Genomics Meets Glycomics—The First GWAS Study of Human N-Glycome Identifies HNF1α as a Master Regulator of Plasma Protein Fucosylation

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Abstract

Over half of all proteins are glycosylated, and alterations in glycosylation have been observed in numerous physiological and pathological processes. Attached glycans significantly affect protein function; but, contrary to polypeptides, they are not directly encoded by genes, and the complex processes that regulate their assembly are poorly understood. A novel approach combining genome-wide association and high-throughput glycobiology analysis of 2,705 individuals in three population cohorts showed that common variants in the Hepatocyte Nuclear Factor 1α (HNF1α) and fucosyltransferase genes FUT6 and FUT8 influence N-glycan levels in human plasma. We show that HNF1α and its downstream target HNF4α regulate the expression of key fucosyltransferase and fucose biosynthesis genes. Moreover, we show that HNF1α is both necessary and sufficient to drive the expression of these genes in hepatic cells. These results reveal a new role for HNF1α as a master transcriptional regulator of multiple stages in the fucosylation process. This mechanism has implications for the regulation of immunity, embryonic development, and protein folding, as well as for our understanding of the molecular mechanisms underlying cancer, coronary heart disease, and metabolic and inflammatory disorders.

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Introduction

Glycosylation is a post-translational modification that enriches protein complexity and function. Over half of all known proteins are modified by covalently bound glycans, which are important for normal physiological processes, including protein folding, degradation and secretion, cell signalling, immune function and transcription [1–4]. Configuration and composition of attached glycans significantly change the structure and activity of polypeptide portions of glycoproteins [5] and since this process is not template driven, complexity of the glycoproteome is estimated to be several orders of magnitude greater than for the proteome itself [6]. Disregulation of glycosylation is associated with a wide range of diseases, including cancer, diabetes, cardiovascular, congenital, immunological and infectious disorders [1,3,7]. Enzymes that are involved in glycosylation may therefore be promising targets for therapy [8]. The most prominent example of the importance of N-glycosylation is the group of rare diseases named congenital disorders of glycosylation [9] where different mutations in the biosynthesis pathway of N-glycans cause significant mortality and extensive motor, immunological, digestive and neurological symptoms [10,11].
HNF1α is a Master Regulator of Fucosylation

Author Summary

By combining recently developed high-throughput glycan analysis with genome-wide association study, we performed the first comprehensive analysis of common genetic polymorphisms that affect protein glycosylation. Over half of all proteins are glycosylated; but, due to difficulties in glycan analysis and the absence of a genetic template for their synthesis, knowledge about the complex processes that regulate glycan assembly is still limited. We demonstrated that HNF1α regulates the expression of key fucosyltransferase and fucose biosynthesis genes and acts as a master regulator of plasma protein fucosylation. Proper protein fucosylation is essential in numerous processes including inflammation, cancer, and coronary heart disease, thus the identification of a master regulator of plasma protein fucosylation has important implications for understanding both normal biological functions and disease processes.

Due to experimental limitations in quantifying glycans in complex biological samples, our understanding of the genetic regulation of glycosylation is currently very limited [12]. However, recent technological advances have allowed reliable, high-throughput quantification of N-glycans [13], which now permits investigation of the genetic regulation and biological roles of glycan structures and brings glycomics into line with genomics, proteomics and metabolomics [14]. Recently we completed the first comprehensive population study of human plasma N-glycome which revealed variability that by far exceeds the variability of proteins and DNA [15]. However, within a single individual composition of plasma glycome is rather stable [16] and environmental factors have limited impact on the majority of glycans [17]. Specific altered glyco-phenotypes that can be associated with specific pathologies were also identified to exist in a population [18].

Variations in glycosylation are of great physiological significance as alterations in glycans significantly change the structure and function of polypeptide parts of glycoproteins [5]. A particularly interesting element of protein glycosylation is the addition of fucose to non-reducing ends of N-glycans. Fucose is a relatively novel sugar in evolutionary terms with two important structural features that distinguishes it from all other mammalian six-carbon monosaccharides; it lacks a hydroxyl group on the carbon at the 6-position and is the only monosaccharide that is in the L-configuration. The conversion of GDP-mannose to GDP-fucose is catalyzed by two enzymes (GMD and FX) that display remarkable evolutionary conservation [19,20]. On the other hand, the large family of genes that add fucose to proteins and lipids (fucosyltransferases, FUTs) has a very complex evolutionary history, including several more recent events specific to primates [21]. In mammals, fucose-containing glycans have important roles in blood transduction reactions, in the selectin-mediated leukocyte-endothelial adhesion that initiates an inflammatory response, in host-microbe interactions, and numerous ontogenic events [10,19]. Acute phase proteins have altered fucosylation in many diseases [22] and changes in the levels of fucosylated glycans have been shown to be associated with several important pathological processes, including cancer [23].

Hepatocyte nuclear factor 1α (HNF1α) and its downstream target HNF4α are transcription factors that regulate gene expression in both the liver and pancreas in a tissue-specific manner and are key regulators of metabolic genes [24]. Mutations in the encoding genes HNF1α and HNF4α cause Maturity Onset Diabetes of the Young (MODY) types 3 and 1 respectively [25,26]. Recently, HNF1α single nucleotide polymorphisms (SNPs) have been associated with plasma C-reactive protein (CRP) [27], LDL cholesterol and gamma glutamyltransferase (GGT) [28], and coronary heart disease [29]. HNF4α variants have been associated with ulcerative colitis [30] and with the plasma concentrations of CRP and apolipoprotein A1 (APOA1) [31]. Currently there is little evidence to link these transcription factors with fucose metabolism and the upstream mechanisms regulating fucosylation pathways are unknown.

Results

Common variants in fucosyltransferase genes affect the relative proportions of plasma N-glycans

We performed the first systematic analysis of the genetic regulation of individual N-glycans in plasma from 2,705 individuals in three population cohorts, from Croatia and Scotland, which have previously been characterized in great detail [32]. Desialylated 2AB-labelled human plasma N-glycans were separated into 13 structurally related groups of glycans, referred to as DG1–DG13 (see Table S1 for a list of specific glycans found within each DG group) [13]. The concentration of plasma N-glycans measured in each of these groups was then expressed as a proportion of the total plasma N-glycome to obtain 13 quantitative variables in each examinee. All N-glycans contain two core N-acetylglucosamine (GlcNAc) residues, to which a “core” fucose can be α1,6-linked to the inner GlcNAc, which is directly linked to an asparagine residue on the protein. Additional fucose residues can be transferred to different positions on antennas that have been added to the core glycan structure (Table S1). Two further traits were derived from the original variables to calculate the percentage of glycan structures containing core (FUC-C) or antennary (FUC-A) fucos, yielding a total of 15 glycan traits for analysis.

We conducted a meta-analysis of genome-wide association study (GWAS) data for the fifteen plasma N-glycan traits measured in three population-based cohorts, CROATIA-VIS (n = 924), CROATIA-KORCUILA (n = 896) and ORCADES (n = 737). Additive SNP effects were tested in each cohort independently and then combined in an inverse-variance weighted meta-analysis. The genome-wide significance threshold for the meta-analysis was set at 5 × 10^{-8}.

Genome-wide significant associations were found for DG1, DG6, DG7, DG9, DG11, as well as FUC-A (Table 1; Figure 1 and Figure 2). Association profiles for DG1, DG7, and DG9 are represented in their genomic context in Figure 1 for the associated region. Quantile–quantile plots for each association were consistent with an excess of true genetic associations, with modest genomic control inflation for each population (inflation factor <1.04 for all traits and each population as well as the meta-analysis), suggesting that the observed results were not due to population stratification (Figure 2A–2C).

Fifteen SNPs located in the region encompassing the fucosyltransferase 8 gene (FUT8, Entrez GeneID: 2530) on chromosome 14 were significantly associated with plasma concentrations of desialylated glycan (DG) 1, the most significant being rs7199838 (p = 3.46 × 10^{-15}) located 5’ of the gene. FUT8 was also associated with DG6, however for this trait only one SNP, rs10483776, reached genome-wide significance (p = 9.58 × 10^{-7}). All SNPs significantly associated with DG1 levels were in high LD (r^2 > 0.5) and located between two recombination hotspots, while no associations were found with SNPs located outside these boundaries nor with other genes located within this association.
interval (Figure 1A). The effect size of the G allele of rs7159888 was \(-0.2617\) (s.e. 0.0301) for DG1 in the meta-analysis of the 3 populations studied (standard deviation units, after adjustment for sex and age; Figure 2D). All significant SNPs in this region had a similar effect size (absolute value of the range: 0.1826–0.3251), accounting for between 1 and 6 percent of the trait variance after adjustment for age and sex. The effect of rs7159888 on DG1 was consistent across populations with similar amplitude and direction of effect (Figure 2D) with the effect for each population plotted separately along with the pooled effect. Haplotype analysis found that a single SNP model performed better than the 3- or 5-SNP haplotype model in every population.

A single SNP located on chromosome 19, rs3760776, was associated with DG7, DG9, DG12 and FUC-A (p = 3.42 \times 10^{-12}, p = 3.51 \times 10^{-17}, p = 9.44 \times 10^{-10}, p = 1.41 \times 10^{-12}) This SNP is located at the 5’ end of the fucosyltransferase 6 gene (FUT6, Entrez GeneID: 2528). The association interval for this SNP contains the NRTN, FUT6 and FUT3 genes (see Figure 1C), of which FUT6 and FUT3 are both biologically plausible candidates to explain the observed associations. The effect size of the G allele of rs3760776 is 0.3387 (s.e. 0.0487) for DG7 (standard deviation units, after adjustment for significant covariates: sex, age and fibrinogen); and 0.4104 (s.e. 0.0487), 0.2974 (s.e. 0.0486), and 0.3446 (s.e. 0.0486) for DG9, DG12 and FUC-A respectively (standard deviation units, after adjustment for age and fibrinogen). These effects account for 2% (DG7), 3% (DG9), 2% (DG12) and 2% (FUC-A) of the trait variance after adjustment for age and fibrinogen). These effects account for 2% (DG7), 3% (DG9), 2% (DG12) and 2% (FUC-A) of the trait variance. A forest plot of the effect size of rs3760776 in every population and for each of these populations studied (standard deviation units, after adjustment for sex, age and fibrinogen) is also within the region, so the causal variant(s) may affect one or both of these genes. The best 5-SNP haplotype contained rs3760776 and encompassed FUT6 but not FUT3 in every population and for every glycan group tested which suggests that the association is with FUT6, not FUT3.

The glycan structures which were significantly associated with genetic variants in the FUT6 and FUT8 genes are summarised in Table 1. Glycan group DG1 consists of a single structure GlcNAc2Man3GlcNAc2 that is known to be a substrate for the 1,6-fucosyltransferase (FUT8) (Table S1) [15,33]. Group DG6 contains three glycan structures, two of which are core fucosylated so the results are consistent with the known biological role of FUT8. In contrast, groups DG7, DG9 and DG12 include glycans containing antenmary fucose while FUC-A was derived as an overall measure of antenmary fucosylation. FUT6 encodes the enzyme fucosyltransferase VI which was reported to be the key enzyme responsible for the \(\xrightarrow{}\)fucosylation of plasma proteins [34]. The association of FUT8 and FUT6 genes with N-glycan structures containing core and antenmary fucosylation is supported by their known biological functions [35] and the fact that they were identified in this study is an effective proof of principle that HPLC measured glycan levels can be used to identify genes that regulate protein glycosylation.

Novel association of HNF1\(\alpha\) with N-glycans

Two SNPs on chromosome 12, rs7953249 and rs733596, showed genome-wide significant associations with DG7 (p = 1.97 \times 10^{-08}, p = 1.75 \times 10^{-08}). The latter SNP was also associated with DG11 (p = 4.44 \times 10^{-08}), with an effect in the opposite direction, and was close to genome-wide significance with DG9 (Table S2). Both SNPs are located in the HNF1\(\alpha\) (Entrez GeneID: 6927) gene region: rs7953249 is located 13 kb 5’ to the gene and rs733596 is in intron 9. Two other genes are found between the recombination hotspots that comprise the boundaries of the association interval, CI2orf43 and OASL (Figure 1B). However, none of the most significantly associated SNPs are located in these genes and all SNPs with suggestive p-values (p < 1 \times 10^{-05}) are located within HNF1\(\alpha\) (Table S2). The effect size of the G allele of rs733596 is –0.1767 (standard deviation units, after adjustment for sex, age and fibrinogen; s.e. 0.0314) for DG7, which only contains glycans with antenmary fucose, and in the opposite direction (0.1699 standard deviation units, after adjustment for age and fibrinogen; s.e. 0.0310) for DG11, which has no antenmary fucose (Table 1). All significant SNPs in this region had a similar effect size (absolute value of the range: 0.1396–0.1767), representing 1–3% of the trait variance. Figure 2E shows the effect size for rs733596 with DG7 for each population separately and the pooled meta-analysis. Comparison of models including rs7953249 and rs733596 separately and combined suggests that the causal variant is located between these two SNPs. This was confirmed by analysis of imputed data based on HapMap release 2 with the most significant SNPs located across intron 1 of HNF1\(\alpha\).

| Trait | Fucosylation status | Chr | SNP | Meta-analysis P-value | Meta-analysis sample size | Associated interval size, kb (no. of genes in interval) | Genes within associated interval | Major allele, minor allele (MAF) | Effect size for minor allele (s.e.) (z-score units)* |
|-------|---------------------|-----|-----|-----------------------|---------------------------|--------------------------------------------------------|---------------------------------|--------------------------------|---------------------------------------------|
| DG1   | NF                  | 1q23.3 | rs7159888 | 3.46 \times 10^{-18} | 2,441 | 650 (1) | FUT8 | G, A (0.44) | 0.26 (0.03) |
| DG6   | CF                  | 14q23.3 | rs10483776 | 9.58 \times 10^{-09} | 2,332 | 650 (1) | FUT8 | A, G (0.21) | 0.22 (0.04) |
| DG7   | AF                  | 12q24.31 | rs7953249 | 1.97 \times 10^{-08} | 2,321 | 118 (3) | HNF1A-CI2orf43-OASL | G, A (0.45) | 0.17 (0.03) |
| DG7   | AF                  | 1p13.3 | rs3760776 | 3.42 \times 10^{-12} | 2,320 | 54 (3) | NRTN-FUT6-FUT3 | G, A (0.11) | 0.34 (0.05) |
| DG9   | AF, CF              | 1p13.3 | rs3760776 | 3.51 \times 10^{-17} | 2,333 | 54 (3) | NRTN-FUT6-FUT3 | G, A (0.11) | 0.41 (0.05) |
| DG11  | NF/CF, AF           | 12q24.31 | rs735396 | 4.44 \times 10^{-08} | 2,332 | 118 (3) | HNF1A-CI2orf43-OASL | G, A (0.38) | 0.17 (0.03) |
| DG12  | AF                  | 1p13.3 | rs3760776 | 9.44 \times 10^{-10} | 2,330 | 54 (3) | NRTN-FUT6-FUT3 | G, A (0.11) | 0.30 (0.05) |
| FUC-A | AF                  | 1p13.3 | rs3760776 | 1.41 \times 10^{-12} | 2,321 | 54 (3) | NRTN-FUT6-FUT3 | G, A (0.11) | 0.34 (0.05) |

*Effect size shown is the \(\beta\)-coefficient, which represents change in glycan levels measured in standard deviation units (after adjustment for covariates) per copy of the allele modelled. NF, non-fucosylated; CF, core fucosylated; AF, antenmary fucosylated; See also Tables S1 and S2.

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A

rs7159888

P=1.5e-18

recombination rate

Observed (logP)

80

60

40

20

0

FUT8

64900

65100

65300

Chromosome 14 position (hg18) (kb)

B

rs735396

P=2.507e-06

recombination rate

Observed (logP)

8

6

4

2

0

SPPL3

HNF1A

C12orf43

119800

119900

Chromosome 12 position (hg18) (kb)

C

rs3760776

P=3.513e-17

recombination rate

Observed (logP)

16

12

8

4

0

TMEM146

DUS3L

NRTN

FUT3

FUT5

NDUFA11

CAPS

RANBP3

5800

Chromosome 19 position (hg18) (kb)
HNF1α regulates multiple stages in protein fucosylation: (1) regulation of GDP-fucose biosynthesis

The shared characteristic of all glycan groups that showed association with HNF1α SNPs was the presence or absence of antennary fucose (Table S2). We hypothesised that HNF1α transcriptionally regulates the expression of genes involved in the separate steps of fucosylation. This is supported by the fact that a functionally related transcription factor, HNF4α, was previously shown to bind the regulatory elements of the GDP-mannose-4,6-dehydrogenase (GMDS) gene in a genome-wide ChIP-ChIP. GMDS is involved in the de novo pathway of L-fucose synthesis to produce GDP-fucose, the substrate used by both core and antennary fucosyltransferases to N-glycosylated proteins [24]. Moreover, HNF1α directly regulates the expression of the hepatic fucosyltransferase VI gene (FUT6) [36]. Therefore, we tested whether HNF1α and/or HNF4α might regulate other genes involved in GDP-fucose biosynthesis. To this end, HNF1α and HNF4α were transiently knocked-down in liver and pancreatic cell lines using RNA interference. Both HNF1α and HNF4α expression levels decreased upon knockdown of either of them in hepatocytes (Figure 3A). In pancreatic cells, HNF1α knockdown up-regulates HNF4α expression but the reverse is not true (Figure S1). This confirms the differential regulation of gene expression downstream of HNFs in liver vis-a-vis pancreas [28,37]. It also corroborated recent findings in murine Hnf1α heterozygote pancreas, where the levels of Hnf4α mRNA increase [38].

As a positive control, the expression of FUT6, a known target of HNF4α in hepatocytes, was first analysed. The ablation of the HNF4α transcript abolished the expression of FUT6 in HepG2 cells confirming that the knockdown was effective. Surprisingly, knockdown of HNF1α resulted in 50% reduction in FUT6 transcript levels suggesting that HNF1α also regulate FUT6 expression in HepG2. This experiment suggested that our hypothesis may potentially explain and provide a direct link between HNF1α and the fucosylation genes. Therefore, we focused on the genes responsible for fucose biosynthesis, a rate limiting step in protein fucosylation. To this end, we analysed the expression of GMDS and L-Fucokinase which regulate de novo and salvage pathways of fucose synthesis, respectively. In HepG2 liver cells, HNF1α and HNF4α knockdown resulted in dramatic down-regulation in the expression of GMDS (91 and 77%, respectively) and L-Fucokinase (92 and 98%, respectively) (Figure 3B). In the pancreatic Panc1 cell lines, HNF4α RNAi resulted in a 70% decrease in GMDS and L-Fucokinase transcript levels (Figure 1). However, HNF1α RNAi led to a 90% reduction in GMDS transcript levels but did not affect L-Fucokinase mRNA abundance (Figure 1). This suggests that HNF1α regulates de novo synthesis of d-fucose in both cell lines tested (liver and pancreas), but only the salvage pathway in the liver cell line tested. HNF4α, on the other hand, regulates both pathways in both cell types tested.

We therefore focused on HNF1α direct transcriptional regulation of HNF4α, GMDS and L-Fucokinase in HepG2 cells. In order to investigate the latter, we performed a bioinformatics analysis to delineate in silico HNF1α and HNF4α binding sites. First, we assessed the conservation of regulatory elements (at the 5’ and 3’ end) between human and other primates as described previously [39]. It was recently shown the sites are not conserved between primates and rodents [40]. Second, the conserved regions were then mined for potential sites using ECR browser and the TRANSFAC database [41]. Finally, the potential sites were analysed manually to ascertain the likely binding sites based on homology to HNF1α and HNF4α consensus binding sites mined using genome-wide ChIP analyses [40,42]. This limited our analysis to 5 sites (primer pairs P16 to P20, Figure 3C) in the GMDS promoter, 3 sites in the promoter (primer pairs P21 to P23, Figure 3D) as well as 2 sites at the 3’end (primer pairs P24 and P25, Figure 3D) of the L-Fucokinase gene and 3 sites in the promoter (primer pairs P34 to P36, Figure 3D) as well as a single site at the 3’ end (primer pair P37) of the HNF4α gene. The primer pairs are less than 1kbps away from each other and some contained both HNF1α and HNF4α binding sites (or half sites) within the 200bps amplifiable regions.

Using these primer pairs, we performed chromatin immunoprecipitation (ChIP) assays to delineate the occupancy of these sites by HNF1α, HNF4α or both proteins as described earlier [43,44]. In HepG2, both HNF1α and HNF4α bind the promoters of GMDS (P17, Figure 3C), L-Fucokinase (P22 although the two factors cannot be re-precipitated, Figure 3D) and HNF4α (P36, Figure 3E).

Also, we show binding of HNF1α and HNF4α at the 3’UTR of L-Fucokinase as well as HNF4α binds the 3’UTR of HNF1α (Figure 3D and 3E, respectively). The interactions of these proteins is not affected by shearing as the primers acts as genomic controls for each other and no signal above background was apparent in the IgG isotype control antibody. Together, the data suggests a complex network of interactions between HNF1α and HNF4α to regulate fucose biosynthesis gene expression and point to a novel and an appreciated role for HNF1α in regulating the two genes studied (GMDS and L-Fucokinase).

HNF1α regulates multiple stages in protein fucosylation: (2) transcriptional regulation of core and antennary fucosyltransferases

After confirming the role of HNF1α in the biosynthesis of GDP-fucose, we analysed the role of HNF1α and HNF4α in the regulation of the expression of fucosyltransferase (FUT) genes (FUT3-11) in HepG2 and Panc1 cell lines to assess whether these hepatic factors regulate other stages of protein fucosylation. In HepG2 cells, HNF1α knockdown down-regulated the expression...
of all FUT genes (Figure 4A and 4B), except FUT8 which was induced upon the loss of HNF1α (Figure 4C). HNF4α knockdown led to a statistically significant downregulation of FUT3, FUT5, FUT6, FUT10, FUT11 but not FUT7 or FUT9 (Figure 4A and 4B). Conversely, FUT8 expression levels increased 10 fold upon the loss of HNF4α (Figure 4C). FUT4 was not expressed in HepG2 cells confirming earlier studies [45]. In the pancreas, all FUT genes were down-regulated (Figure 2) pointing to a key role for HNF1α in the regulation of fucosylation in the pancreas. Knockdown of HNF4α in liver cells reduced the expression of all FUT genes analysed except FUT7 or FUT9, but to a lesser extent than HNF1α knockdown (Figure 4A and 4B), however, FUT8 was again up-regulated (Figure 4A and 4B). The data supports a wider effect of HNF1α on the expression of the fucosyltransferase genes compared to HNF4α. The data also suggests that HNF1α and HNF4α downregulate FUT8, which adds fucose to the core glycan, in contrast to all other FUTs that add fucose to the antennary arms of glycans [33]. We observed a rather high correlation between concentrations of antennary and core fucosic in our population samples (r = 0.574, p = 4.01 × 10^{-85}), indicating that the availability of the common substrate of both core and antennary FUTs, GDP-fucose, is a rate-limiting factor in protein fucosylation. It therefore appears that HNF1α not only enhances the activity of antennary FUTs but also, by down-regulating FUT8, increases the amount of GDP-fucose available for antennary fucosylation.

FUT3, FUT6 and FUT5 were the only FUTs to be highly repressed (more than 3-fold) upon the loss of both HNF1α and HNF4α in liver cells (Figure 4A), suggesting a co-regulation of the three genes. In pancreatic cells, FUT3 and FUT6, but not FUT5 followed the same dynamics (Figure 2), FUT3 and FUT6 expression was not repressed upon HNF4α loss (Figure 2). This could be explained by a differential role for HNF4α in regulating FUT3 but not FUT3 or FUT6. These data suggest that HNF1α is the major regulator of the fucosylation pathway in both liver and pancreatic cell lines. While HNF4α also regulates the expression of these genes, its role is probably secondary to HNF1α. However, none of the genes studied here have previously been shown to be regulated in vivo by HNFs. Only the GMDS promoter has previously been shown to be chromatin immunoprecipitated with HNF4α antibody [28].

Bioinformatic analysis showed that FUT3, FUT5 and FUT6 are clustered in one locus in the human genome (see and Figure S3) [39]. This also corroborated our findings that FUT3, FUT5 and FUT6 are co-regulated downstream of HNF4α and HNF1α (Figure 4A). However, the FUT3/5/6 cluster was neither syntenic nor conserved in the mouse genome. We therefore focused on primate conservation only.

The promoter, intergenic and 3′ regulatory element conserved regions were analysed for HNF binding sites as detailed above for GMDS and L-Fucokinase. This analysis identified a limited number of sites in regulatory regions of FUT3, FUT5, FUT6, and FUT10. It did not identify any binding sites in silico in the FUT11 promoter, but a highly conserved long range enhancer was found within the ADK gene, that is 650 kb upstream and rich in HNF binding sites. We were unable to detect any HNF binding sequences within the FUT8 regulatory elements analysed.

Using ChIP, the binding of HNF1α and HNF4α to the putative response elements identified in silico was analysed. ChIP analysis showed that HNF1α and HNF4α bound multiple sequences within the predicted regulatory regions of multiple FUT genes, including FUT3, FUT5, FUT6, FUT10, FUT11 (Figure 4D–4I). HNF4α, and not HNF1α, bound the promoter of FUT5 (P13 and P15, Figure 4G). The unique binding of HNF4α to the promoter of FUT5 corroborated our findings that knockdown of HNF4α in pancreatic Panc1 cells abolished the expression of FUT5 but not FUT3 or FUT6 (Figure 2).

Using re-precipitation (reChIP), we confirmed that both HNF transcription factors bound (i) the promoters of FUT3, FUT6 and FUT10 (Figure 4E, 4F, 4D, and 4I respectively); (ii) 3′UTRs of FUT6 (Figure 4D); and (iii) the long range enhancer 650 kb upstream of FUT11 (Figure 4H). This shows that HNF1α and HNF4α are potential regulators of the expression of these genes in vivo.

Discussion

By performing the first genome-wide association analysis (GWAS) of protein glycosylation we have taken the first steps towards mapping the complex network of genes that regulate protein N-glycosylation. We also identified common variants in three genes which exert a relative strong influence on N-glycans in plasma (1–6% of variance explained). Importantly, all of the identified genes (FUT6, FUT11 and HNF1α) are involved in fucosylation, indicating that the addition of this unusual sugar may be a rate-limiting step in N-glycan synthesis. A gene encoding the transcription factor HNF1α, with previously unknown biological links to glycosylation, is shown to be strongly associated with the relative proportions of plasma N-glycans. The possible function(s) of HNF1α are a focus of intense current interest following its recently reported associations in GWAS with plasma C-reactive protein (CRP) [27], gamma-glutamyl transerase (GGT) [28], LDL cholesterol and apolipoprotein [29,31] and coronary artery disease [29,46]. Our analysis of gene knockdowns (RNAi) showed that HNF1α is an upstream regulator of several key genes involved in different stages of the fucosylation pathway. We have demonstrated that HNF1α binds the promoters in vivo, and is necessary and sufficient for the in vitro expression, of two genes, fucokinase and GMDS, required for de novo and salvage pathways of fucose synthesis, respectively (Figure 5C). Fucose synthesis is the rate limiting step for fucosylation in eukaryotes and prokaryotes [35] and, by up-regulating its synthesis, HNF1α increases the availability of fucose to the glycosylation machinery. In addition, HNF1α directly regulates the expression of several fucosyltransferase (FUT) genes (Figure 5D). Our results also demonstrate that HNF1α reciprocally regulates core versus antennary fucosylation; while activating FUTs involved in antennary fucosylation, HNF1α represses FUT8, which adds fucose to the core-GlcNAc. In this way, HNF1α decreases the consumption of GDP-fucose for core-fucosylation, and further increases the pool of fucose available for antennary fucosylation.

Having shown this novel regulation of fucosylation genes, we scanned earlier genome wide studies for HNF factors to identify whether these genes were picked up. In fact, other genome wide
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A

Fold enrichment over GAPDH normalised to gene levels in control cells (1)

|          | HNF1α | HNF4α | FUT6 |
|----------|-------|-------|------|
| Scrambled| -     | +     | -    |
| HNF1α RNAi| -     | +     | -    |
| HNF4α RNAi| -     | +     | -    |

B

GMDS

Fold enrichment over GAPDH normalised to gene levels in control cells (1)

|          | GMDS | L-fucokinase |
|----------|------|--------------|
| Scrambled| +    | -            |
| HNF1α RNAi| -    | +            |
| HNF4α RNAi| -    | +            |

C

ChIP

|      | Input | IgG | HNF1α | HNF4α | HNF1α/4α reChIP |
|------|-------|-----|-------|-------|-----------------|
| P16  |       |     |       |       |                 |
| P17  |       |     |       |       |                 |
| P18  |       |     |       |       |                 |
| P19  |       |     |       |       |                 |
| P20  |       |     |       |       |                 |
| P21  |       |     |       |       |                 |
| P22  |       |     |       |       |                 |
| P23  |       |     |       |       |                 |
| P24  |       |     |       |       |                 |
| P25  |       |     |       |       |                 |

GMDS promoter

L-Fucokinase promoter

D

ChIP

|      | Input | IgG | HNF1α | HNF4α | HNF1α/4α reChIP |
|------|-------|-----|-------|-------|-----------------|
| P21  |       |     |       |       |                 |
| P22  |       |     |       |       |                 |
| P23  |       |     |       |       |                 |
| P24  |       |     |       |       |                 |
| P25  |       |     |       |       |                 |

L-Fucokinase 3’UTR

E

ChIP

|      | Input | IgG | HNF1α | HNF4α | HNF4α reChIP |
|------|-------|-----|-------|-------|--------------|
| P34  |       |     |       |       |              |
| P35  |       |     |       |       |              |
| P36  |       |     |       |       |              |
| P37  |       |     |       |       |              |

HNF4α promoter

HNF4α 3’UTR

F

Fold luciferase activity

|      | HNF1α RNAi |
|------|------------|
|      | -          |
|      | +          |

G

Fold luciferase activity

|      | HNF1α RNAi |
|------|------------|
|      | -          |
|      | +          |

100ng pcDNA3 HNF1α
the reported association between common variants in HNF1α involved in fucosylation may be the molecular mechanism behind regulation of antennary fucosyltransferases and down-regulation of artery disease, inflammatory bowel disease and cancer.

Increasing acute-phase response requires a rapid increase in the concentration of fucose, which are also highly glycosylated and have high content of antennary-fucose [50]. Mounting a successful acute phase response [15], which are also highly glycosylated and have high content of antennary-fucose [50].

We hypothesize that the role of HNF1α and its transcriptional co-factor HNF4α in the regulation of fucosylation is an essential part of mounting an acute phase response to infection in humans. Antennary fucosylation of their glycoprotein ligands is needed for binding of E-, L- and P-selectins to their target cells and the initiation of inflammation [48]. The decrease in fucosylation in the rare Leukocyte Adhesion Deficiency II (LAD II) impairs neutrophil function, which can be restored by oral administration of fucose [49]. Recently, we have reported moderate correlations between fucosylated plasma N-glycans and components of the acute phase response [15], which are also highly glycosylated and have high content of antennary-fucose [50]. Mounting a successful acute-phase response requires a rapid increase in the concentration of acute-phase proteins and this in turn is dependent on their efficient fucosylation. Our results indicate that fucosylation is a rate-limiting step in plasma protein glycosylation, and by both increasing de novo and salvage synthesis of GDP-fucose, up-regulation of antennary fucosyltransferases and down-regulation of core-fucosyltransferase, HNF1α appears to be a master regulator of this process. Variants in HNF1α and HNF4α genes were previously reported to be associated with concentrations of acute phase proteins in human plasma [24,27]. Plasma protein fucosylation plays an important role in inflammation [22] and the central role of HNF1α in the regulation of multiple genes involved in fucosylation may be the molecular mechanism behind the reported association between common variants in HNF1α and inflammatory markers (such as CRP) as well as several diseases in which inflammation plays a key pathogenic role (such as coronary artery disease, inflammatory bowel disease and cancer).

Materials and Methods

Study populations and genotyping

All three populations recruited adult individuals within a community irrespective of any specific phenotype. The CROATIA-VIS and CROATIA-KORCULA studies are both cohorts from the Croatian Dalmatian islands recruited in 2003–2004 and 2007 respectively. The ORCADES study is ongoing with participants recruited from the Orkney islands in Scotland. Fasting blood samples were collected, biochemical and physiological measurements taken and questionnaires of medical history as well as lifestyle and environmental exposures collected following similar protocols.

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 [32].

The CROATIA-KORCULA study includes 969 Croatians between the ages of 18 and 98 [32]. The field work was performed in 2007 in the eastern part of the island, targeting healthy volunteers from the town of Korčula and the villages of Lumbarda, Žrnovo and Račišće.

The Orkney Complex Disease Study (ORCADES) is an ongoing study in the isolated Scottish archipelago of Orkney [32]. Data for participants aged 18 to 100 years, from a subgroup of ten islands, were used for this analysis.

DNA samples were genotyped according to the manufacturer’s instructions on Illumina Infinium SNP bead microarrays (HumanHap300v1 for the CROATIA-VIS cohort, HumanHap300v2 for the ORCADES cohort and HumanCNV370v1 for the CROATIA-KORCULA cohort). Genotypes were determined using Illumina BeadStudio software. Genotyping was successfully completed on 991 individuals from CROATIA-VIS, 953 from CROATIA-KORCULA and 761 from ORCADES.

Ethics statement

All studies conformed to the ethical guidelines of the 1975 Declaration of Helsinki and were approved by appropriate ethics boards with all respondents signing informed consent prior to participation.

Glycan release and labelling

The N-glycans from plasma sample (5 μl) proteins were released and labelled with 2-aminobenzamide (LudgerTag 2-AB labelling kit Ludger Ltd., Abingdon, UK) as described previously [13]. Labelled glycans were dried in a vacuum centrifuge and redissolved in known volume of water for further analysis.

Sialidase digestion

After initial HPLC quantification sialidase digestion was performed to improve measurement precision. Aliquots of the 2-AB-labeled glycan pool were dried down in 200-μl microcentrifuge tubes. To these, the following was added: 1 μl of 500 mM sodium acetate incubation buffer (pH 5.5), 1 μl (0.005 units) of ABS, Arthrobacter ureafaciens sialidase (releases α2-3, 6, 8 sialic acid, Prozyme) and H2O to make up to 10 μl. This was incubated overnight (16–18 h) at 37°C and then passed through a Micropure-EZ enzyme remover (Millipore, Billerica, MA, USA) before applying to the HPLC.

Hydrophilic interaction high-performance liquid chromatography (HILIC)

Released glycans were subjected to hydrophilic interaction high performance liquid chromatography (HILIC) on a 250×4.6 mm i.d. 5 μm particle packed TSKgel Amide 80 column ( Tosoh Bioscience, Stuttgart, Germany) at 30°C with 50 mM formic acid adjusted to pH 4.4 with ammonia solution as solvent A and acetonitrile as solvent B. 60 min runs were on a 2795 Alliance separations module (Waters, Milford, MA). HPLCs were equipped with a Waters temperature control module and a Waters 2475 fluorescence detector set with excitation and emission wavelengths of 330 and 420 nm, respectively. The system was calibrated using
Figure 4. HNF1α is major regulator of the fucosyltransferase genes. Real time PCR for FUT3, FUT5 and FUT6 RNAs (A); FUT7, FUT9, FUT10 and FUT11 (B); FUT8 (C) after HNF1α and HNF4α RNA interference (RNAi) in HepG2 cells. H) analyses show that Hnf1α (HNF1) and Hnf4α (HNF4) bind predicted regulatory elements for key fucosylation genes in liver cells. D), E), F), G), H) and I) Binding of HNF1α and HNF4α to predicted regulatory elements of the FUT3, FUT5, FUT6, FUT10 and FUT11. Abbreviations: the gene names are detailed in Figure 1; P1–P37: Primer sets 1 to 37; 3’UTR: 3’ untranslated region. reChIP: re-precipitation (or sequential) ChIP.

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Figure 5. HNF1α is at the heart of fucosylation regulation. A) Diagram showing HNF1α and HNF4α regulation of pathways involved in fucose synthesis. B) Pathways of core (tri-mannosyl-chitobiose) and antennary fucosylation and their proposed regulation by the transcription factors Hnf1α and Hnf4α, encoded by the HNF1α and HNF4α genes respectively. Red arrows indicate genes that are targets of Hnf1α/Hnf4α. Solid lines show direct regulation and dashed lines show indirect regulation of the FUT genes shown. Genes shown in blue depict GWAS identified genes (HNF1α, FUT6 and FUT8). Abbreviations: Neu5Ac, N-Acetylneuraminic (sialic) acid; GlcNAc, N-acetylgalactosamine; Man, mannose; Gal, galactose; poly-LacNAc, Galβ1,4GlcNAC (LacNAC) antennary chains; Asn, asparagine residue on glycosylated protein. 

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an external standard of hydrolyzed and 2-AB-labeled glucose oligomer from which the retention times for the individual glycans were converted to glucose units (GU) [51]. Glycans were analyzed on the basis of their elution positions and measured in glucose units then compared to reference values in NIBRT’s “GlycoBase v3.0 ” database available at http://glycobase.nibrt.ie) for structure assignment [52].

HPLC analysis was performed partly in the National Institute for Biotechnology and Training (NIBRT) in Dublin, Ireland, and partly in the Glycobiology laboratory of Genos Ltd in Zagreb, Croatia. Both laboratories used the same columns and separation conditions. Duplicate analysis of a number of samples was performed and confirmed full reproducibility of the analytical results both within and between laboratories.

Glycan structural features

Levels of glycans sharing the same structural features were approximated by adding the structures having same characteristics: Core fucosylated glycans (FUC-C) = DG6/DG5+DG6*100; Antennary fucosylated glycans (FUC-A) = DG7/DG5+DG7*100.

Genotype and phenotype quality control

Genotyping quality control was performed using the same procedures for all cohorts. Individuals with a call rate less than 97% were removed as well as SNPs with a call rate less than 98% (95% for CROATIA-VIS), minor allele frequency less than 0.02% or Hardy-Weinburg equilibrium p-value less than 1 x 10^-10. Differences in SNP call rate threshold were used to account for observed differences between genotyping arrays. 924 individuals passed all quality control thresholds from CROATIA-VIS, 568 in CROATIA-KORCULA and 263 in ORCADES. An EM based algorithm was used to infer haplotypes from genotypic data. The “scan.haplo” function of the GenABEL package for R [53], which calls the “haplo.score.slide” function of the haplo.stats package for R [57], was used to test a sliding window of 3- and 5-SNP haplotypes across the associated interval. These results were compared to a single SNP model across the same region obtained using the “qtscore” function of the GenABEL package for R. A significant difference between haplotype and single-SNP analysis was determined using the Akaike information criterion [58].

ChiP identification of HNF binding sites in FUT genes

To establish whether HNF1α and HNF4α bind the regulatory elements of the fucosylation genes, their genomic loci were analysed using bioinformatics to identify HNF response elements. Conserved elements between human and mouse genomes [39] were analysed initially to delineate the binding sites of HNF1α and HNF4α using the TRANSFAC database and the ECR browser (http://ecrbrowser.dcode.org/). Primers for ChiP, reChiP and real-time PCR are listed in Text S1.

RNA interference

Production of the RNA duplexes for RNA interference was described in details earlier (Kittler et al., 2005). The target sequences (see Text S1) against HNF1α and HNF4α were designed using the siDESIGN Center (Dharmacon). The Transfection of HepG2 and PANC1 cells was carried out as described by (Yu et al., 2002).

Luciferase assays

The PCR products of ChiP primers (Sequences are detailed in the Text S1) were cloned into pGEM-T easy vector (Promega) and subcloned into pGL4 vectors (Promega) as described earlier (Essafi et al., 2005). pGL4-luc vectors (100 ng) and internal control of pRL-TK (20 ng) renilla plasmid were transiently co-transfected into HepG2 and PANC1 cells (10^5) using the calcium phosphate co-precipitation. Cells were harvested 48 hr post-transfection for luciferase reporter assay using the Dual-Luciferase reporter assay system (Promega). The luciferase activity was normalized by Renilla luciferase activity. All assays were performed in three separate experiments done in triplicate.

ChiP

ChiP was carried out on HepG2 cells essentially as detailed earlier (Essafi et al., 2005). The antibodies used were HNF1α (sc-6547) and HNF4α (sc-6556) from Santa Cruz Biotechnology. The corresponding control IgG antibodies were from Sigma-Aldrich.
Real-time PCR
RNA isolation, cDNA synthesis and Real time PCR were performed as described earlier (Birkenkamp et al., 2007). PCR primer sequences are listed in Text S1.

Supporting Information
Figure S1 PANC1 cells were treated with RNAi against HNF1α and HNF4α. The expression levels of the indicated genes were analysed by real time SYBR Green PCR.
Found at: doi:10.1371/journal.pgen.1001256.s001 (0.28 MB TIF)

Figure S2 PANC1 cells were treated with RNAi against HNF1α and HNF4α. The expression levels of the indicated genes were analysed by real time SYBR Green PCR.
Found at: doi:10.1371/journal.pgen.1001256.s002 (0.41 MB TIF)

Figure S3 The clustering of FUT3, FUT5 and FUT6 on chromosome 19. The position of primer pairs used for ChiP are indicated at the bottom of the figure.
Found at: doi:10.1371/journal.pgen.1001256.s003 (0.39 MB TIF)

Table S1 (A) Main structures present in glycan groups for which Genome-wide significant associations were found (SNPs in FUT8, FUT6 and HNF1α genes). (B) Glycan structures present in different HPLC peaks.
Found at: doi:10.1371/journal.pgen.1001256.s004 (0.10 MB DOC)

Table S2 Desialylated glycan traits (DG1-13, FUC-A, FUC-C) and their associations with FUT6, FUT8 and HNF1α SNPs showing their effect sizes (Beta) in standard deviation units (with standard errors) and p-values. The fucosylation status of each trait is also shown.
Found at: doi:10.1371/journal.pgen.1001256.s005 (0.06 MB DOC)

Text S1 Supplemental Materials and Methods.
Found at: doi:10.1371/journal.pgen.1001256.s006 (0.05 MB DOC)

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Author Contributions
Conceived and designed the experiments: G Lauc, A Essafi, C Hayward, A Knezevic, O Gornik, U Gyllensten, JF Wilson, AF Wright, ND Hastie, H Campbell, PM Rudd, I Rudan. Performed the experiments: A Essafi, JE Huffman, A Knezevic, JJ Kattla, O Polasek, V Vitart, JL Abrahams, M Pucic, M Novokmet, I Redzic, SH Wild, F Borovecki, I Kolcic, L Zgaga. Analyzed the data: G Lauc, A Essafi, JE Huffman, C Hayward, A Knezevic, JJ Kattla, O Polasek, O Gornik, V Vitart, JL Abrahams, M Pucic, M Novokmet, I Redzic, S Campbell, SH Wild, F Borovecki, W Wang, I Kolcic, L Zgaga, U Gyllensten, JF Wilson, AF Wright, ND Hastie, H Campbell, PM Rudd, I Rudan. Contributed reagents/materials/analysis tools: G Lauc, C Hayward, O Polasek, V Vitart, S Campbell, F Borovecki, W Wang, I Kolcic, L Zgaga, U Gyllensten, JF Wilson, AF Wright, ND Hastie, H Campbell, PM Rudd, I Rudan. Wrote the paper: G Lauc, A Essafi, JE Huffman, C Hayward, O Gornik, V Vitart, U Gyllensten, JF Wilson, AF Wright, ND Hastie, H Campbell, PM Rudd, I Rudan. Led the writing of the paper: I Rudan, G Lauc. Performed functional genomic studies: A Essafi. Carried out the genotyping and performed statistical analyses: JE Huffman, C Hayward, V Vitart, S Campbell, SH Wild, F Borovecki. Performed laboratory analyses of glycans: A Knezevic, JJ Kattla, O Gornik, JL Abrahams, M Pucic, M Novokmet, I Rudan. Performed field work and constructed genealogies in Croatia: O Polasek, I Kolcic, L Zgaga. Assisted in writing of the paper and checked the manuscript for important intellectual content: W Wang, U Gyllensten, JF Wilson, AF Wright, ND Hastie, H Campbell, PM Rudd. Designed the study: U Gyllensten, JF Wilson, AF Wright, ND Hastie, H Campbell, PM Rudd, I Rudan, G Lauc.

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