GLYCOSYLATION PATTERN CHARACTERIZATION OF NATURAL AND RECOMBINANT GLYCOPROTEINS USING A TWO-DIMENSIONAL MAPPING TECHNIQUE

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Introduction

Several glycan profiling methods have been recently published. They are based on measurement of the oligosaccharide effective size on gel filtration chromatography (1), High Performance Anion Exchange Chromatography retention times and electrophoretic mobilities on High Performance Capillary Electrophoresis (2, 3); and 2-3 dimensional (2D-3D) carbohydrate mapping database of pyridylaminated-oligosaccharide derivatives using C18 and amide-silica HPLC and an extra anion exchange column, respectively (4, 5).

The methodology that will be described is based on the separation of derivatized oligosaccharides on Fluorophore Assisted Carbohydrate Electrophoresis (FACE) (6) and Amine Adsorption-HPLC (7). The first report about the suitability of separation of ANTS (8-anime-1,3,6-naphthalene tri sulfonic acid) oligosaccharide derivatives using NH2-HPLC under ion suppression conditions is advanced. We propose a two-dimensional (2D) sugar-mapping technique for (ANTS-derivatives) of neutral and sialyl oligosaccharides as a simple and sensitive technique for structural characterization of N-linked oligosaccharides from natural and recombinant glycoproteins using only picomoles of samples. In addition, the contribution of each monosaccharide residue was determined which facilitate the understanding of the behavior of asialo and sialo complex oligosaccharides. The proposed methodology includes: i) reductive amination with ANTS of enzymatically released oligosaccharides, ii) simultaneous separation of derivatized oligosaccharides by FACE and NH2-HPLC column under ion suppression conditions, iii) plotting of the relative migration indexes (RMI with respect to a mixture of maltoligosaccharides of different degree of polymerization) (X-axis) and relative retention times (tRMan7 relative to Man-GlcNAc2 oligosaccharide) (Y-axis) in a two coordinates graphic. This methodology fulfill almost all the requirements for a complete characterization of neutral and charged oligosaccharides released from N-glycosylated glycoprotein as it is demonstrated with several examples of natural and DNA-recombinant glycoproteins.

Results and Discussion

The analysis of oligomannosides and complex oligosaccharides using FACE and NH2-HPLC gave structural complementary information. Individual monosaccharide contributions to RMI and tRMan7 were determined. Oligomannose series shows a constant increment per mannone residue added of 0.7 GU and 0.09 on RMI and tRMan7 respectively. The deletion of a Gal residue in asialo complex oligosaccharide decreases the RMI and tRMan7 in 1 GU and 0.08 respectively. Furthermore, a larger effect was observed when asialo and sialo complex oligosaccharides are compared. The introduction of a sialic acid determines faster migration on FACE while the same sialic acid produces a remarkable increase on HPLC retention time. Changes on sialic acid linkage configuration are also detected on the RMI and tRMan7.

A two-dimensional plotting of RMI vs tRMan7

More than 40 different standard oligosaccharides were studied in terms of electrophoresis and HPLC behavior. All these compounds were plotted into a 2D graphic of RMI referred as GU and retention time relative to Man7 (tRMan7). Figure 1 shows a family of straight lines corresponding to oligomannosides, asialo and sialylated di, tri and tetraantennas. Incorporation of a new member of the oligomannosides or asialo complex series will meet their own curves due to the constant contribution of the corresponding monosaccharides. Analysis of sialo oligosaccharides, showed that, to each antennary structure corresponds one line e.g. the di, tri or tetraantennary lines are composed by the elements of the same structural motif but varying on the number and configuration linkages of the attached sialic acids. All this elements determines straight lines with negative slopes. Since the addition of a Fuc unit to the inner core is depicted by a positive shift of the corresponding point, then, the fucosylated isomers generate a new set of points and thus a new line parallel to the unfucosylated structures. As well as Fuc addition generates a new line, some other factors does, e.g. the deletion of galactose or N-Acetyl glucosamine units and the introduction of lactosamine extensions (data not shown).

Apart from the already described straight lines another set of parallel lines with positive slopes can be detected. All these ones are characterized by a parameter "L" which is a function of the "effective charge" disclosed by the molecule. Thus, this parameter "L" is, among other factors, dependent of the number of functional carboxylic groups and on the configuration linkages of the sialic acid at the non-reducing end of the sialo oligosaccharide. Each new line is constituted by the RMI and \( t_{\text{Man}^7} \) values of oligosaccharides which display the same "effective charge" determined by the equation:

\[
L = 2 \times \# \text{Sialic Acid (} \alpha 2,6 \text{ linked) } + 1 \times \# \text{Sialic Acid (} \alpha 2,3 \text{ linked)}
\]

The cross-linking of these straight lines facilitate the preliminary structural characterization of sialylated oligosaccharides. Therefore a sialylated oligosaccharide released from N-glycosylated protein could be characterized, at least, in terms of the number of antennae, the presence or not of inner core fucosylation and the number and type of sialic acids.

**N-linked glycan profiling of glycoproteins**

IgG 2a produced in vivo and in vitro gave a typical N-linked glycosylation profile of the IgG family showing small asialo oligosaccharides as has been reported before. The major structures corresponds to the monogalacto-core fucosylated diantennae (in two positional isomers) followed by the galacto-core fucosylated diantennae in proximal ratios of 0.5:1 and 0.8:1. Other two minor species were detected, the digalacto-core fucosylated and the unfucosylated galacto diantennas (Figure 2-4).

Other examples were also studied and the corresponding pair of values of each oligosaccharide were plotted in the 2D graphic (Figure 1). These include the glycosylation profiles of several heterologous proteins expressed in the methylotrophic yeast *Pichia pastoris*, where Man\(_3\)GlcNAc\(_2\) was the common oligosaccharide. An structural modification of this oligosaccharides was possible to be characterized in conjunction with exoglycosidase digestion. Fungal proteins were also analyzed as was the case of *Trichoderma reesei* endoglucanase 1 "glycoforms". The presence of charged species was considered due to low RMI values and strong retention in HPLC. When plotting the RMI and \( t_{\text{Man}^7} \) in the 2D graphic, the corresponding points lay on parallel lines to the oligomannosides standards and proximal to \( L = 3 \) and \( L = 7 \) lines of sialylated oligosaccharides.
This behavior suggested phosphorylation of oligomannosides oligosaccharides. This was further confirmed by ESI-MS in combination with mild acid hydrolysis and alkaline phosphatase digestion.

**Conclusions**

- The most relevant conclusion is the achievement of a 2D analysis through the electrophoretic (RMI) and chromatographic (t<sub>Man<sup>7</sup></sub>) parameters of the same ANTS-oligosaccharide derivatives.
- The existence of a complex network which in conjunction with the defined contribution of each monosaccharide gives a predictive character to this web.
- Linear dependence within each group of compounds (asialo, mono, di, tri or tetrasialylated oligosaccharides), greatly facilitate the structural characterization of an isolated oligosaccharide.
- The developed methodology serve to analyze N-linked glycosylation pattern of natural and recombinant glycoproteins.

![Figure 3. FACE separation of N-linked ANTS-derived oligosaccharides of monoclonal IgG 2a. G5 represents the position of a malto pentasaccharide; lane A) in vitro sample oligosaccharides; lane B) in vivo sample oligosaccharides and lane C) mixture of standard asialo complex diantennary oligosaccharides.](image)

![Figure 4. Asialo diantennary structures identified from monoclonal IgG 2a; 1 - 1', 2 - 2', 3 - 3' and 4 - 4' corresponds to the peak numbers from the HPLC chromatogram.](image)