Global site-specific analysis of glycoprotein N-glycan processing

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N-glycans contribute to the folding, stability and functions of the proteins they decorate. They are produced by transfer of the glycan precursor to the sequon Asn-X-Thr/Ser, followed by enzymatic trimming to a high-mannose-type core and sequential addition of monosaccharides to generate complex-type and hybrid glycans. This process, mediated by the concerted action of multiple enzymes, produces a mixture of related glycoforms at each glycosite, making analysis of glycosylation difficult. To address this analytical challenge, we developed a robust semiquantitative mass spectrometry (MS)-based method that determines the degree of glycan occupancy at each glycosite and the proportion of N-glycans processed from high-mannose type to complex type. It is applicable to virtually any glycoprotein, and a complete analysis can be conducted with 30 µg of protein. Here, we provide a detailed description of the method that includes procedures for (i) proteolytic digestion of glycoprotein(s) with specific and nonspecific proteases; (ii) denaturation of proteases by heating; (iii) sequential treatment of the glycopeptide mixture with two endoglycosidases, Endo H and PNGase F, to create unique mass signatures for the three glycosylation states; (iv) LC-MS/MS analysis; and (v) data analysis for identification and quantitation of peptides for the three glycosylation states. Full coverage of site-specific glycosylation of glycoproteins is achieved, with up to thousands of high-confidence spectra hits for each glycosite. The protocol can be performed by an experienced technician or student/postdoc with basic skills for proteomics experiments and takes ~7 d to complete.

INTRODUCTION

Asparagine-linked-glycans (N-glycans) are among the most common and important post-translational modifications of proteins. They have critical roles in the folding, conformation and stability of the proteins themselves1,2, and participate as ligands in intra- and intercellular recognition and host-pathogen interactions3,4. Altered biosynthesis of N-glycans has been associated with many diseases, such as cancer5,6, influenza7–9, and AIDS10–14. For example, increased levels of underprocessed high-mannose-type glycans have been reported to occur during breast cancer progression in both mice and humans6. Some broadly neutralizing antibodies (bnAbs) of the highly glycosylated HIV envelope glycoprotein include underprocessed high-mannose glycans in their epitopes, whereas others require fully processed complex-type structures containing sialic acid11,15. As the structures of N-glycans affect the activity and pharmacodynamics of glycoprotein biotherapeutics16, careful choice of cell lines used to express proteins10, growth conditions17, and purification methods18–20 to control the consistency of N-glycosylation is needed.

Given the importance of N-glycans to the structure and functions of glycoproteins, there is increasing need for robust methods for analysis of N-linked glycosylation that can be integrated with state of the art proteomics. Complicating the analysis is the inherent diversity of N-glycan structures present on any glycoprotein, which is a consequence of the non-template-driven biosynthesis. It begins with the en bloc transfer of Glc3Man9GlcNAc2 from a lipid-linked glycosyl donor to the nascent polypeptide by an oligosaccharyltransferase (OST) to a sequence-defined glycosite, Asn-X-Thr/Ser (where X can be any amino acid residue except proline). Although none of the proteins analyzed in the present study contains an atypical glycosylation site (N-X-Cys/Val), these sites have been verified in previous studies by the presence of glycans on intact glycopeptides21,22. As illustrated in Figure 1, the glycan is then subjected to trimming by removal of the glucose residues to a high-mannose-type glycan, then further trimming to the conserved Mann9GlcNAc2 core, followed by addition of sugars by the sequential action of glycosyltransferases that produce a highly related set of complex-type structures (Fig. 1). Moreover, for glycoproteins with more than one glycosite, processing at each site may differ based on the access of the glycans to the processing enzymes. Indeed, well documented examples of proteins that contain high-mannose-type glycans at one or more glycosites, and highly processed complex-type glycans at other sites are IgM23,24, influenza hemagglutinin8,25,26, and the HIV envelope glycoprotein10,27–29.

Analysis of N-glycans of glycoproteins

Over the past two decades, several strategies have emerged to characterize N-glycans of glycoproteins and identify the glycosites that are recognized and used by the cellular glycosylation machinery. Several methods employ endoglycosidases, such as PNGase F and PNGase A, to release the glycans from the protein, followed by analysis of glycoforms using MS30–35 or high-performance liquid chromatography (HPLC)36,37 with or without prior derivatization38. The MS-based methods provide a composition for each molecular ion, which is annotated as a high-mannose or complex-type glycan consistent with biosynthetic principles. Furthermore, tandem mass spectra of derivatized N-glycans generated by both matrix-assisted laser desorption/ionization
(MALDI) and electrospray ionization (ESI) MS have been widely used for characterization of detailed structural information of N-glycans expressed in different biological systems.30,34. Freely available software tools such as GRITS Toolbox (http://www.grits-toolbox.org/) are able to automatically process, annotate, and archive glycomics data in a high-throughput manner. The HPLC methods rely on retention times of N-glycan standards for identification of the glycan species. With both methods, supporting experiments using glycosidase digestions and/or permethylation analysis can provide additional support for structure assignments. Such methods find wide utility in characterizing glycosylation of highly studied glycoproteins such as immunoglobulins (e.g., IgG), or detecting individual differences in N-glycans of serum glycoproteins12,15,59,60. Although these methods provide key information about the nature of the glycoforms present on a glycoprotein and their relative abundance, they do not reveal which glycoforms are utilized or the extent to which each glycosite is occupied.

A number of proteomics-based methods focus on identification of the glycosites that are utilized by the OST and the degree to which the site is occupied by a glycan.40-44. The basic strategy is to immobilize glycopeptides by using lectins or by coupling periodate-treated glycopeptides to hydrazide-activated beads. After protease digestion, endoglycosidase H (Endo H) is used to remove all high-mannose- and hybrid-type N-glycans that have at least 5 mannose residues (Fig. 1). This enzyme leaves a GlcNAc-Asn residue that adds +203 to the peptide mass. Subsequently, the remaining complex N-glycans are removed with PNGase F in the presence of H2O18, which both removes the glycan and converts Asn to Asp, with a +3 addition to the peptide mass. Peptides carrying glycosites (Asn-X-Thr/Ser) with Asn (unoccupied), GlcNAc, N-acetylgalcosamine; Man, mannose; Sia, sialic acid.

Figure 1 | N-linked glycan processing in the endoplasmic reticulum and Golgi apparatus. N-glycans that are Endo H-sensitive are shown in the red box; these include high-mannose and hybrid glycans that have two terminal mannose residues, which is required for recognition by Endo H (red oval). Asn, asparagine; ER, endoplasmic reticulum; Fuc, fucose; Gal, galactose; Glc, glucose; GlcNAc, N-acetylgalacosamine; Man, mannose; Sia, sialic acid.

Major limitations of this approach are the relatively low abundance of glycopeptides in the protein digest mixture, as well as the inherent low ionization efficiency of glycopeptides during MS analysis.40,41,49,50. For these reasons, glycopeptides are typically enriched from peptide digests prior to MS analysis with different purification methods, such as hydrophilic interaction chromatography (HILIC)29,51,52 and hydrazide chemistry.40. Despite the use of different fragmentation methods, including collision-induced dissociation (CID), high-energy collision dissociation (HCD), and electron-transfer dissociation (ETD), the inherent heterogeneity of glycoforms at each glycosylation site, as well as difficulties in obtaining good peptide backbone fragmentations for peptides with large glycans, impedes comprehensive identification of intact glycopeptides28,30. Annotation of LC-MS/MS data from such experiments is possible by using commercial algorithms such as Byonic (http://www.proteinmetrics.com/products/byonic/). However, quantitative assessment of the relative abundance of the glycan structures detected at each glycosite is still a substantial challenge due to unknown ionization efficiencies for peptides with each glycoform, especially for those with sialic acid-containing structures.38. Moreover, if the glycopeptides are enriched prior to analysis, peptides with no glycan are lost, and the degree of glycosite occupancy cannot be assessed.

Development of the protocol and applications of the method

The protocol described here arose from the need of the HIV vaccine effort for robust semiquantitative glycoproteomics methods that would: (i) establish the occupancy of N-glycans at each glycosite and (ii) assess the degree to which glycans were processed from high-mannose type to complex type. The need stems from the fact that the primary candidate for a vaccine is the HIV envelope glycoprotein trimer (Env) that contains 75–90 N-glycans, creating a glycan shield to protect against attack by the immune system.53,54. Despite the dense cover of N-glycans, bnAbs that bind to HIV Env do occur in 10–30% of HIV-1-infected patients.55,56. Importantly, some bnAbs show interactions with high-mannose glycans57,58, whereas others show dependence on complex-type glycans12,15,59,60. Thus, to support the rational design of an HIV Env vaccine, we developed a method that could assess the proportion of high-mannose-type and complex-type N-glycans at each glycosite on the Env glycoprotein.40. However, as demonstrated here, the method is broadly applicable to any glycoprotein.

One of the key aspects of our method is the sequential use of two endoglycosidase treatments to introduce unique mass signatures for glycosites that carry no glycan, high-mannose/hybrid-type glycans, and complex-type glycans, as illustrated in Figure 2a. After protease digestion, endoglycosidase H (Endo H) is used to remove all high-mannose- and hybrid-type N-glycans that have at least 5 mannose residues (Fig. 1). This enzyme leaves a GlcNAc-Asn residue that adds +203 to the peptide mass. Subsequently, the remaining complex N-glycans are removed with PNGase F in the presence of H2O18, which both removes the glycan and converts Asn to Asp, with a +3 addition to the peptide mass. Peptides carrying glycosites (Asn-X-Thr/Ser) with Asn (unoccupied), GlcNAc-Asn (Endo H–treated), or Asp (PNGase F–treated), display similar ionization efficiencies during MS analysis.40,42, allowing us to use ion intensity peak area to quantify the relative distribution of the three glycosylation states at each glycosite detected10,61. Another distinguishing feature of the method is the use of multiple...
proteases, which can generate up to thousands of spectra hits for each glycosite, resulting in >95% sequence coverage. This simple strategy effectively converts the glycoproteomics analysis to a proteomics analysis, allowing the use of robust proteomics software to analyze data in a high-throughput manner.

A major advantage of this method is that it provides a glimpse into the site occupancy and processing of N-glycans at each glycosylation site. It provides a semiquantitative analysis of the three glycosylation states, and a complete analysis can be conducted on only 30 µg of protein, even for complex glycoproteins such as HIV Env, comprising up to 30 glycosites per monomer. It should be noted that hybrid structures, which can potentially contain sialic acid (see Fig. 1), are included in the high-mannose/hybrid-type category because they are cleaved by Endo H. However, hybrid structures typically have low abundance.

One major limitation of the method is that glycan structures are removed before LC-MS/MS, and the class of the glycan is inferred by the specificities of the endoglycosidases. MS/MS methods based on analysis of intact glycopeptides provide more information on the spectra of glycans at individual glycosites, and are complementary to this protocol, which provides semi-quantitative information on site occupancy and glycan processing. However, neither method provides the precise structure complete with glycosidic linkages between monosaccharides. Although this protocol is designed for site-specific glycosylation analysis of purified glycoproteins, it is, in principle, applicable to more-complex protein samples, such as membrane-enveloped viruses (e.g., HIV, influenza virus, coronavirus) that comprise only 10–12 proteins. However, as described in the ‘Experimental design’ section, modification of the protocol would be needed to survey glycosites in more complex samples such as cells or human serum.

Overview of the procedure

The procedure for site-specific analysis of glycoprotein N-glycan processing is summarized in Figure 2b and consists of the following key stages.

- In the first stage, buffer exchange for glycoproteins is needed if they are dissolved in the buffers that contain nonvolatile salts (Steps 1–12). Glycoproteins are then denatured and alkylated at pH 6 (Steps 13–17) to minimize non-enzymatic deamidation while retaining protein sequence coverage. Proteins are digested with several different protease treatments, including ‘triple digestion’, chymotrypsin, and a combination of trypsin and chymotrypsin, in order to maximize sequence coverage and increase the confidence of detecting each glycosite (Step 18).
- In the second stage (Steps 19–25), proteases are denatured by heating to prevent incorporation of 18O-water into the C termini of peptides during the subsequent PNGase F treatment.
- In the third stage (Steps 26–36), sequential endoglycosidase treatment is employed to create unique mass signatures relative to the predicted amino acid sequence, for peptide glycosites that are not occupied (+0 Da) or that are occupied by high-mannose/hybrid- (+203 Da) or complex-type glycans (+3 Da) (Fig. 2a).
- The resulting samples are subjected to LC-MS/MS analysis (Step 37).
- Data processing for identification and quantitation of peptides for the three glycosylation states is done using the Integrated Proteomics Pipeline (IP2) software package (stage 5, Steps 41–53). The MS1 and MS2 data are extracted from MS raw files with RawConverter and processed using multiple components from the IP2 software package (Steps 38–53).

Alternative methods

Over the past decade there has been substantial progress in site-specific glycosylation analysis of purified glycoproteins. Exemplary, and perhaps the most relevant to this protocol, is the work on HIV-1 Env by the Desaire and Cripsin groups. The methods developed by the two groups focus on characterizing intact glycopeptides with or without enrichment of glycopeptides prior to MS analysis, and thus can provide complementary information about individual glycoforms at each glycosite. However, milligram quantities of material may be required for their methods, which is attributed in part to the relatively low ionization efficiencies of glycopeptides and in part to the fact that each glycopeptide is actually a mixture of many different glycoforms. Although the complexity is reduced by using only two proteases, chymotrypsin and trypsin, identification of all forms of a glycopeptide with multiple N-glycans is still a challenge, even when using a combination of CID and ETD for fragmentation. Quantitative measurements are also challenging, due to unknown ionization of glycopeptides with different N-glycan species during MS analysis. Moreover, there is limited information about site occupancy, due to markedly different ionization efficiencies between peptides and the corresponding glycopeptides, and/or enrichment of glycopeptides prior to MS analysis.

Other methods that use endoglycosidases to identify glycosites have long been in use. However, typically, glycopeptides are selectively captured using treatment with periodate followed...
by binding to hydrazide-activated beads, or are enriched using lectins. The peptides are then released using PNGase F, in which glycosylated asparagine is converted to aspartic acid (+1 Da mass shift), allowing indirect identification of glycosites in the released peptides\(^{40,41}\). Alternatively, for a purified protein, PNGase F can be applied directly, and site occupancy of glycosylation sites can be determined by measuring the ratio of peptides with glycosites containing asparagine to aspartic acid during ESI-MS analysis\(^{28}\). By comparison with the protocol described here, addition of a second endoglycosidase, Endo H, provides additional information about the extent of glycan processing, in addition to the degree of site occupancy.

Experimental design

**Proteolysis with a combination of proteases.** Glycoproteins are denatured and alkylated at pH 6 instead of a mildly alkaline pH to minimize the non-enzymatic deamidation of Asn to Asp, which can complicate data analysis\(^{65}\). To maximize sequence coverage, a number of different protease digestions are used, including digestion by chymotrypsin, a combination of trypsin and chymotrypsin, and combinations of Arg-C, trypsin, elastase and subtilisin (‘triple digestion’)\(^{66}\) (Step 18). Of note, a combination of all proteolytic digestions is essential for detecting all glycosites on heavily glycosylated proteins with large molecular weights, such as the HIV Env trimer and the spike glycoprotein of Middle East respiratory syndrome coronavirus (MERS-CoV S-2P protein). Triple digestion steps times more MS/MS spectra for a given sample than an Orbitrap Fusion or Orbitrap Lumos, is more sensitive and generates several measurement of site-specific glycosylation of purified proteins, multidimensional protein identification technology (MudPIT), in which peptides are systematically separated based on the charge in the first dimension and hydrophobicity in the second, will accelerate site-specific N-glycan processing of glycoproteins in these complex protein samples\(^{68}\).

**Sequential endoglycosidase treatment.** Sequential treatment of glycopeptides with Endo H followed by PNGase F is employed to generate different mass signatures for glycosites that contain no glycan (+0), high-mannose/hybrid-type glycan (+203), and complex-type glycan (+3) (Steps 26–36, Fig. 2a). The endoglycosidase digestions are highly efficient and progress rapidly to completion, so the deglycosylation reactions are conducted for 1 h to minimize non-enzymatic deamidation that can occur during this step. Removal of N-glycans increases the ionization efficiencies of peptides and allows us to localize glycosylation sites on peptides with multiple modifications, which is challenging for analysis of intact glycopeptides (Supplementary Fig. 1).

**LC-MS/MS analysis.** In principle, the deglycosylated peptides can then be analyzed by any type of LC-MS/MS (Step 37). A high-resolution mass spectrometer, such as the Orbitrap Elite, provides satisfactory sequence coverage for most glycoproteins. An instrument with a higher scan speed, such as an Orbitrap Fusion or Orbitrap Lumos, is more sensitive and generates several times more MS/MS spectra for a given sample than an Orbitrap Elite. Thus, they are able to provide site-specific glycan-processing information for those sites that may be missed in the results generated by the Orbitrap Elite as a result of heavy glycosylation and the large molecular weights of the glycoproteins. Although single-dimension separation is sufficient for characterization of site-specific glycosylation of purified proteins, multidimensional protein identification technology (MudPIT), in which peptides are systematically separated based on the charge in the first dimension and hydrophobicity in the second, will accelerate measurement of site-specific N-glycan processing of glycoproteins in these complex protein samples\(^{68}\).

**Replicates and controls.** Ideally, each glycoprotein is digested in at least two technical replicates and analyzed by the same MS instrument. Invertase produced by the yeast *Saccharomyces cerevisiae* and α-1-acid glycoprotein from bovine serum are known to be occupied by high-mannose-type and complex-type glycans, respectively, and can be used as controls to test completion of endoglycosidase treatment. As a check on the overall success of the protocol, non-enzymatic deamidation of Asn residues and/or O\(^{18}\)-incorporation at the C terminus should be seen in <5% of the total peptides.

**Materials and equipment**

**Reagents**

- Urea (MilliporeSigma, cat. no. US128)
- Ammonium acetate (CH\(_3\)COONH\(_4\); MilliporeSigma, cat. no. A1542)
- Iodoacetamide (IAA; MilliporeSigma, cat. no. I1149)
- DTT (Fisher BioReagents, cat. no. BP172-5)
- Water (purified by the GenPure Pro water purification system; Thermo Fisher Scientific)

**Equipment**

- Hydrazide-activated beads
- Chymotrypsin
- Trypsin
- Elastase
- Subtilisin
- PNGase F
- Endo H
- Invertase
- O\(^{18}\)-incorporation
- MS instrument
**PROTOCOL**

eyes, skin, and intestines irreversibly. Handle it in a hood while wearing personal protective equipment, including lab coat, gloves, and safety glasses.

- Arg-C (Promega, cat. no. V1881)
- Trypsin (Promega, cat. no. V5111)
- EDTA disodium salt dihydrate (MilliporeSigma, cat. no. E5134)
- Ethylene bridged hybrid (BEH) 1.7-µm C18 resin (Waters, cat. no. 186003556)
- 4-µm Jupiter C18 (Phenomenex, cat. no. 04A-4396)
- Acetonitrile (ACN; VWR, cat. no. 200004-350) ! CAUTION ACN has modest toxicity in small doses and should be handled in a hood while wearing gloves.
- Tris base (Fisher Scientific, cat. no. BP152-10)
- Calcium chloride dihydrate (CaCl₂·2H₂O; MilliporeSigma, cat. no. C3306)
- Elastase (Promega, cat. no. V1891)
- Subtilisin (MilliporeSigma, cat. no. P5380)
- Chymotrypsin (Promega, cat. no. V1061)
- Endo H (New England Biolabs, cat. no. P0702L)
- PNGase F (New England Biolabs, cat. no. P07055)
- Urea (8 M) Dissolve 0.048 g of urea in 100 µl of 100 mM ammonium acetate (pH 6). Prepare fresh solution for each experiment.

**EQUIPMENT SETUP**

**EQUIPMENT**

- EASY 1000 system (Thermo Fisher Scientific)
- Orbitrap Fusion Tribrid mass spectrometer equipped with the EASY-nLC system (Thermo Fisher Scientific) or an Orbitrap Fusion Tribrid mass spectrometer equipped with the EASY-nLC system (Thermo Fisher Scientific)
- Freezer set to −80 °C (Forma Scientific)
- Water bath set to 30 °C (Fisher Scientific, cat. no. 15-462-5Q)
- Incubator set to 37 °C (Fisher Scientific, cat. no. 15-103-0514)
- Incubator set to 56 °C (Fisher Scientific, cat. no. 15-103-0514)
- Incubator set to 100 °C (Fisher Scientific, cat. no. 05-412-500)
- NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, cat. no. ND-2000)
- Eppendorf microcentrifuge 5415R (MilliporeSigma, cat. no. Z065212)
- Microcentrifuge (Santa Cruz Biotechnology, cat. no. sc-358765)
- Lyophilizer (Labconco)
- Desiccator (Hach)
- Fisher Vortex Genie 2 (Fisher Scientific, cat. no. 12-812)
- Parafilm (VWR, cat. no. 52858-000)
- Kimtech Science Kimwipes tissues (Kimberly-Clark)
- Freezer set to −80 °C (Forma Scientific)
- Mass spectrometer: the protocol below is optimized for either (i) an Orbitrap Elite Hybrid Ion Trap-Orbitrap mass spectrometer equipped with the EASY-nLC system (Thermo Fisher Scientific) or (ii) an Orbitrap Fusion Tribrid mass spectrometer equipped with the EASY 1000 system (Thermo Fisher Scientific)

**Software**

- RawConverter, v1.1.0.19 (http://fields.scripps.edu/rawconv/)
- Notepad++, v7.5.4 (https://notepad-plus-plus.org/download/v7.5.3.html)
- glyco_motif_filter, v1.0 (http://fields.scripps.edu/yates/wp/?page_id=687)
- IP2 (Integrated Proteomics Pipeline, v5.0.1; http://goldfish.scripps.edu/ip2/login.jsp)

**REAGENT SETUP**

**Ammonium acetate (100 mM, pH 6)** Dissolve 0.077 g of ammonium acetate in 10 ml of water. Adjust the buffer pH to 6 by adding acetic acid. Dispense into aliquots and store the buffer at −20 °C for up to 2 months.

**DTT (500 mM)** Dissolve 0.0077 g of DTT in 100 µl of water. Prepare fresh solution for each experiment.

**Iodoacetamide (300 mM)** Dissolve 0.00925 g of iodoacetamide in 100 µl of water. Prepare fresh solution for each experiment.

**Ammonium bicarbonate (100 mM, pH 8)** Dissolve 0.079 g of ammonium bicarbonate in 10 ml of water. Measure the buffer pH. Dispense into aliquots and store the buffer at −20 °C for up to 2 months.

**100 mM Tris-Cl (pH 7.8)** Dissolve 12.114 g of Tris in 800 ml of water. Adjust the pH to 7.8 with concentrated HCl. Bring the final volume to 1 liter with water. Store the buffer at 4 °C for up to 2 months.

**Acetonitrile (ACN; VWR, cat. no. 200004-350)** Dissolve 0.077 g of ammonium acetate in 1 ml of water. Adjust the buffer pH to 6 by adding acetic acid. Dispense into aliquots and store the buffer at −20 °C for up to 2 months.

**EDTA (1 M, pH 8)** Add 186.12 g of EDTA·2H₂O to 300 ml of H₂O. Stir vigorously on a magnetic stirrer. Adjust the pH to 8.0 with NaOH pellets. The disodium salt of EDTA will not go into solution until the pH of the solution is adjusted to −8.0 by the addition of NaOH. Bring the final volume to 500 ml with water. Store the buffer at 4 °C for up to 2 months.

**Ammonium bicarbonate (100 mM, pH 5.5)** Dissolve 0.077 g of ammonium acetate in 10 ml of water. Adjust the buffer pH to 5.5 by adding acetic acid. Dispense into aliquots and store the buffer at −20 °C for up to 2 months.

**18O-water (97 atom% of 18O; MilliporeSigma, cat. no. 329878)**

▲ CRITICAL Store the reagent in a desiccator at room temperature (20–25 °C) for up to 1 year.

- Fetalin (MilliporeSigma, cat. no. F3385)
- Invertase (MilliporeSigma, cat. no. I0408)
- IgG (MilliporeSigma, cat. no. A4506)
- IgM (MilliporeSigma, cat. no. I8260)
- α-1-Acid glycoprotein (AGP; MilliporeSigma, cat. no. G3643)
- Transferrin (MilliporeSigma, cat. no. T8158)
- NaOH pellets (MilliporeSigma, cat. no. 221465)
- Glycoprotein of interest: In this protocol, we use three example proteins: BG505 SOSIP.664 trimer (laboratory-made, expressed in HEK 293-F cells as described in a previous study) MERS-CoV S-2P protein (laboratory-made, expressed in HEK 293-F cells as described in a previous study) and influenza virus hemagglutinin (HA, laboratory-made, expressed in HEK 293-F cells as described in a previous study).

**EQUIPMENT**

- Eppendorf Research Plus pipette (Eppendorf)
- Biotix microcentrifuge tubes, 1.5 ml (VWR, cat. no. MTI-0150-BC)
- Low-binding pipette tips, 10 µl (Corning, cat. no. 4150)
- Low-binding pipette tips, 200 µl (Corning, cat. no. 4151)
- Centrifugal filter, 10 kDa (MilliporeSigma, cat. no. MRCPRT010)
- Centrifugal filter, 30 kDa (MilliporeSigma, cat. no. MRCH0030)
- Incubator set to 37 °C (Fisher Scientific, cat. no. 15-103-0514)
- Incubator set to 56 °C (Fisher Scientific, cat. no. 15-103-0514)
- Incubator set to 100 °C (Fisher Scientific, cat. no. 05-412-500)
- Water bath set to 30 °C (Fisher Scientific, cat. no. 15-462-5Q)
- pH meter (MilliporeSigma, cat. no. Z283037)
- NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, cat. no. ND-2000)
- Eppendorf microcentrifuge 5415R (MilliporeSigma, cat. no. Z605212)
- Microcentrifuge (Santa Cruz Biotechnology, cat. no. sc-358765)
- Lyophilizer (Labconco)
- Desiccator (Hach)
- Fisher Vortex Genie 2 (Fisher Scientific, cat. no. 12-812)
- Parafilm (VWR, cat. no. 52858-000)
- Kimtech Science Kimwipes tissues (Kimberly-Clark)
- Freezer set to −80 °C (Forma Scientific)
- Mass spectrometer: the protocol below is optimized for either (i) an Orbitrap Elite Hybrid Ion Trap-Orbitrap mass spectrometer equipped with the EASY-nLC system (Thermo Fisher Scientific) or (ii) an Orbitrap Fusion Tribrid mass spectrometer equipped with the EASY 1000 system (Thermo Fisher Scientific)

**Software**

- RawConverter, v1.1.0.19 (http://fields.scripps.edu/rawconv/)
- Notepad++, v7.5.4 (https://notepad-plus-plus.org/download/v7.5.3.html)
- glyco_motif_filter, v1.0 (http://fields.scripps.edu/yates/wp/?page_id=687)
- IP2 (Integrated Proteomics Pipeline, v5.0.1; http://goldfish.scripps.edu/ip2/login.jsp)
**Mass spectrometer setup** The instrument should be mass-calibrated prior to use. The detailed settings below were used with the Orbitrap Fusion and the Orbitrap Elite.

| Parameter               | Orbitrap fusion | Orbitrap elite |
|-------------------------|-----------------|----------------|
| Ionization mode         | Positive        | Positive       |
| Electrospray voltage    | 2.5 kV          | 2.5 kV         |
| MS1 resolution          | 120 k           | 120 k          |
| MS2 resolution          | 15 k            | 15 k           |
| MS1 scan range          | 400–1,500       | 400–2,000      |
| MS1 maximum injection time | 50 ms        | 200 ms         |
| MS2 maximum injection time | 250 ms        | 1,000 ms       |
| MS1 AGC target          | $4 \times 10^5$ | $1 \times 10^6$ |
| MS2 AGC target          | $5 \times 10^4$ | $5 \times 10^4$ |
| Isolation               | $2 \text{ m}/\text{z}$ with quadrupole | $2 \text{ m}/\text{z}$ with ion trap |
| Fragmentation           | CID at 35%      | CID at 35%     |
| Dynamic exclusion       | 5 s             | 60 s           |
| Charge states           | 2–7 selected    | +1 and unassigned are rejected |
| Intensity threshold     | $5e4$           | 500            |
| DDA method              | 3 s cycle time  | Top ten        |

AGC, automatic gain control.

**PROCEDURE**

**Buffer exchange for glycoproteins** ● **TIMING** ~11 h

▲ **CRITICAL** To minimize sample loss, low-protein-binding microcentrifuge tubes and pipette tips should be used during sample preparation. Only volatile salts are used in the protocol and these can be removed by lyophilization, resulting in higher sensitivity due to minimal protein loss. High-purity reagents should be used for sample preparation and MS analysis to minimize signals derived from contaminants.

▲ **CRITICAL** If the protein is a lyophilized powder, then proceed to the ‘Denaturation and alkylation of glycoprotein’ section (Step 13).

1. Insert a 10-kDa centrifugal filter into a tube.

2. Rinse the filter by adding 100 µl of 100 mM ammonium acetate immediately before use. Seal the tube with the attached cap.
3| Centrifuge the device (the tube and the filter) at 8,000 g for 50 min at 4 °C.

4| Pipette the solution containing ~30 µg of target protein into the filter. Seal the tube with the attached cap. If the volume of the solution is <100 µl, bring the final volume to 100 µl with water. ▲ CRITICAL STEP Protein concentration is estimated with the NanoDrop A280 assay.

5| Centrifuge the device at 8,000 g for 50 min at 4 °C.

6| Pipette 100 µl of 100 mM ammonium acetate into the filter.

7| Centrifuge the device at 8,000 g for 50 min at 4 °C.

8| Repeat Steps 6 through 7 at least two times. ▲ CRITICAL STEP To complete the buffer exchange, it is important to extensively wash the filter that contains glycoproteins (at least three times) with 100 mM ammonium acetate (pH 6). ▲ CRITICAL STEP To minimize non-enzymatic deamidation, the acidic buffer, 100 mM ammonium acetate (pH 6), should be used instead of mildly alkaline buffers during buffer exchange.

9| Wash the filter membrane with 100 µl of 100 mM ammonium acetate (pH 6).

10| Collect the solution and store it in a low-protein-binding microcentrifuge tube.

11| Repeat Steps 9 and 10 at least four times and combine the fractions. ▲ CRITICAL STEP Wash the filter membrane with the buffer at least five times after buffer exchange in order to achieve maximum recovery of proteins. ■ PAUSE POINT The solution can be stored up to 1 week at −80 °C or in liquid nitrogen.

12| Lyophilize 500 µl of the solution at room temperature for at least 5 h. ▲ CRITICAL STEP If volatile salts such as ammonium acetate are added to the buffer, it is important to completely remove them with the lyophilizer.

Denaturation and alkylation of glycoprotein ● TIMING ~7 h

13| Dissolve the glycoprotein in 100 µl of 8 M urea in 100 mM ammonium acetate (pH 6) and place the solution at room temperature for 1 h. ▲ CRITICAL STEP The acidic buffer, 100 mM ammonium acetate, should be used for sample preparation instead of mildly alkaline buffers, if possible, in order to keep non-enzymatic deamidation to a minimum.

14| Add 2 µl of 500 mM DTT (to a final concentration of ~10 mM) and incubate the solution at 56 °C for 1 h. ▲ CRITICAL STEP Spin down (1,000 g, 25 °C, 1 min) all the solution from the walls of microcentrifuge tube before adding DTT.

15| Add 11 µl of freshly prepared 500 mM iodoacetamide (to a final concentration of ~50 mM) and incubate the solution in the dark at room temperature for 45 min. ▲ CRITICAL STEP To preserve activity of iodoacetamide, prepare iodoacetamide solution immediately before use, as it is unstable. Perform the alkylation step in the dark, as iodoacetamide is light-sensitive.

16| Buffer-exchange to 100 mM ammonium bicarbonate (pH 8) by using a centrifugal filter with a membrane nominal molecular weight limit of 10 kDa. ▲ CRITICAL STEP To complete buffer exchange, it is important to wash the filter at least three times with 100 µl of 100 mM ammonium bicarbonate at 8,000 g for 50 min at 4 °C.

17| Dispense the sample into five equal aliquots for the following proteolytic digestion. ▲ CRITICAL STEP Do not dry glycoproteins after denaturation, as they may not be fully soluble in the buffer after lyophilization. ■ PAUSE POINT The samples can be stored at −80 °C or in liquid nitrogen for at least 2 weeks.
Protease treatments ● TIMING ~24 h
18] Five aliquots (containing 6 µg of denatured glycoproteins each) are subjected to treatments with multiple proteases and combinations of proteases involving: Arg-C followed by trypsin (option A), elastase (option B), and subtilisin (option C). Aliquots A–C will later be combined into a ‘triple digestion’ sample (Step 21). The remaining two aliquots are digested with chymotrypsin (option D) or a combination of trypsin and chymotrypsin (option E).

(A) Arg-C followed by trypsin
(i) Add Arg-C to one aliquot that contains ~6 µg of denatured glycoproteins at an enzyme/protein ratio of 1:20 (wt/wt). Bring the final volume to 100 µl with 100 mM ammonium bicarbonate (pH 8). Add DTT and EDTA to final concentrations of 5 mM and 0.2 mM, respectively.
   ▲ CRITICAL STEP Arg-C is able to cleave at the C terminus of arginine residues, including sites next to proline, resulting in increased sequence coverage when combined with trypsin digestion.
(ii) Incubate the solution at 37 °C for 4 h.
(iii) Lyophilize the resulting peptide mixture for at least 3 h to remove water and volatile salt.
(iv) Redissolve the peptide mixture in 500 µl of 100 mM ammonium acetate (pH 6).
(v) Add sequencing-grade modified trypsin to the solution at a trypsin/protein ratio of 1:10 (wt/wt).
   ▲ CRITICAL STEP Sequencing-grade modified trypsin is able to digest glycoproteins at pH 6 while preserving adequate digestion efficiency.
(vi) Incubate the reaction at 37 °C for 16 h.

(B) Elastase
(i) Add elastase to the second aliquot of the denatured glycoprotein at an elastase/protein ratio of 1:20 (wt/wt).
   ▲ CRITICAL STEP The utility of triple digestion can generate higher sequence coverage than digestion with any of the single enzymes alone.
(ii) Bring the final volume to 500 µl with 100 mM ammonium bicarbonate (pH 8).
(iii) Incubate the reaction at 37 °C for 16 h.

(C) Subtilisin
(i) Add subtilisin to the third aliquot of the denatured glycoprotein at a subtilisin/protein ratio of 1:20 (wt/wt).
(ii) Bring the final volume to 500 µl with 100 mM ammonium bicarbonate (pH 8).
(iii) Incubate the reaction at 37 °C for 4 h.
   ▲ CRITICAL STEP Do not incubate the reaction longer than 4 h, in order to obtain appropriate lengths of peptides for MS detection.

(D) Chymotrypsin
(i) Add chymotrypsin to the fourth aliquot of the denatured glycoprotein at a chymotrypsin/protein ratio of 1:13 (wt/wt).
   ▲ CRITICAL STEP Do not add too much chymotrypsin to the solution, as it can be self-digested, which will suppress the ESI signals of analytes.
(ii) Bring the final volume to 500 µl with 100 mM ammonium bicarbonate (pH 8).
(iii) Incubate the reaction in a 30 °C water bath for 10 h.

(E) A combination of trypsin and chymotrypsin
(i) Add trypsin and chymotrypsin to the fifth aliquot of the denatured glycoprotein at enzyme/protein ratios of 1:20 (wt/wt) and 1:13 (wt/wt) respectively.
(ii) Bring the final volume to 500 µl with 100 mM ammonium bicarbonate (pH 8).
(iii) Incubate the reaction at 37 °C for 16 h.
   ■ PAUSE POINT The peptide mixtures derived from combination proteolytic digestion can be stored at −80 °C or in liquid nitrogen for at least 2 weeks.

Denaturation of proteases ● TIMING ~10 h
19] Lyophilize the peptide mixtures derived from each of the five protease digestions (Step 18A–E) for at least 5 h.
20] Redissolve each sample in 100 µl of water.
21] Combine the peptide mixtures derived from Step 18A–C into a ‘triple digestion’ sample.
22] Separately, heat the combined triple digestion sample (Step 21) and the samples generated from digestion with chymotrypsin (Step 18D), and a combination of trypsin and chymotrypsin (Step 18E) at 100 °C for 30 s.
23] Cool the samples at room temperature for 30 s.
**PROTOCOL**

24| Repeat Steps 22 and 23 at least five times.  
    ▲ CRITICAL STEP To completely deactivate the proteases used, Steps 22 and 23 should be repeated at least five times. Any remaining active proteases will accelerate incorporation of $^{18}$O-water into the C termini of peptides during the following PNGase F treatment conducted in $^{18}$O-water. 

25| Lyophilize the samples for at least 3 h to remove any remaining volatile salts from the samples.  
    ▲ CRITICAL STEP Complete removal of volatile salts from the samples is important for the following Endo H treatment, which has optimal activity at pH 5.5.  
    ■ PAUSE POINT The peptide mixtures can be stored at −80 °C or in liquid nitrogen for at least 2 weeks. 

Sequential endoglycosidase treatment ● TIMING ~7 h 
26| Separately, redissolve each of the three samples in 20 µl of 100 mM ammonium acetate (pH 5.5). 

27| Add Endo H to the peptide mixtures at a minimum enzyme/glycoprotein ratio of 250 NEB units per 10 µg.  
    ▲ CRITICAL STEP The quantity of Endo H needed for deglycosylation may vary from protein to protein. A set of control experiments should be done before using this protocol (see Reagent Setup). 

28| Incubate the reaction at 37 °C for 1 h.  
    ■ PAUSE POINT The Endo H–treated peptides can be stored up to 1 week at −80 °C or in liquid nitrogen. 

29| Lyophilize the Endo H–treated samples for at least 3 h immediately before use. In the meantime, proceed to Step 30.  
    ▲ CRITICAL STEP Complete removal of ammonium acetate in the samples ensures that the following PNGase F treatment will proceed to completion. 

30| Lyophilize the PNGase F solution for at least 1 h immediately before use and redissolve the PNGase F enzyme in the same volume of $^{18}$O-water (see Reagent Setup). 

31| Add 20 µl of 100 mM ammonium bicarbonate (pH 8) prepared with $^{18}$O-water to the PNGase F solution.  
    ▲ CRITICAL STEP Steps 31 and 32 should be done as quickly as possible to reduce contact of the reaction mixture with air. 

32| Add PNGase F to the Endo H–treated peptide mixtures at a minimum enzyme/glycoprotein ratio of 500 NEB units per 10 µg.  
    ▲ CRITICAL STEP The quantity of PNGase F that is needed for deglycosylation can be determined by treating a defined amount of the desired glycoprotein with a series of PNGase F concentrations. Gel shifts of N-glycans on SDS-PAGE can be used to determine completion of deglycosylation. 

33| Seal the microcentrifuge tube that contains the mixture. 

34| Incubate the reaction at 37 °C for 1 h. 

35| Dispense the reaction mixture into aliquots that contain ~2 µg of peptides and store them in liquid nitrogen immediately.  
    ■ PAUSE POINT The samples can be stored in liquid nitrogen for at least 1 month. 

**LC-MS/MS analysis ● TIMING 6–48 h per glycoprotein** 
36| Set up the LC-MS/MS system to characterize the deglycosylated peptides as described in the ‘Equipment Setup’ section.  
    ▲ CRITICAL STEP Each glycoprotein is digested in two or three technical replicates and analyzed by the same MS instrument. 

? TROUBLESHOOTING 

Data analysis ● TIMING variable; hours to 1 d per glycoprotein 
37| Extract the MS1 and MS2 spectra from the MS raw files using the spectrum-converting software RawConverter. 

38| Add the MS raw files to the ‘files to covert’ window of RawConverter. 

39| Set ‘Experiment Type’ as data dependent. Enable ‘Select monoisotopic m/z in DDA’ ‘Output formats’ should be set to ‘MS1, MS2, and MS3’, in which both MS1 and MS2 data are extracted from the MS raw files.
40| Use a text editor such as Notepad++ (https://notepad-plus-plus.org/) to prepare a file containing the sequences of the target glycoproteins in FASTA format.

41| Add the resulting file to a predefined database, such as the European Bioinformatic Institute Bos Taurus protein database, with the ‘database’ of IP2.

42| Set the parameters as: Source: Uniprot; Organism Name: Bos Taurus; Generate reverse (decoy) sequences: yes; Add contaminant proteins: No.

**CRITICAL STEP** Reverse (decoy) sequences should be generated and included in the final database in order to estimate peptide probabilities and false-discovery rates.

43| Upload the resulting file that contains the sequences of the target glycoproteins to the ‘database’ window of IP2.

44| Upload the database to IP2.

45| Upload the MS1 and MS2 files to IP2.

46| Start a ProLuCID search in the IP2 (v5.0.1) software package.

47| Set the mass tolerance at 50 p.p.m. for precursor ions and 20 p.p.m. for fragment ions (MS2 spectra are detected in the Orbitrap instrument). No enzyme specificity is considered for searching. Set carboxyamidomethylation (+57.02146 C) as a fixed modification. Set oxidation (+15.9994 M), deamidation (+2.988261 N), GlcNAc (+203.079373 N), and pyroglutamate formation from N-terminal glutamine residue (−17.026549 Q) as variable modifications.

**CRITICAL STEP** Sequence coverage of target proteins should be >95%.

**TROUBLESHOOTING**

48| Filter the results generated by the ProLuCID search by using DTASelect (v2.0, another component of the IP2 software kit). The parameters are set as: minimum number of peptides per protein: ≥2, spectrum false-positive rate: ≤0.05, and precursor delta mass cutoff: ≤10 p.p.m.

**CRITICAL STEP** All peptides with N+203 modifications that are not located at the consensus motif should be manually checked before they are removed. Asparagine residues that are not located at the motif should be considered as potential glycosylation sites when multiple spectra hits with N+203 modifications that do not contain the motif are consistently detected. Further verification is needed for these potential glycosylation sites.

**TROUBLESHOOTING**

49| Filter the results generated by DTASelect with the software ‘Glyco_motif_filter’ to remove those peptides with N+3 and/or N+203 modifications that are not located at the motif (N-X-S/T, X can be any amino acid residue except proline).

**CRITICAL STEP** All peptides with N+203 modifications that are not located at the consensus motif should be manually checked before they are removed. Asparagine residues that are not located at the motif should be considered as potential glycosylation sites when multiple spectra hits with N+203 modifications that do not contain the motif are consistently detected. Further verification is needed for these potential glycosylation sites.

50| Start a label-free analysis using Census (another component of the IP2 software package). The parameters are set as: ‘find missing peptide’: enabled, mass tolerance: ≤10 p.p.m., retention time tolerance: ≤0.1 min.

**CRITICAL STEP** Ion injection time is used to further normalize the resulting peak area.

51| Determine the abundance of each peptide from each raw file by the sum of the ion intensity peak area over all identified charge states.

**CRITICAL STEP** To improve the accuracy of the method, a set of peptides with N+0, N+3, and N+203 modifications is considered only when at least one of the three has a peak area of at least $5 \times 10^8$. Of note, ~2 µm of purified glycoproteins are loaded onto the column in the present study. This value was empirically determined as optimal to distinguish information from spectral noise and will vary from instrument to instrument. A control experiment should be done by using well-characterized model glycoproteins such as invertase (occupied by high mannose glycans) and α-1-acid glycoprotein (occupied by complex-type glycans) before setting the peak area threshold. Peak area values will be dependent on the type of LC system and mass spectrometer used, and the appropriate threshold will need to be determined for other instrument types.

**TROUBLESHOOTING**

52| Combine the data derived from two or three technical replicates after analysis of each MS run separately.
## Troubleshooting

Troubleshooting advice can be found in Table 1.

| Step | Problem | Possible reason | Solution |
|------|---------|----------------|----------|
| 36   | No peptides detectable | The LC-MS instrument is not performing properly | Make sure that the LC-MS instrument is well-calibrated and working properly according to specifications |
| 48   | Low sequence coverage of target glycoproteins | Glycoproteins may not have been washed out completely after buffer exchange (Step 16) | Wash the filter membrane at least five times after buffer exchange to maximize recovery of target glycoproteins |
|      | No peptides detected after combination proteolytic digestion | Proteases stored in the buffer at −80 °C for too long may not be active (Step 18) | Make fresh protease solution |
|      | Too many peaks that belong to non-enzymatic deamidation are detected | The microcentrifuge tube that contains 18O-water is not completely sealed or was exposed to air for too long (Steps 30–33) | Seal the microcentrifuge tube completely and place it into a desiccator. Perform Steps 30–33 as quickly as possible to reduce contact of the reaction mixture with air |
| 51   | The output file generated by Census is empty or cannot be downloaded | The MS1 file was not extracted correctly by RawConverter or the names of the target glycoproteins are not recognized by Census | Extract the MS1 spectra from raw files again using RawConverter. Delete spaces that are not recognized by Census from the names of the target glycoproteins |

**TIMING**

Steps 1–12, buffer exchange for glycoproteins: ~11 h
Steps 13–17, denaturation and alkylation of glycoprotein: ~7 h
Step 18, protease treatments: ~24 h
Steps 19–25, denaturation of proteases: ~10 h
Steps 26–35, sequential endoglycosidase treatment: ~7 h
Step 36, LC-MS/MS analysis: 6–48 h per glycoprotein
Steps 37–52, data analysis: variable; hours to 1 d per glycoprotein, depending on complexity

**ANTICIPATED RESULTS**

This protocol is used to determine site-specific N-glycan processing of glycoproteins. In the most widely used strategy, glycoproteins are digested with specific proteases such as trypsin, resulting in (glyco)peptides that are suitable for LC-MS/MS analysis. Glycosylated peptides, however, have much lower ionization efficiency during MS analysis relative to peptides, and thus milligram quantities of materials are generally used for typical glycoproteomics methods. Characterization of glycopeptides with multiple glycosites by MS/MS is still challenging, even with combination of different types of fragmentation techniques. Quantitative measurement of glycopeptides is complicated by the fact that ionization efficiencies of glycopeptides differ with variable glycoforms. This protocol describes an alternative way to overcome these problems by the use of combination proteolytic digestion followed by sequential endoglycosidase treatment.

**Validation of sequential endoglycosidase treatment**

One of the distinguishing features of this protocol is the use of sequential treatment of glycopeptides with endoglycosidases (Endo H followed by PNGase F) to create unique mass signatures for glycosites that have no N-glycan, high-mannose-type glycan, or complex-type glycan. This strategy converts the glycoproteomics analysis to a proteomics analysis, resulting in higher sensitivity of the protocol (only 30 µg of sample is needed for a complete analysis). The key to success is that the sequential endoglycosidase treatments proceed to completion, avoiding mis-assignment of N-glycan processing status. To this end, we applied the protocol to assess site-specific N-glycan processing of two well-characterized model glycoproteins, invertase produced by the yeast *S. cerevisiae* and α-1-acid glycoprotein from bovine serum. Glycosites on invertase are occupied by underprocessed oligomannose, and those on α-1-acid glycoprotein are fully processed complex-type glycosylation...
All N-glycosites on both glycoproteins were identified with multiple MS/MS spectra ranging from 54 to >10,000 per site (Supplementary Tables 2 and 3). Low percentages of spectra hits that contain non-enzymatic deamidation or 18O-incorporation into the C termini of peptides were found among all spectra hits identified, which is attributed to the complete denaturation of proteases used (Supplementary Tables 4 and 5). As described in the procedure section, as well as the previous study\textsuperscript{10}, a set of peptides with N\textsuperscript{+0}, N\textsuperscript{+3}, and N\textsuperscript{+203} modifications was considered only when at least one of the three had a peak area of at least >5 × 10\textsuperscript{8}. As expected, the 14 N-glycosites of invertase were identified as entirely high-mannose-type glycosylation, and site occupancy was >90% for all glycosites except the sites N64 and N275 (Fig. 3a, b). By contrast, the five N-glycosites of α-1-acid glycoprotein were completely complex-type glycosylation, and site occupancy was >98% for all five sites (Fig. 3c, d). These results indicated that sequential endoglycosidase treatment reached completion.

Validation of MS detection of glycotyoes

Another major assumption is that the endoglycosidase-treated peptide glycosites that are unoccupied or occupied by high-mannose glycans or complex-type glycans are detected equally during MS analysis. To test this assumption, the HIV-1 Env trimer BG505 SOSIP.664, expressed in the presence of kifunensine (high-mannose only), was selected as a model protein (hereafter referred to as Kif_BG505). N-glycans on Kif_BG505 were first removed by sequential endoglycosidase treatment, and as expected, glycosites comprised >95% high-mannose-type glycosylation, and site occupancy was >90% for all glycosites except the sites N64 and N275 (Fig. 3a, b). By contrast, the five N-glycosites of α-1-acid glycoprotein were completely complex-type glycosylation, and site occupancy was >98% for all five sites (Fig. 3c, d). These results indicated that sequential endoglycosidase treatment reached completion.

Figure 3 | Validation of sequential endoglycosidase treatment. (a) Scatter plot of the site-specific N-glycan processing of invertase produced by the yeast Saccharomyces cerevisiae. (b) Color-coded bar graph of the site-specific N-glycan processing of invertase. (c) Scatter plot of the site-specific N-glycan processing of α-1-acid glycoprotein. (d) Color-coded bar graph of the site-specific N-glycan processing of α-1-acid glycoprotein. A set of peptides with N\textsuperscript{+0}, N\textsuperscript{+3}, and N\textsuperscript{+203} modifications was displayed only when at least one of the three had a peak area of at least >5 × 10\textsuperscript{8}. Data were obtained from six independent experiments. Mean ± s.e.m. were plotted. a, b adapted from ref. 10, Nature Publishing Group.
Examples of the protocol

Although the protocol was initially developed for analysis of site-specific N-glycan processing of the HIV Env trimer, it is applicable to analysis of site-specific N-glycan processing of recombinant glycoprotein therapeutics (Fig. 5a,b), serum glycoproteins (Fig. 5c,d), and soluble or membrane-bound envelope glycoproteins from viruses (Fig. 6). It is also likely to be useful in characterization of glycoprotein processing in more complex systems such as whole cells due to its high sensitivity.

Site-specific N-glycan processing of recombinantly produced therapeutic glycoproteins, including IgG and IgM, was determined (Fig. 5a,b). Human serum IgG, which contains IgG1-4 with IgG1 and IgG2 as the major isotypes, was found to be entirely complex-type glycosylation, consistent with previous studies (Fig. 5a)32,49,52. Of the five N-glycosites on IgM, which is the major antibody produced in the primary immune response, three sites were shown to be completely occupied by complex-type structures (N171, N332, and N395), whereas the other two sites, N402 and N563, were primarily high-mannose-type glycosylation (Fig. 5b). The glycosite N563 that is proximal to the CH4 domain and thus is a poor substrate for OST was found to be partially glycosylated, in agreement with previous studies24,71.

Data were obtained from at least six independent experiments. Mean ± s.e.m. were plotted. Image adapted from ref. 10, Nature Publishing Group.
Abnormal glycosylation of serum glycoproteins is a common feature in various human diseases, such as cancers and congenital disorders of glycosylation (CDGs). In particular, serum transferrin was first used to diagnose abnormal glycosylation in CDG patients in the 1990s. Normal transferrin has two N-glycosylation sites, each of which is fully occupied, whereas in type I CDGs, an increase of mono-glycosylated transferrin was found due to defects of oligosaccharide assembly and transfer to glycoproteins in those patients. Analysis of site-specific N-glycan processing of commercially available human serum transferrin revealed that the two glycosites of this protein were entirely complex-type glycosylation, in line with the previous studies. Another serum glycoprotein, fetuin, was also found to be entirely complex-type glycosylation, with full occupancy of two sites (N99 and N156) and partial glycosylation of the third glycosite (N176, 89% of site occupancy).

Membrane-bound envelope glycoproteins of various viruses, such as MERS-CoV S-2P protein and HIV Env trimer, are the target of neutralizing antibodies, and thus are the focus of vaccine development. N-linked glycans on those envelope glycoproteins serve as a shield to protect the underlying protein from immune surveillance and thus confound development of effective vaccines for those viruses. Importantly, some neutralizing antibodies to those viruses have glycan-dependent epitopes, suggesting that vaccine design efforts would benefit greatly from understanding the N-glycan processing status at each glycosylation site. MERS-CoV S-2P protein is a large trimer (~600 kDa) with ~25 N-linked glycans per monomer, each of which comprises two noncovalently associated subunits, S1 and S2. Characterization of site-specific N-glycan processing of a prefusion-stabilized MERS-CoV S-2P protein ectodomain (MERS S-2P) revealed that all 23 glycosites on the protein were fully occupied, except for the N104 site. High-mannose glycans were predominantly found in the S1-NTD (residues 18–353), whereas other regions of the S protein, including the RBD (367–588), the two subdomains (589–751), and S2 (752–1,291), contained glycosites occupied largely by complex-type glycans. The glycan N1176, which is in the epitope for antibody G4 and was reported to mask antibody recognition, was found to have a complex-type structure. Of note, we did not observe that the proteases used had biases on specific glycotytes.

Figure 6 | Application of the protocol for characterization of site-specific N-glycan processing of virus envelope glycoproteins. (a–c) Site-specific N-glycan processing of recombinant envelope glycoproteins derived from viruses, including the prefusion-stabilized spike glycoprotein ectodomain of Middle East respiratory syndrome coronavirus (MERS-CoV S-2P protein, a), influenza virus hemagglutinin from H3N2 strain A/Victoria/361/2011 (b), and HIV-1 envelope glycoprotein (c). The proportions of high-mannose and complex-type glycans at the glycosites highlighted in yellow were assigned based on the proportion of spectra hits because peak area did not reach the threshold of $5 \times 10^8$. Data were obtained from at least six independent experiments. Mean ± s.e.m. were plotted. b,c adapted from ref. 10, Nature Publishing Group.
(Supplementary Fig. 2). Of the 12 glycosites on the recombinant influenza haemagglutinin (HA) of A/Victoria/361/2011, three were >85% high-mannose (N45, N165, and N285), four were fully complex-type glycosylation (N22, N38, N63, and N483), and the rest were occupied by a mixture of high-mannose and complex-type glycans (Fig. 6b). It is striking to observe that the site N122 was not occupied and another site N144 was only 32% occupied on this HA glycoprotein. We also applied the protocol to the benchmark HIV Env trimer BG505 SOSIP.664, resulting in identification of all 28 glycosites with up to 2,000 spectra hits per glycosite (Fig. 6c). All 28 glycosites were >90% occupied, except the sites N185e, N197, N618, and N625. Of those that were largely occupied, 14 were >75% high-mannose, four were >75% complex-type glycosylation, and six other sites had a mixture of high-mannose and complex-type glycans. In particular, the glycosites, N295, N332, N339, N386, and N392, in the high-mannose patch region, were found to be occupied predominantly by underprocessed oligomannose, consistent with previous studies. The N160 glycan, which is critical for binding of the bnAbs PG9 and PG16 to HIV Env, was composed predominantly of high-mannose structures, occupied predominantly by underprocessed oligomannose, consistent with previous studies. Interestingly, the high-mannose and complex-type glycans identified at each glycosylation site of BG505 SOSIP.664 matched the pathway of N-glycan processing, in which high-mannose structures are first trimmed from Man₉ to Man₅ before addition of the terminal mannose and complex-type glycans identified at each glycosylation site of BG505 SOSIP.664 matched the pathway of N-glycan processing, in which high-mannose structures are first trimmed from Man₉ to Man₅ before addition of the terminal monosaccharides that define complex-type/hybrid glycans, as compared to the results of the same protein obtained on intact glycopeptide level. Thus, the sites were predominantly occupied by Man₉ if they were 100% high-mannose glycosylation, whereas other sites were occupied by mixtures of processed high-mannose structures (Man₉ to Man₅) and simple complex-type structures if they were occupied by a mixture of high-mannose and complex-type glycans.

We believe this protocol will be of wide interest to the proteomics and glycomics fields, and will be used by many outside those fields who want to gain high-level information about the glycoproteins they investigate.

Data availability
The data that support the findings of this study are available from the corresponding author upon reasonable request.

Further information on experimental design is available in the Life Sciences Reporting Summary.
68. Wolters, D.A., Washburn, M.P. & Yates, J.R. An automated multidimensional protein identification technology for shotgun proteomics. *Anal. Chem.* 73, 5683–5690 (2001).

69. Pallesen, J. et al. Immunogenicity and structures of a rationally designed prefusion MERS-CoV spike antigen. *Proc. Natl. Acad. Sci. USA* 114, E7348–E7357 (2017).

70. Imre, T. et al. Glycosylation site analysis of human alpha-1-acid glycoprotein (AGP) by capillary liquid chromatography-electrospray mass spectrometry. *J. Mass Spectrom.* 40, 1472–1483 (2005).

71. Pabst, M. et al. A microarray-matrix-assisted laser desorption/ionization-mass spectrometry approach for site-specific protein N-glycosylation analysis, as demonstrated for human serum immunoglobulin M (IgM). *Mol. Cell Proteomics* 14, 1645–1656 (2015).

72. Tang, N.L.S. Chemical diagnostics from bench to bedside preface. Top. Curr. Chem. 336, V–VI (2014).

73. Satomi, Y., Shimonishi, Y., Hase, T. & Takao, T. Site-specific carbohydrate profiling of human transferrin by nano-flow liquid chromatography/electrospray ionization mass spectrometry. *Rapid Commun. Mass Spectrom.* 18, 2983–2988 (2004).

74. Nilsson, J. et al. Enrichment of glycopeptides for glycan structure and attachment site identification. *Nat. Methods* 6, 809–811 (2009).

75. Hulsmeier, A.J., Tobler, M., Burda, P. & Hennet, T. Glycosylation site occupancy in health, congenital disorder of glycosylation and fatty liver disease. *Sci. Rep.* 6, 33927 (2016).

76. Guttman, M. et al. CD4-induced activation in a soluble HIV-1 Env trimer. *Structure* 22, 974–984 (2014).

77. Pancera, M. et al. Structural basis for diverse N-glycan recognition by HIV-1-neutralizing V1-V2-directed antibody PG16. *Nat. Struct. Mol. Biol.* 20, 804–813 (2013).

78. McLellan, J.S. et al. Structure of HIV-1 gp120 V1/V2 domain with broadly neutralizing antibody PG9. *Nature* 480, 336–343 (2011).
Experimental design

1. Sample size
   Describe how sample size was determined. Not applicable

2. Data exclusions
   Describe any data exclusions. Not applicable

3. Replication
   Describe whether the experimental findings were reliably reproduced. Each glycoprotein is digested in at least two technical replicates and analyzed by the same MS

4. Randomization
   Describe how samples/organisms/participants were allocated into experimental groups. Not applicable

5. Blinding
   Describe whether the investigators were blinded to group allocation during data collection and/or analysis. Not applicable

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.

6. Statistical parameters
   For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

   n/a | Confirmed
   ☑ | The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)
   ☑ | A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
   ☑ | A statement indicating how many times each experiment was replicated
   ☑ | The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section)
   ☑ | A description of any assumptions or corrections, such as an adjustment for multiple comparisons
   ☑ | The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted
   ☑ | A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)
   ☑ | Clearly defined error bars

See the web collection on statistics for biologists for further resources and guidance.

Software

Policy information about availability of computer code

7. Software
   Describe the software used to analyze the data in this study. Freely available software RawConverter is applied to exact the MS1 and MS2 data.
For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). Nature Methods guidance for providing algorithms and software for publication provides further information on this topic.

### Materials and reagents

Policy information about availability of materials

8. Materials availability
   - Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company. No

9. Antibodies
   - Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species). Not applicable

10. Eukaryotic cell lines
   - a. State the source of each eukaryotic cell line used. Not applicable
   - b. Describe the method of cell line authentication used. Not applicable
   - c. Report whether the cell lines were tested for mycoplasma contamination. Not applicable
   - d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use. Not applicable

### Animals and human research participants

Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals
   - Provide details on animals and/or animal-derived materials used in the study. Not applicable

Policy information about studies involving human research participants

12. Description of human research participants
   - Describe the covariate-relevant population characteristics of the human research participants. Not applicable