Polymorphisms in $B3GAT1$, $SLC9A9$ and $MGAT5$ are associated with variation within the human plasma $N$-glycome of 3533 European adults

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The majority of human proteins are post-translationally modified by covalent addition of one or more complex oligosaccharides (glycans). Alterations in glycosylation processing are associated with numerous diseases and glycans are attracting increasing attention both as disease biomarkers and as targets for novel therapeutic approaches. Using a recently developed high-throughput high-performance liquid chromatography (HPLC) analysis method, we have reported, in a pilot genome-wide association study of 13 glycan features in 2705 individuals from three European populations, that polymorphisms at three loci ($FUT8$, $FUT6$/ $FUT3$ and $HNF1A$) affect plasma levels of $N$-glycans. Here, we extended the analysis to 33 directly measured and 13 derived glycosylation traits in 3533 individuals and identified three novel gene association ($MGAT5$, $B3GAT1$ and $SLC9A9$) as well as replicated the previous findings using an additional European cohort. $MGAT5$ (meta-analysis association $P$-value $= 1.80 \times 10^{-12}$ for rs1257220) encodes a glycosyltransferase which is known to synthesize the associated glycans. In contrast, neither $B3GAT1$ (rs7928758, $P = 1.66 \times 10^{-08}$) nor $SLC9A9$ (rs4839604, $P = 3.50 \times 10^{-13}$) had previously been associated functionally with glycosylation of plasma proteins. Given the glucuronyl transferase activity of B3GAT1, we were able to show that glucuronic acid is present on antennae of plasma glycoproteins underlying the corresponding HPLC peak. $SLC9A9$ encodes a proton pump which affects pH in the endosomal compartment and it was recently reported that changes in Golgi pH can impair protein sialylation, giving a possible mechanism for the observed association.

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INTRODUCTION

Glycosylation is a post-translational modification that influences protein and lipid structural complexity and function. It is essential for multicellular life and its complete absence is embryonic lethal (1). Genetic defects that affect glycosylation biosynthetic or degradation pathways have been associated with a wide range of congenital disorders (2,3). Multiple proteins are modified by covalently bound glycans, which has implications for protein folding, degradation, cell signalling, secretion, immune function and transcription (4,5). Misregulation of glycosylation has been associated with a wide spectrum of common diseases including cancer, diabetes, cardiovascular, immunological and infectious disorders (4,6,7). Terminal variability in glycans is common (e.g. ABO blood groups) and helps to create diversity that allows our bodies to evade pathogens and adapt to changing environments (8). Glycosylation is not directly template-driven, which gives rise to the extensive complexity of the glycoproteome, estimated to be several orders of magnitude more complex than the proteome itself (9).

Owing to experimental limitations in quantifying glycans in biological samples, our understanding of the genetic regulation of glycosylation is still very limited (2). Recent technological advances have allowed reliable, high-throughput quantification of N-glycans (10), which now allows investigation into the genetic regulation and biological role of these glycan structures and brings glycomics into line with genomics, proteomics and metabolomics (11). In contrast to other analytical techniques, high-performance liquid chromatography (HPLC) separates glycans into chromatographic peaks which contain mostly similar structures (Fig. 1). Therefore, even without complete separation of all glycan structures, many glycome features of an individual sample can be deduced from ratios of different HPLC peaks (12).

An initial pilot study was performed using three cross-sectional population samples [CROATIA-Vis, CROATIA-Korcula and ORCADES (The Orkney Complex Disease Study)] on a subset of directly measured [desialylated glycans] measured (13) and derived glycan species [antennary fucosylated glycans (FUC-A), core fucosylated glycans (FUC-C)] to investigate whether a genome-wide association study (GWAS) approach could be applied to the identification of genetic variants affecting N-glycans in human plasma. Single-nucleotide polymorphisms (SNPs) located within fucosyltransferase 8 (FUT8), fucosyltransferase 6 (FUT6) and hepatic nuclear factor 1 alpha (HNF1a) genes were found to be significantly associated with levels of several glycan groups (13).

In this study, a similar analysis was performed to include additional glycan species and an additional population sample (the Northern Swedish Population Health Study: NSPHS). We were able to strengthen our previous results as well as describe additional associations of these genes with new glycan traits and novel associations with mannosyl (α-1,6-)glycoprotein β-1,6-N-acetyl-glucosaminyltransferase V (MGAT5), β-1,3-glucuronyltransferase 1 (B3GAT1), and solute carrier family 9, member 9 (SLC9A9).

RESULTS

A meta-analysis of genome-wide association results was conducted for 46 plasma N-glycan traits (Table 1) measured in four population-based samples, CROATIA-Vis, CROATIA-Korcula, ORCADES and NSPHS. Sex-specific analyses were also performed for each trait, and descriptive statistics and heritabilities for all traits are presented in Supplementary Material, Table S1. Heritability of traits (adjusted for sex and age) was variable between populations, as might be expected given the isolated nature of the populations. However, all traits had at least one population displaying a heritability >0.2, with most having at least one population in which heritability was >0.4. DG10 showed a fairly consistent heritability across populations (h² = 0.31–0.40), whereas GP10 was much more variable (h² = 0.21–0.65). Meta-analysis summary data for each significantly associated gene region are presented in Table 1. Summary data for all genome-wide significant SNPs are presented in Supplementary Material, Tables S2 (all participants), S3 (women only) and S4 (men only).

The most statistically significant association was found with a pair of SNPs located on chromosome 19. rs3760776 and rs778885 were both significantly associated with DG7, DG9, GP14, and only rs3760776 reached genome-wide significance with DG12 and A-FUC. The association of rs3760776 with DG7, DG9, DG12 and A-FUC was reported in the previous study, and the addition of another population sample improved the association P-value for each trait (13). The most significant association was found with DG9 for SNP rs3760776 (P = 3.2 × 10⁻²⁰), located at the 5' end of the fucosyltransferase 6 (FUT6, Entrez GeneID: 2528). The effect size of the G allele of rs3760776 was the largest of all traits, 0.44 (SE 0.04) for DG9 [z-score units, after adjustment for sex, age and principal components (PC)]. The association region for this SNP contains the NRTN, FUT6 and FUT3 genes between two recombination hotspots (Supplementary Material, Fig. S1). FUT6 encodes the enzyme fucosyltransferase VI, which was reported to be the key enzyme responsible for the antennary fucosylation of plasma proteins (14), thus the causal variant(s) probably affect this gene. SNP rs3760776 explained 1.1, 4.8, 5.7 and 3.0% of the variance of DG9 (adjusted for sex, age and PC) in CROATIA-Vis, CROATIA-Korcula, ORCADES and NSPHS, respectively.

An association peak containing 22 SNPs was located in the region encompassing the FUT8 (Entrez GeneID: 2530) gene on chromosome 14 (Supplementary Material, Fig. S2). We previously reported associations of this gene with plasma concentrations of DG1 and DG6. The most significant signal for DG1 was still located 5' of the gene with the addition of the NSPHS population sample; however, the SNP showing the strongest association changed from rs7159888 in the previous study to rs11621121 whose P-value decreased from 1.0 × 10⁻¹⁷ to 1.7 × 10⁻²³. The association with DG6 produced the same top SNP (rsi0483776), with its P-value decreasing from 9.6 × 10⁻⁰⁶ to 9.3 × 10⁻¹⁰. SNPs in this region were also associated with DG10, GP1, GP10, C-FUC and A2 in a manner consistent with the biological role of FUT8. The SNP rs11621121 explained 3.0, 5.9, 1.0 and 2.8% of the variance of DG1.
Five SNPs on chromosome 12, the most significant being rs735396, showed genome-wide significant associations with DG7 ($P = 7.8 \times 10^{-12}$). SNPs within this region were also associated with GP13 ($P = 3.8 \times 10^{-10}$), GP15 ($P = 4.3 \times 10^{-10}$), DG9 ($P = 3.2 \times 10^{-11}$), DG11 ($P = 4.2 \times 10^{-10}$) and FUC-A ($P = 5.0 \times 10^{-10}$). Associations of DG7 and DG11 with SNPs in this region were previously reported but the significance of the association has increased with the addition of NSPHS, and the number of SNPs with significant association has also increased (13). All SNPs are located within or 5′ of the HNF1 homeobox A (HNF1a, Entrez GeneID: 6927) gene region, with rs735396 found in intron 9. Three other genes are located between the recombination hotspots that comprise the boundaries of the association interval, C12orf27, C12orf43 and OASL (Supplementary Material, Fig. S3); however, functional studies reported in the previous paper have demonstrated that HNF1a plays a role in the regulation of several genes which play key roles in the fucosylation of glycan chains (13). SNP rs735396 explained 2.9, 0.9, 0.8 and 2.6% of the variance of sex-, age- and PC-adjusted DG7 in CROATIA-Vis, CROATIA-Korcula, ORCADES and NSPHS, respectively.

An SNP located on chromosome 2, rs1257220, was associated with DG11 and TA, a derived trait quantifying tetra-antennary glycans, ($P = 1.9 \times 10^{-10}$, $P = 1.8 \times 10^{-10}$). This SNP is located in the intron between exons 1 and 2 of the gene encoding mannosyl (α1,6)-glycoprotein β-1,6-N-acetylglucosaminyltransferase (MGAT5, Entrez GeneID: 4249). There is an area of high recombination in the middle of the MGAT5, indicating that the causal variant is most likely in the region 5′ of this hotspot (Fig. 2). The effect size of the A allele of rs1257220 is 0.19 (SE 0.03) for both DG11 and TA ($z$-score units, after adjustment for sex, age and PC). SNP rs1257220 explained 1.1, 1.6, 0.8 and 2.0% of the variance of sex-, age- and PC-adjusted TA in CROATIA-Vis, CROATIA-Korcula, ORCADES and NSPHS, respectively.

A single SNP located on chromosome 11, rs7928758, was associated with DG13 ($P = 1.7 \times 10^{-08}$). This SNP is located in the final intron of the β-1,3-glucuronosyltransferase1 (B3GAT1, Entrez GeneID: 27087) gene. There is a spike in the recombination rate towards the end of B3GAT1; therefore, the causal variant is most likely located 5′ of this spot (Fig. 3). The effect size of the C allele of rs7928758 is −0.23 (SE 0.04) ($z$-score units, after adjustment for sex, age and PC). SNP rs7928758 explained 0.2, 0.4, 2.4 and 1.6% of the variance of sex-, age- and PC-adjusted DG13 in CROATIA-Vis, CROATIA-Korcula, ORCADES and NSPHS, respectively.

Given this association, we performed a mass spectrometric analysis of the 2AB-labeled glycans of three fractions of
DG13 and confirmed the presence of glucuronic acid on large, complex-type N-glycans (Supplementary Material, Fig. S4). Two examples of the obtained fragment spectra which revealed the diagnostic ion at mass-to-charge ratio (m/z) 542 corresponding to an N-glycan antenna decorated with glucuronic acid are shown in Supplementary Material, Figure S5.

A pair of SNPs on chromosome 3 were significantly associated to tetrasialylated glycans, rs1372288 (P = 3.6 × 10⁻¹⁰⁻¹) and rs4839604 (P = 3.5 × 10⁻¹³). Both SNPs are found 3′ of the solute carrier family 9, member 9 (SLC9A9) Entrez GenID: 285195). The effect size for the minor allele for both SNPs is 0.22 (SE 0.03) (z-score units, after adjustment for sex, age and PC). All significant SNPs fall between the 3′ end of the SLC9A9 gene and just before the 3′ end of CHST2 but not within CHST2, which is oriented tail-to-tail with SLC9A9 (Fig. 4). There is a large spike in the recombination rate separating the region of association from the 3′ end of CHST2 so the causal variant most likely affects falls in the region closer to SLC9A9. SNP rs4839604 explained 3.0, 1.5, 0.7 and 2.3% of the variance of tetrasialylated glycans.

### Table 1. Genetic markers associated with plasma N-glycan levels

<table>
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<tr>
<th>Trait</th>
<th>Chr</th>
<th>Lowest P-value SNP in associated interval</th>
<th>Meta-analysis P-value</th>
<th>Meta-analysis sample size</th>
<th>Associated interval size, kb (number of significant SNPs in interval)</th>
<th>Genes within associated interval[^a]</th>
<th>Major allele, minor allele (MAF)</th>
<th>Effect size for minor allele (SE[^b])</th>
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<tr>
<td>All participants</td>
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<td>G, A (0.13)</td>
<td>-0.36 (0.04)</td>
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<td>118 (5)</td>
<td>HNF1A; C12orf43; OASL</td>
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<td>MGA-T5</td>
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<td>SLC9A9</td>
<td>G, A (0.23)</td>
<td>0.22 (0.03)</td>
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[^a]: Results are reported for NCBI Build 36 and all SNPs are on ‘+’ strand.
[^b]: z-Score units for trait adjusted for sex, age and PC.
For the top SNPs reported for the combined analysis, there were no significant differences observed for effect size between men and women (Supplementary Material, Table S5). Any discrepancy between sexes appears to be due to difference in sample size rather than difference in effect.

**DISCUSSION**

We have performed the first comprehensive genome-wide association analysis of the human plasma glycome as represented by 46 glycosylation traits analysed in the plasma of 3533 individuals from four isolated European populations. The predominance of loci involved in fucosylation displaying the strongest association with glycan levels in plasma (13) was confirmed.

FUT8 is the fucosyltransferase responsible for the $\alpha_{1,6}$-fucosylation of the core $N$-acetyl-glucosamine (GlcNAc) structure of $N$-glycans (15). Glycan groups associated with the gene region containing $FUT8$ contain fucose attached to their core, so the results are consistent with the known biological role of FUT8. In contrast, groups DG7, DG9 and DG12 include glycans containing antennary fucosylation, and FUC-A was derived as an overall measure of antennary fucosylation on biantennary glycans. $FUT6$ encodes the enzyme fucosyltransferase VI, which was reported to be the key enzyme responsible for the $\alpha_{3}$-fucosylation of plasma proteins (14). The association of $FUT8$, $FUT6$ and $FUT3$ with $N$-glycan structures containing core and antennary fucosylation is supported by their known biological functions (16).
various fucosyltransferases act to add fucose in a generic N-glycan is presented in Figure 5.

In our previous study, HNF1a was associated with several glycan traits but the mechanism for this association was not obvious. Functional studies showed that HNF1a and HNF4a act to co-regulate the expression of most fucosyltransferase (FUT3–11) genes in HepG2 cells as well as gene expression levels of key enzymes needed for synthesis of GDP-fucose, the substrate for fucosyltransferases, thereby regulating both core and antennary fucosylation (13). Rare mutations within this gene cause Maturity Onset Diabetes of the Young 3 [MODY3 (MIM:600496)]. Common polymorphisms in HNF1a have been associated with variations of the plasma levels of C-reactive protein (17–22), low-density lipoprotein (23) and gamma-glutamyl transferase (24), and found to be susceptibility loci for type 2 diabetes (25,26), coronary heart disease (27) and pancreatic cancer (28). Although none of these associations reported the same top SNP as our study, none could be discounted from tagging the same region. There is strong support for the gamma-glutamyltransferase, coronary heart disease and C-reactive protein loci to also influence N-glycan levels with the reported associated SNPs in complete linkage disequilibrium (LD) with our top SNP ($r^2 = 1, D' = 1$). The top SNP in the Voight et al. (25) type 2 diabetes study (rs7957197, $r^2 = 0.102, D' = 1$) is most likely not tagging the same pleiotropic signal but due to the very high $D'$ value this cannot be completely ruled out without further investigation.

Three novel non-fucosylation-related associations were identified in this expanded analysis. The MGAT5 gene codes for the enzyme mannosyl (α1-6)-glycoprotein β1,6-N-acetyl-glucosaminyltransferase V (GnT-V), which adds GlcNAc residues to mannose in a β1,6 orientation on the antennary structure of N-glycans (Fig. 5). This is a decisive step
in the generation of tetra-antennary glycans. The majority of structures contained within the DG11 peak are tetra-antennary glycans (29); therefore, the associations with DG11 and TA (a derived trait quantifying total tetra-antennary glycans) are biologically plausible. Tetra-antennary glycans are important regulators of membrane function (30) since they affect the half-life of numerous receptors on the cell membrane (31). This has important implications for many dynamic processes, from immunity to cancer progression and metastasis. MGAT5 synthesizes cell-surface ligands for galectins—proteins involved in the proliferation of T-cells and apoptosis. Loss of MGAT5 expression lowers the threshold needed for T-cell activation, and as Mgat5-deficient mice displayed several autoimmune phenotypes it was postulated that MGAT5 expression might be implicated in autoimmune disorders in humans (32). Recently, polymorphisms within MGAT5 have been tentatively associated with the severity of multiple sclerosis (33,34) in two small studies which may support this theory; however, the SNPs reported in these studies do not appear to be tagging the same LD blocks as our glycan-associated SNPs. Expression of MGAT5 is upregulated in oncogenic cells, and an increased production of galectin ligands on the cell surface allows the tumour cell to retain growth factors such as epidermal growth factor and transforming growth factor-β (35).

B3GAT1 is a member of the glucuronyltransferase gene family. This gene product functions as the key enzyme in a glucuronyl transfer reaction during the biosynthesis of the carbohydrate epitope HNK-1. It acts to add a glucuronic acid (GlcA) to the terminal N-acetyllactosamine (Lac) disaccharide to form the HNK-1 epitope precursor (36,37). The HNK-1 epitope is expressed on a subset of human

Figure 4. Significance plot (A) and forest plot (B) for the most significant SNP for regions of interest from the meta-analysis of tetrasialylated glycans. (A) –Log 10 of the P-values are plotted against chromosome positions. The most significant SNP is labelled along with its P-value and information on SNP function (see annotation legend in the upper right corner). Gene information and regional LD (r²) are also shown. (B) The estimate of the effect size of the best SNP for each individual population and the pool along with its standard error is shown, in which the size of the square for individual cohorts represents the proportion that it is contributing to the estimate of pooled effect size. The effect size presented is the β-coefficient, which represents a change in glycan levels measured in standard deviation units (adjusted for age, sex and PC) per copy of the allele modelled.
lymphocytes, including natural killer cells, but it was not previously reported to exist on plasma proteins. We were able to show through mass spectrometry (MS) analysis that glucuronic acid is present on some glycans which make up the DG13 plasma glycan pool, explaining this association.

SLC9A9, solute carrier family 9 (sodium/hydrogen exchanger), is a proton pump which affects pH in the endosomal compartment (38). This gene was not previously linked to glycosylation, but it was recently reported that changes in Golgi pH can impair protein sialylation (39), thus the association between SLC9A9 and tetrasialylated glycans makes biological sense. Sialic acids are found in cell secretions and are usually the terminal component of glycoproteins and glycolipids on the outer cell surface and therefore are involved in cell communication and defence. They act to shield recognition sites that may be antigenic in order to prevent autoimmunity but also function as ligands for many molecules such as hormones, inorganic cations and antibodies (40). Microorganisms are able to exploit the prominent role of sialic acids within the human body by either coating themselves with or binding to sialic acid in order to penetrate and infect the cell. A better understanding of processes and pathways underlying sialylation could lead to new avenues for the treatment of infection and disease (41).

Polymorphisms in SLC9A9 have recently been suggestively associated with attention-deficit hyperactive disorder [ADHD (MIM:143465)] (42,43). The first study reported only a gene-based P-value so it cannot be determined if the variant underlying the reported signal could be the same as for the glycosylation signal, whereas in the latter study the reported SNP tagging the disease signal is not in strong LD with our top SNP (rs1372288 and rs9810857, r² = 0.063, D' = 0.575; rs4839604 and rs9810857, r² = 0.106, D' = 0.727). Glycosylation analysis in ADHD patients also revealed differences in tetrasialylated glycans between children with ADHD and matching controls (44). Although disease causality cannot be ascertained by these data, the associations still provide a novel set of molecules which could act as clinical markers of disease.

Recent advances in high-throughput methods of analysing N-glycans have now made it possible to measure these traits in large cohorts. Analysing these traits with GWAS reveals several genome-wide significant associations illuminating the genetic control of distinct biological pathways, including fucosylation, sialylation and glucuronyl transfer, even in a modest sample size. Some of these biological processes are known to be altered in several disease states. For example, fucosylation of acute phase proteins is modified in many diseases, such as acute inflammation (45,46), rheumatoid arthritis (47) and diabetes (48), and changes in the levels of fucosylated glycans have been shown to be associated with several important pathological processes, including cancer and inflammation.

Figure 5. Pathways of core (tri-mannosyl-chitobiose) and antennary fucosylation. Genes highlighted indicate genome-wide significant genes identified from GWAS (FUT6/FUT3, FUT8 and MGAT5). Structures are given using Oxford notation (62). Asn, asparagine residue on glycosylated protein.
between the ages of 18 and 98 (52). The field work was performed in 2007 and 2008 in the eastern part of the island, targeting healthy volunteers from the town of Korčula and the villages of Lumbarda, Zrnovo and Račišće. The CROATIA-Korčula study includes 969 Croatians between the ages of 18 and 98 (52). The field work was performed in 2007 and 2008 in the eastern part of the island, targeting healthy volunteers from the town of Korčula and the villages of Lumbarda, Zmovo and Račišće.

The CROATIA-Vis study includes 1008 Croatians, aged 18–93 years, who were recruited from the villages of Vis and Komiza on the Dalmatian Island of Vis during 2003 and 2004 within a larger genetic epidemiology program (50,51).

The CROATIA-Korčula study includes 969 Croatians between the ages of 18 and 98 (52). The field work was performed in 2007 and 2008 in the eastern part of the island, targeting healthy volunteers from the town of Korčula and the villages of Lumbarda, Zmovo and Račišće.

ORCADES was performed in the Scottish archipelago of Orkney, which collected data between 2005 and 2011 (53). Data for 889 participants aged 18–100 years from a subgroup of 10 islands were used for this analysis.

NSPHS is a family-based population study including a comprehensive health investigation and collection of data on family structure, lifestyle, diet, medical history and samples for laboratory analyses from peoples living in the north of Sweden (54). Data were available from 700 participants aged 14–91 years.

DNA samples were genotyped according to the manufacturer's instructions on Illumina Infinium SNP bead microarrays (HumanHap300v1 for CROATIA-Vis, HumanHap300v2 for ORCADES and NSPHS and HumanCNV370v1 for CROATIA-Korčula). Genotypes were determined using the Illumina BeadStudio software. Genotyping was successfully completed on 991 individuals from CROATIA-Vis, 953 from CROATIA-Korčula, 889 from ORCADES and 656 from NSPHS. The number of participants available for meta-analysis prior to phenotype quality control was 3367 (1941 women, 1426 men).

Extreme outliers (those with values more than three times the interquartile distances away from either the 75th or the 25th percentile values) were removed for each glycan measure to eliminate individuals not representative of normal variation within the population.

**Genotype and phenotype quality control**

Genotyping quality control was performed using the same procedures for all cohorts. Individuals with a call rate <97% were removed as well as SNPs with a call rate <98% (95% for CROATIA-Vis), minor allele frequency <0.02 or Hardy–Weinberg equilibrium P-value less than $1 \times 10^{-10}$. In total, 924 individuals passed all quality control thresholds from CROATIA-Vis, 898 from CROATIA-Korčula, 889 from ORCADES and 656 from NSPHS. The number of participants available for meta-analysis prior to phenotype quality control was 3367 (1941 women, 1426 men).

**Genome-wide association analysis**

Genome-wide association analysis was performed for each population and then combined using an inverse-variance weighted meta-analysis for all traits. Each trait was adjusted for sex, age and the first three PC obtained from the population-specific identity-by-state (IBS)-derived distance matrix (described below) and the residuals transformed to ensure their normal distribution using quantile normalization. Sex-specific analyses were adjusted for age and PC only. The residuals expressed as z-scores were used for association analysis. The ‘mmmscore’ function of the GenABEL package (56) for the R statistical software was used for the association test under an additive model. This score test for family-based association takes into account pedigree structure and allows unbiased estimations of SNP allelic effect when relatedness is present between examinees (57). The relationship matrix used in this analysis was generated by the ‘ibs()’ function of GenABEL (using weight = ‘freq’ option), which utilizes IBS genotype sharing to estimate the realized pairwise kinship coefficient. All lambda values for the population-specific analyses were <1.05, showing that this method efficiently accounts for family structure. Inverse-variance weighted meta-analysis was performed using the MetABEL package (56) for R. An association was considered statistically significant at the genome-wide level if the $P$-value for an individual SNP was robust and reproducible across measurement centres (13,29). Chromatography yielded a total of 33 measured traits. From the HPLC analysis, undigested N-glycans were separated into 16 structurally related groups of glycans, referred to as glycan peak (GP)1–16. Desialylated 2AB-labelled N-glycans were also separated into 13 structurally related groups of glycans, referred to as DG1–DG13. WAX chromatography resulted in four peaks: monosialylated, disialylated, trisialylated and tetrasialylated structures. The amount of N-glycans measured in each of these groups was quantified relatively, as a proportion of the total plasma N-glycome. An additional 13 parameters were gained by calculating some structural determinants from measured traits. A description of each glycan trait is presented in Table 2 along with equations for the additional 13 derived traits.
less than $5 \times 10^{-8}$. The effect of the most significant SNP in each gene region expressed as a percentage of the variance explained was calculated for each glycan trait adjusted for sex, age and first three PC in each cohort individually using the 'polygenic' function of the GenABEL package for R.

**Result interpretation**

Regions of association were visualized using the web-based software LocusZoom (58) to display the LD of the region. A preliminary literature search for each associated gene was performed using GWAS Catalog (59) for associations reaching genome-wide significance within the study or reasonably suggestive if sample size was large ($\geq 10,000$). Additional associations were reported if they were replicated by another population either within the original publication or in another study. SNAP (60) was used to look at LD between SNP associations from the literature and this study. 1000 Genomes Pilot 1 data from the CEU population was used to calculate LD, and SNPs were accepted to be tagging the same region if $D’ = 1$ and $r^2 > 0.8$.

**Mass spectrometry**

Nano-LC-ESI-MS/MS. MS analysis of the collected glycan fractions was performed using an Ultimate 3000 nano-LC system (Dionex/LC Packings, Amsterdam, The Netherlands) equipped with a reverse-phase trap column ($C_{18}$ PepMap 100 Å, 5 μm, 300 μm × 5 mm; Dionex/LC Packings) and a nano column ($C_{18}$ PepMap 100 Å, 3 μm, 75 μm × 150 mm; Dionex/LC Packings). The column was equilibrated at room temperature with eluent A (0.1% formic acid in water) at a pH of 8.0.
flow rate of 300 nl/min. After injection of the samples, a gradient was applied to 25% eluent B (95% acetonitrile) in 15 min and to 70% eluent B at 25 min, followed by an isocratic elution with 70% eluent B for 5 min. The eluate was monitored by UV absorption at 214 nm. The LC system was coupled via an online nanospray source to an Esquire HCTultra ESI-IT-MS (Bruker Daltonics, Bremen, Germany) operated in the positive ion mode. For electrospray (1100–1250 V), stainless steel capillaries with an inner diameter of 30 μm (Proxeon, Odense, Denmark) were used. The solvent was evaporated at 170°C, employing a nitrogen stream of 71/min. Ions from m/z 500 to 1800 were recorded. Automatic fragment ion analysis was enabled, resulting in MS/MS spectra of the most abundant ions in the MS spectra. Glycan structures were assigned using GlycoWorkbench (61).

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

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REFERENCES

42. Brookes, K., Xu, X., Chen, W., Zhou, K., Neale, B., Lowe, N., Anney, R.,
38. Roxrud, I., Raiborg, C., Gilfillan, G.D., Stromme, P. and Stenmark, H.
24. Yuan, X., Waterworth, D., Perry, J.R.B., Lim, N., Song, K., Chambers,
220
520–528.
54. Kolz, M., Johnson, T., Sanna, S., Teumer, A., Vitart, V., Perola, M.,
58. Pruim, R.J., Welch, R.P., Sanna, S., Teslovich, T.M., Chines, P.S., Gliedt,
220
43. Mick, E., Todorov, A., Smalley, S., Hu, X., Loo, S., Todd, R.D.,
44. Pivac, N., Knezevic, A., Gornik, O., Pucic, M., Iglw, W., Peeters, H.,
45. Mick, E., Todorov, A., Smalley, S., Hu, X., Loo, S., Todd, R.D.,
51. Vitart, V., Rudan, I., Hayward, C., Gray, N.K., Floyd, J., Palmer, C.N.,
52. Kolz, M., Johnson, T., Sanna, S., Teumer, A., Vitart, V., Perola, M.,
53. McQuillan, R., Leutenegger, A.L., Abdel-Rahman, R., Franklin, C.S.,
54. Mick, E., Todorov, A., Smalley, S., Hu, X., Loo, S., Todd, R.D.,
58. Pruim, R.J., Welch, R.P., Sanna, S., Teslovich, T.M., Chines, P.S., Gliedt,
60. Johnson, A.D., Handsaker, R.E., Pulit, S.L., Nizzari, M.M., O’Donnell,
64. Pruim, R.J., Welch, R.P., Sanna, S., Teslovich, T.M., Chines, P.S., Gliedt,
66. Roxxrud, I., Raiborg, C., Gilfillan, G.D., Strompe, P. and Stenmark, H.