Analysis of zwitterionic and anionic N-linked glycans from invertebrates and protists by mass spectrometry

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Abstract Glycomic analyses over the years have revealed that non-vertebrate eukaryotes express oligosaccharides with inorganic and zwitterionic modifications which are either occurring in different contexts as compared to, or are absent from, mammals. Examples of anionic N-glycans (carrying sulphate or phosphate) are known from amoebae, fungi, molluscs and insects, while zwitterionic modifications by phosphorylcholine, phosphoethanolamine and aminoethylphosphonate occur on N-, O- and lipid-linked glycans from trichomonads, annelids, fungi, molluscs, insects, cestodes and nematodes. For detection of zwitterionic and anionic glycans, mass spectrometry has been a key method, but their ionic character affects the preparation and purification; therefore, as part of a glycomic strategy, the possibility of their presence must be considered in advance. On the other hand, their ionisation and fragmentation in positive and negative ion mode mass spectrometry as well as specific chemical or enzymatic treatments can prove diagnostic to their analysis. In our laboratory, we combine solid-phase extraction, reversed and normal phase HPLC, MALDI-TOF MS, exoglycosidase digests and hydrofluoric acid treatment to reveal N-glycans modified with anionic and zwitterionic moieties in a wide range of organisms. It is to be anticipated that, as more species are glycomically analysed, zwitterionic and anionic modifications of N-glycans will prove rather widespread. This knowledge is - in the longer term - then the basis for understanding the function of this cornucopia of glycan modifications.

Keywords Glycomics · Mass spectrometry · Phosphate · Sulphate · Glucuronate · Phosphorylcholine · Phosphoethanolamine · Aminoethylphosphonate

Introduction Glycoconjugates are not merely the product of ‘ligation’ of an oligosaccharide with the underlying protein or lipid, but can be further modified with non-sugar anionic components such as sulphate and phosphate. Due to their charged character, these modifications have a profound effect on recognition events in vivo. For example, mannose-6-phosphate is well known as being necessary for recognition and targeting of mammalian lysosomal enzymes [1], the clearance of mammalian pituitary glycoprotein hormones by Kupffer cells is mediated by lectins binding sulphated GalNAc [2] and the sulpho-glucuronate-based HNK-1 epitope has functions in neural development [3]. However, the biological significance of glycan phosphorylation and sulphation (other than proteoglycans) in non-mammalian systems remains obscure, but this is in part due to a probable underestimation of their occurrence. Nevertheless, previous reports indicate that methyl and mannose phosphodiesters occur respectively, on slime mould and yeast glycoproteins [4, 5] and that sulphate is a component of fucans and galactans from marine organisms [6], invertebrate glycosaminoglycans [7], sea urchin sialoconjugates [8], plant chloroplast sulfoquinovosyldiacetylglycerol [9] and Dictyostelium N-glycans [4, 10]. More recent data (see also Fig. 1) using mass spectrometry mean that this list can be extended to include sulphate on mollusc and insect N- and O-glycans [12–14]. Additionally, rather well known from mammalian systems are the ‘organic’ anionic modifications of glycans including sialic and glucuronic acids; in protists, the occurrence of sialic
Acid is probably restricted to trypanosomatids which steal this unit from host glycoproteins via their trans-sialidase [20]. In contrast, the presence of sialic acid in insects was, for some years, a controversial topic, but low levels of sialylated N-glycans and a relevant sialyltransferase were found in Drosophila [21, 22]. Sialylation may have roles in neural development in Drosophila [23] and fertilisation of sea urchins [24]. Glucuronic acid, on the other hand, is present as a component of chondroitin and heparan sulphates from all animals, including invertebrates [25] but is now known to be present on insect N-, O- and lipid-linked glycans [12, 26, 27]; it has been recently found (also in methylated form) on the antennae of N-glycans of two mollusc species [14, 28]. On the other hand, glucuronic acid (but not sialic acid) has been found on O-glycans from both Schistosoma mansoni and Caenorhabditis elegans [29, 30], but neither have been detected, to date, as modifications of nematode or trematode N-linked glycans.

In terms of the zwitterionic modifications, phosphorylcholine and phosphoethanolamine are of immunological relevance as targets of mammalian pentraxins (e.g., C-reactive protein) and antibodies [31, 32]; furthermore, phosphorylcholine is an immunomodulatory modification of N-glycans and glycolipids from nematode parasites which may interact directly or indirectly with Toll-like receptors of immune cells and so effect the balance of host immune systems [33]. However, other than the phosphoethanolamine on glycosylphosphatidylinositol anchors [34], zwitterionic modifications of glycans are unfamiliar to those who specialise in mammalian glycomics. In 'lower' organisms (for some examples, see Fig. 1), the zwitterion phosphorylcholine has been reported as a phosphodiester modification of fungal cell walls [35], of nematode and cestode N-glycans [15, 36, 37] and of glycolipids from fungi, annelids and nematodes [38–40], whereas phosphoethanolamine, the non-methylated version of phosphorylcholine, is present on trichomonad and fungal N-glycans as well as on insect O-glycans and glycolipids [16, 17, 41, 42]. The ‘variant’ of phosphoethanolamine lacking the oxygen ‘bridge’, thereby possessing a C-P bond, is known as aminoethylphosphonate and has been detected (also in methylated form) on mollusc, cnidarian and locust glycoconjugates [14, 18, 43, 44].

For glycan analysis, the presence of anionic moieties is known to pose problems, as their physicochemical properties
alter characteristics during purification or result in altered ionisation [45]. Some of the same problems and solutions also apply to zwitterionically-modified oligosaccharides as their polarity means that such glycans can be lost upon organic or solid phase extraction. Also, other than well-characterised sialidases and phosphatases, there are no (pure) enzymes known to cleave any of these modifications, but some chemical treatments (hydrofluoric acid or methanolic HCl) are possible. Certainly, a pre-requisite for understanding the evolution and function of anionic and zwitterionic glycans in non-vertebrate systems is the adequate detection and structural elucidation of the relevant glycomodifications, which have variable effects on chromatographic separations or are of low abundance leading to them being easily overlooked; here, we summarise ways to analyse such glycans in non-mammalian systems, also in a historical context, but with a focus on N-glycomic methodologies used in our laboratory. Naturally, such approaches are also applicable to, or in part first used with, anionic glycans from vertebrates.

Purification of zwitterionic and anionic glycans

A first step in a detailed global glycomic analysis is to consider the means of releasing the oligosaccharide chains from the proteins or lipids. For cleavage from polypeptides, both chemical and enzymatic approaches are possible. Whereas the former (e.g., hydrazinolysis or β-elimination) can prove harsh and can affect labile residues on the analytes, the latter are restricted by enzyme specificities, which mean that some classes of glycans will not be released. For N-glycans, peptide:N-glycosidases (PNGases) are known from both bacterial and eukaryotic sources; most used are the commercially-available ones from Flavobacterium (PNGase F) and almond (PNGase A). The familiar PNGase F is unable to cleave core α1-3-fucosylated N-glycans [46] while being less affected by the peptide length; the opposite is true for PNGase A, which can remove such glycans, but optimally only from small peptides. Therefore, as core α1-3-fucose is common in invertebrates and in some protists, best results are obtained if both enzymes are employed sequentially. Indeed, we generally proteolyse samples and then treat the glycopeptides first with PNGase F to release the bulk of the N-glycans (e.g., oligomannosidic and core α1-6-fucosylated structures) prior to using PNGase A to cleave the remaining ones carrying the core α1-3-fucose. This selective release results in a further deconvolution of complex glycomes even before the enrichment into neutral and anionic pools (Fig. 2). Whereas new PNGases have been recently discovered with broader substrate specificities [47–49], a universal O-glycanase is yet to be found.

Although glycans are often analysed as mixtures, it is undoubtedly optimal if a suitable fractionation method is used to overcome suppression of glycans of low abundance. Normal phase, reversed phase and graphitised carbon materials are commonly used in glycan purification, either to initially ‘clean-up’ a glycan preparation, crudely separate glycan pools or purify into fractions with each containing a low number of glycan species. Graphitised carbon has proven to be an invaluable solid phase for the enrichment of anionic oligosaccharides, whether the anionic moiety be sialic acid, glucuronate, sulphate or phosphate [4, 12, 50, 51]; however, dominant monoanionic glycans are also found in the ‘neutral’ fraction. Typically, we elute the neutral-enriched fraction with 40 % acetonitrile prior to the ‘acidic-enriched’ pool with 0.1 % trifluoroacetic acid and these pools are often then ready for mass spectrometric analyses, although occasionally a subsequent reversed-phase step may be required as in a recent study on a marine snail [14]. The next option is whether to analyse free, reduced, permethylated or fluorescently-labelled N-glycans. In our own studies, we observe that labelling with 2-aminopyridine, which facilitates the use of reversed phase HPLC columns, also improves the ionisation of the oligosaccharides and the fragmentation of the core region (thus allowing direct demonstration of core fucosylation; Fig. 2).

During the analysis of fluorescently-labelled N-glycans, classical C18 columns (e.g., Hypersil) are commonly employed [52], but in a study on the free-living nematode Pristionchus pacificus we compared Hypersil C18 with two fused-core reversed-phase resins (RP-amide and Kinetex XB-C18). Thereby, differences in the retention of phosphorylcholine-modified glycans were observed; for the standard C18 material, these glycans were rather highly-retained, whereas on the fused core columns they eluted earlier and, in general, isomers were more finely separated [19]. This means that, for a typical polishing step with C18 reversed-phase resin, sufficient organic solvent must be applied to elute phosphorylcholine-modified glycans; otherwise these glycans may be lost. In contrast, N-glycans modified with phosphoethanolamine, methylphosphate or sulphate elute early on a standard C18 column [16,53]. While glycans with multiple such residues are poorly resolved/retained on reversed phase columns, the power of this method is to resolve isomers of neutral and mono-anionic glycans; this is especially useful when the target is to distinguish antennal or core modifications.

Normal phase resins are also frequently used for N-glycan analyses and their application for size fractionation is reminiscent of the gel filtration methods (e.g., Bio-Gel P4) common in the 1980’s and 1990’s for exoglycosidase-based glycan sequencing [54]. Interestingly, sulphate not only leads to earlier retention on reversed phase columns, but also on normal phase [13]. On the other hand, a ‘mixed’ hydrophilic interaction/anion exchange resin (HIAX; Dionex AS11) is an excellent means of separating by size as well as by increasing numbers of anionic residues, regardless of type [4, 12, 55].

For samples particularly complex in terms of numbers of glycans of different mass, 2D-HPLC (e.g., with normal phase as a first or second separation step) can also be used to reduce...
Historically, the use of FAB-MS was extremely useful in N-glycan analysis; in the 1990’s, this method was used to analyse oligosaccharides from nematodes carrying phosphorylcholine [36, 56] as well as glycosylphosphatidylinositol anchors from eukaryotes modified with phosphoethanolamine [57]. As the glycans were commonly derivatised by permethylation, subsequent extraction procedures can result in the loss of the polar ionically-modified glycans. Thus, the zwitterionic glycans were only detected either by permethylation after hydrofluoric acid (HF) treatment, which removes the phosphorylcholine moieties, or by peracetylation which is suitable for lower mass glycans: both approaches were employed on N-glycans from nematode parasites [36, 56]. Thus, in part, the positions of the phosphorylcholine modifications could only be inferred.

On the other hand, the presence of aminoethylphosphonate on locust N-glycans was fully compatible with standard FAB-MS [18], with the 6-substitution of mannose or GlcNAc being shown by displacement with a perdeuteromethyl group ($^{31}$P and $^1$H-NMR were also employed in that study). Positive and negative ion mode FAB-MS also revealed the presence of aminoethylphosphonate or its N-methylated derivative on mollusc glycolipids [43, 58]. As for studies on phosphorylcholine-modified glycoconjugates, HF also removed the phosphate to yield the underlying neutral structure. Seemingly sulphated glycans have been less studied using FAB-MS, but examples include repeating units of the dermatan sulphate of a marine ascidian [59] and of the chondroitin sulphate from squid [60] as well as detection of sulphated N-glycolylneuraminic acid in sea urchin eggs [61]. Furthermore, FAB-MS was one of a number of methods used to show the presence of methylphosphate on Dictyostelium N-glycans [62].

**Fast atom bombardment spectrometry**

Currently ESI-MS and MALDI-TOF MS are the most widely-used mass spectrometric methods for glycan analysis. ESI-MS is often considered ‘softer’ than MALDI-TOF MS [45] and is frequently used ‘on-line’ with liquid chromatography, but tends to produce multiply-charged ions [63]. It appears that there is only one study on the analysis of phosphorylcholine-modified N-glycans by LC-ESI-MS; on-line porous-grafted carbon separation of pyridylaminated oligosaccharides from Ascaris suum thereby revealed a set of eleven zwitterionic N-glycans which were all sensitive to hydrofluoric acid treatment; as a signature MS/MS fragment ion for nematode antennal phosphorylcholine modifications, $m/z$ 369 was detected which corresponds to PC$_1$HexNAc$_1$ [64]. ESI-MS was also used to analyse an unusual phosphoethanolamine-modified N-glycan originating from Campylobacter [65]. In terms of O-glycans, ESI-MS has revealed the presence of phosphorylcholine and phosphoethanolamine linked to insect O-glycans [12, 66] as well as of aminoethylphosphonate on a jellyfish O-glycan [44].

Examples of invertebrate phosphorylated and sulphated protein-linked glycans analysed, at least in part, by ESI-MS include unusual phosphoglycans from Trypanosoma cruzi and Leishmania [67–69], a phosphoester-linked disaccharide from Dictyostelium [70] and fragments of a sulphated polysaccharide from a starfish [71]. Whereas underivatised sulphated O-glycans from insects and an example sulphated N-glycan from an oyster were analysed by PGC-ESI-MS/MS, a modified post-permethylation clean-up procedure allowed detection of sulphated N-glycans in mosquito using nanospray mass spectrometry [12, 13]. NSI-MS following permethylation was also used to detect sialic acid on N-glycans from Drosophila and glucuronic acid on O-glycans from both Drosophila and mosquitoes [12, 21], while ESI-IT-MS was used to define methylated glucuronic acid substitutions of antennal fucose on N-glycans derived from the shell-forming fluid of a mussel [28]. In the case of another mollusc, aspects of the structure of the GlcA(MeHex)Fuc branches and the methylaminoethylphosphonate-modified N-glycans, as well as zwitterionic and anionic O-glycans, of a marine snail were revealed by PGC-ESI-MS/MS [14].

**Matrix-assisted laser-desorption/ionisation mass spectrometry**

MALDI-TOF MS is arguably the most robust, versatile and flexible method for glycomic analysis, although the ionisation is often considered relatively harsh; it tends to produce singly-charged ions in the mass range relevant to glycan analysis, which eases interpretation. It can either be performed on free glycans or ones in permethylated form or those derivatised.
with a fluorescent label at the reducing terminus [72]. While modified post-derivatisation strategies allow MALDI-TOF MS of sulphated glycans in their permethylated form [73], fluorescent labelling of glycans with, e.g., 2-aminoypyridine or 2-aminobenzamidine, facilitates HPLC methods which can then be used ‘off-line’ with MALDI-TOF MS. As mentioned above, in the case of 2-aminoypyridine, labelling also increases sensitivity in the mass spectrometer and yields intense Y-fragments of protonated quasimolecular ions, which aid identification of core modifications without affecting the ability to observe B-ions resulting from antennal losses. Over the years various matrices have been proposed for glycan analysis and currently we use 6-aza-2-thiothymine (ATT) in both positive and negative ion modes.

Considering the individual types of modifications, the zwitterionic character of phosphorylcholine ensures that glycans which carry this moiety are rather easily detected in the positive ion mode, with the m/z 328 or 369 ions (PC1Hex1 and PC1HexNAc1) being strong characteristic antennal MS/MS fragments as observed, respectively, for fungal and nematode N-glycans [17, 19]. Indeed, even with pyridylaminated glycans for which the core region is generally well observed in MS/MS, antennal PC1Hex(NAc)1 fragments dominate the masses for which the core region is generally well observed in the positive ion mode, with the m/z 328 or 369 ions (PC1Hex1 and PC1HexNAc1) being strong characteristic antennal MS/MS fragments as observed, respectively, for fungal and nematode N-glycans [17, 19]. Indeed, even with pyridylaminated glycans for which the core region is generally well observed in MS/MS, antennal PC1Hex(NAc)1 fragments dominate the spectrum and so information regarding the core is only inferred by fragment ions corresponding to the loss of the core (e.g., loss of 299, 445 or 607 for GlcNAc1Fuc0,1Gal0,1-PA; see Fig. 3e and f and Refs. [19, 74]). In a marine snail, we have also found phosphorylcholine substituting galactose linked to core α1-6-fucose residues [14].

Although hydrofluoric acid treatment, which removes phosphorylcholine (amongst other phosphoethers as well as galactofuranose and some fucose linkages), tends to reduce the signal of low abundance glycans, in part due to the loss of the well-ionisable zwitterionic modification, it does then aid identification of core modifications without affecting the ability to observe B-ions resulting from antennal losses. Over the years various matrices have been proposed for glycan analysis and currently we use 6-aza-2-thiothymine (ATT) in both positive and negative ion modes.

Fig. 3 Comparative mass spectrometry of N-glycans modified by phosphoethers and sulphate. Example positive and negative ion mode MALDI-TOF MS and MS/MS (also after hydrofluoric acid treatment) of N-glycans from invertebrate and protist organisms. (a-c) Comparison of the positive (+) and negative (−) ion mode MALDI-TOF MS spectra before and after hydrofluoric acid (HF) treatment of isobaric N-glycans modified with either phosphate (P), sulphate (S) or methylaminomethylphosphate (MAEP or *) from two marine organisms together with a relevant MS/MS spectrum of either the m/z 1270 or 1272 [M-H]− or [M + H]+ quasimolecular ions. (d) Positive (+) and negative (−) ion mode MALDI-TOF MS spectra before and after hydrofluoric acid treatment of a Trichomonas vaginalis N-glycan (Tv2 strain) with the corresponding MS/MS spectrum of m/z 1274. (e, f) Positive ion mode MALDI-TOF MS and MS/MS of a nematode N-glycan modified with phosphorylcholine (PC) before and after hydrofluoric acid treatment. (g, h) Comparison of the positive (+) and negative (−) ion mode MALDI-TOF MS spectra before and after hydrofluoric acid (HF) treatment of isobaric N-glycans modified with methylphosphate (PMe) and phosphate or sulphate from Dictyostelium with relevant MS/MS spectra of either the m/z 1485 or 1487 quasimolecular ions. Further structural details were in each case proven by enzymatic digestion. The data in a are unpublished, whereas those in panels b–h underlie assignments in previous publications on Volvarina rubella (b and c; marine snail), Trichomonas vaginalis (d; protist), Caenorhabditis elegans (e and f; nematode) and Dictyostelium discoideum (g and h; slime mould) [4, 14, 16, 74].

Also, in our recent studies, we have revealed sulphation on a range of invertebrate N-glycans, relying on a number of approaches to show that the 80 Da modification of these oligosaccharides is indeed sulphate and not phosphate; generally, mass accuracies of many instruments (other than FT-MS) are insufficient to distinguish phosphate and sulphate (Δ = 0.01 Da), but there are a number of differences regarding their properties. First, phosphorylated glycans ‘fly’ well in both positive and negative ion modes, whereas the detection of sulphated glycans is often impeded by ‘in source’ loss of sulphate, an effect most obvious in the positive ion mode. This can, though, depend on the choice of matrix. Using 2,5-dihydrobenzoic acid (DHB), sodiated adducts of intact sulphated glycans are detected; with ATT, traces of these [M + Na]+ ions can be observed in positive mode, whereas very good MS and MS/MS spectra can be obtained for the intact glycan in negative mode [13]. Second, phosphate (also methylphosphate, but not sulphate) is sensitive to hydrofluoric acid treatment resulting in a shift of 80 or 94 Da (Fig. 3a, b, g and h), whereas methanolic HCl can be used to remove sulphate at least to some extent under conditions which do not result in loss of other residues. We have applied these methods to example slime mould, insect and mollusc N-glycans [4, 12, 14]. Although there are commercially-available sulphatases, we have yet to find a source which is compatible with a MALDI-TOF-based digestion assay; in contrast, standard phosphatases are ‘MALDI-compatible’. Both phosphate and sulphate (like...
methylphosphate 6-linked to mannose or sulphated oyster N-glycans [4, 12, 13, 53]. This overcame the problem that such glycans were early-eluting and not well-resolved when fractionated by RP-HPLC, particularly when they contained more than one anionic moiety. Interestingly, a phosphorylated HexHexNAc2PMeP glycan (for MS data, see Fig. 3h) eluted slightly earlier than the corresponding isobaric sulphated form on HIAX [4]. Furthermore, in the case of oyster plasma N-glycans, sulphation was observed to reduce retention time on both RP- and NP-HPLC; however, the sulphated glycans accounting for about half the structures, some also carrying the sulphate in the context of a blood group A modification, could be analysed by MS and MS/MS in negative ion mode with the monosulphated being detected as [M-H]- and disulphated as [M-2H + Na]+ ions [13]. A mix of NP-HPLC, MALDI-TOF MS and ESI-MS has been recently applied by others to demonstrate sulphation of N-glycans from a lepidopteran species [75].

In the case of glucuronic acid, its mass is the same as a methylated hexose (Δm/z 176) and so it is important to distinguish these units from each other. Fortunately, not only are glucuronylated N-glycans enriched in the acidic fraction eluted from graphitised carbon, they ionise well in both positive and negative ion modes of MALDI-TOF MS; fragmentation in positive ion mode is effective as shown for mosquito N-glycans [12] as well as for glucuronylated antennae from a marine snail [14]. While sialic acid modifications can be detected by MALDI-TOF MS, with the potential to stabilise these residues in a linkage-specific manner [76], we have not yet detected sialic acids in our own studies on N-glycans derived from insects, nematodes, molluscs or protists other than in obvious food or media components.

A key part of our strategy is that exoglycosidases or chemical treatments are used to gain extra information, in addition to the retention time and MS/MS data, regarding the structure of an N-glycan. Thereby, we have also partly extended the tools available by having our own in-house recombinant exoglycosidases, such as galactosidases and hexosaminidases [77, 78], in addition to those from commercial sources. However, the background contaminations in some enzyme preparations (e.g., β-glucuronidase or α-galactosidase) can make subsequent MS measurements difficult and so low-abundance structures can be refractory to further analyses (only with reasonable amounts can reinjection onto HPLC be performed). A further challenge is that steric hindrance, a lack of standards or the inability to remove certain modifications (e.g., methylation) can complicate the interpretation. Nevertheless, even pyridylaminated glycans of low abundance (e.g., 2 pmol in an HPLC fraction) can be structurally defined using our approach by MS/MS before and after enzymatic and chemical incubation without further purification.

Conclusion

Glycan analysis remains a challenge, especially for unknown glycans in low amount from novel sources; current glycoinformatic tools are very much mammalian-centered and ‘in silico’ annotations are not appropriate for the analysis of N-glycans from invertebrate or protist sources. Furthermore, analytical methods requiring a larger amount of material (NMR or GC-MS) cannot be performed on low abundance isomers and so one must rely on robust, sensitive procedures in which complex mixtures of glycans are avoided. This may, depending on the purity of the glycan sample, require one or two steps of solid-phase extraction followed by on-line or off-line fractionation in combination with mass spectrometry. Ideally, orthogonal methods (more than one type of mass spectrometry, chemical or enzymatic treatments) are used. These concepts have formed our own analytical workflows and the results of our recent studies highlight the extreme variability of N-glycan structures in invertebrates and protists; thereby, we observe that anionic or zwitterionic modifications are often significantly present. Thus, tailoring the methods to take this into account, one should always expect the unexpected when exploring the glycoiniverse.

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