercellular adhesion molecule-1; sVCAM-1, soluble vascular cell adhesion molecule-1.

VEGF levels in 6p21.1 near *VEGFA* (vascular endothelial growth factor A) and in 9p24.2 near *VLDLR* (very-low-density lipoprotein receptor).⁷

Proinflammatory cytokines

Locus near DLEU1 shows a large effect on TNFa

We identified a novel variant with a large effect on TNF α levels (β =2.13 SD) in 13q14.3 near *DLEU1* and *DLEU7* (deleted in lymphocytic leukaemia 1 and 7) in the meta-analyses.

The human leukocyte antigen (HLA) locus shows a small effect on $\mbox{IL}1\beta$

A novel variant at 6p22.1 in the human leucocyte antigen locus associating with IL1 β level was identified in the meta-analyses. In the conditional analyses, we observed two independent association signals at this locus (table 1, online supplementary figure S2J). The same locus and the same lead variant rs6917603 showed also a suggestive effect on IL4 level (online supplementary figure S2L), but the meta-analysed result was not significant after multiple correction (p= 5.56×10^{-09}).

Chemokines

We replicated previously reported loci near CXCL10 (C-X-C motif chemokine ligand 10) and ACKR1 (atypical chemokine receptor 1) associating with IP10 levels and with MCP1 levels, respectively.⁷

Supplemental genome-wide tests in NFBC1966

Altogether 236 individuals having fever or CRP > 10 mg/L were excluded from the supplemental genome-wide tests performed in the NFBC1966 population. The results of the supplementary analyses were congruent with the original findings (online supplementary table S4).

Comparisons of genetic effects on inflammatory phenotypes versus other traits

Elevated circulating concentrations of inflammatory markers increase the risk for CVD.^{19 20} We used the gwas-pw method²¹ to evaluate the presence of shared genetic determinants between inflammatory phenotypes showing significant genetic association in the present study and other cardiovascular health-related traits (LDL-C, HDL-C, TotC, TG, CAD risk, ischaemic stroke risk) obtained from open-access sources.²²⁻²⁴ Altogether 56 genomic regions showed robust statistical evidence for containing a genetic variant influencing one or more of the inflammatory phenotypes and at the same time one or more of the other traits studied (model 3 posterior probability greater than 0.99; online supplementary figure S3). The ABO locus was one of the loci harbouring variants influencing multiple traits. In this locus, we observed negative linear relationships between the SNP effects on sE-selectin and sICAM-1 levels and CAD risk, stroke risk, as well as LDL-C, HDL-C and TotC levels (figure 3). The results in the other loci are provided in online supplementary figure S3.

DISCUSSION

The present study examines the genetic determinants of 16 circulating inflammatory phenotypes in 5284 individuals from Northern Finland with a subsequent meta-analysis of 10 phenotypes in 3 other Finnish populations, adding up to a total of 13 577 participants. We report seven novel and replication of six previously published genetic associations.

We identified a novel association for sVCAM-1 concentration at the ABO locus. This locus was also associated with sE-selectin and sICAM-1 levels as observed previously.^{18 26} The present GWAS suggested two distinct association signals in the ABO locus for the sE-selectin and sICAM-1 levels versus sVCAM-1 level, and the supplementary tests provided further support for at least two mechanisms contributing to circulating concentrations of CAMs in this locus. The two mechanisms include the blood type A subtype A1, which has a robust lowering effect on sE-selectin and sICAM-1 levels,^{18 26} and the blood type B which seems to have an increasing effect on sVCAM-1 level. The lowering effect of the A1 subtype on sE-selectin and sICAM-1 could arise from increased glycosyltransferase activity that possibly modifies the shedding of the CAMs from the endothelium and/or their clearance rate from circulation.^{18 26} The underlying mechanism explaining the association between the blood type B and higher sVCAM-1 concentration remains unknown and warrants research. VCAM-1-mediated adhesion involves interaction with galectin-3, a protein that has a specificity for galactosides.³¹ As the B antigen holds an additional galactose monomer compared with the A and O antigens, and galectins are known to recognise blood type antigens,³² it raises the speculation that the amount of unbound sVCAM-1 in the

TG -	0.2	0.19		-0.24	-0.2	-0.14	-0.24	-0.06	1
TotC -	-0.76	-0.72	-0.45	0.57	0.34	0.93	0.33	1	-0.06
HDL-C -	-0.27	-0.24	0.26	0.31	0.08	0.1	1	0.33	-0.24
LDL-C -	-0.74	-0.72	-0.6	0.51	0.35	1	0.1	0.93	-0.14
stroke -	-0.31	-0.26	-0.07	0.43	1	0.35	0.08	0.34	-0.2
CAD -	-0.59	-0.55	-0.21	1	0.43	0.51	0.31	0.57	-0.24
sVCAM-1 -	0.49	0.54	1	-0.21	-0.07	-0.6	0.26	-0.45	
sICAM-1 -	0.9	1	0.54	-0.55	-0.26	-0.72	-0.24	-0.72	0.19
sE-selectin -				-0.59					
SE-Selectin SUCANN' CAD STOKE LOL-C HOL-C TO TO									

Figure 3 SNP effects on soluble adhesion molecule levels versus other cardiovascular health-related traits in the *ABO* locus. The Pearson's r of the genetic effects (Z-scores) were estimated using a set of SNPs that located in the *ABO* locus (defined as the LD block²⁵ containing *ABO* gene in the gwas-pw²¹ analyses) and that were available in both the present study and open-access data sets.^{22–24} Positive correlations are indicated with red color, negative correlations are indicated with blue color, and correlations with p≥0.05 are left blank. The scatter plot representations as well as correlations in the other loci are shown in online supplementary figure S3. CAD, coronary artery disease; HDL-C, high-density lipoprotein cholesterol; LD, linkage disequilibrium; LDL-C, low-density lipoprotein cholesterol; sE-selectin; soluble E-selectin; sICAM-1, soluble intercellular adhesion molecule-1; TG, total triglycerides; TotC, total cholesterol.

circulation could be influenced by a possible competitive binding of galectin-3 with sVCAM-1 and the B antigen.

To evaluate the shared genetic mechanisms, we compared the correspondence of genetic effects on inflammatory phenotypes versus cardiovascular health-related traits. We observed a negative relationship between the genetic effects on CAM levels and the genetic effects on LDL-C and TotC levels, as well as lower risk for CAD and ischaemic stroke, in the ABO locus. This denotes that the genotypes in the ABO locus associating with higher levels of soluble CAMs tended to associate with lower circulating cholesterol levels as well as lower risk of cardiovascular outcomes. This was unexpected since previous evidence suggests that increased soluble CAM levels are linked with atherosclerosis progression and vascular outcomes.^{20 33} Possible explanations unravelling the negative correlation advocate that soluble CAMs may compete with leucocyte adhesion to the endothelial molecules or that enhanced ectodomain shedding may contribute to the reduced recruitment of leucocytes to the subendothelial space, thereby promoting cardioprotective effects.³⁴ Our results suggesting a negative relationship between the genetic effects on soluble CAM and circulating cholesterol levels advocate that altered cholesterol metabolism could contribute to the CAD risk associated with the ABO locus; the genetic effects of the same SNPs on LDL-C or TotC show positive correlation with CAD risk. Nevertheless, further studies are warranted to understand the exact mechanisms.

Another novel association with sVCAM-1 level was detected in chr12. The lead SNP of this locus is in LD with rs117468318 (r²=1 in NFBC1966) that locates in the 5'UTR of *HSP90B1* encoding heat shock protein gp96 and, according to RegulomeDB,²⁸ is likely to affect transcription factor binding, suggesting a possible regulatory mechanism for the detected association. HSP90B1/gp96 is a chaperone that is essential for assembly of 14 of 17 integrin pairs in the haematopoietic system.³⁵ Integrin α 4 β 1 is an important ligand of VCAM-1; if altered transcription of *HSP90B1* had a downstream effect on integrin α 4 β 1 level, this could further modify the level of unbound sVCAM-1 in circulation.

The third novel locus showing association with sVCAM-1 level was identified in chr17 near ABCA8. The lead SNP rs112001035 is an expression quantitative trait locus (eQTL) for ABCA8 in multiple tissue types.³⁶ If ABCA8 is involved in the regulation of HDL level via interaction with ABCA1³⁷ and if plasma HDL levels contribute to VCAM-1 expression,³⁸ then altered expression of the ABCA8 could influence circulating levels of sVCAM-1 by modulating HDL particle concentration. However, this hypothesis is not supported by the fact that the effect of the lead SNP on HDL particle concentration is negligible in a metabolomics GWAS $(\beta = -0.043 \text{ SD}, p = 0.049)$.³⁹ There is evidence suggesting that ABCA8 may be involved in sphingolipid metabolism,⁴⁰ and it has been hypothesised that ABCA8 may be involved in the formation of specific membrane domains during ApoA-I lipidation.³⁷ Thus, the association between the ABCA8 and sVCAM-1 level could be related to altered HDL composition possibly contributing to endothelial homeostasis rather than absolute particle concentration. However, more evidence is needed to draw conclusions.

We detected a novel effect of rs11220471 in chr11 near *ST3GAL4* on sE-selectin levels in the NFBC1966 population. *ST3GAL4* encodes a member of the glycosyltransferase 29 family of enzymes involved in protein glycosylation. In mice, St3Gal4 is needed for synthesis of functional selectin ligands.²⁹ The altered levels or structure of selectin ligands due to variation in *ST3GAL4* could contribute to the levels of unbound sE-selectin in circulation, providing a biologically rational mechanism for the detected association.

In the meta-analyses, we detected a novel large-effect locus for VEGF in chr4 (β =-2.38 SD) near *STK32B*. Mutations in this locus have been associated previously with coeliac disease,⁴¹ CAD⁴² and Ellis-van Creveld syndrome.⁴³ *STK32B* may play a role in the hedgehog signalling pathway, which has been implicated in metastasis and angiogenesis in cancer⁴⁴ and downregulated in coeliac disease.⁴⁵ The hedgehog signalling has shown to be involved in the regulation of VEGF expression during developmental angiogenesis in avian embryo.⁴⁶ Thus, previous literature and our results advocate that *STK32B* may be involved in the regulation of VEGF levels possibly via hedgehog signalling-related mechanism.

The other novel findings obtained in meta-analysis include a large-effect locus on TNF α level in chr13 (β =2.13 SD). The locus in 13q14.3 associating with TNF α locates near *DLEU1* and *DLEU7*. This region is recurrently deleted in tumours and haematopoietic malignancies.⁴⁷ *DLEU1* is a part of a transcriptionally coregulated gene cluster that modulates the activity of the nuclear factor kappa B (NF-kB) pathway,⁴⁸ which is also modulated by TNF α .⁴⁹ It is largely unknown how the *DLEU1* and related *DLEU2* regulate NF-kB activity⁵⁰; our result suggests that TNF α signalling might be involved in this mechanism.

At last, we identified a small-effect locus in chr6 harbouring two independent association signals on IL1 β and showing suggestive association also on IL4 level. This association signal is in the region coding the human leucocyte antigen proteins, and

further experimental evidence would be needed to identify the exact mechanism how the locus contributes to IL levels.

The strengths and limitations of our study should be considered. The sample size of the present study should provide adequate power for detecting genetic associations with circulating markers of systemic inflammation.⁸ The use of genetically isolated populations, such as inhabitants of Northern Finland, should further enhance the power for locus identification in GWAS settings. We were able to perform meta-analyses only for 10 out of the total of 16 inflammatory phenotypes, and the novel findings are largely based on NFBC1966 population only. Thus, replication of the present findings in other populations would be helpful. In particular, the associations of the novel rare, large-effect variants need to be interpreted with caution until the associations are validated in other populations. The interassay coefficient of variability measures for sE-selectin and VEGF in particular are notably larger than 15%, which is considered to be the limit for acceptable values (online supplementary table S1). However, to our consideration, all the findings identified in the present study locate on genome regions with biologically relevant genes. Furthermore, the replications of the previously reported loci speak for the data adequacy and add confidence to the novel associations. Finally, as we have not included functional experiments in this work, we are limited to previous literature when explaining the potential biological mechanism behind the identified associations.

The present results provide novel information on genetic mechanisms influencing levels of inflammatory phenotypes in circulation. The evident role of the *ABO* locus in the regulation of the soluble CAM levels likely encompasses at least two distinct mechanisms influencing sE-selectin, sICAM-1 and sVCAM-1 levels. Our findings provide evidence that increased soluble CAM concentrations per se may not be a risk factor for cardiovascular outcomes. In particular, genetic variation associating with increased sE-selectin or sICAM-1 levels at the *ABO* locus seems to contribute to lower cardiovascular risk. Furthermore, genetic effects at the *ICAM1* locus providing a direct molecular link to sICAM-1 concentration do not correlate with the genetic effects on CAD risk nor stroke risk. Overall, the present study extends the knowledge about the molecular pathways involved in inflammatory load.

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Contributors ES, SS and JK conceptualised the study. ES, MK and AA-O performed the statistical analyses. TK, K-HH, MS, KS and SJ performed the quantifications of the inflammatory markers. OR, MP, VS, TL, HV, MS, SJ, JJ, SK-K, MM, K-HH, M-RJ, SS and JK provided funding or other resources to conduct the study. SS and JK supervised the study. ES, SS and JK wrote the original manuscript. All authors contributed to revising the content and approved the final version.

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