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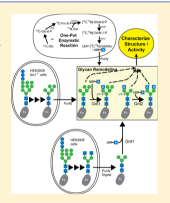
Intramolecular N-Glycan/Polypeptide Interactions Observed at Multiple N-Glycan Remodeling Steps through [13C,15N]-N-Acetylglucosamine Labeling of Immunoglobulin G1

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Supporting Information

ABSTRACT: Asparagine-linked (N) glycosylation is a common eukaryotic protein modification that affects protein folding, function, and stability through intramolecular interactions between Nglycan and polypeptide residues. Attempts to characterize the structure-activity relationship of each N-glycan are hindered by inherent properties of the glycoprotein, including glycan conformational and compositional heterogeneity. These limitations can be addressed by using a combination of nuclear magnetic resonance techniques following enzymatic glycan remodeling to simultaneously generate homogeneous glycoforms. However, widely applicable methods do not yet exist. To address this technological gap, immature glycoforms of the immunoglobulin G1 fragment crystallizable (Fc) were isolated in a homogeneous state and enzymatically remodeled with [13C, 15N]-N-acetylglucosamine (GlcNAc). UDP-[13C, 15N]GlcNAc was synthesized enzymatically in a one-pot reaction from [13C]glucose and [15N-amido]glutamine. Modifying Fc with recombinantly expressed glycosyltransferases (Gnt1 and Gnt2) and UDP-[13C, 15N]-GlcNAc resulted in complete glycoform conversion as judged by mass spectrometry. Twodimensional heteronuclear single-quantum coherence spectra of the Gnt1 product, containing a



single [13C,15N]GlcNAc residue on each N-glycan, showed that the N-glycan is stabilized through interactions with polypeptide residues. Similar spectra of homogeneous glycoforms, halted at different points along the N-glycan remodeling pathway, revealed the presence of an increased level of interaction between the N-glycan and polypeptide at each step, including mannose trimming, as the N-glycan was converted to a complex-type, biantennary form. Thus, conformational restriction increases as Fc N-glycan maturation proceeds. Gnt1 and Gnt2 catalyze fundamental reactions in the synthesis of every glycoprotein with a complex-type N-glycan; thus, the strategies presented herein can be applied to a broad range of glycoprotein studies.

Protein asparagine-linked (N) glycosylation is a cotranslational event that confers a wide range of properties to the underlying polypeptide, including enhanced folding and stability, favorable pharmacokinetic properties, decoration with specific epitopes central to function (reviewed in ref 1), and allosteric modulation of protein function.² Many of these properties can be attributed to intramolecular interactions between N-glycan and polypeptide epitopes.³ Investigations of the structure-activity relationships of glycoprotein glycans must surmount two challenges: glycan compositional and conformational heterogeneity.

Unlike the template-dependent biosynthesis of nucleic acids or proteins, N-glycans are synthesized, ligated, and remodeled by glycosylhydrolases and glycosyltransferases that operate without a template. This complex biosynthesis generates significant compositional heterogeneity (reviewed in refs 1 and 4). Immature N-glycans with a high mannose (Man) content are transferred to the nascent polypeptide chain during import into the lumen of the endoplasmic reticulum from lipidlinked donor molecules. For glycans destined to be complextype, a single Man residue is removed followed by export to the Golgi, where three more Man residues are removed to form a Man5 structure (Figure 1). In the next step, an N-acetylglucosamine (GlcNAc) residue forms the base of the first N-glycan branch upon addition to the C2 hydroxyl of the $Man\alpha 1-3$ residue by the glycosyltransferase enzyme Gnt1. The final hydrolysis step removes two of the five remaining Man residues. Gnt2 then adds a GlcNAc residue to the C2 hydroxyl of the remaining $Man\alpha 1-6$ residue to form the base of a second branch. The steps that follow do not proceed to completion for every glycan, resulting in significant heterogeneity. The most common modifications are fucosylation, synthesis of additional GlcNAc branches, and transfer of galactose (Gal) and sialic acid residues. Lastly, the glycoprotein is exported to the cell surface where glycosyltransferase- or glycosylhydrolase-mediated modifications may occur.

Glycoengineered proteins, including antibodies and antibody fragments, are of great interest because of the potential to enrich them with glycoforms with elevated therapeutic properties⁵ or to obtain homogeneous preparations for detailed studies. Many routes that show promising results have been reported, including, but not limited to, purely synthetic

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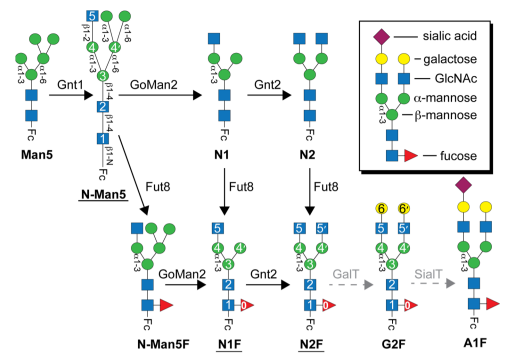


Figure 1. Native IgG1 Fc *N*-glycan processing in the Golgi. Conversions catalyzed by the enzymes indicated above the solid arrows and labeled with black type largely proceed to completion. Reactions catalyzed by enzymes denoted with gray type a dashed arrow modify some but not all of the secreted IgG1. Glycoforms studied here by nuclear magnetic resonance are underlined. Carbohydrate residues are numbered according to ref 30 and represented using the CFG convention and shown in the inset⁴⁹ (GlcNAc, *N*-acetylglucosamine). Glycosidic linkages of the human IgG1 Fc *N*-glycan are indicated.

methods, $^{6-8}$ enzymatic remodeling of the glycan termini *in vitro* with sugar nucleotides, $^{9-12}$ approaches to enzymatically transfer a synthetic carbohydrate, $^{13-16}$ and genetic manipulation of expression host organisms. $^{17-24}$ Each method faces its unique challenges, and a single method for each situation has not been found.

N-Glycans are mobile moieties that often inhibit crystallization. Thus, N-glycans are enzymatically removed for studies of protein structure as a matter of routine. Solution nuclear magnetic resonance (NMR) spectroscopy offers a significant advantage in that proteins, highly concentrated in a buffered solution, are not required to crystallize and can be studied with complete carbohydrates (reviewed in ref 3). Solution NMR spectroscopy offers another advantage over solid-state methods: the entire ensemble of interconverting conformations contributes to the observable signals. While X-ray crystallography is adept at capturing high-resolution images of low-energy conformations at low temperatures, NMR can capture subtle shifts in conformational equilibria. NMR methods for studying N-glycans often incorporate ¹³C-labeled sugars into N-glycans with glycosyltransferase enzymes after purification and have utilized terminal galactose and sialic acid transfer reactions, ^{25–27} though an expression-based method to label the entire glycan is also available. 28,29

This work describes a novel *in vitro* route to UDP-[\(^{13}\text{C}\),\(^{15}\text{N}\)]GlcNAc and stepwise, site-specific labeling of the immunoglobulin G1 (IgG1) fragment crystallizable (Fc) *N*-glycan. One significant advantage of this approach is facile assignment of NMR resonances due to the stepwise labeling approach. This permits an assessment of *N*-glycan structures isolated at multiple crucial points along the *N*-glycan remodeling pathway as it would occur in the Golgi. The nomenclature used to describe carbohydrate residues of the Fc

N-glycan is highlighted in Figure 1.³⁰ For example, "(5')-GlcNAc" refers to the *N*-acetylglucosamine residue at the nonreducing end of the GlcNAc β 1–2Man α 1–6Man β 1–4GlcNAc β 1–4GlcNAc β -N moiety.

IgG1 is a critical defense protein that recognizes specific pathogen epitopes through tight binding, antigen recognition domains and triggers a pro-inflammatory, pathogen-destroying immune response through the Fc region. Interactions between IgG1 Fc and certain cell surface receptors, the lowaffinity Fc γ receptors, require Fc N-glycosylation at Asn297 of the Fc C γ 2 domain. Transient, intramolecular interactions between the Fc N-glycan and polypeptide are central to this requirement, though the structure—activity relationship of this phenomenon remains undefined. This report covers new methods for addressing this relationship that are broadly applicable to structure—function studies of N-glycans.

■ EXPERIMENTAL PROCEDURES

Materials. All materials were purchased from Sigma-Aldrich unless otherwise noted. Structure figures were prepared using PyMOL (Schrödinger LLC).

Protein Expression. An expression plasmid encoding the GlmS enzyme (glutamine:fructose-6-phosphate transaminase, EC 2.6.1.16) from *Escherichia coli* was prepared by amplifying and cloning the GlmS open reading frame from the pMA1 phagemid³⁶ into the *Nco*I and *Xho*I restriction sites of the pET21d plasmid (Merck Millipore). The final cloned open reading frame encoded GlmS with N- and C-terminal tags: M +A+C2-E609+LEHHHHHHH. Plasmid preparation was verified by DNA sequencing (Iowa State University DNA Facility). GlmS was expressed in transformed *E. coli* BL21star(DE3) cells carrying the GlmS:pET21d vector in the presence of ampicillin (50 mg/L). Expression was induced with 0.5 mM isopropyl β-

D-1-thiogalactopyranoside (IPTG) once the culture density reached an OD₆₀₀ of 0.7, and cells were incubated for 20 h at 18 °C in an orbital shaking incubator. Cells were harvested in 50 mL aliquots by centrifugation; the spent medium was decanted and the pellet frozen and stored at -80 °C. Cells from a single frozen aliquot were lysed in 10 mL of 25 mM 4morpholinepropanesulfonic acid (MOPS), 100 mM sodium chloride, 5 mM β -mercaptoethanol, and 1 mM ethylenediaminetetraacetic acid (EDTA) (pH 7.2), with four or five passages through an EmulsiFlex-C5 homogenizer (Avestin) operating at 15000 psi. Insoluble debris was removed by centrifugation at 25000g for 1 h. The clarified lysate was washed extensively in a 10 kDa cutoff Amicon centrifugal filter unit in the same buffer (without EDTA) to remove salts and concentrated to ~1 mL. GlmS-containing washed cell lysate was prepared fresh for each reaction. GlmS proved to be unstable during purification, and removing the C-terminal six-His tag failed to improve stability.

A vector containing the GlmM (phosphoglucosamine mutase, EC 5.4.2.10) open reading frame from Bacillus anthracis cloned into the pDEST17 plasmid³⁷ was transformed into E. coli BL21star(DE3) cells. Protein expression was induced after a culture grown at 37 °C in Luria-Bertani medium reached an OD₆₀₀ of 0.6 with 0.5 mM IPTG. Cells were incubated for 20 h at 18 °C in an orbital shaking incubator and then separated from the growth medium by centrifugation. Cells were lysed using the homogenizer described above in a buffer containing 50 mM 2-amino-2-hydroxymethylpropane-1,3-diol (Tris), 200 mM sodium chloride, and 10 mM imidazole (pH 8.2) and then centrifuged at 25000g for 1 h to remove insoluble debris. Clarified lysate containing GlmM was loaded directly on a Ni²⁺-NTA column (Qiagen) using a Biologic LP chromatography system (Bio-Rad) and eluted with a linear gradient from 10 to 250 mM imidazole in the same buffer. Fractions containing GlmM were pooled, concentrated, and loaded on a Superdex 200 column (GE Healthcare) equilibrated with 25 mM Tris and 100 mM sodium chloride (pH 8.2). GlmM eluted as a sharp peak. Positive fractions were pooled and concentrated to 250 μ M GlmM as judged by A_{280} , diluted to 25% (v/v) glycerol, and frozen at −80 °C.

A vector containing the open reading frame encoding the bifunctional GlmU [glucosamime-1-phosphate-N-acetyltransferase (EC 2.3.1.157) and UDP-N-acetylglucosamine diphosphorylase (EC 2.7.7.23)] from E. coli cloned into the pET21b plasmid³⁸ was transformed into E. coli BL21star(DE3) cells. Protein expression was performed largely as described from GlmM, except cells were lysed with a buffer containing 50 mM Tris, 500 mM sodium chloride, and 10 mM imidazole (pH 8.2); gel-filtration chromatography was performed in a buffer containing 10 mM Tris, 100 mM sodium chloride, and 5 mM β -mercaptoethanol (pH 8.2). GlmU eluted as a sharp peak. Positive fractions were pooled and concentrated to 190 μ M GlmU as judged by A_{280} , diluted to 25% (v/v) glycerol, and frozen at -80 °C.

IgG1 Fc was expressed in HEK293F and HEK293S($lec1^{-/-}$) cells and purified as previously described.² The luminal domains of Gnt1 and Gnt2 were expressed as GFP fusions at the N-terminus using the pGen2 vector¹² in HEK293F or HEK293S($lec1^{-/-}$) cells in the same manner. Crude expression medium was passed over a Ni²⁺-NTA column using gravity. Next, the column was washed with 12 column volumes of 50 mM Tris, 500 mM sodium chloride, and 30 mM imidazole (pH 8.0) and then eluted with the same buffer containing 250 mM

imidazole. Fractions containing the desired protein were washed extensively with 50 mM 2-(N-morpholino)-ethanesulfonic acid¹⁰ and 100 mM potassium chloride (pH 6.25), concentrated with a 10 kDa cutoff centrifugal filter to 25–40 μ M protein as judged by A_{280} , diluted to 50% (v/v) glycerol, and frozen at -80 °C.

One-Pot UDP-[13C,15N]GlcNAc Synthesis and Partial **Purification.** Buffer, small molecule, and protein components were combined in a single tube to final concentrations of 100 mM MOPS (pH 7.2), 100 mM sodium chloride, 5 mM magnesium chloride, 1 mM dithiothreitol, 1 mM α -D-glucose 1,6-bisphosphate, 5 mM $_{D}$ -[$^{13}C_{U}$]glucose, 10 mM adenosine triphosphate, 25 mM [15N-amido]glutamine, 10 mM acetylcoenzyme A, 10 mM uridine triphosphate, 17 units of hexokinase (from Saccharomyces cerevisiae), 18 units of phosphoglucose isomerase (from S. cerevisiae), a 10% (v/v) final concentration of washed cell lysate containing recombinant EcGlmS, 4 µM BaGlmM, and 2.4 µM EcGlmU in water. The reaction mixture was incubated at room temperature for 24 h in a closed tube in the dark. Reaction purification was performed on the basis of a published method.³⁹ Briefly, reaction products were applied to a diethylaminoethylenecellulose column equilibrated in 10 mM triethylammonium bicarbonate (TEAB) (pH 7.0). The column was washed with 2 column volumes of 10 mM TEAB and 4 column volumes of 50 mM TEAB and eluted with 4 column volumes of 100 mM TEAB. Fractions from the 100 mM elution were pooled, lyophilized, resuspended in H2O, and lyophilized again to remove the remaining TEAB. The concentration of the purified material was determined by comparing ¹H signals of the compound to that of an internal 0.5 mM 4,4-dimethyl-4silapentane-1-sulfonic acid (DSS) standard.

Preparing N-Man5 *N***-Glycans.** IgG1 Fc (13 mg/mL) expressed in HEK293S($lec1^{-/-}$) cells was incubated with 50 mM MES (pH 6.25), 100 mM potassium chloride, 20 mM manganese chloride, 1 mM UDP-[13 C, 15 N]GlcNAc, and 5 μ M Gnt1 at room temperature for 24 h in the dark. The reaction mixture was exchanged into an NMR buffer [10 mM MOPS (pH 7.2), 100 mM potassium chloride, and 0.5 mM DSS in >98% deuterium oxide] using a 10 kDa cutoff centrifugal concentrator.

Preparing N1F and N2F N-Glycans. IgG1 Fc (14 mg/ mL) expressed in HEK293F cells was incubated with 50 mM sodium citrate and 40 units of an N-acetylglucosaminidase (New England Biolabs) for 48 h at 37 °C. IgG1 Fc was then purified using a Protein A column as described previously.³⁰ Next, a Gnt1-catalyzed reaction was performed as described above to incorporate a single [13C,15N]GlcNAc residue. The second residue was added using 6 mg/mL IgG1 Fc, 50 mM MES (pH 6.25), 100 mM potassium chloride, 20 mM manganese chloride, 1 mM UDP-[13C, 15N]GlcNAc, and 4 μM Gnt2 at room temperature for 48 h in the dark. The reaction mixture was exchanged into the same NMR buffer described above using a 10 kDa cutoff centrifugal concentrator. Reactions were monitored by permethylation and matrixassisted laser desorption ionization mass spectrometry (MALDI-MS) analysis at each step as previously described 40 using a Voyager-DE PRO instrument (Applied Biosystems).

Endoglycosidase Treatments To Cleave N-Glycans. A pET:GFP-EndoF1 plasmid for expression in *E. coli* was provided by K. Moremen (University of Georgia, Athens, GA) and expressed and purified using standard protocols (Qiagen). Purified EndoF1 ($10~\mu M$) was added to $60~\mu M$ IgG1

Fc in a 50 mM phosphate buffer (pH 6.0) and incubated for 12 h at 37 $^{\circ}$ C.

NMR Spectroscopy. NMR spectra were recorded using 5 mm Shigemi NMR tubes in a spectrometer equipped with a cryogenically cooled probe and an Avance II console (Bruker) and operating at 50 °C and 16.4 T. Fc dimer concentrations were between 60 and 100 μ M in a final volume of 300 μ L. The pulse sequence for the $^{1}H-^{13}C$ heteronuclear single-quantum coherence (HSQC) spectra of Fc did not include a sensitivity enhancement element or coherence selection gradients to minimize the loss of broad peaks. Data were analyzed using Topspin (version 3.2), NMRviewJ (One Moon Scientific), and NMRPipe. Chemical shifts were referenced directly (^{1}H) and indirectly (^{13}C and ^{15}N) to the internal DSS methyl peak at 0.07 ppm (^{1}H).

■ RESULTS AND DISCUSSION

One-Pot Synthesis of UDP-[13C,15N]-N-Acetylglucosamine. Synthesis of UDP-GlcNAc from glucose by eukaryotic and prokaryotic organisms proceeds along similar metabolic pathways; however, the prokaryotic system includes a bifunctional enzyme (GlmU) that catalyzes the last two steps (Figure 2A).^{42–44} Because of its relative simplicity, the prokaryotic

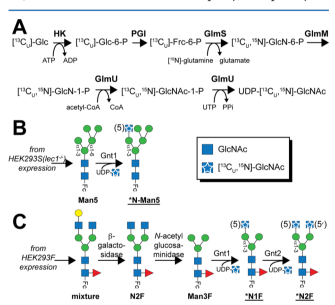


Figure 2. Schemes for *in vitro* enzymatic conversions described in this study. (A) A one-pot synthesis of UDP-[\begin{small}^{13}\text{C},\begin{small}^{15}\text{N}\end{small} GlcNAc utilizes enzymes from bacterial pathways and [\begin{small}^{13}\text{C}]glucose. Carbohydrate remodeling started with Fc bearing either a mannose-type (B) or a complex-type (C) *N*-glycan. [\begin{small}^{13}\text{C},\begin{small}^{15}\text{N}\end{small} GlcNAc is shown as a blue square with a white star in the cartoon figures and by "*N" in the glycan name; residue numbers corresponding to the convention introduced in Figure 1 are given in parentheses.

pathway was recapitulated *in vitro* using a combination of commercially available and laboratory-expressed enzymes with off-the-shelf metabolites, including $[^{13}\mathrm{C}_{\mathrm{U}}]$ glucose and $[^{15}\mathrm{N}-amido]$ glutamine. Recovery of the starting $[^{13}\mathrm{C}_{\mathrm{U}}]$ glucose in the form of UDP- $[^{13}\mathrm{C},^{15}\mathrm{N}]$ GlcNAc, following purification, was $\sim 18\%$ (moles of UDP- $[^{13}\mathrm{C},^{15}\mathrm{N}]$ GlcNAc per mole of $[^{13}\mathrm{C}_{\mathrm{U}}]$ glucose). A higher conversion would likely be achieved by increasing the concentrations of certain donor substrates, including $[^{15}\mathrm{N}-amido]$ glutamine and acetyl-CoA; however, the cost of greater metabolite concentrations is unlikely to be offset

by the increased yield. A 1 mL reaction mixture, starting with 1 mg of $[^{13}C_U]$ glucose, produced 0.55 mg of UDP- $[^{13}C,^{15}N]$ -GlcNAc. This is enough UDP- $[^{13}C,^{15}N]$ GlcNAc for the Gnt1-catalyzed labeling of at least 13 mg of IgG1 Fc [a typical sample for two-dimensional (2D) NMR analysis contains 1 mg of IgG1 Fc].

A ¹H–¹³C heteronuclear single-quantum coherence (HSQC) spectrum of the purified material revealed chemical shifts that were identical to database values for UDP-GlcNAc⁴⁵(Figure S1 of the Supporting Information). A ¹H–¹⁵N HSQC spectrum also showed a single, intense peak for the amide moiety (Figure S1B of the Supporting Information). This preparation proved to be sufficiently pure for glycosyltransferase-catalyzed sugar additions even though minor impurities were observed in the NMR spectra. This one-pot synthetic route permits an additional labeling strategy to incorporate a ¹³C-labeled acetate moiety from acetyl-CoA. Methyl groups have beneficial spin relaxation properties in magnetic resonance applications that permit measurements on very large (~1 MDa) or dynamic systems. ^{46,47} Furthermore, the synthesis of the methyl donor in this reaction, (¹³C-acetyl)-CoA, is easily achieved. ⁴⁸

Beyond the applications to *N*-glycans, UDP-[¹³C,¹⁵N]-GlcNAc is a "gateway" nucleotide sugar that can be converted to many other products, too diverse to completely describe here. Potential immediate applications include using UDP-[¹³C,¹⁵N]GlcNAc as a substrate in glycosaminoglycan or sialic acid biosynthesis.⁴⁹ Alternatively, a simple single enzymecatalyzed epimerization leads to UDP-[¹³C,¹⁵N]-*N*-acetylgalactosamine⁵⁰ that serves as the basis of eukaryotic *O*-GalNAc glycans (including mucins and many others) and is critical for glycosaminoglycan and glycosphingolipid biosynthesis.⁴⁹

The enzymatic method for preparing UDP-GlcNAc described here offers marked benefits over previously described methods with respect to isotope labeling for NMR and MS-based studies. Other enzymatic methods for synthesizing UDP-GlcNAc that start from GlcNAc or GlcN have been described. However, isotope-labeled GlcNAc and GlcN can be cost prohibitive and are available with only limited labeling patterns, unlike the scheme presented here that can be used to produce a wide array of custom labeling patterns starting with inexpensive starting materials, including glucose and glutamine. Numerous chemical methods are also available and can be adapted for the synthesis of GlcNAc analogues; however, these methods are less efficient than the one-pot enzymatic method presented here (not limited to refs 56–58).

Gnt1-Catalyzed Conversion of the Man5 *N*-Glycan. Addition of a GlcNAc residue in a β 1–2 linkage to the (3)Man residue, catalyzed by Gnt1 (also known as lec1 or MGAT1), is a crucial step in hybrid and complex-type *N*-glycan maturation (Figure 1). A Gnt1-deficient HEK293 cell line (HEK293S or $lec1^{-/-}$) halts this process and produces glycoproteins with nearly homogeneous Man5 *N*-glycans. Glycans on IgG1 Fc expressed using this cell line are thus auspicious substrates for investigating Gnt1 activity *in vitro*. As expected on the basis of published reports, S1,62 MS analysis of IgG1 Fc incubated with UDP-GlcNAc and Gnt1 revealed complete conversion of Fc-Man5 to the Fc-N-Man5 glycoform (Figure 3; experimental mass of 1825.3 Da, observed mass of 1824.9 Da).

Enzymatic Conversion to a Complex-Type Fc N-Glycan. HEK293F cells, unlike the HEK293S ($lec1^{-/-}$) cells, have the capacity to generate complex-type polysaccharides and express Fc with a fucosylated biantennary N-glycan that varies with respect to the amount of terminal galactose incorporated

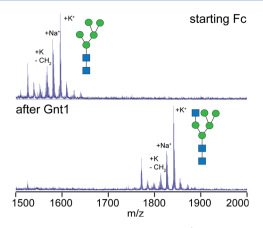


Figure 3. Gnt1-catalyzed remodeling of $lec1^{-/-}$ -expressed IgG1 Fc with a Man5 N-glycan as monitored by MALDI-MS. Enzymatic remodeling occurs when the N-glycans are attached to the Fc polypeptide; however, the analysis shown here includes N-glycan removal followed by permethylation.

(Figure 4). Exoglycosidase treatment removed terminal sugars and resulted in Fc with a primarily Man3F glycan (Figure 4; experimental mass of 1345.7 Da, observed mass of 1345.5). This material was then used as a substrate for modification by Gnt1, the result being Fc containing the (5)-[¹³C,¹⁵N]GlcNAc residue (*N1F; Figures 2C and 4; experimental mass of 1597.8 Da, observed mass of 1597.3 Da). The fact that Gnt1 could

recognize a Man3F substrate was unknown; Gnt1 was shown to modify a polysaccharide with an identical display of three Man residues tethered through the β -Man to a GlcNAc, ^{61,62} but because the Fut8 core fucosyltransferase modifies N-glycans only after Gnt1 *in vivo* ⁶³ (Figure 1), it was unknown if the presence of the (0)fucose(Fuc) residue would prevent this modification *in vitro*.

The *N2F glycoform was prepared from the *N1F material following a reaction catalyzed by Gnt2, which likewise forms a GlcNAc β 1–2Man linkage, except Gnt2 modifies the (3')Man residue rather than the (3)Man modified by Gnt1 (Figures 2C and 4; experimental mass of 1850.0 Da, observed mass of 1850.1 Da). On the basis of MALDI-MS analysis, the Gnt1-and Gnt2-catalyzed reactions proceeded nearly to completion [>95% conversion (Figure 4)].

This approach represents a significant advance to *in vitro* enzymatic *N*-glycan remodeling. In contrast to the methods described in the introductory section, this method rebuilds *N*-glycans from a paucimannose (Man3) core *N*-glycan that is present in all eukaryotic *N*-glycans and permits incorporation of ¹³C or ¹⁵N labels at each step. Thus, it is expected that all eukaryotic *N*-glycans could be remodeled in this manner with suitable exoglycosidases, many of which are commercially available. The robust nature of this approach is reflected in the high conversion of the IgG1 Fc glycoprotein, a notably difficult protein to remodel enzymatically. ^{11,12,64} Furthermore, this method utilizes commercially available sugar nucleotides (if stable isotope-enriched sugars are not required) in place of

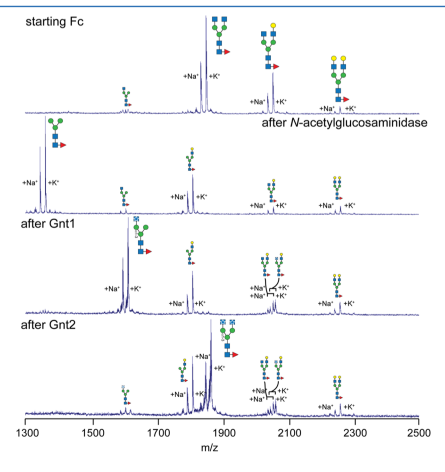


Figure 4. Generation of *N1F and *N2F Fc glycoforms from HEK293F-expressed Fc was confirmed using MALDI-MS analysis. These steps are shown in Figure 2C. Enzymatic remodeling occurs when the *N*-glycans are attached to the Fc polypeptide; however, the analysis shown here includes *N*-glycan removal followed by permethylation.

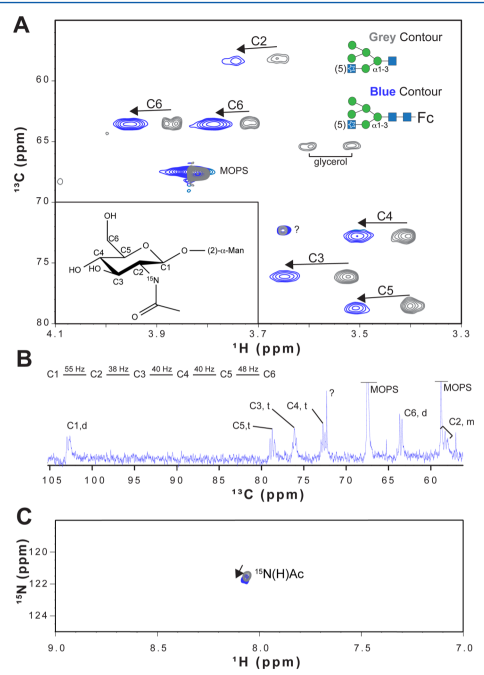


Figure 5. $^{1}H-^{13}C$ HSQC spectra of IgG1 Fc with a Man5 N-glycan following addition of $[^{13}C,^{15}N]$ GlcNAc. (A) A 2D $^{1}H-^{13}C$ HSQC spectrum of the *N-Man5 N-glycan following EndoF1-catalyzed hydrolysis is shown as gray contours. Blue contours show the positions of peaks from IgG1 Fc bearing a *N-Man5 N-glycan. Arrows indicate the direction of peak movement because of interactions with the Fc polypeptide. Peak labels that correspond to a figure of β-linked GlcNAc are shown (inset) and refer to the carbon position of the $^{1}H-^{13}C$ peak. $^{1}J_{C-C}$ couplings are not resolved because of the limited resolution in the ^{13}C dimension. (B) 1D ^{13}C -observe NMR spectrum of *N-Man5 Fc. $^{1}J_{C-C}$ values are indicated. (C) 2D $^{1}H-^{15}N$ HSQC spectra before and after N-glycan hydrolysis with the same colors used in panel A.

synthetic oligosaccharide precursors for the case of transglycosylation (see ref 64).

NMR Analysis of [¹³C,¹⁵N]GlcNAc-Labeled Fc Glycoforms. NMR spectra of IgG1 Fc following Gnt1-catalyzed remodeling of the Man5 glycan using UDP-[¹³C,¹⁵N]GlcNAc revealed peaks for each ¹H-¹³C and ¹H-¹⁵N moiety (Figure 5). The peaks were relatively intense and narrow, considering the glycans are part of the ~55 kDa Fc. This property indicates the presence of significant GlcNAc motion relative to the polypeptide domains and is consistent with similar measurements of galactose and sialic acid residues on the 3-4-5

branch of the *N*-glycan [as opposed to the 3'-4'-5' branch (Figure 1)]. 12,25,34 $^{1}J_{\mathrm{C-C}}$ couplings from a one-dimensional (1D) $^{13}\mathrm{C}$ -observe experiment agreed with the resonance assignments based on an assignment of β -GlcNAc (Figure 5B). Similar spectra were observed at 25, 37, and 50 °C, and peak positions in duplicate samples were reproduced.

The Fc polypeptide influences the position and line shapes of GlcNAc peaks. Endoglycosidase F1 treatment hydrolyzed the glycosidic linkage between the (1)GlcNAc and (2)GlcNAc residues, releasing the glycan from its covalent attachment to Fc. As a result, resonances of the released *N*-glycan were

different from those of the Fc-conjugated material [average Δ^{1} H = 0.098 ppm (Figure 5)], even though the Fc polypeptide was still present in the experiment. It was previously shown that the N-glycan termini (residues 6 and 6') exchange between a restricted and free state on a microsecond time scale, which results in a single observable peak in NMR spectra that represents the population-weighted average of the two states.³⁴ Mutating the polypeptide surface to prevent interaction resulted in the predominance of the free state.² On the basis of these reports, it is likely that peaks corresponding to the (5)GlcNAc residue also represent the population-weighted average of two distinct states. Thus, the farther the peak is shifted from that of the free state (seen with the hydrolyzed Nglycan), the greater the restriction mediated by intramolecular interactions between N-glycan and polypeptide residues. It is not clear how the Fc polypeptide influences the frequencies of (5)GlcNAc resonances in the restricted state, though electric field effects likely contribute. 65 Multiple charged surface residues lie in the proximity of the (5)GlcNAc residue, including lysine 334, which is within 8 Å (Figure S2 of the Supporting Information).

A 2D ¹H-¹³C spectrum of the *N1F glycoform reveals a pattern of peaks similar to that observed with *N-Man5 (Figure 6, red and blue contours, respectively). A spectrum of the *N2F

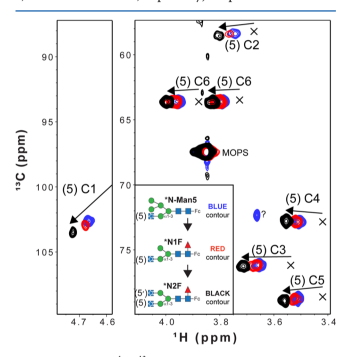


Figure 6. Overlay of $^{1}H^{-13}C$ HSQC spectra collected with purified IgG1 Fc glycoforms that reveals shifts of peaks away from that of a hydrolyzed N-glycan. The positions of peaks from an Fc N-glycan following EndoF-catalyzed hydrolysis are indicated with X's. Arrows show the direction of peak movement as the N-glycan matures. The C1 peak for the hydrolyzed glycan was obscured by residual water in the sample and was not observed.

glycoform was likewise similar, but not identical, with respect to the position of peaks corresponding to the (5)GlcNAc residue (Figures 6 and 7, black contour). As the glycoform advanced from *N-Man5 to *N1F to *N2F, the deviation of the peak positions as compared to the released glycan (shown with an "X" in Figure 6) likewise increased. A small shift in peak positions was observed between spectra of *N-Man5 and *N1F

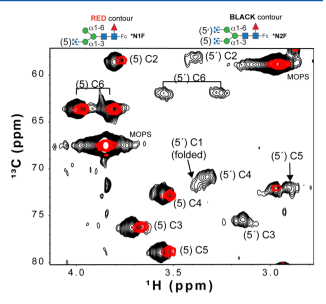


Figure 7. Broad, low-intensity peaks appear in a ${}^{1}H-{}^{13}C$ HSQC spectrum following Gnt2-catalyzed addition of a second $[{}^{13}C, {}^{15}N]$ -GlcNAc. The resonance assignments for the (5')GlcNAc C3, C4, and C5 peaks were obtained by comparison to a report by Yamaguchi et al. 28

(average $\Delta^1 H = 0.016$ ppm), though a larger shift was seen between *N1F and *N2F (average $\Delta^1 H = 0.034$ ppm). This indicates the (5')GlcNAc residue (*N1F \rightarrow *N2F) had a greater effect on resonance frequency than removing the two Man residues [*N-Man5 \rightarrow *N1F (see Figure 6)]. Changes in resonance frequencies of the (5)GlcNAc residue likely reflect changes in structure of the *N*-glycan. Peaks observed with the *N-Man5, *N1F, and *N2F Fc glycoforms shifted away, in a stepwise manner, from peaks observed in spectra of a hydrolyzed *N*-glycan. This shift indicated that *N*-glycan conformation becomes more restricted as maturation, mediated by glycan-modifying enzymes in the Golgi, proceeds. This conclusion is supported by previous work showing similar directions of chemical shift changes for galactose and sialic acid resonances upon temperature changes. ^{12,34}

Peak line widths also reflected changes in the N-glycan. NMR line widths are a direct reflection of transverse relaxation rates of nuclear orientations in an NMR experiment. These values decrease with the tumbling rate of the nucleus and can also be increased by conformational exchange (R_{ex}) on the microsecond to millisecond time scale (contributions from magnetic field inhomogeneity and chemical shift anisotropy likewise increase line widths but are not considered here).66 Line widths of (5)GlcNAc resonances from the ¹H-¹³C HSQC spectra, measured without applying a line-broadening function during processing, increased following attachment of the N-glycan to Fc and indicated structural differences between the different glycoforms. The line width of (5)GlcNAc H6, after removing the effect of ^{2&3}J_{HH} coupling, showed a representative response and increased from 7 Hz (hydrolyzed *N-Man5) to 14 Hz (*N-Man5-Fc) to 19 Hz (*N1F-Fc) to 14 Hz (*N2F-Fc) as shown in Figure S3 of the Supporting Information. These values are much smaller than the expected value of 35 Hz for the same nucleus at the same magnetic field strength tumbling with an apparent molecular mass of 55 kDa. The value of 35 Hz is based on contributions of nearby nuclei to the line width and excludes effects of chemical shift anisotropy (CSA) and $R_{\rm ex}$. ⁶⁶ It

is unclear why the values measured from the *N1F glycoform are larger than those of the *N-Man5 and *N2F forms, but this may reflect changes in exchange rates.

Interactions between the N-glycan and polypeptide surface can explain both the peak position and line width differences in the spectra of Fc glycoforms. Phe241 is located adjacent to the β -Man residue in Fc models determined by X-ray crystallography 67,68 (Figure 8) and was shown to restrict N-

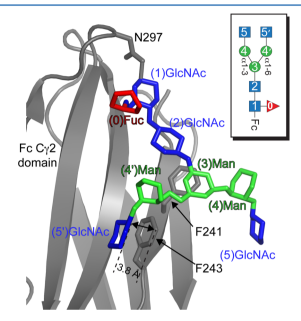


Figure 8. Structural model of the IgG1 Fc–*N*-glycan interface (based on Protein Data Bank entry 4ku1⁷¹). F243 directly contacts the (5')GlcNAc residue. The (5)GlcNAc does not appear to make a direct contact with the polypeptide surface. Carbohydrate residues are shown in a stick model with the color of each residue corresponding to the key for each residue shown in Figure 1. Only ring atoms of the carbohydrates are shown for the sake of simplicity, and galactose residues present in the original Protein Data Bank model are not shown for the sake of clarity.

glycan motion.² Covalent attachment of the *N*-glycan through Asn297 places the glycan in a prime position to make this crucial contact. A smaller shift is observed between the *N-Man5 and *N1F glycoforms as two Man residues are removed and a (0)Fuc is added (Figure 1). The Fuc is not expected to influence movement of the *N*-glycan branch termini greatly because it does not appear to interact with the Fc polypeptide.⁶⁹ Man residues interfere with contacts along the polypeptide surface and likely result in a relatively small stabilization upon removal. Bowden et al.⁷⁰ observed the C2 hydroxyl of the Man residue at the nonreducing terminus of the Man α 1-6Man β moiety in the Man5 *N*-glycan (see Figure 1) was positioned to prevent an interaction with F243 and shifted the entire *N*-glycan away from the polypeptide surface.

The (5')GlcNAc, however, can adopt a conformation suitable for contact with F243^{2,67,68} (Figure 8). This is supported by the relatively greater shift of (5)GlcNAc peaks following Gnt2-catalyzed glycan modification, and in the appearance of the (5')GlcNAc resonance that appeared to be broad and weak (¹H6 line width of ~67 Hz) >0.5 ppm (¹H) upfield of the (5)GlcNAc peaks (Figure 7). A ¹H–¹⁵N HSQC spectrum of the *N2F glycoform likewise shows two distinct peaks corresponding to acetamide moieties of the 5 and 5'

[¹³C,¹⁵N]GlcNAc residues (Figure S4 of the Supporting Information). This dramatic shift of all (5')GlcNAc resonances is consistent with close contact of the (5')GlcNAc residue with the polypeptide for at least some period of time. NMR spectra of the (6')-¹³C-Gal residue of Fc with a G2F glycoform (Figure 1) showed similar peak displacements and reduced intensities, which was due to a transient interaction with the polypeptide surface.³⁴ Furthermore, line widths and positions of these peaks from Fc with double F241S and F243S mutations overlapped with an *N*-glycan on a trypsinized Fc glycopeptide.²

SUMMARY

A one-pot, purely enzymatic method was used to produce UDP-[\$^{13}C,\$^{15}N]GlcNAc efficiently from [\$^{13}C]glucose and [\$^{15}N]glutamine. IgG1 Fc \$N\$-glycans were enzymatically remodeled from a natural immature form and an unnatural core \$N\$-glycan, common to all eukaryotic \$N\$-glycans, to high conversion using UDP-[\$^{13}C,\$^{15}N]GlcNAc. NMR spectra recorded at different points along the \$N\$-glycan remodeling pathway reveal enhanced interactions of the carbohydrate with the polypeptide surface. These small changes in equilibria cannot be observed by \$X\$-ray crystallography. These methods are broadly applicable beyond IgG1 Fc and provide spectroscopic probes for characterizing \$N\$-glycan structure and motion.

ASSOCIATED CONTENT

S Supporting Information

2D $^{1}H-^{13}C$ and $^{1}H-^{15}N$ HSQC spectra of UDP- α -D- $[^{13}C,^{15}N]$ GlcNAc, Fc model showing the location of charged residues, representative line shapes and peak information extracted from 2D NMR measurements, and a $^{1}H-^{15}N$ HSQC-TROSY spectrum of ($^{15}N-Y;2\times[^{13}C,^{15}N]$ -GlcNAc)-IgG1 Fc (*N2F glycoform). This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

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