Glycoproteomic Analysis of Antibodies*‡

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Antibody glycosylation has been shown to change with various processes. This review presents mass spectrometric approaches for antibody glycosylation analysis at the level of released glycans, glycopeptides, and intact protein. With regard to IgG fragment crystallizable glycosylation, mass spectrometry has shown its potential for subclass-specific, high-throughput analysis. In contrast, because of the vast heterogeneity of peptide moieties, fragment antigen binding glycosylation analysis of polyclonal IgG relies entirely on glycan release. Next to IgG, IgA has gained some attention, and studies of its O- and N-glycosylation have revealed disease-associated glycosylation changes. Glycoproteomic analyses of IgM and IgE are lagging behind but should complete our picture of glycosylation’s influence on antibody function. Molecular & Cellular Proteomics 12: 10.1074/mcp.R112.026005, 856–865, 2013.

BIOLOGICAL ROLE OF IMMUNOGLOBULIN GLYCOSYLATION

Immunoglobulins (Igs)1 are produced by the adaptive immune system in order to identify and neutralize foreign antigens and pathogens to which the host has been exposed. In humans, five known classes of Igs (IgG, IgM, IgA, IgE, and IgD) are secreted in variable amounts by B cells during an immune response. Although these Ig classes are built from Ig domains and are thus structurally related, they differ considerably in several aspects, such as their glycosylation (1). Over the past 30 years, numerous studies have explored the structural, biological, and clinical roles of Ig glycosylation, focusing mainly on IgG molecules, which are the most abundant serum Ig, occurring at 10 to 15 mg/ml (value for IgG1) in human circulation (1). Each IgG molecule consists of two heavy and two light chains that together form two fragment antigen binding (Fab) portions and one fragment crystallizable (Fc) portion (Fig. 1). Two N-glycans are linked to the heavy chains at Asn 297 in the Cγ2 domain of the protein backbone (Fc part). These Fc glycans are in part located in a cavity between the two heavy chains and influence the conformation of the protein (2, 3). Their removal by glycosidases or via mutation of the glycosylation sites reduces the binding of IgG to Fc-gamma receptors (FcγRI) (4–6). The Fc-linked carbohydrates are complex-type biantennary N-glycans with a high level of core-fucosylation and a variable number of galactoses (Gal) resulting in the prevalent glycoforms G0F (no Gal), G1F (one Gal), and G2F (two Gal). A minor proportion of these glycans might contain a bisecting N-acetylgalactosamine (GlcNAc) residue and/or terminal sialic acids substituting antenna Gal (7) (see Fig. 1).

Many reports have described variations of IgG Fc glycosylation, especially of the degree of galactosylation, related to age, sex, heritability, and pregnancy, as well as to autoimmune diseases, infectious diseases, and cancers (e.g. Refs. 8–15). For instance, an increase in IgG G0F is observed in the serum of patients with rheumatoid arthritis (7) and correlates with disease progression and severity (16, 17). These clinical observations have led researchers to examine in detail the relationship between Fc glycan structures, the biological properties of IgG, and the degree of inflammation. It was found that an absence of sialic acids and low levels of galactosylation might confer important pro-inflammatory properties to IgG by facilitating the formation of immune complexes and favoring the binding of IgG to activating FcγRI (18–20). Similarly, the absence of core-fucose or the presence of bisecting GlcNAc improved the affinity of the Fc tail to FcγRIIa, thereby enhancing antibody-dependent cellular cytotoxicity (21–23). On this basis, new glycoengineered anti-cancer antibodies carrying afucosylated Fc glycans are currently in clinical development, such as the anti-CD20 monoclonal antibody (mAb) obinutuzumab (GA101) for use against B-cell lymphoma (24, 25).

In addition, Fc-linked glycans appear to modulate the activation of the comple ment system. Whereas the classical complement pathway can be triggered by the preferential binding of C1q to fully galactosylated IgG, the lectin pathway is recruited through the recognition ofagalactosylated IgG by mannose-binding lectin (26, 27). In contrast, the presence of terminal galactose and/or sialic acid residues on Fc glycans

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Review

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1 The abbreviations used are: ECD, electron capture dissociation; ESI, electrospray ionization; Fab, fragment antigen binding; Fc, fragment crystallizable; FTICR, Fourier transform ion cyclotron resonance; Gal, galactoses; GlcNAc, N-acetylgalactosamine; HILIC, hydrophilic interaction liquid chromatography; HR, hinge region; Ig, immunoglobulin; mAb, monoclonal antibody; RP, reversed-phase.

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might confer anti-inflammatory properties to IgG via interaction with the human lectins Dectin-1 (28) and dendritic cell-specific intercellular adhesion molecule-3-grabbing non-integrin (19, 29, 30). Thus, variations in the structure of IgG Fc glycans might skew the immune system toward a pro- or an anti-inflammatory response by modulating the interaction of IgG with several immune components, including FcγR, complement factors, and lectins. Interestingly, it was recently established that IgG Fc glycosylation may be modulated by factors such as hormones (e.g. estradiol and progesterone), cytokines (e.g. IFN-γ and IL-21), bacterial DNA (CpG oligodeoxynucleotide), and food metabolites (e.g. all-trans retinoic acid and drugs) (31–33).

The influence of glycosylation on the biological properties of other Ig classes has been poorly explored. Some reports have established that variations in the glycosylation of IgA and IgE modulate the affinity for their respective receptors, FcαR and FcεR (1). Results from clinical studies also support the idea that there is some structural and functional role of glycosylation in all classes of Ig. An example is IgA1, which exhibits O-glycosylation at various sites of its hinge region peptide (see Fig. 1). In nephropathy, lowered levels of IgA1 O-glycan sialylation and galactosylation have been observed (34). These abnormally glycosylated IgA1s were shown to have a longer half-life, to self-aggregate, and to form complexes with other molecules of the immune system, including IgG and mannose-binding lectin, thereby promoting IgA deposition in the kidney mesangium and exacerbating inflammation (1).
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tural analysis of chemically or enzymatically released glycans (15, 35, 36). Mass spectrometric analysis of glycoproteins at the glycopeptide or released glycan level are currently the methods of choice for obtaining sensitive and comprehensive glycosylation information from complex biological samples (37).

Analysis at the glycopeptide level is the most favorable approach, as site-specific glycan heterogeneity can be characterized and glycan compositions can be correlated to their attachment sites on the protein (35). In particular, liquid chromatography–mass spectrometry (LC/MS) has been widely used for glycopeptide analysis. The advantage of LC-electrospray ionization (ESI)-MS analysis is the up-front chromatographic separation of the (glyco)peptides prior to MS analysis. Obviously the choice of an efficient chromatographic separation method for a glycopeptide mixture after proteolytic digestion is crucial. For this purpose, C18 reversed-phase chromatography is the up-front chromatographic separation of the (glyco)peptides prior to MS analysis. A very convenient approach for IgG Fc glycosylation analysis is the measurement of (tryptic) Fc glycopeptides, which is generally performed via RP-LC-MS/MS (37, 48–50). Chromatographic separation is observed on the basis of small structural differences in a single amino acid side chain. Tryptic Fc glycopeptides of IgG1 carrying tyrosine residues in position 296 and 300 elute in front of tryptic IgG3/4 glycopeptides (F296 and Y300), which again elute in front of tryptic IgG2/3 Fc glycopeptides (F296 and F300). In contrast, IgG3 from Asian donors was reported to exhibit a tyrosine (Y) in position 296, resulting in identical peptide moieties for IgG3 and IgG4 (45). Thus, allotypic variations have to be taken into account when comparing subclass-specific IgG Fc-glycosylation profiles of genetically different groups.

A very convenient approach for IgG Fc glycosylation analysis is the measurement of (tryptic) Fc glycopeptides, which is generally performed via RP-LC-MS/MS (37, 48–50). Chromatographic separation is observed on the basis of small structural differences in a single amino acid side chain. Tryptic Fc glycopeptides of IgG1 carrying tyrosine residues in position 296 and 300 elute in front of tryptic IgG3/4 glycopeptides (F296 and Y300), which again elute in front of tryptic IgG2/3 Fc glycopeptides (F296 and F300). In contrast, changes in the glycan structure with regard to galactosylation, fucosylation, and bisecting affect RP retention times. Consequently, IgG1, IgG2/3, and IgG3/4 glycopeptide clusters are observed in distinct retention time windows. Isomeric tryptic Fc glycopeptide species belonging to different IgG subclasses (i.e., fucosylated IgG1 and non-fucosylated IgG3/4, or fucosylated IgG3/4 and non-fucosylated IgG2/3) are consistently separated by RP-LC, allowing their unambiguous assignment to specific IgG subclasses upon mass spectrometric detection. Sialic acid, however, can have a strong influence on IgG Fc glycopeptide retention, depending on the solvent system. The use of an acetonitrile gradient in aqueous 0.1% formic acid results in greater retention of sialylated glycopeptides eluted together with their non-sialylated counterparts. Long-term stable and robust mass spec-
Trometric analysis was achieved by employing a sheath-flow ESI sprayer with isopropanol:water:propionic acid (50:30:20; v:v:v) as a sheath liquid. The relative standard deviations for the eight major observed glycopeptide species remained less than 4% over a time range of several months, thereby allowing the analysis of thousands of samples with high precision.

HILIC-LC-MS also has been reported as a versatile tool for the separation of glycans and glycopeptides (52–54). Tryptic IgG1 Fc glycopeptides experience more retention than IgG2 Fc glycopeptides as a result of the additional oxygen atoms presented by the tyrosine residues at positions 296 and 300 (54). Furthermore, greater retention is observed with increasing glycan size/complexity, and chromatographic distinction between the 3-arm and 6-arm isomers of monogalactosylated species is often possible because of the slightly greater retention of the 3-arm isomer (54, 55). The high organic modifier content applied in HILIC mobile phases makes this separation technique particularly well suited for MS interfacing.

Fast and straightforward analysis of IgG Fc glycosylation is achieved by enriching the tryptic Fc glycopeptides using HILIC solid phase extraction followed by direct-infusion ESI-MS(/MS) (56, 57). Alternatively, MALDI-MS of purified Fc-glycopeptides can be performed with either positive- or negative-mode ionization (37, 58–60). When combined with delayed-extraction TOF detection, MALDI analysis of sialylated Fc-glycopeptides might result in a vast degree of in-source decay, largely dependent on the matrix chosen for sample preparation. When α-cyano-4-hydroxycinnamic acid is used, sialylated species are almost completely degraded. In contrast, the analysis of sialylated Fc glycopeptides is possible with 2,5-dihydroxybenzoic acid and α-cyano-4-hydroxycinnamic acid, especially when combined with negative-mode ionization (61). Interestingly, MALDI Fourier transform ion cyclotron resonance (FTICR) MS, which features an intermediate-pressure ion source, allows the registration of sialylated glycopeptides with both 2,5-dihydroxybenzoic acid and α-cyano-4-hydroxycinnamic acid (59). This may be attributed to the efficient cooling of nascent ions, which limits in-source decay (62). Although direct infusion-ESI-MS and MALDI-MS have superior throughput relative to LC/MS approaches, the accurate relative quantification of polyclonal human IgG Fc glycoforms might be compromised by the presence of isomeric tryptic glycopeptides of different IgG subclasses. However, this is not an issue with mAbs. These high-throughput
approaches show particularly high potential for biopharmaceutical quality control and fermentation monitoring (57). It has to be taken into account, however, that unlike human polyclonal IgG, which is over 95% glycosylated in the Fc moiety, biotechnologically produced IgG might contain Fc peptides holding the consensus N-glycosylation sequence but lacking glycosylation. As most non-glycosylated peptides will be lost upon HILIC solid phase extraction, RP and porous graphitized-carbon-based sample preparations might be advantageous for such samples in order to allow the simultaneous analysis of glycosylated and non-glycosylated versions of the Fc peptide covering the N-glycosylation site.

Another strategy for analyzing IgG Fc glycosylation on biopharmaceuticals involves ESI-high-resolution-MS(−M/MS) of intact mAbs or Fc portions prepared via reduction or enzymatic digestion (57, 63, 64). The high mass accuracy obtained with current high-resolution mass spectrometers allows one to determine the glycoform composition on intact monoclonal antibodies based on accurate mass with a typical 15-ppm (63) to 2-ppm (64) mass accuracy error. Moreover, up to 33% peptide sequence coverage has been reported for an intact commercial recombinant IgG using an LC-ESI-electron transfer dissociation high-resolution MS/MS approach in which time-domain transients recorded in different LC-MS/MS experiments were averaged prior to Fourier transform signal processing (64). Although intact glycoprotein analysis works well to profile Fc glycoforms on mAbs, it might not be applicable for highly complex samples such as human polyclonal IgG.

To elucidate the role of IgG Fc glycosylation in autoimmunity, inflammatory diseases, and cancer, many studies use murine disease models. Fc glycosylation of murine IgGs, however, considerably differs from that of human IgGs with regard to sialylation, fucosylation, and bisection (Fig. 2). The sialic acid on murine IgG appears to be exclusively N-glycolyl-neuraminic acid, whereas human IgG exclusively exhibits N-acetylenuraminic acid (27, 65). Moreover, serum-derived murine IgG1 and IgG2a/b both show high levels of disialylated Fc glycopeptides (signal at m/z 1127.41; Fig. 2A) (65–67). On human IgG, disialylated Fc N-glycopeptides have only recently been reported for recombinantly expressed mAb at a low relative abundance (57), but they can also be found on polyclonal IgG from human circulation, albeit at a low relative intensity (signal at m/z 1180.79; Fig. 2B). Fucosylation on murine IgG is even higher than on human IgG, with non-fucosylated glycoforms being almost completely missing. Also, bisected species are lacking on murine IgG Fc portions, making the overall glycoform repertoire of murine IgG much more restricted than that of human IgG. Thus, IgG Fc glycosylation variation observed in murine models might not directly translate to the human situation.

IgG samples that are biotechnologically produced or derived from human circulation are generally available in relatively large amounts (often microgram quantities), and the sensitivity of MS methods is therefore not an issue. It has been demonstrated, however, that IgG subpopulations might diverge considerably from total serum IgG in terms of Fc glycosylation profiles (68–71). Thus, the analysis of specific subpopulations of IgG has been found to be rewarding and has repeatedly revealed skewed glycosylation profiles that might have a profound influence on the biological activity of, for example, pathogenic autoantibodies and alloantibodies (68, 69, 71). Notably, these antibodies often may be obtained in only minute amounts by means of affinity purification, and conventional nano-LC/MS has in some cases been found to have insufficient sensitivity to analyze their Fc glycosylation. A recently reported transient-isotachophoresis separation in neutrally coated capillaries with a porous sheathless sprayer interfaced with an ultra-high-resolution TOF mass spectrometer addressed this issue, bringing the lower limit of detection down to ~20 amol (72). This high sensitivity was reached as a result of reduced ion suppression, which is typical of ESI at very low flow rates such as those used with capillary electrophoresis sheathless ESI-MS (73).

**Fab Glycosylation Analysis**—Besides the conserved N-glycosylation sites on the Fc portion, additional carbohydrate chains can be linked to the hypervariable regions of Ig. For instance, between 15% and 25% of IgG molecules isolated from the serum of healthy human subjects have been reported to carry N-glycans on their variable domains (74–76). IgG populations with Fab glycans have been called asymmetric antibodies and were found to be bound by the lectin concanavalin A (77, 78). Interestingly, the amount of asymmetric IgG was found to increase during pregnancy, as well as after the treatment of antibody-producing cells with hormones (e.g. progesterone, estrogen) and cytokines (e.g. IL-6) (79–81). More recently, HPLC and MS analyses of Fab-linked glycans from human serum IgG have revealed primarily complex-type biantennary N-glycans with high contents of core-fucose (~80%), bisecting GlcNAc (>50%), and sialic acid (~80%) (74, 75, 82). Depending on their structures and locations, the Fab glycans may influence IgG effector functions by increasing or decreasing the affinity for the antigen (1). One report furthermore suggests that Fab glycosylation could modulate antibody half-life (83). Therefore, a better understanding of IgG functionality requires a detailed analysis of Fab specific glycosylation.

The choice of an appropriate strategy for the analysis of IgG Fab glycosylation is determined by the biological source (monoclonal IgG versus polyclonal IgG antibodies). Monoclonal IgGs, which exhibit well-defined Fab glycosylation sites, can be analyzed at the level of glycopeptides and IgG portions (Fc, Fab, heavy and light chains), as well as after the selective release of Fab-glycans using glycosidases. LC/MS allows the analysis of both Fc- and Fab-glycopeptides at the same time, thereby revealing site-specific N-glycan microheterogeneity on therapeutic antibodies (84). Alternatively, the glycosylation of heavy and light chains of IgG mAbs can be studied via...
The separation of Fab and Fc fragments of IgG is generally accomplished using the enzymes papain (75, 83, 86, 87) or pepsin (75). Papain cleaves IgG just above the disulfide bridges between the two heavy chains, resulting in two Fab portions and an Fc portion of similar molecular weight (~50 kDa each). Pepsin cleaves below the disulfide bridges, generating a F(ab')2 (±100 kDa) and two ½ Fc portions (±25 kDa each). In 2002, a streptococcal cysteine proteinase, IdeS, was reported to cleave IgG specifically at a unique site below the hinge region, leading to the formation of F(ab')2 fragments with great yield and specificity (88). A recombinant version of this enzyme is now commercially available under the brand name FabRICATOR (Genovis, Lund, Sweden). A multitude of approaches have been used to purify F(ab) and F(ab')2 fragments. After pepsin digestion, Fc glycopeptides and F(ab')2 portions were separated using size exclusion (75). Papain digestion was followed by ion exchange chromatography (75) or affinity chromatography using Protein A (82, 87). In all cases, the Fab-linked N-glycans were released using PNGase F and analyzed via HPLC or MS.

Another way to separately analyze Fc and Fab glycans of a mAb is to release them from the entire IgG molecule using discriminating glycosidases and/or enzymatic conditions. For example, PNGase F and endoglycosidase F2 were reported to selectively release, in native condition, the Fc and Fab glycans, respectively (86).

Polyclonal IgGs exhibit a vast diversity of amino acid sequences of the variable regions created during somatic hypermutation, resulting in a multitude of Fab-glycosylation sites differing in number and location, as well as in the nature of their glycan chains. This enormous heterogeneity complicates, if not precludes, Fab glycosylation analysis at the glycopeptide level. Consequently, Fab glycosylation analysis of polyclonal IgG has hitherto relied on the analysis of released glycans from parts of IgG or from entire IgG molecules.

Recently, a method using sequential enzymatic release of Fc glycans and Fab glycans has been reported (74). Fab glycans, but not Fc sugars, were found to be resistant to PNGase F cleavage under native conditions (74). Therefore, IgG Fc glycans were first released under native conditions, and after IgG isolation, denaturing conditions allowed the liberation of Fab glycans.

For all techniques that use released glycans, a major drawback is that samples have to be extremely pure. Fc glycosylation is close to 100%, whereas Fab glycosylation is found on a only minor portion of polyclonal IgGs. Minor Fc contamination in Fab samples can bias the results. Fab and Fc glycosylation analysis at the released glycan level might be similarly compromised by the presence of other glycoprotein contaminants. This underlines the importance of highly specific purification methods.

**Immunoglobulin A Glycosylation Analysis**—Immunoglobulin A has several N- (IgA1 and 2) and O-glycosylation sites (IgA1 only; see Fig. 1), and both N- and O-glycosylation have been analyzed at the released glycan level (89, 90). In secretory fluids, such as mucosa and milk, two IgA molecules are dimerized by the N-glycosylated secretory component and the joining (J-)chain (91).

Site-specific N-glycosylation analysis of IgA has been done at the glycopeptide level after employing Asp-N endoprotease (92). Two N-glycopeptides were identified, and the peptide sequences were obtained by means of Edman degradation. Based on the calculated masses of these sequences, different glycan compositions were deduced from MALDI-TOF-MS of desialylated glycopeptides. Differential treatment with galactosidase and fucosidase, as well as two-dimensional HPLC on released glycans using C18 and amide columns, revealed fully galactosylated complex-type biantennary structures with or without bisecting GlcNAc and fucose (92). More recently, tryptic glycopeptides of size-exclusion chromatography-purified IgA1 have been analyzed using LC-FTICR-MS, with sequence confirmation using electron capture dissociation (ECD)-FTICR-MS/MS (93). Glycan compositions and linkages were established via gas-liquid chromatography. Interestingly, bi-, tri- and tetra-antennary complex type glycans were observed (93). N-glycosylation analysis of secretory IgA from human colostrum has recently been performed at the glycopeptide level using in-gel trypsin digestion and subsequent LC/MS and LC-MS/MS (91), revealing pronounced site-specific differences in glycosylation.

Also, the O-glycosylation of IgA has been extensively studied at the glycopeptide level (91, 94–98). Specific O-glycosylation changes were found in IgA nephropathy (34). More specifically, aberrantly glycosylated IgA1, with Gal-deficient hinge region (HR) O-glycans, plays a pivotal role in the pathogenesis of IgA nephropathy (95, 96, 99). Renfrow and coworkers showed IgA1 O-glycan heterogeneity via the use of FTICR-MS and LC-FTICR-MS to obtain accurate mass profiles of IgA1 HR glycopeptides from three different IgA1 myeloma proteins (95). Additionally, in that study, the first ECD fragmentation approach on an individual IgA1 O-glycopeptide from an IgA1 HR preparation that was reproducible for each IgA1 myeloma protein was obtained (Fig. 3). These results suggest that future analyses of IgA1 HRs from IgA nephropathy patients and healthy controls should be feasible.

Novel strategies for the analysis of clustered O-glycans involve the use of a combination of IgA-specific proteases and trypsin and ECD-FTICR-MS/MS. They provide a variety of IgA1 HR fragments that allow the unambiguous localization of all O-glycosylation sites for the six most abundant glycoforms, leading to the identification of Gal-deficient sites (96). Additionally, the published protocol was adapted for on-line LC-ECD-MS/MS and LC–electron transfer dissociation–MS/MS analysis. This work appears to be a relevant clinical approach for defining the molecular events leading to the pathogenesis of a chronic kidney disease, and at the same
time it might be generally applicable for the analysis of clustered sites of O-glycosylation (96).

PERSPECTIVES

As demonstrated extensively for IgG, as well as for some IgA, a detailed structural analysis of N- and O-glycosylation is required in order for one to understand their three-dimensional structures and immune functions. To our knowledge, the glycosylation of other Igs (IgD, IgE, IgM) has hitherto not been addressed at the glycoproteomic level. The numerous O-glycosylation sites in the IgD HR and N-glycosylation sites (≥5 N-glycosylation sites) in IgM and IgE make their comprehensive glycosylation analysis at the glycopeptide level challenging. Additionally, the analysis of IgE and IgD from human circulation is particularly demanding, as these antibodies are generally present only in minute amounts (100). For IgE, glycoproteomic analysis would be needed in order to allocate its complex type and oligomannosidic glycans to their specific site(s) and analyze the NxS site on position 383, which has been predicted to be unoccupied (101). A particular analytical challenge will be the analysis of the variable region glycosylation of Ig subclasses other than IgG.

Ig glycosylation studies are routinely done in many different labs, and thus the amount of data produced is increasing tremendously. A recent approach combining genome-wide association and high-throughput glycomics analysis of plasma samples from 2705 individuals in three population cohorts showed that common variants in certain genes can influence N-glycan levels in human plasma (102). Based on a follow-up study, a high-throughput isolation and glycosylation analysis of IgG variability and heritability of the IgG glycome in three different populations was published (10). Although a variety of associations of clinical and physiological parameters with Ig glycosylation have been established, we believe that many more processes and diseases are marked by Ig glycosylation changes and that we have seen only the tip of the iceberg. Future Ig glycosylation profiling at the site-specific level by means of the mass spectrometric analysis of glycopeptides, when applied to human disease cohorts as well as in vitro and in vivo models of immunological processes, is expected to provide valuable new insights into the modulatory role of Ig glycosylation.

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