Negative Ion Fast-Atom Bombardment Tandem Mass Spectrometry To Determine Sulfate and Linkage Position in Glycosaminoglycan-Derived Disaccharides

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Negative ion fast-atom bombardment tandem mass spectrometry has been used in the analysis of monosulfated disaccharides. These commercially obtained disaccharides have been enzymatically prepared from glycosaminoglycans using polysaccharide lyases. Three disaccharides from chondroitin sulfate and dermatan sulfate and two disaccharides from heparan sulfate and chemically derivatized heparin were analyzed. All five disaccharides were isomeric, with differences in sulfate position and linkage position. The full-scan mass spectra are useful in differentiating isomers when the sulfate group resides on different saccharide units. This structural information was obtained from fragment ions produced through cleavage at the glycosidic linkage. The full-scan mass spectra of each monosulfated disaccharide also produced intense molecular anions having long lifetimes. Collisional activation of these resulted in tandem mass spectra rich in significant product ions. Some of these fragment ions were formed through ring cleavage and were useful in the determination of both sulfate and linkage position. (J Am Soc Mass Spectrom 1992, 3, 797–803)

Glycosaminoglycans (GAGs) are sulfated, linear polysaccharides that are found in virtually all living tissues [1]. The GAGs differ in types of saccharide constituents, sulfation, and linkage configuration and position. GAGs can be separated into two major families, those with (1 → 4) linkages (i.e., heparin and heparan sulfate) and those with alternating (1 → 3, 1 → 4) linkages (i.e., chondroitin and dermatan sulfates). Theses GAGs are composed of alternating, glycosidically linked uronic acid and hexosamine residues. Although the uronic acid residue is only sulfated at C-2, the hexosamine residue can be sulfated at C-2, C-3, C-4, and C-6. Both of these families play an important role in cell–cell interaction [2], as well as in a wide variety of other biological functions [3]. Our laboratory is involved in developing new methods to study the structure of these complex polysaccharides [4–6]. We rely on polysaccharide lyases [7, 8] to break down these GAGs into smaller oligosaccharides for structure determination. The aim is to relate features in the fine structure of these GAGs, such as the position of sulfate groups, to specific biological activities [9].

Carbohydrate structure determination has increasingly come to rely on mass spectrometry primarily because of its high sensitivity [10]. Mass spectral approaches for the analysis of nonvolatile, highly sulfated oligosaccharides prepared from GAGs often require prior derivatization [11, 12]. One method to analyze underivatized sodium salts of sulfated oligosaccharides, prepared using polysaccharide lyases, involves the use of negative ion fast-atom bombardment mass spectrometry (FAB-MS). This results in both molecular and fragment anions, from the cleavage of glycosidic linkages, that afford sequence information [13, 14]. The fragmentation observed in the negative ion FAB-MS of underivatized samples, however, does not permit the assignment of sulfate position within a saccharide residue nor does it afford information on the linkage position. These are important structural features, since a complex mixture of GAGs is often found in a single biological sample.

Negative ion FAB tandem mass spectrometry (MS/MS) has recently been applied to oligosaccharide analysis and its fragmentation mechanisms and nomenclature discussed [15, 16]. FAB MS/MS of neutral oligosaccharides has shown utility in determining

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linkage position [17, 18] and studies on acidic monosaccharides have shown FAB MS/MS to be useful for determining sulfate position [19]. In this paper we report the negative ion FAB-MS and FAB MS/MS of a series of commercially obtained, GAG-derived disaccharides demonstrating the utility of this method to obtain ions regarding both linkage and sulfate position.

Experimental

Materials. UUA2S(1 → 4)-d-GlcNAc and UUA(1 → 4)-d-GlcNAc6S (where UUA is a 4-deoxy-α-L-threo-hex-4-eno-pyranosyluronic acid, GlcNAc is N-acetylgalactosamine, and S is sulfate) were purchased from Pharmacia Enzymes (Aberdeen, Scotland). UUA2S(1 → 3)-d-GalNAc, UUA(1 → 3)-d-GalNAc4S, and UUA(1 → 3)-d-GalNAc6S were also received from Seikagaku America (Rockville, MD). These disaccharides could also be prepared by treating GAG or chemically derivatized GAG with the appropriate polysaccharide lyase and purifying the desired disaccharide using high-performance liquid chromatography (HPLC) [8, 20]. Triethanolamine (TEA) and sodium dodecylsulfate (SDS) were from Sigma Chemical Co. (St. Louis, MO). The Bio-Gel P2 desalting column was from BioRad (Richmond, CA). Thioglycerol, deuterium oxide (99.99 atom %) and 3-(trimethylsilyl)propionic acid-2,2,3,3-2H4 sodium salt (TSP-2H4) were from Aldrich Chemical Co. (Milwaukee, WI). Sodium borate was from Fisher Chemical Co. (Fair Lawn, NJ) and borate acid was from Mallinckrodt (Paris, KY).

Characterization of the Disaccharide Standards by 1H-NMR Spectroscopy. Each disaccharide was dissolved in 2H2O and exchanged three times by freeze drying. The 1H-NMR spectra of each disaccharide (1.5 mg/0.5 mL) was obtained at 300 MHz using a Bruker WM 360 NMR spectrometer (Spectrospin AG, Switzerland) at 25 °C with TSP-2H4 as the internal standard. The spectra were assigned on the basis of reported literature values and were consistent with the structures of each disaccharide [14, 21]. The purity of each disaccharide was estimated by 1H-NMR based on the absence of signals corresponding to contaminants.

Characterization of the Disaccharide Standards by Capillary Zone Electrophoresis (CZE). CZE analysis was performed using a Dionex capillary electrophoresis system (Sunnyvale, CA) equipped with a variable wavelength ultraviolet detector set at 220 nm. The experiments used a fused silica (externally coated except where the tube passed through the detector) capillary tube (75 μm i.d., 375 μm o.d., 70 cm long) from Dionex. The capillary tube was washed extensively and activated with 0.1 M phosphoric acid, 0.5 M sodium hydroxide, and deionized water then filled with the operating buffer. The UUA2S(1 → 3)-d-GalNAc, UUA(1 → 3)-d-GalNAc4S, and UUA(1 → 3)-d-GalNAc6S were analyzed using 10 mM sodium borate and 50 mM boric acid at pH 8.8 [22] and 10 kV and the UUA2S(1 → 4)-d-GlcNAc and UUA(1 → 4)-d-GlcNAc6S were analyzed using 50 mM SDS, 10 mM sodium borate at pH 8.5 and 20 kV [21]. The samples were injected by gravity injection (1 nL containing 1 ng) and electrophoresis was performed at 25 °C using the above operating buffers.

Mass Spectrometry. All mass spectra were acquired using a VG ZAB-HF mass spectrometer (VG Analytical Ltd., Manchester, UK) equipped with an Ion Tech sapphire-field FAB gun and a standard VG source. The gun was operated at 8 keV with a 2-μA current. The samples were desalted at least twice using a Bio-Gel P2 column (35 cm × 2.5 cm) eluted with deionized water. The samples (approximately 1 μL of a 20 mg/mL solution) were dissolved in 2 μL TEA (or thioglycoler) and added to a standard VG stainless steel probe tip.

For FAB-MS and FAB MS/MS experiments, ions were produced by bombarding the sample with 8 keV xenon atoms. Precursor ions were selected using the magnetic sector at a mass resolution of 500 and were collisionally activated by introducing helium into the collision cell to reduce the beam of precursor ions by approximately 50%. Mass-analyzed ion kinetic energy (MIKE) spectra were obtained by scanning the electrostatic analyzer and averaging 8 scans with the use of the multichannel analysis software on the VG 11-250 data system. The detector dynode was set at 7 kV and positioned for optimal sensitivity across the entire energy range [23].

Results and Discussion

The two families of GAGs found in tissues that are most often associated with important biological activities are the heparin/heparan sulfate and the chondroitin/dermatan sulfates [1]. This study used commercially available disaccharide standards prepared from GAGs and chemically modified GAGs that had been depolymerized using various polysaccharide lyases [7], fractionated, and purified by HPLC [8, 20]. Each disaccharide was dissolved in deionized, distilled water to give a concentrated (20 mg/mL) stock solution. A portion of this stock solution was used directly for CZE and FAB-MS analysis, while a second portion was exchanged with 2H2O for 1H-NMR spectroscopy. NMR analysis at 360 MHz resulted in spectra consistent with structure for each of the five monosulfated disaccharides [14, 21]. The purity of each was > 90% by NMR. CZE analysis gave electropherograms that showed a single symmetrical peak for each of the disaccharides indicating a purity of > 95% [21, 22]. The purity of these disaccharides had also been previ-
ously established by strong anion exchange HPLC [24] and reversed phase ion pairing HPLC analysis [25].

Studies by our group have demonstrated that it is possible to analyze the underderivatized sodium salts of GAG-derived disaccharides by FAB-MS [13, 14] and to obtain both molecular anion and fragment ions resulting in both structure and sequence information. The analysis of thoroughly desalted samples using triethanolamine as the liquid FAB matrix gave more intense spectra and more fragmentation than other commonly used matrices, such as thioglycerol [13]. All five of the GAG-derived, monosulfated disaccharides gave negative ion FAB spectra with very intense molecular anion peaks at m/z 458 and 480 ([M – H]− and [M + Na – 2H]−) with a low intensity molecular anion at m/z 502 ([M + 2Na – 3H]−) (where M is the fully protonated molecule).

Two difficulties remain for this analytical approach: one is the presence of matrix ions that can confuse the interpretation of spectral data; the second is the low amount of fragmentation that occurs, making full structural assignment difficult. Ions arising from the triethanolamine matrix can be subtracted out, however, giving spectra that are easier to interpret (Figure 1). The ring-cleavage fragments, crucial for the assignment of structure, are generally not observed. Our previous studies [13, 14] identified several GAG-derived oligosaccharides that were difficult to distinguish when analyzed by FAB-MS. These problems are particularly apparent when two isomers are analyzed. Isomeric disaccharides can differ in both sulfate and linkage position yet show very similar FAB mass spectra. It is possible to distinguish between monosulfated disaccharides only when the sulfates are on different saccharide units (i.e., 2S from 4S and 6S, Figure 1). No significant differences are observed in the spectra of monosulfated disaccharides isomers having sulfates on the same saccharide unit (i.e., 4S from 6S). For example, while the spectra of ∆UA2S(1 → 3)-d-GalNAc and ∆UA2S(1 → 4)-d-GlcNAc contained an ion at m/z 277 corresponding to [M + Na – H – GalNAc]− or [M + Na – H – GlcNAc]−, ∆UA(1 → 3)-d-GalNAc4S, ∆UA(1 → 3)-d-GalNAc6S, and ∆UA(1 → 4)-d-GlcNAc6S contained ions at m/z 300 corresponding to [M-∆UA] (Figure 1). In addition, differences in linkage positions (i.e., 1 → 3 and 1 → 4) can not be distinguished from the full-scan FAB mass spectra of monosulfated disaccharides (Figure 1).

Because all the monosulfated disaccharides produce very intense [M – H]− and [M + Na – 2H]− ions having long lifetimes, their metastable [26] and collision-induced dissociation (CID) MIKE spectra were recorded. Collision gas was not required to obtain tandem mass spectra rich in structurally significant product ions; however, collisional activation resulted in some structurally significant ions that were not observed in the metastable MIKE spectra.

The CID-MIKE spectra of monosaccharidic sulfated glucosamines have been reported to contain fragment ions from which the sulfate position can be determined [19]. Thus, we obtained the CID-MIKE spectra of [M – H]− ions (m/z 458) of monosulfated disaccharides in an effort to distinguish sulfate position (Figure 2). Whenever possible, we use the nomenclature recently proposed by Domon and Costello [16] to describe the fragment ions drawn in each figure.

The 92X2 ion observed at m/z 342 and the [HSO4]− ion at m/z 97 are common to the [M – H]− CID-MIKE spectra for each isomeric disaccharide (Figure 2). Characteristic ions for both ∆UA2S(1 → 3)-d-GalNAc and ∆UA2S(1 → 4)-d-GlcNAc (Figure 2a and d) are the B1, m/z 237 and 157 ions corresponding to a cleavage of

![Figure 1. Full-scan negative-ion FAB spectra of (a) ∆UA(1 → 4)-d-GlcNAc6S, (b) ∆UA(1 → 3)-d-GalNAc4S, and (c) ∆UA2S(1 → 3)-d-GalNAc. The structures are presented in their polyanionic form (M - 2H). Matrix ions have been subtracted in (a) whereas the asterisks in (b) and (c) mark matrix ions and sodium ion clusters not removed by matrix subtraction.](image-url)
the glycosidic linkage. Although fragment ions corresponding to cleavage through the glycosidic linkage are also observed in the CID-MIKE spectra of ΔUA(1 → 3)-d-GalNAc4S, ΔUA(1 → 3)-d-GalNAc6S, and ΔUA(1 → 4)-d-GlcNAc6S disaccharides at m/z 175, 282, and 300 from [M – H]+, it is possible to differentiate the ΔUA(1 → 3)-d-GalNAc4S and ΔUA(1 → 3)-d-GalNAc6S isomers by the dramatic differences in ion intensities. For example, in the spectrum of ΔUA(1 → 3)-d-GalNAc4S (Figure 2c), glycosidic cleavage gives the most abundant Y₁ product ion at m/z 300 ([M – ΔUA]+). For ΔUA(1 → 3)-d-GalNAc6S, the fragmentation without retention of the glycosidic oxygen gives Z ion formation corresponding to m/z 282 ([M – ΔUA]−) as the most abundant process (Figure 2b). This is not surprising because the mechanism for the formation of the Z ion requires an oxygen ion at the 4 position to undergo epoxide formation prior to expulsion of the nonreducing end sugar and H transfer [27]. It should be noted that ΔUA(1 → 3)-d-GalNAc6S, ΔUA(1 → 3)-d-GalNAc4S, and ΔUA(1 → 4)-d-GlcNAc6S fragment to give C and Z ions at m/z 175 ([M – GalNAc6S or 4S]− or [M – GlcNAc6S]−) and m/z 282 ([M – ΔUA]−) that arise from a sul-
fated saccharide present at the disaccharide's reducing-end sugar (Figure 2b, c, and e). The absence of the same C and Z ions in the spectra of ΔUA2S(1 → 3)-d-GalNAc and ΔUA2S(1 → 4)-d-GlcNAc (Figure 2a and d) offers an additional feature to distinguish these monosulfated disaccharides.

A peak at m/z 309 that corresponds to the loss of 149 μ was observed in several of the CID-MIKE mass spectra (Figure 2a–e). In thioglycerol the m/z 309 peak is not observed, suggesting that this peak was due to the TEA (Mₚ = 149) liquid matrix.

The [M + Na – 2H]⁻ ion at m/z 480 gives product ions in the FAB CID-MIKE spectra that can also be used to deduce sulfate position (Figure 3). Positive ion FAB analysis of carbohydrates has shown that the alkali metal cationized species undergo fragmentations that are more structurally informative than the fragmentation of the protonated molecular ion [28]. Negative ions of carbohydrates containing sodium cations have not been as thoroughly studied as nonsodium ions, so that a well-defined nomenclature is not available. A nomenclature similar to that proposed by Domon and Costello [16] is used for these fragment ions arising from the m/z 480 molecular anion. Each

![Figure 3. The negative-ion CID-MIKE spectra of m/z 480, ([M + Na – 2H]⁻) (a) ΔUA2S(1 → 3)-d-GalNAc, (b) ΔUA(1 → 3)-d-GalNAc6S, (c) ΔUA(1 → 3)-d-GalNAc4S, (d) ΔUA2S(1 → 4)-d-GlcNAc, and (e) ΔUA(1 → 4)-d-GlcNAc6S.](image-url)
CID-MIKE spectrum arising from the [M + Na – 2H]− precursor ion at m/z 480 is unique. For example, ring cleavage results in fragment ions that can be used to distinguish the position of sulfate groups. In these spectra, the presence of an ion at m/z 153 is only observed for ∆UA25(1 → 3)-d-GalNAc6S (Figure 3c) while an ion at m/z 139 is characteristic of ∆UA25(1 → 3)-d-GalNAc6S and ∆UA25(1 → 4)-d-GlcNAc6S (Figure 3b and e). It is interesting to note that these two peaks, which contain the sulfate group and require ring cleavage, do not appear in the corresponding metastable spectra [26]. The FAB CID-MIKE spectra of the ∆UA25(1 → 3)-d-GalNAc and ∆UA25(1 → 4)-d-GlcNAc6S are simple, containing few fragment ions (Figure 3a and e). The ∆UA25(1 → 3)-d-GalNAc gave C1-type fragmentation with proton and sodium addition, resulting in an abundant product ion at m/z 277 ([M – GalNAc]−) (Figure 3a). The remaining product ions obtained from the [M + Na – 2H]− ion of other isomers are quite complex. Many contain sodium and appear to arise from subsequent fragmentation of Z ions. The ion at m/z 244 can be assigned to ring cleavage of the Z1 ion and loss of HOCH2CH=O and the ion at m/z 274 to loss of CH2O by cleavage of the C5–C6 bond of the hexosamine residue in the Z1 ion (Figure 3c).

When all the GAGs present within a tissue are depolymerized using a mixture of polysaccharide lyases, disaccharides containing both (1 → 3 and 1 → 4) glycosidic linkages are obtained. When these are isomeric they are often difficult to separate and analyze [21, 22]. Therefore a tandem mass spectral method capable of distinguishing between such isomeric disaccharides would be very useful. Tandem CID positive-mode FAB-MS has been useful in distinguishing linkage position in neutral oligosaccharides (see ref; 28 and references therein).

It is not surprising, therefore, that similar results for sulfated sugars in the negative-ion mode are observed. The 0.2A2 product of deprotonated ions is characteristic of the (1 → 4) linkage [18]. This is true of the CID-MIKE spectra of either the [M – H]− or the [M + Na – 2H]− ion of monosulfated disaccharides (Figures 2 and 3). The 0.2A2 product ion indicative of a (1 → 4) linkage is at m/z 357 in the [M – H]− spectra and at m/z 379 in the [M + Na – 2H]− spectra (Figures 2 and 3, d and e). These ions at m/z 357 and 379 are not found in the spectra of any of the (1 → 3) monosulfated disaccharides.

Conclusions

In conclusion, FAB CID-MIKE analysis can be used to distinguish both sulfate position and linkage position in GAG-derived monosulfated disaccharides. There are several areas that require future attention. First, the mechanism of sodium interaction with these oligosaccharides as they undergo fragmentation requires additional study. Second, the effect of sulfation on fragmentation also requires further investigation. To answer this question the isomeric disulfated disaccharides of the structure ∆UA25(1 → 4)-d-GlcNAc6S, ∆UA25(1 → 3)-d-GalNAc6S, and ∆UA25(1 → 3)-d-GalNAc4S are currently under investigation in our laboratory using FAB CID-MIKE scanning. Third, the use of chromatographic [29] and electrophoretic [30] methods combined with tandem FAB-MS, such as that successfully applied to nucleosides and peptides, needs to be explored for acidic carbohydrates. Such approaches are particularly useful when only minor differences are observed in the spectra of isomers. Finally, application of this technique to larger oligosaccharides needs to be examined. We have reported the use of FAB-MS to determine structures as large as a dodecasulfated octasaccharide [14]. The low-intensity molecular anion of this octasaccharide will probably require larger amounts of this highly purified sample for analysis. The application of MS/MS to oligosaccharides, such as this octasaccharide, might be very important in helping to determine the sequence of the GAGs.

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References


See also: 802 LAMB ET AL.