ASSOCIATION STUDIES ARTICLE

Defining the genetic control of human blood plasma N-glycome using genome-wide association study

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Abstract

Glycosylation is a common post-translational modification of proteins. Glycosylation is associated with a number of human diseases. Defining genetic factors altering glycosylation may provide a basis for novel approaches to diagnostic and pharmaceutical applications. Here we report a genome-wide association study of the human blood plasma N-glycome composition in up to 3811 people measured by Ultra Performance Liquid Chromatography (UPLC) technology. Starting with the 36 original traits measured by UPLC, we computed an additional 77 derived traits leading to a total of 113 glycan traits. We studied associations between these traits and genetic polymorphisms located on human autosomes. We discovered replicated 12 loci. This allowed us to demonstrate an overlap in genetic control between total plasma protein and IgG glycosylation.

Introduction

Glycosylation—addition of carbohydrates to a substrate—is a common and structurally diverse cotranslational and posttranslational modification of proteins that can affect their physical properties (solubility, conformation, folding, stability, trafficking etc.) (1–4) as well as biological functions, including protein–protein, cell–cell, cell–matrix and host–pathogen interactions (2,3,5,6). Carbohydrate units can be attached to the protein by N- or O-glycosidic bonds with the rare exceptions of C-glycosidic attachment (7). N-glycosylation is the most abundant type of glycosylation (8,9) and, unlike other types, is specific to a consensus asparagine-containing sequence in the primary structure of the protein. Glycoproteins comprise various enzymes, hormones, cytokines, receptors, immunoglobulins, structural, adhesion and other protein molecules, and altered glycosylation is increasingly recognized to be implicated in human pathologies. In particular, association with changes in total plasma protein N-glycome composition or immunoglobulin G (IgG) glycosylation has been found for Parkinson’s disease (10), low back pain (11), rheumatoid arthritis (12), ulcerative colitis, Crohn’s disease (13) and type 2 diabetes (14). Beyond that, aberrant glycosylation is involved in key pathological steps of tumor development and is even considered a new hallmark of cancer (15–17). Glycans are considered as potential therapeutic targets (18) and biomarkers for early diagnosis and disease prognosis (19–22), which makes glycobiology a promising field for future clinical applications. An example of glycoprotein biomarker is AFF-L3—the fucosylated fraction of alpha-fetoprotein—that was approved by the U.S. Food and Drug Administration as a diagnostic marker of primary hepatocellular carcinoma (21).

Protein glycosylation is an extremely complex process depending on the interplay of multiple enzymes catalyzing glycan transfer, glycosidic linkage hydrolysis as well as glycan biosynthesis. Abundance of specific protein glycoforms can be influenced by a variety of parameters including activity of enzymes and availability of substrates, accessibility of a glycosylation site, protein synthesis and degradation. No surprise that, overall, protein glycosylation is a complex process that is controlled by genetic, epigenetic and environmental factors (23–25). Mechanisms of regulation of this process are only started to be understood. Genome-wide association studies (GWASs) can expand our knowledge on this topic by a hypothesis-free search of candidate genes involved in regulation of glycosylation. Their role can be clarified in subsequent functional follow-up studies.

Previous GWAS of total plasma protein N-glycome measured with high-performance liquid chromatography (HPLC) discovered six loci associated with protein glycosylation (26,27). Four of these loci contained genes that have well-characterized roles in glycosylation—the fucosyltransferases FUT6 and FUT8, glucuronyltransferase B3GAT1 and glucosaminyltransferase MGAT3. Other two loci—near the SLC9A9 gene on chromosomes 3 and near the HNF1A gene on chromosome 12—did not contain any genes known to be involved in glycosylation processes. A functional in vitro follow-up study in HepG2 cells showed that the HNF1A gene product acts as a co-regulator of expression of most fucosyltransferase genes (26). In addition, it co-regulates the expression of genes encoding key enzymes required for the synthesis of GDP-fucose, the substrate of these fucosyltransferases. It is noteworthy that identification of HNF1A as one of the master regulators of protein fucosylation enabled to propose a new diagnostic tool for discrimination between HNF1A-MODY monogenic diabetes and type 1 and type 2 diabetes based on the ratio of fucosylated to nonfucosylated triantennary glycans (19). The locus on chromosome 3 contains SLC9A9 gene, which encodes a proton...
pump affecting pH in the endosomal compartment, reminiscent of recent findings that changes in Golgi pH can impair protein sialylation (27).

Since 2011, when the latest GWAS of plasma N-glycome was published, new technologies for glycome profiling have been developed (28). Ultra-performance liquid chromatography (UPLC) became a widely used technology for accurate analysis of plasma N-glycosylation due to its superior sensitivity, resolution, speed and capability to provide branch-specific information of glycan structures (29). Moreover, new imputation panels (such as 1000 Genomes (30) and HRC (31)) became available, increasing the resolution and power of genetic mapping.

In this work, we aimed to advance our understanding of the genetic control of the human plasma N-glycome and to establish a public resource that will facilitate future studies linking glycosylation and complex human diseases. For that, we performed and reported results of GWAS on 113 plasma glycome traits measured by UPLC and genotypes imputed to the 1000 Genomes reference panel in 2763 participants of TwinsUK. Further, we replicated our findings in 1048 samples from three independent and genetically diverse cohorts—PainOR, SOCCS and Qatari Metabolomics Study on Diabetes (QMDiab).

Results

Replication of previously reported loci

We started with replication of six loci that were reported previously. Huffman and colleagues (27) analyzed four independent cohorts with total sample size of 3533, using plasma N-glycome measured with HPLC. Because of technological differences, there is no one-to-one correspondence between HPLC and UPLC traits and exact replication is not possible. Therefore, we analyzed association of single-nucleotide polymorphisms (SNPs) reported by (27) with all 113 UPLC traits measured in this study and considered a locus replicated if we observed P < 0.05/(6 × 30) = 2.78 × 10⁻⁴ (where 30 is a number of principal components, explaining 99% of the variation of the 113 studied traits) in the TwinsUK cohort (N = 2763). Using this procedure, we replicated five of the six previously reported SNPs (Table 1, Fig. 1). For more details, see Supplementary Material, Table 1.

These results not only confirm previous and establish five plasma glycome loci as replicated but also demonstrate that our study is well powered (among replicated loci, all P-values were less than 4 × 10⁻⁴).

Discovery and replication of new loci

In the next step, we searched for new loci via genome-wide association scan with subsequent replication of the revealed association signals.

Table 1. Replication of six previously reported loci [Huffman et al. (27)]

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<tr>
<th>SNP</th>
<th>CHR-POS</th>
<th>Gene</th>
<th>Eff/Ref</th>
<th>EAF</th>
<th>N Trait</th>
<th>BETA (SE)</th>
<th>P-value</th>
<th>EAF</th>
<th>Trait</th>
<th>BETA (SE)</th>
<th>P-value</th>
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<td>rs1257220</td>
<td>2:135015347</td>
<td>MGAT5</td>
<td>A/G</td>
<td>0.67</td>
<td>3263</td>
<td>Tetra-antennary glycan</td>
<td>0.19 (0.03)</td>
<td>1.80E-10</td>
<td>0.25</td>
<td>FASG5(3,3,3)</td>
<td>0.16 (0.032)</td>
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<td>SLCO5A</td>
<td>C/T</td>
<td>0.77</td>
<td>3320</td>
<td>Tetra-antennary glycan</td>
<td>-0.02 (0.03)</td>
<td>3.50E-13</td>
<td>0.83</td>
<td>FRS2/FST2 + FRS2</td>
<td>-0.11 (0.038)</td>
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<td>rs7928758</td>
<td>11:134265967</td>
<td>B3GAT1</td>
<td>T/G</td>
<td>0.88</td>
<td>3233</td>
<td>A4G2G4 (DG13)</td>
<td>0.23 (0.04)</td>
<td>1.66E-08</td>
<td>0.84</td>
<td>A4G13S,62</td>
<td>0.24 (0.038)</td>
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<td>HNF1A</td>
<td>T/C</td>
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<td>3236</td>
<td>A2FG2G2 (DG7)</td>
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<td>0.18 (0.031)</td>
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<td>FUT8</td>
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<td>3234</td>
<td>A2 (DG1)</td>
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<td>1.69E-23</td>
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<td>G/A</td>
<td>0.87</td>
<td>3262</td>
<td>A3F3G3 (DG9)</td>
<td>0.44 (0.04)</td>
<td>3.18E-29</td>
<td>0.91</td>
<td>A4G4S,3,3,3</td>
<td>0.56 (0.050)</td>
</tr>
</tbody>
</table>

Replicated loci are in bold. CHR-POS—chromosome and position of SNP according to GRCh37 human genome build; Eff/Ref—effective and reference alleles; gene—candidate gene for the locus reported in Huffman et al. (27); EAF—effective allele frequency; N—sample size; Trait—glycan trait with statistically strongest association with the SNP; DG (desialylated peak)—HPLC peak after sialidase treatment; BETA (SE)—beta coefficient (in SD units) and standard error of effect; P-value—P-value of association.

The discovery cohort comprised 2763 participants of the TwinsUK study with genotypes available for 8557543 SNPs. The genomic control inflation factor varied from 0.99 to 1.02, suggesting that influences of residual population stratification on the test statistics were small (see Supplementary Material, Table 2; QQ-plots in Supplementary Fig. 1). In total, 906 SNPs located in 14 loci were significantly associated (P < 5 × 10⁻⁴/30 = 1.66 × 10⁻⁴, where 30 is a number of principal components, explaining 99% of the variation of the 113 studied traits) with at least one of 113 glycan traits (in total 5052 SNP-trait associations, see Fig. 2, Table 1). Out of 113 traits, 68 were significantly associated with at least one of the 14 loci. For more details, see Supplementary Material, Table 3.

Among 14 loci, four were previously reported as associated with the plasma N-glycome (Fig. 1). Three loci—on chromosome 12 at 121 Mb (leading SNP: rs1169303, intronic variant of the HNF1A gene), on chromosome 14 at 105 Mb (leading SNP: rs7147636 located in the intron of the FUT8 gene) and on chromosome 19 at 58 Mb (leading SNP: rs7255720, upstream variant of the FUT6 gene)—were replicated to be associated with the plasma N-glycome in two previous GWAS (26,27), while association of the locus on chromosome 11 at 126 Mb (leading SNP: rs1169303, intronic variant of the HNF1A gene) on chromosome 19 at 58 Mb (leading SNP: rs7255720, upstream variant of the FUT6 gene)—were reported to be associated with the plasma N-glycome in two previous GWAS (26,27), while association of the locus on chromosome 11 at 126 Mb (leading SNP: rs1866767 located in the intron of B3GAT1 gene) was reported only in the latest GWAS meta-analysis of plasma N-glycome (27).

Ten further loci that have not been reported before were found here. In order to replicate our findings, we have performed association analysis of these 10 SNPs in three independent cohorts—PainOR, SOCCS and QMDiab.

Functional annotation in silico

For 12 replicated loci, we performed an in silico functional annotation in order to prioritize potentially causal genes. Prioritization used multiple lines of evidence, such as presence of predicted damaging variants in a gene, pleiotropic effects of glycan-associated SNPs on gene expression and the results of a DEPICT (Data-driven Expression Prioritized Integration for Complex Traits) analysis, which employs predicted gene functions and reconstituted gene sets (32). Although most of the observed loci contain genes encoding proteins with a known role in glycosylation (FUT3/FUT6, FUT8, B3GAT1, ST6GAL1, B4GALT1, ST3GAL4, MGAT3, MGAT5 and HNF1A), we still included them in the
Figure 1. Schematic overview of the novel and known loci associated with plasma protein N-glycosylation. Chr—chromosome. Genes, prioritized in this study, represent corresponding loci. *The role of MAN1C1 (mannosidase alpha class 1C member 1) gene in glycosylation is known, but the association of MAN1C1 locus was not replicated in the present study.

Figure 2. Manhattan plot of discovery GWAS (after correction for genomic control). Horizontal line corresponds to the genome-wide significance threshold of $1.7 \times 10^{-9}$. For each SNP the lowest $P$-value among 113 traits is shown. Only SNPs with $P < 1 \times 10^{-5}$ are shown. Points with $-\log_{10}(P) > 20$ are depicted at $-\log_{10}$ ($P$-value) = 20. Gene labels mark loci that were found in previous GWAS (27); underlined gene labels mark novel loci; *loci replicated either in the TwinsUK cohort (for previously reported loci) or in the replication meta-analysis of three cohorts PainOR, QMDiab and SOCCS (novel loci identified in the present study).
Next, we performed gene ontology (GO) gene-set enrichment analysis using Multi-marker Analysis of GenoMic Annotation (MAGMA) approach (37) for the loci with \( P < 1 \times 10^{-5} \) (Supplementary Material, Table 7A). We observed strong enrichment of genes involved in the glycan synthesis pathways—such as ‘glycosaccharide metabolic process’ (\( P \)-value of enrichment after correction for multiple testing, \( 1.56 \times 10^{-29} \)), ‘protein N-linked glycosylation’ (\( P \)-value, \( 4.45 \times 10^{-24} \)) and ‘N-glycan biosynthesis’ (\( P \)-value, \( 1.69 \times 10^{-20} \)). After exclusion of 14 genome-wide significant loci, the significance of enrichment of glycan-related pathways has reduced (corresponding \( P \)-values without correction for multiple testing were 0.57, 0.097 and 0.0063; see Supplementary Material, Table 7B).

Pleiotropy with expression quantitative trait loci

We next attempted to identify genes whose expression level could potentially mediate the association between SNPs and plasma N-glycome. To do this, we performed a summary data-based Mendelian randomization (SMR) analysis followed by heterogeneity in dependent instruments (HEIDI) test (38) using a collection of eQTL (expression quantitative trait loci) data for blood (39), 44 tissues provided in the GTEx database version 6p (40) and six blood cell types collected in the CEDAR study (see Supplementary Material, Table 3) for 40 samples of leukocyte types, including platelets and monocytes. In short, SMR tests the association between gene expression in a particular tissue/cell type and a trait using the top associated SNP as a genetic instrument. Significant SMR test may indicate that the same functional variant influences both expression and the trait of interest (causality or pleiotropy) but may also indicate that functional variants underlying gene expression are in linkage disequilibrium with those controlling the traits. Inferences about functional variants may be shared between plasma glycan trait and expression were made based on HEIDI test: \( F_{\text{HEIDI}} > 0.05 \) (likely shared), \( 0.05 > F_{\text{HEIDI}} < 0.001 \) (possibly shared) and \( F_{\text{HEIDI}} < 0.001 \) (sharing is unlikely).

We applied SMR/HEIDI analyses for replicated loci that demonstrated genome-significant association in our discovery data (11 loci). In total, we included in the analysis
expression levels of 20,448 transcripts (probes). For 15 probes located in 7 loci associated with plasma glycosylation, we observed significant \( P_{\text{HEIDI}} < 0.05/20448 = 2.445 \times 10^{-9} \) association with the top SNPs associated with plasma N-glycome (see Supplementary Material, Table 8). Subsequent HEIDI test showed that the hypothesis of shared functional variant between plasma glycan traits and expression was most likely showed that the hypothesis of shared functional variant (see Supplementary Material, Table 8). Subsequent HEIDI test showed that the hypothesis of shared functional variant between plasma glycan traits and expression was most likely shown to be true for four probes: ST6GAL1 in whole blood [from Westra et al. (39)], TMEM121 in whole blood [GTEx (40)], MGAT3 in CD19+ cells [CEDAR (41)] and CHCHD10 in whole blood [from Westra et al. (39)]. For the other five probes, we conclude that the functional variant is possibly shared (0.001 \( < P_{\text{HEIDI}} < 0.05 \)) between glycan traits and expression of ST3GAL4 [in two different tissues: muscle skeletal and pancreas; GTEx (40)], B3GAT1 [in two tissues: whole blood from Westra et al. (39) and lung tissue from GTEx (40)] and SYNGR1 [in tibial nerve tissue from GTEx (40)].

**Overlap with complex traits**

We next investigated the potential pleiotropic effects of our loci on other complex traits and diseases using PhenoScanner v1.1 database (42). For 12 replicated SNPs (Tables 1 and 2), we looked up traits that were genome-wide significantly \( (P < 5 \times 10^{-8}) \) associated with the same SNP or SNP in a strong \( (r^2 > 0.7) \) linkage disequilibrium. The results are summarized in Supplementary Material, Table 9. For 8 out of 12 loci, we observed associations with a number of complex traits. Four loci (near IKZF1, FUT8, MGAT3 and DERL3) were associated with levels of glycosylation of IgG (43). Two loci (on chromosome 12 at 121 Mb and on chromosome 11 at 126 Mb, containing HNF1a and ST3GAL4 genes, respectively) were associated with LDL and total cholesterol levels (44,45). The locus-containing HNF1a was additionally associated with the level of plasma C reactive protein (46,47) and gamma glutamyl transferase (48). Locus on chromosome 22 at 39 Mb (containing MGAT3) was associated with adult height (49). Locus on chromosome 14 at 65/66 Mb (near FUT8) was associated with age at menarche (50). Note, however, that PhenoScanner analysis does not allow distinguishing between pleiotropy of a variant shared between the traits and linkage disequilibrium between different functional variants affecting separate traits.

**Summary of in-silico follow-up**

We compared the genes suggested by our in silico functional investigation with the candidate genes suggested previously for five known loci (see Table 3). For three out of five loci (B3GAT1, FUT8 and FUT6/FUT3), we selected the same genes as suggested by the authors of the previous study (27). All three genes are known to be involved in the glycan synthesis pathways. B3GAT1 gene encodes beta-1,3-glucuronoyltransferase 1. According to SMR/HEIDI analysis, the same functional variant possibly mediates the association between B3GAT1-containing loci on chromosome 11 with glycan trait as well as with the level of B3GAT1 expression in whole blood and lung tissue. The FUT8 locus was associated mostly with core-fucosylated biantennary glycans, which are known to be linked to the immunoglobulins (51). Since the FUT8 gene encodes fucosyltransferase 8, an enzyme responsible for the addition of core fucose to glycans, this gene is the most biologically plausible in this locus. Evidence for prioritization of this gene in our study was also provided by DEPICT.

FUT3 and FUT6 encode fucosyltransferases 3 and 6 that catalyze the transfer of fucose from GDP-beta-fucose to alpha-2,3 sialylated substrates. The FUT3/FUT6 locus was associated with antenatally fucosylated of tri- and tetra-antenatally sialylated glycanes, and therefore we consider these genes as good candidates. Moreover, in the FUT6 gene (chromosome 19, 58 Mb), we found the missense variant rs17855739 (substitution G \( \rightarrow \) A) that leads to the amino acid change from negatively charged glutamic acid to positively charged lysine. FATHMM-XF predicted this variant as pathogenic for transcripts of FUT6 gene. Thus, we can consider this SNP as a potentially causal functional variant. DEPICT prioritized the FUT3 gene.

For two other loci (on chromosome 2 at 135 Mb and on chromosome 12 at 121 Mb), we were not able to prioritize genes by the DEPICT and eQTL analyses. However, the first locus contained the MGAT5 gene encoding mannosyl-glycoprotein-N-acetyl glucosaminyl-transferase that is involved in the glycan synthesis pathways. The second locus contained several genes including HNF1a, which was previously shown to co-regulate the expression of most fucosyltransferase (FUT3, FUT5, FUT6, FUT8, FUT10 and FUT11) genes in a human liver cancer cell line (HepG2 cells) as well as to co-regulate expression genes encoding key enzymes needed for synthesis of GDP-fucose, the substrate for fucosyltransferases, thereby regulating multiple stages in the fucosylation process (26). Thus, we considered HNF1a as the candidate gene for this locus.

Four of the seven novel loci contain genes that are known to be involved in glycansynthesis pathways—ST6GAL1, ST3GAL4, B4GALT1 and MGAT3 (see Table 3 and Fig. 1). Moreover, SMR and HEIDI analyses have shown that expression of ST6GAL1, ST3GAL4 and MGAT3 genes may mediate the association between corresponding loci and plasma N-glycome. ST6GAL1 and ST3GAL4 genes encode sialyltransferases, enzymes that catalyze the addition of sialic acid to various glycoproteins. The locus-containing ST6GAL1 was associated with ratio of sialylated and non-sialylated galactosylated biantennary glycans. The locus containing ST3GAL4 was associated with galactosylated sialylated tri- and tetra-antenatally glycans. The locus containing MGAT3 was associated with core-fucosylation of bisected glycans, which is in line with the known effect of GfT-III (product of MGAT3 gene) on FUT8 activity (52). For this locus, we have found two possible functional variants: non-coding variant rs7423 and in-frame deletion rs149306472 that was predicted to be pathogenic for the product of the SYNGR1 gene. The SYNGR1 gene encodes an integral membrane protein associated with presynaptic vesicles in neuronal cells. Since MGAT3 has a known role in glycan biosynthesis, we choose MGAT3 as the candidate gene for this locus. The B4GALT1 gene encodes galactosyltransferase, which adds galactose during the biosynthesis of different glycoconjugates. This gene was associated with galactosylation of biantennary glycans. Thus, we observe consistency between known enzymatic activities of the products of selected candidate genes and the spectrum of glycans that are associated with corresponding loci.

The other three novel loci do not contain genes that are known to be directly involved in glycan synthesis (Fig. 1). Variant rs9624334 (chromosome 22 at 24 Mb) is located in the intron of SMARCB1 gene that is known to be important in antiviral activity, inhibition of tumor formation, neurodevelopment, cell proliferation and differentiation (53). However, gene prioritization analysis (DEPICT) showed that the possible candidate gene is DERL3, which encodes a functional component of endoplasmic reticulum (ER)-associated degradation for misfolded luminal glycoproteins (54) (see Table 3). Additionally,
This indicates that core-fucosylation of bisected glycans is under with similar glycan traits (core-fucosylation of bisected glycans). Expression in heart and liver and the lowest expression in spleen of those influenced by the monogalactosylated structures that are biochemically close to the gene that encodes immunoglobulin heavy chains. This locus was associated with sialylation of core-fucosylated biantennary glycans (43), and the authors suggested that this might be a candidate gene based on our eQTL pleiotropy analysis. Thus, we consider this locus. Interestingly, this and the second gene that encodes interferon regulatory factor 1 (IFNAR1—also overlapping with IgG glycosylation—was suggested as candidate gene for previous GWAS on HPLC plasma glycome reported the leading SNP rs9624334 of lymphocyte differentiation. Taking into account that IgG is the most abundant glycoprotein in the blood plasma (51) is secreted by B cells (57), IKZF1 seems to be a plausible candidate gene.

### Gene network regulating N-glycosylation

To identify possible clusters in the gene network of plasma protein N-glycosylation, we draw a graph in which 11 genome-wide significant loci and genome-wide significantly (P < 1.66 × 10^-8) associated glycan traits were presented as nodes and edges represent observed associations (Fig. 3). We labeled each glycan trait as ‘immunoglobulin-linked’ (Ig-linked), ‘non-immunoglobulin-linked’ (non-Ig-linked) or mixed depending on the contribution of Ig- and non-Ig-linked glycans to the trait value (Supplementary Material, Table 10), which was inferred based on the information about protein-specific glycosylation reported previously (51). For more details about the procedure of Ig/non-Ig/mixed assignment, see Supplementary Material, Note 4.

The resulting network shows that candidate genes and glycan traits cluster into two major subnetworks or hubs (Fig. 3). The first subnetwork contained the six loci: FUT8, DERL3/CHCHD10, IKZF1, TMEM121, ST6GAL1 and MGAT3, with FUT8 as a hub. These loci, as well as the loci containing B4GALT1, were associated with core-fucosylated biantennary glycans. It is known that the majority of plasma core-fucosylated biantennary glycans are linked to immunoglobulins (51). Moreover, in previous studies, these seven genes were found to be associated with N-glycosylation of IgG (43,56). At the same time, these genes were associated with non-Ig-linked glycans. We can consider this cluster (7 genes out of 11) as related to both IgG and non-IgG glycosylation. Taking into account that IgG is the most prevalent glycosylated plasma protein, it is not surprising that more than a half of replicated loci are actually associated with immunoglobulins glycosylation. However, previous GWAS on HPLC plasma N-glycome reported only one locus—FUT8—as overlapping with IgG glycosylation-associated loci.

### Table 3. Summary of in-silico functional annotation for 12 replicated loci

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<td>I, IgG</td>
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<td>+</td>
<td>I</td>
<td></td>
<td></td>
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<tr>
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<td>I</td>
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<td>Nerve</td>
<td>rs17855739</td>
<td>Whole</td>
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</table>

For each locus, we report the gene nearest to the top SNP and plausible candidate genes. CV—variant with predicted (by FATHMM-XF or FATHMM-INDEL) pathogenic impact on the gene; SMR/HEIDI—evidence for pleiotropy with expression demonstrated by SMR/HEIDI analysis; D—evidence provided by DEPICT analysis; Funct. studies—evidence from functional studies; Glycan synth.—known genes involved in glycan synthesis or its regulation are present in the locus; Prev. annot.—the region has previously been revealed in glycome GWAS, and the gene was suggested as candidate gene (Pl—gene was reported as affecting plasma protein N-glycome by Huffman et al. (27)).
Figure 3. A network view of associations between loci and glycan traits. Square nodes represent genetic loci labeled with the names of candidate gene(s), circle nodes represent glycan traits. Squares with polka dot pattern represent candidate genes, located in genomic regions that were previously found to be associated with IgG N-glycome. Grey squares represent candidate genes, located in genomic regions associated with plasma N-glycome. Grey/white squares represent candidate genes, located in genomic regions associated with plasma N-glycome, that were previously found to be associated with IgG N-glycome. Circles with polka dot pattern highlights glycan traits mostly containing glycans that are linked to immunoglobulins. Grey circles represent traits that are mostly formed by glycans linked to other (not immunoglobulin) proteins. Grey/white circles represent glycan traits, formed by a mixture of glycans that are linked to immunoglobulin and non-immunoglobulin proteins. Arrows represent genetic association ($P < 1.66 \times 10^{-9}$) between gene and specific glycan.

The second subnetwork contained four loci (ST3GAL4, HNF1A, FUT3/FUT6 and B3GAT1, with FUT3/FUT6 as a hub) associated with tri- and tetra-antennary glycans. It is known that these types of glycans are linked to plasma proteins other than IgG (51). Thus, we attributed this cluster to non-IgG plasma protein N-glycosylation. Among these four loci, we report ST3GAL4 as the novel locus controlling N-glycosylation of non-IgG plasma proteins. We attribute it to non-immunoglobulins plasma protein N-glycosylation owing to its association with tetra-antennary glycans.

Discussion
We conducted the first GWAS of total plasma N-glycome measured by UPLC technology. Our efforts brought the number of loci significantly associated with total plasma N-glycome from 6 (26,27) to 16, of which 12 were replicated in our study. This allowed us to use a range of in silico functional genomics analyses and identify candidate genes in the established loci.

Compared to the HPLC glycan measurement technology used in previous GWAS of plasma N-glycome (26,27), UPLC technology provides better resolution and quantification of glycan
structures, resulting in increased power of association testing. Thus, despite the reduced sample size in our study (2763 samples here vs. 3533 samples in the study reported by Huffman et al. (27)), we have detected 14 vs. 6 plasma N-glycome QTLs. In particular, we revealed and replicated novel loci containing ST3GAL4 and ST6GAL1. An interesting question is why these two strongly associated loci were not detected in previous HPLC-based GWAS. In our study, ST3GAL4 locus showed the strongest association with FGPI7 peak (which corresponds to A2G2S[3,6,3] trait; \(P = 8.6 \times 10^{-29}\)). On an average UPLC chromatogram, a nearby FGPI9 peak has 8× and 28× bigger area than FGPI7 and FGPI8 peaks, respectively. On the HPLC chromatogram, these three peaks are merged into one FG9 peak. Thus, FG9 HPLC peak is mostly formed by FGPI9. In our study, we revealed association of ST3GAL4 locus neither with FGPI8 (\(P = 6.14 \times 10^{-1}\)) nor with FGPI9 (\(P = 8.25 \times 10^{-6}\)) peaks. We can therefore assume that association in the previous study (27) was not detected due to a small signal-to-noise ratio. We suggest the same reasoning for the ST6GAL1 locus. In our study it was associated with FGPI3 peak (\(P = 3.12 \times 10^{-22}\)), which together with FGPI2 forms GP6 HPLC peak. ST6GAL1 locus showed association neither with GP6 in Huffman et al. study (\(P = 0.012\)) (27) nor with GP12 in our study (\(P = 3.19 \times 10^{-7}\)). FGPI2 peak has 1.5× bigger area than FGPI3 Thus, we propose that ST6GAL1 has an exclusive effect on FGPI3 [F2A[3]G1S[3-6\(^1\)] trait].

It should be noted that we used a new imputation panel (1000 Genomes instead of HapMap in the previous studies) that more than tripled the number of polymorphisms analyzed (from 2.4 M SNPs to 8 M). That may have contributed to the higher power of our study as well.

In addition to detecting novel loci, we were able to replicate five (HNF1A, FUT6, FUT8, B3GAT1 and MGAT5) out of six loci that were reported previously to be associated with human plasma N-glycome measured using the HPLC technology (26,27). However, 3 out of 10 novel loci have not been replicated. For two of them, associated SNPs had relatively low (~2%) MAF. For rare variants, multiple testing burden is increased compared to common variants, which may lead to higher than expected false positive rate (58) in discovery. At the same time, relative effects of drift that are more pronounced for rare variants (59,60) may increase false negative rate in replication. The third unreplicated locus is located in HLA region known to have very complex structure (61). This locus showed highest (although not significant) heterogeneity of association (\(P\)-value of Cochran’s Q-test = 0.07) among loci in the replication meta-analysis.

Among six plasma glycome loci that were identified as genome-wide significant previously (26,27), only one (region of FUT8) had overlap with a locus identified as associated with IgG glycome composition (43). A recent multivariate GWAS study of plasma IgG glycome composition (56) identified five new loci, including the region of FUT3/FUT6, thus bringing the overlap between plasma and IgG glycome loci to two. In our study, among 12 replicated loci, the majority (8 loci) overlapped with loci that were reported to be associated with IgG glycome composition (43,56) (Fig. 3). We therefore established a strong overlap between IgG and plasma glycome loci.

In a way, this overlap is to be expected. It is known that majority of serum (and therefore plasma) glycoproteins are either immunoglobulins produced by B-lymphocytes or glycoproteins secreted by the liver (62). We thus expected overlap between IgG and total plasma glycome loci, and we expected that loci associated with the plasma N-glycome would be enriched by genes with tissue specific expression in liver and B cells. Indeed, we find that plasma N-glycome loci are enriched by genes expressed in plasma cells, antibody producing-cells and B-lymphocytes, and we also find overlap between plasma N-glycome loci and CD19+ eQTLs. However, we neither find enrichment of genes that are expressed in liver (Supplementary Material, Table 5C) nor overlap between plasma N-glycome loci and liver eQTLs. In the future, it will be important to achieve better resolution and separation of loci that are related to glycosylation of non-immunoglobulin glycoproteins. This could be achieved either technologically (e.g. performing analyses of IgG-free fractions of proteins), or this could be attempted via statistical modeling.

The genetic variation in the FUT3/FUT6 locus is a major (in terms of proportion of variance explained and number of glycans affected) genetic factor for non-immunoglobulin glycosylation. According to current knowledge, these enzymes catalyze fucosylation of antennary GlcNAc52, resulting in glycan structures that are not found on IgG (51,63). This is consistent with the spectrum of glycan traits associated with FUT3/FUT6 locus in our work (Fig. 3). However, this locus was recently found to be associated with IgG glycosylation (56). The authors could not explain this finding because at that time IgG glycans were not known to contain antennary fucose. Two explanations could have been proposed for this surprising finding: either enzymes encoded by FUT3/FUT6 locus exhibit non-canonical activity of core fucosylation or some IgG glycans actually do contain antennary fucose. Recently, the latter was demonstrated in the study by Russell et al. (10). Based on our results showing no evidence for association between FUT3/FUT6 locus and core fucosylation, we can speculate that association of this locus with IgG glycosylation could be explained by the presence of antennary fucose on some IgG-linked glycans.

An interesting pattern starts emerging out of study of genetic control of plasma glycosylation. We now see a clear overlap in genetic control between plasma and IgG glycosylation, which calls for future studies that would help distinguish among global, cell-specific, tissue-specific and protein-specific pathways of protein glycosylation. Many (8 out of 12) replicated loci contained genes that encode enzymes directly involved in glycosylation (FUT3/FUT6, FUT8, B3GAT1, ST6GAL1, B4GALT1, ST3GAL4, MGAT3 and MGAT5). However, glycosylation results from a complex interplay not only of enzymes responsible for transfer of monosaccharides to a growing glycan chain but also of enzymes involved in biosynthesis of individual monosaccharides and mechanism ensuring timely localization of all components involved in the process. Moreover, a number of environmental factors, such as diet and smoking, were found to be associated with plasma N-glycome composition (25). Studies in mouse models suggested complex compensatory mechanisms to pay role in glycosylation, which can complicate interpretation of the observed associations (64,65). We now start seeing loci and genes, which are likely to reflect complex aspects of plasma protein glycosylation, such as regulation of fucosylation by HNF1a (26). Such regulatory genes, in our view, are plausible candidates that will help linking glycans with complex human diseases. The results of our study provide evidence for potential role of genes DERL3, CHCHD10, TMEM121, IGH and IKZF1, although for some loci we could not prioritize only one particular gene (e.g. for locus containing DERL3 and CHCHD10). These genes can be considered candidates for future experimental research. To facilitate further studies of glycosylation and of the role of glycome in human health and diseases we have made full results of our plasma N-glycome GWAS (almost 1 billion of trait-SNP associations) freely available to the scientific community via GWAS archive.
Previous GWAS of HPLC measured plasma N-glycome (27) identified six genes controlling plasma N-glycosylation of which four implicated genes with obvious links to the glycosylation process. Here, using a smaller sample but more precise UPLC technology and new GWAS imputation panels, we confirmed the association of five known loci and identified replicated additional seven loci. Our results support the idea that genetic control of plasma protein N-glycosylation is a complex process, which is under control of genes that belong to different pathways and are expressed in different tissues. Further studies with larger sample size are warranted to further decrypt the genetic architecture of the glycosylation process and explain the relations between glycosylation and mechanisms of human health and disease.

Materials and Methods

Study cohort description

This work is based on analysis of data from four cohorts—TwinsUK, PainOR, SOCCS and QMDiab. Sample demographics can be found in Supplementary Material, Table 11.

TwinsUK. The TwinsUK cohort (66) (also referred to as the UK Adult Twin Register) is a nationwide registry of volunteer twins in the UK, with about 13 000 registered twins (83% female, equal number of monozygotic and dizygotic twins, predominantly middle-aged and older). The Department of Twin Research and Genetic Epidemiology at King’s College London hosts the registry. From this registry, a total of 2763 subjects had N-linked total plasmaglycan measurements that were included in the analysis.

QMDiab. The QMDiab is a cross-sectional case–control study with 374 participants. QMDiab has been described previously and comprises male and female participants in near equal proportions, aged between 23 and 71 years, mainly of Arab, South Asian and Filipino descent (67,68). The initial study was approved by the institutional review boards of Hamad Medical Corporation (HMC) and Weill Cornell Medicine—Qatar (WCM-Q) (research protocol #11131/11). Written informed consent was obtained from all participants. All study participants were enrolled between February 2012 and June 2012 at the Dermatology Department of HMC in Doha, Qatar. Inclusion criteria were a primary form of type 2 diabetes (for cases) or an absence of type 2 diabetes (for controls). Sample collection was conducted in the afternoon, after the general operating hours of the morning clinic. Patient and control samples were collected in a random order as they became available and at the same location using identical protocols, instruments and study personnel. Samples from cases and controls were processed in the laboratory in parallel and in a blinded manner. Data from five participants were excluded from the analysis because of incomplete records, leaving 176 patients and 193 controls. Of the 193 control participants initially enrolled, 12 had Hba1C levels above 6.5% (48 mmol/mol) and were subsequently classified as cases, resulting in 188 cases and 181 controls.

SOCCS. SOCCS study (69,70) comprised 2057 (colorectal cancer) CRC cases (61% male; mean age at diagnosis, 65.8 ± 8.4 years) and 2111 population controls (60% males; mean age, 67.9 ± 9.0 years) as ascertained in Scotland. Cases were taken from an independent, prospective, incident CRC case series and aged <80 years at diagnosis. Control subjects were population controls matched by age (±5 years), gender and area of residence within Scotland. All participants gave written informed consent and study approval was from the MultiCentre Research Ethics Committee for Scotland and Local Research Ethics committee. Sample collection is described in (69,70).

PainOR. The PainOR (71) is the University of Parma cohort of patients of a retrospective multicenter study (ClinicalTrials.gov Identifier: NCT02037789) part of the PainOMICS project funded by European Community in the Seventh Framework Programme (Project ID: 602736). The primary objective is to recognize genetic variants associated with chronic low back pain (CLBP); secondary objectives are to study glycomics and Activomics profiles associated with CLBP. Glycomic and Activomic approaches aim to reveal alterations in proteome complexity that arise from post-translational modification that varies in response to changes in the physiological environment, a particularly important avenue to explore in chronic inflammatory diseases. The study was firstly approved by the institutional review boards of Istituto di Ricovero e Cura a Carattere Scientifico (IRCCS) Foundation San Matteo Hospital Pavia and then by the institutional review boards of all clinical centers that enrolled patients. Copies of approvals were provided to the European Commission before starting the study. Written informed consent was obtained from all participants. In the period between September 2014 and February 2016, 1000 patients (including 38.1% male and 61.9% female, averaging 65 ± 14.5 years) were enrolled at the Anesthesia, Intensive Care and Pain Therapy Department of University Parma Hospital. Inclusion criteria were adult Caucasian patients who were suffering of lower back pain (pain between the costal margins and gluteal fold, with or without symptoms into one or both legs) more than 3 months who were admitted at Pain Department of University Parma Hospital. We exclude patients with recent history of spinal fractures or lower back pain due to cancer or infection. Sample collection was performed in all patients enrolled, according to the standard operating procedures published in PlosOne in 2017 (72). Samples were processed in PainOMics laboratory in a blinded manner in University of Parma.

Genotyping

For full details of the genotyping and imputation see Supplementary Material, Table 12.

TwinsUK. Genotyping was carried out using combination Illumina SNP arrays: HumanHap300, HumanHap610Q, 1 M-Duo and 1.2M-Duo 1 M. Standard quality control of genotyped data was applied, with SNPs filtered by sample call rate > 98%; MAF > 1%; SNP call rate, > 97% (for SNP with MAF ≥ 5%) or < 99% (for SNPs with 1% ≤ MAF ≤ 5%); and HWE, P < 1 × 10^{-6}. In total 275 139 SNPs passed criteria. Imputation was done using IMPUTE2 software with 1000G phase 1 version 3 and mapped to the GRCh37 human genome build. Imputed SNPs were filtered by imputation quality (SNPTEST proper-info) > 0.7, MAF ≥ 1% and MAC ≥ 10, leading to 8 557 543 SNPs passed to the GWAS analysis.

QMDiab. Genotyping was carried out using Illumina Omni array 2.5 (version 8). Standard quality control of genotyped data was applied, with SNPs filtered by sample call rate > 98%; MAF > 1%; SNP call rate, > 98%, and HWE P × c < 1 × 10^{-6}. In total 1 223 299 SNPs passed criteria. Imputation was done using SHAPEIT software with 1000G phase 3 version 5 and mapped to the GRCh37 human genome build. Imputed SNPs were filtered
by imputation quality > 0.7, leading to 20483276 SNPs passed to the GWAS analysis.

**SOCSS.** Details of the genotyping procedure can be found here (73). Genotyping was carried out using illumina SNP arrays: HumanHap300 and HumanHap240S. Standard quality control of genotyped data was applied, with SNPs filtered by sample call rate > 95%; MAF > 1%; SNP call rate, >95%; and HWE, $P < 1 \times 10^{-6}$. In total 514 177 SNPs passed criteria. Imputation was done using SHAPEIT and IMPUTE2 software with 1000 Genomes, phase 1 (Integrated haplotypes, released June 2014) and mapped to the GRCh37 human genome build. Imputed SNPs were not filtered, leading to 37 780 221 SNPs passed to the GWAS analysis.

**PainOR.** Genotyping was carried out using illumina HumanCore BeadChip. Standard quality control of genotyped data was applied with SNPs filtered by sample call rate > 98%; MAF > 0.625%; SNP call rate, >97%; and HWE, $P < 1 \times 10^{-6}$. In total 253 149 SNPs passed criteria. Imputation was done using Eagle software with HRC r1.1 2016 reference and mapped to the GRCh37 human genome build. Imputed SNPs were not filtered, leading to 39 127 685 SNPs passed to the GWAS analysis.

**Phenotyping**

**Plasma N-glycome quantification**

Plasma N-glycome quantification of samples from TwinsUK, PainOR and QMDiab were performed at Genos by applying the following protocol. Plasma N-glycans were enzymatically released from proteins by PNGase F, fluorescently labeled with 2-aminobenzamide and cleaned up from the excess of reagents by hydrophilic interaction liquid chromatography solid phase extraction (HILIC-SPE), as previously described. (74). Fluorescently labeled and purified N-glycans were separated by HILIC on a Waters BEH Glycan chromatography column, 150 $\times$ 2.1 mm, 1.7 $\mu$m BEH particles, installed on an Acquity UPLC instrument (Waters, Milford, MA, USA) consisting of a quaternary solvent manager, sample manager and a fluorescence detector set with excitation and emission wavelengths of 250 and 428 nm, respectively. Following chromatography conditions previously described in details (74), total plasma N-glycans were separated into 39 peaks for QMDiab, TwinsUK and PainOR cohorts. The amount of N-glycans in each chromatographic peak was expressed as a percentage of total integrated area. Glycan peaks (GPs)—quantitative measurements of glycan levels—were defined by automatic integration of intensity peaks on chromatogram. The number of defined GPs varied among studies from 36 to 42 GPs.

Plasma N-glycome quantification for SOCSS samples were done at the National Institute for Bioprocessing Research and Training (NIBRT) by applying the same protocol as for TwinsUK, PainOR and QMDiab, with the only difference in the excitation wavelength (330 nm instead of 250 nm).

**Harmonization of GPs**

The order of the GPs on a UPLC chromatogram was very similar among the studies. However, depending on the cohort some peaks located near one another might have been indistinguishable. The number of defined GPs varied among studies from 36 to 42. To conduct GWAS on TwinsUK following by replication in other cohorts, we harmonized the set of peaks (or GPs). According to the major glycostructures within the GPs we manually created the table of correspondence between different GPs (or sets of GPs) across all cohorts, where plasma glycome was measured using UPLC technology. Then, based on this table of correspondence, we defined the list of 36 harmonized GPs (Supplementary Material, Table 13) and the harmonization scheme for each cohort. We validated the harmonization protocol by comparing with manual re-integration of the peaks on chromatogram level using 35 randomly chosen samples from three cohorts: TwinsUK, PainOR and QMDiab. We show the full concordance between two approaches (Pearson correlation coefficient $R > 0.999$, see Supplementary Material, Table 14 for the details). We applied this harmonization procedure for the four cohorts: TwinsUK, QMDiab, CRC and PainOR, leading to the set of 36 glycan traits in each cohort.

**Normalization and batch correction of GPs**

Normalization and batch correction were performed on harmonized UPLC glycan data for four cohorts: TwinsUK, PainOR, SOCSS and QMDiab. We used total area normalization (the area of each GP was divided by the total area of the corresponding chromatogram). Normalized glycan measurements were log10-transformed due to right skewness of their distributions and the multiplicative nature of batch effects. Prior to batch correction, samples with outlying measurements were removed. Outlier was defined as a sample that had at least one GP that is out of three standard deviations from the mean value of GP. Batch correction was performed on log10-transformed measurements using the ComBat method (75), where the technical source of variation (batch and plate number) was modeled as a batch covariate. Again, samples with outlying measurements were removed.

From the 36 directly measured glycan traits, 77 derived traits were calculated (see Supplementary Material, Table 10). These derived traits average glycosylation features such as branching, galactosylation and sialylation across different individual glycan structures, and consequently, they may be more closely related to individual enzymatic activity and underlying genetic polymorphism. As derived traits represent sums of directly measured glycans, they were calculated using normalized and batch-corrected glycan measurements after transformation to the proportions (exponential transformation of batch-corrected measurements). The distribution of 113 glycan traits can be found in Supplementary Material, Figure 2.

Prior to GWAS, the traits were adjusted for age and sex by linear regression. The residuals were rank transformed to normal distribution [intransform function in GenABEL (76,77) R package].

**Genome-wide association analysis**

Discovery GWAS was performed using TwinsUK cohort (N = 2763) for 113 GP traits. Genome-wide Efficient Mixed Model Association algorithm (GEMMA) (78) was used to estimate the kinship matrix and to run linear mixed model regression on SNP dosages assuming additive genetic effects. Obtained summary statistics were corrected for genomic control inflation factor $\lambda_{GC}$ to account for any residual population stratification. An association was considered statistically significant at the genome-wide level if the P-value for an individual SNP was less than $5 \times 10^{-8}/(29 + 1)=1.66 \times 10^{-8}$, where 29 is an effective number of tests (traits) that was estimated as the number of principal components that jointly explained 99% of the total plasma glycome variance in the TwinsUK sample.
Locus definition

In short, we considered SNPs located in the same locus if they were located within 500 Kb from the leading SNP (the SNP with lowest P-value). Only the SNPs and the traits with lowest P-values are reported (leading SNP-trait pairs). The detailed procedure of locus definition is described in Supplementary Material, Note 1.

Replication

We have used TwinsUK cohort for the replication of six previously described loci (27) affecting plasma N-glycome. From each of six loci we have chosen leading SNP with the strongest association as reported by authors (27). Since there is no direct trait-to-trait correspondence between glycan traits measured by HPLC and UPLC technologies we tested the association of the leading SNPs with all 113 glycan traits in TwinsUK cohort. We considered locus as replicated if its leading SNP showed association with at least one of 113 glycan traits with replication threshold of \( P < 0.05/6(\times 50) = 2.78 \times 10^{-4} \), where six is number of loci and 30 is a number of principal components that jointly explained 99% of the total plasma N-glycome variance.

For the replication of novel associations, we used data from three cohorts: PainOR (N = 294), QMDiab (N = 327) and SOCCS (N = 472) with total replication sample size of \( N = 1048 \) samples that have plasma UPLC N-glycome and genotype data (for details of genotyping, imputation and association analysis; see Supplementary Material, Table 12). We used the leading SNPs and traits for the replication that were identified in the discovery step. For these SNPs we conducted a fixed-effect meta-analysis using METAL software (79) combining association results from three cohorts. The replication threshold was set as \( P < 0.05/10 = 0.005 \), where 10 is the number of replicated loci. Moreover, we checked whether the sign of estimated effect was concordant between discovery and replication studies.

Functional annotation in silico

**Functional annotation of associated variants.** All SNPs and indels in high LD (\( r^2 > 0.6 \)) with the 12 lead variants at replicated loci were selected. LD was calculated using genotype data for 503 samples with European descent from 1000 Genomes phase 3 version 5 data and Plink tool (80) (version 1.9) using –show-tags option. Additionally, for each of the 12 replicated loci we have selected the set of SNPs that had strong associations, defined as those located within \(+/−250 \text{ kbp}\) window from the strongest association, and having \( P\text{-value} < T \), where \( \log(10(T)) = \log(10(P\text{min}) + 1) \), where \( P\text{min} \) is the \( P\)-value of the strongest association in the locus. This additional inclusion criteria was applied since genotype data for TwinsUK samples was imputed using 1000 Genomes phase 1 version 3 panel and some of the SNPs from this panel are not exists in 1000 Genomes phase 3 version 5 panel. The list of selected variants can be found in Supplementary Material, Table 4A. Next all selected variants passed to functional annotation using Ensembl Variant Effect Predictor (VEP) method (33). Then, we used FATHMM-XF (34) and FATHMM-INDEL (35) methods to predict the impact of SNPs and small indels. Predictions are given as P-values in the range [0, 1]; values above 0.5 are predicted to be deleterious, while those below 0.5 are predicted to be neutral or benign. P-values close to the extremes (0 or 1) are the highest-confidence predictions that yield the highest accuracy.

**Gene-set and tissue/cell enrichment analysis.** To prioritize genes in associated regions, gene set enrichment and tissue/cell type enrichment analyses were carried out using DEPICT software v. 1 rel. 194 (32). For the analysis we have chosen independent variants (see ‘Locus definition’) with \( P < 5 \times 10^{-4}/30 \) (14 SNPs) and \( P < 1 \times 10^{-5}/30 \) (93 SNPs). We used 1000G dataset for calculation of LD (81). GO enrichment analysis was performed using FUMA GENE2FUNC (82) analysis based on MsigDB c5 (83) and MAGMA (37) with default parameters and ‘All genes’ as background genes.

**Pleiotropy with complex traits.** We have investigated the overlap between associations obtained here and elsewhere, using PhenoScanner v1.1 database (42). For 12 replicated SNPs (Table 1; Table 2) we looked up traits that have demonstrated genome-wide significant (\( P < 5 \times 10^{-8} \)) association at the same or at strongly (\( r^2 > 0.7 \)) linked SNPs.

**Pleiotropy with eQTLs.** To identify genes whose expression levels could potentially mediate the association between SNPs and plasma glycan traits we performed a SMR analysis followed by HEIDI method (38). In short, SMR test aims at testing the association between gene expression (in a particular tissue) and a trait using the top associated eQTL as a genetic instrument. Significant SMR test indicates not only evidence of causality or pleiotropy but also the possibility that SNPs controlling gene expression are in linkage disequilibrium with those associated with the traits. These two situations can be disentangled using the HEIDI test.

The SMR/HEIDI analysis was carried out for leading SNPs that were replicated and were genome-wide significant (\( P < 1.7 \times 10^{-8} \)) on discovery stage (11 loci in total, see Table 1). We checked for overlap between these loci and eQTLs in blood (39), 44 tissues provided by the GTEX database (40) and in nine cell lines from CEDAR dataset (41), including six circulating immune cell types (CD4+ T-lymphocytes, CD8+ T lymphocytes, CD19+ B lymphocytes, CD14+ monocytes, CD15 granulocytes and platelets). Technical details of the procedure may be found in Supplementary Material, Note 2. Following Bonferroni procedure, the results of the SMR test were considered statistically significant if \( P_{\text{SMR}} < 2.445 \times 10^{-6} \) (0.05/20448, where 20448 is a total number of probes used in analysis for all three data sets). Inferences whether functional variant may be shared between plasma glycan trait and expression were made based on HEIDI test: \( P_{\text{HEIDI}} > 0.05 \) (likely shared), \( 0.05 < P_{\text{HEIDI}} < 0.001 \) (possibly shared) and \( P_{\text{HEIDI}} < 0.001 \) (sharing is unlikely).

Data availability

Summary statistics from our plasma N-glycome GWAS for 113 glycan traits are available for interactive exploration at the GWAS archive (http://gwasarchive.org). The dataset was also deposited at Zenodo (http://doi.org/10.5281/zenodo.1298406) (84). The data generated in the secondary analyses of this study are included with this article in the supplementary tables.

Supplementary Material

Supplementary Material is available at HMG online.

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Conflict of Interest statement. Y.A. is a founder and co-owner of Maatschap PolyOmica, a private organization, providing services, research and development in the field of computational and statistical (gen)omics, and PolyKnomics, one of the organization providing the “GWASArchive”. G.L. is a founder and owner of Genos Ltd, a biotech company that specializes in glycay analysis and has several patents in the field. All other authors declare no conflicts of interest. Other authors declare no competing financial interests.

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Author Contributions

S.Z.S. and Y.A.T. contributed to the design of the study, carried out statistical analysis and produced the figures; S.Z.S., Y.A.T. and A.S.S. contributed to interpretation of the results; S.Z.S., Y.A.T., I.K.S., K.S. and Y.A. wrote the first version of the manuscript; S.Z.S., A.S.S., Y.A.T., I.K.S. and Y.A. wrote the revised second version of the manuscript; L.K., F.V., S.Z.S. and J.K. contributed to data harmonization and quality control; M.S., M.V., F.V., T.P., J.Stambuk, I.T.-A., J.K., J.Simunovic, M.P.-B. and G.L. contributed to plasma N-glycome measurements; M.M. and T.S. analyzed TwinsUK dataset and contributed to interpretation of the results; L.K., A.M.H., M.D. and S.M.F. analyzed SOCCS dataset and contributed to interpretation of the results; M.A., F.M.K.W. and C.D. designed PainOR study and contributed to interpretation of the results; K.S. and G.T. analyzed QMDiab dataset and contributed to interpretation of the results; E.L., J.D. and M.G. designed CEDAR study and contributed to interpretation of the results; Y.A. and G.L. conceived and oversaw the study, contributed to the design and interpretation of the results; all co-authors contributed to the final manuscript revision.

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