SHORT COMMUNICATION

Glycoform-resolved FcγRIIIa affinity chromatography–mass spectrometry

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ABSTRACT

Determination of the impact of individual antibody glycoforms on FcγRIIIa affinity, and consequently antibody-dependent cell-mediated cytotoxicity (ADCC) previously required high purity glycoengineering. We hyphenated FcγRIIIa affinity chromatography to mass spectrometry, which allowed direct affinity comparison of glycoforms of intact monoclonal antibodies. The approach enabled reproduction and refinement of known glycosylation effects, and insights on afucosylation pairing as well as on low-abundant, unstudied glycoforms. Our method greatly improves the understanding of individual glycoform structure–function relationships. Thus, it is highly relevant for assessing Fc-glycosylation critical quality attributes related to ADCC.

Fcγ receptors (FcγR) are key elements in many immunological responses. FcγRIIIa has an important role in mediating antibody-dependent cell-mediated cytotoxicity (ADCC). Thus, investigating antibody–FcγRIIIa interactions is highly relevant for fundamental and clinical research, as well as (bio)therapeutic innovations. The Fc glycosylation of immunoglobulin G (IgG) has large effects on its FcγRIIIa interaction. Most significantly, IgG fucosylation reduces FcγRIIIa affinity by up to 100-fold. This is in part attributable to a unique glycan–glycan interaction between the IgG Fc and the FcγRIIIa N162 glycan. Knowledge of the effects of IgG glycosylation led to the successful development and approval of FcγRIIIa-affinity liquid chromatography (AC) and functional-FcγRIIIa affinity chromatography (SPR) or AC necessitated high IgG glycoform purity. Advantageous features of multiple IgG glycoforms of a therapeutic mAb. This was achieved by AC hyphenated to mass spectrometry (MS). MS allowed molecular resolution while the separation dimensionality provides FcγRIIIa (V158) affinity. Several glycosylation traits are therefore critical quality attributes. Analytical methods for assessing effector functions and Fc/Fc-receptor interactions in therapeutic antibodies were recently reviewed. Functional cellular activity assays have the advantage of high relevance towards the in vivo situation. The downstream signaling is determined by a complex interplay of immune complexes binding to activating and inhibitory FcγR. The interaction of FcγRIIIa with the Fc portion occurs in an asymmetrical 1:1 stoichiometry that precludes cellular activation in vivo by monomeric antibodies. Thus, the translation of functional cellular assays is more straightforward compared to cellular or cell-free binding assays studying monomeric antibodies. However, the fundamental interactions of Fc and FcγRIIIa do not depend on whether IgG is present in monomeric form or as an immune complex, and it is important to study them. Several cell-free physico-chemical assays are well established and generally, the first choice for assessing the potential impact of monomeric IgG Fc-attributes on receptor binding due to their increased analytical performance with regard to assay complexity, affinity resolution, and robustness. Moreover, these binding assays strongly linked in vitro FcγRIIIa affinity and ADCC activity. Explicitly, retention time differences in FcγRIIIa affinity liquid chromatography (AC) were linked to ADCC. A common disadvantage of all previous methods is the averaged and potentially biased output because of the naturally occurring IgG glycoform heterogeneity. Therefore, unraveling the impact of individual IgG glycoforms on FcγRIIIa affinity previously required laborious glycoengineering. The lack of molecular resolution of established in vitro affinity assessment techniques, such as surface plasmon resonance (SPR) or AC necessitated high IgG glycoform purity. Here, we present the simultaneous assessment of FcγRIIIa affinity of multiple IgG glycoforms of a therapeutic mAb. This was achieved by AC hyphenated to mass spectrometry (MS). MS allowed molecular resolution while the separation dimensionality provides FcγRIIIa (V158) affinity. Advantageous features are: (1) affinity assessment of individual, previously unstudied glycoforms from biosynthetic mixtures, omitting the need for advanced glycoengineering; and (2) increased affinity differentiation compared to established in vitro techniques due to the simultaneous assessment. MS has become an important technique for characterizing intact proteins. The combination with AC (AC-MS) proved its potential for functional characterization of
therapeutic mAbs recently. AC-MS based on the fetal/neonatal Fc receptor (FcRn) showed decreased FcRn affinity, and by extension IgG half-life, for a mAb oxidized at M255. However, FcRn affinity is only very weakly influenced by glycosylation. Therefore, resolving complex glycosylation heterogeneity molecularly on an intact protein level is necessary for FcγRIIIa-AC-MS, which makes it more challenging and more powerful at the same time. For method development, a good balance between MS response and separation efficiency is crucial. In contrast to previous AC-UV studies, we employed a simple ammonium acetate buffer. We optimized ammonium acetate concentration and linear pH gradient (Supplemental Figure 1), aiming at an at least equal separation efficiency compared to the previously reported FcγRIIIa AC-UV conditions (Figure 1(a)). This was achieved using 50 mM ammonium acetate and a pH gradient from pH 5 – pH 3 (Figure 1(b)). We used the same column for which FcγRIIIa AC retention times were previously linked to ADCC and obtained comparable profiles (Figure 1). A 15 T Fourier transform ion cyclotron resonance (FT-ICR)-MS instrument provided high sensitivity and mass accuracy for determining even low-abundant glycoforms (Figure 1(c,d), Supplemental Table 3 and 4). We detected 21 compositions, reflecting 27 partially isomeric glycoforms (Figure 1(b) and Supplemental Table 4). Typical charge state distributions ([M + 24H]24+ to [M + 29H]29+) of folded protein conformations were observed for the entire pH gradient (inserts Figure 1(c,d)). The method showed very good intra- and inter-day variability of relative peak areas for the resolved glycoforms of the mAb (Supplemental Figure 3 and Table 5). Further,
relative peak areas of glycoforms of the intact mAb showed good agreement compared to the glycoform profile based on released glycan analysis (Supplemental Figure 5). This demonstrated the feasibility of performing relative quantitation and provides additional proof of correct glycoform assignment.

AC-MS enables the simultaneous in vitro comparison of FcγRIIIa affinity of antibody glycoform mixtures, in contrast to SPR or AC-UV. Previously, FcγRIIIa affinities of seven glycoengineered mAbs were assessed using AC-UV. More recently, AC-UV and subsequent glycan analysis were used to assess the heterogeneity of Fc-glycosylation of a therapeutic antibody, using a non-glycosylated FcγRIIIa. However, glycosylation has been shown to be critical for differential recognition of IgG-glycoforms by wild-type FcγRIIIa. Thus, no significant differences for individual glycoforms, except for glycoform levels, were observed. With our method, we can directly compare the affinity of 21 glycoforms present in a therapeutic mAb without the need for fractionation or glycoengineering. Herein, we use a human embryonic kidney (HEK) cell-produced FcγRIIIa. This may lead to altered glycosylation and affect affinity/selectivity compared to the natural receptor. Currently, available natural FcγRIIIa data is missing site-specificity or is inferred (and incomplete). While the majority of CHO and HEK cell glycoforms seems to qualitatively overlap with natural glycosylation, neither seems good at producing triantennary structures. HEK cells can produce antennary fucosylation, but also unnatural LacDiNAc structures. At the same time, the functional impact of individual FcγRIIIa glycoforms is unclear.

Fucosylation is known to have the highest impact on FcγRIIIa affinity. Single afucosylation is sufficient to drastically increase binding, owing to the asymmetric 1:1 binding between IgG and FcγRIIIa. Consequently, glycoforms containing fucosylation on both heavy chains (2x fucosylated) populate the earlier eluting peak in the AC chromatogram (Figure 1(b,c)). The 1x (a)fucosylated and 2x afucosylated species are found in the later eluting AC peak (Figure 1(b,d)). However, the AC-MS system achieves a much more detailed differentiation of affinity than AC-UV, because it can rank FcγRIIIa affinity of multiple (partially) co-eluting glycoforms in one run (Supplemental Table 6). In contrast, AC-UV requires comparison of multiple runs, making retention time stability a limiting factor. Other assays only give an average value for all glycoforms present, resulting in a potentially significant impact of even low-abundant glycoforms of vastly higher affinity. As a result, we could demonstrate that 2x afucosylated complex glycoforms (G0(N)/G0(N), G0(N)/G1(N)) show an increased FcγRIIIa affinity over 1x (a)fucosylated (Figures 1(b) and 2(a)). Furthermore, AC-MS can distinguish the positive influence of galactosylation on FcγRIIIa affinity in all fucosylation variants (Figures 1(b), 2(a), Supplemental Figure 4). Previously, this was only observed for 2x fucosylated glycoforms by AC-UV and mainly 1x afucosylated glycoforms by AC-UV and SPR. More recently, a comprehensive binding study applying AC-UV, SPR and ADCC analysis showed the positive influence of afucosylation and galactosylation for highly homogeneous 2x fucosylated glycoforms and 0x fucosylated glycoforms.

A further challenge in glycoengineering is biosynthetic interferences between the different glycosylation features. For example, bisection correlates with other glycosylation features depending on the expression system, which complicates independent affinity assessment. Furthermore, glycoengineered mAbs often still exhibit a high degree of glycosylation heterogeneity. The high molecular resolution of our method allows the study of subtle FcγRIIIa affinity differences by dissecting the contribution of individual glycostructure features. For example, we could demonstrate that bisection increases FcγRIIIa affinity independent of afucosylation and galactosylation. This was observed regardless of the presence of galactoses (Figure 2(b)).

In addition, FcγRIIIa affinity of typically low-abundant glycoforms (=0.1% relative abundance, Supplemental Table 5) could be studied (Figure 1(b)). For example, mono-antennary glycoforms lacking N-acetylgalactosamine (-N) showed decreased affinity compared to its complex bi-antennary glycoforms (e.g., G0F/G0F-N vs. G0F/G0F, Figure 1(b), Supplemental Figure 4 and Table 6). Glycoengineering, and therefore classical affinity assessment, of low-abundant species, is often deemed too tedious, a limitation efficiently addressed by our method.

High mannose species have been shown to increase ADCC activity via FcγRIIIa binding that was attributed to the lack of core-fucose. A minor amount of high mannose glycans,
paired with either another high mannose (0.8% M5/M5) or the complex glycan (0.2% M5/G0F), was detected in the therapeutic mAb (Figure 1(b)).^28 Largely, these showed higher affinity compared to 2x fucosylated glycoforms. However, the affinity of M5-containing glycoforms was reduced compared to 1x (a)fucosylated complex glycoforms. These findings are supported by previous studies describing increased binding affinities^12,23,27 and ADCC activities^12,23 of high mannose glycoforms over 2x fucosylated complex glycoforms, and similar or decreased affinities and activities compared to afucosylated complex glycoforms.\(^{12,13,27}\) Identity and affinity of the M5/M5 were confirmed by glycoengineering (Supplemental Figure 6 and 7). Moreover, the isolated analysis of the glycoengineered M5/M5 mAb showed comparable retention as observed in the therapeutic mAb, giving no indication for a measurable bias in retention behavior caused by competitive receptor binding of other glycoforms.

Glycoforms with one non-glycosylated heavy chain (X/NG) are normally also found to a low extent in therapeutic antibodies.\(^{29}\) Decreased binding affinities and ADCC activities were observed for these glycoforms in previous studies.\(^{30}\) We observed elution of fucosylated glycoforms (G0F/NG, G1F/NG, G2F/NG) already before the pH gradient (for initial pH of 5) started, whereas G0/NG showed the lowest affinity of the afucosylated glycoforms, but it exhibited a higher affinity compared to the 2x fucosylated glycoforms (Supplemental Table 6, Supplemental Figure 4).

In conclusion, FcyRIIIa AC-MS represents a great advance in characterizing the impact of antibody Fc glycosylation on FcyRIIIa affinity. It omits the need for high purity glycoengineering, which is notoriously difficult, especially for biosynthetic pathway intermediates,\(^{24}\) and has hitherto complicated and, in part, prohibited the affinity assessment of various glycoforms. We were able to perform an FcyRIIIa affinity assessment of individual glycoform pairings present in a therapeutic mAb, including so far unstudied, low-abundant glycoforms (e.g., G0F-N/G0F). Due to its high molecular and high FcyRIIIa affinity resolution, the method allows independent assessment of individual glycosylation features. For example, we could show, for the first time, the increased affinity of afucosylated complex glycoforms over singly (a)fucosylated complex glycoforms and high mannose glycoforms. Furthermore, we could support the independent, positive influence of bisection. FcyRIIIa AC-MS will allow a fast and reliable assessment of FcyRIIIa affinity of novel (synthetic) glycoforms, even in the presence or excess of other glycoforms.

FcyRIIIa AC-MS may be highly useful in other applications. For example, other post-translational modifications as well as protein backbone differences may be analyzed in addition to glycosylation, again separating independent effects. Thus, FcyRIIIa AC-MS will be an enormously valuable tool in studying and tuning ADCC activation of (therapeutic) antibodies. Largely by the omission of high purity glycoengineering, proteoform-related critical quality attributes and their interactions could be defined earlier in therapeutic mAb development. In the future, glycoengineering of FcyRIIIa towards natural glycosylation would be highly desirable. In general, AC-MS approaches have a high potential for studying biomolecular interactions in a proteoform-resolved manner and are extremely valuable in unraveling structure–function relationships.

**Materials and methods**

**Chemicals**

All chemicals were purchased from Sigma-Aldrich (Steinheim, Germany) and had at least analytical grade quality. Deionized water was used from a Purelab ultra (ELGA Labwater, Ede, The Netherlands). Ammonium acetate solution (7.5 M, Sigma-Aldrich) and glacial acetic acid (Fluka-Honeywell) were used for preparing MS-compatible mobile phases.

**Antibodies**

In this study, four different monoclonal antibodies (mAbs) differing in glycosylation were used. First, a standard therapeutic mAb covering a typical degree of glycosylation heterogeneity was investigated. In addition, a glycoengineered high mannose-containing variant of the first mAb and a mAb containing a high level of afucosylation and bisection were analyzed. Lastly, a highly fucosylated and highly bisected anti-D mAb were investigated (reported as +B).\(^{12}\) The first three mAbs were provided by Roche Diagnostics (Penzberg, Germany) and the last one by Sanquin (Amsterdam, The Netherlands).

**FcyRIIIa column preparation**

FcyRIIIa affinity column was provided by Roche Diagnostics (Penzberg, Germany). The preparation of the FcyRIIIa affinity column is reported elsewhere.\(^{9}\) In brief, biotinylated human FcyRIIIa_V158 (HEK cell-produced with glycosylation profile as reported\(^{22}\)) was incubated with streptavidin sepharose for 2 h upon mild shaking and the receptor-derivatized sepharose was packed in a Tricorn 5/50 Column housing (inner diameter 5 mm x length 50 mm, GE Healthcare).

**FcyRIIIa affinity liquid chromatography**

For FcyRIIIa affinity liquid chromatography with non-MS compatible conditions and UV detection (absorbance measured at 280 nm), previously reported pH gradient and mobile phase composition were used.\(^{13}\) Equilibration buffer/mobile Phase A consisted of 20 mM sodium citrate pH 6.0 and 150 mM NaCl. Mobile phase B contained 20 mM citrate pH 3.0 and 150 mM NaCl. Final MS-compatible conditions contained 50 mM ammonium acetate pH 5.0 (mobile phase A) and 50 mM ammonium acetate pH 3.0 (mobile phase B). All experiments were performed using a biocompatible Thermo Ultimate3000 instrument at 25 C. The samples were diluted with mobile phase A, if necessary (conc. 1–5 µg/µL). Injection volume was set to 10 µL. The flow rate was set to 500 µL/min. For hyphenation with MS, the flow rate was reduced from 500 µL/min to 30 µL/min via flow-splitting prior to the
electrospray ionization (ESI) source. Upon injection, the column was washed with 5 column volumes (10.5 min) of mobile phase A, and the sample was eluted in a linear gradient of 15 column volumes (31.5 min) ending with 100% mobile phase B. In addition, a washing step using 5 column volumes (10.5 min) of mobile phase B and a linear pH gradient of 7.5 min ending with 100% mobile phase A was applied after the elution gradient. Re-equilibration using 15 column volumes (31.5 min) of mobile phase A was applied between runs.

**Fourier transform ion cyclotron resonance mass spectrometry**

Online ESI-MS coupling of FcγRIIIa AC was performed on a Bruker 15 T solariX FT-ICR-MS (Bruker Daltonics, Bremen, Germany). Instrument tuning and calibration were performed using direct infusion (2 µL/min) of the therapeutic mAb (0.1 µg/µL in mobile phase A) (Supplemental Figure 2 and Table 2). The m/z of all charge states of the main glycoform (G0F/G1F) were used for calibration. By using the charge state envelope of the most abundant glycoform, the relevant mass range was sufficiently covered. External calibration was not needed for the intended purpose of the study because the masses and glycoforms of the analyzed mAbs were known. ESI capillary voltage was set to 4000 V and endplate offset to −500 V. Nebulizer gas pressure was set to 0.8 bar, dry gas to 3 L/min and source temperature to 200 C. The ion funnel 1 was operated at 150 V, radio frequency amplitude at 300 Vpp and skimmer 1 at 125 V. Spectra were acquired in an m/z-range of 506–20,000. Accumulation time was set to 1 s. The amount of data points was set to 128 k. Each spectrum in serial mode analysis resulted from the summation of 20 spectra. This resulted in the acquisition of 2.6 data points per minute in the chromatogram. This acquisition rate was chosen as a compromise of sensitivity (for low-abundant glycoforms) and resolution, and it was sufficient for the generally broad eluting peaks observed in this study. All mass spectra were visualized and processed using DataAnalysis 5.0 (Bruker Daltonics). For deconvolution, the Maximum Entropy tool was used (145,000–150,000 Da, datapoint spacing = 1, instrument resolving power = 3000). Further information regarding data processing are described in the Supplemental Material.

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**Disclosure statement**

A. Heidenreich, D. Reusch, and M. Habenger are employees of Roche Diagnostics, the manufacturer of the column and the therapeutic antibodies used in this research.

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