Assays for determining heparan sulfate and heparin

$O$-sulfotransferase activity and specificity

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Abstract $O$-sulfotransferases (OSTs) are critical enzymes in the cellular biosynthesis of the biologically and pharmacologically important heparan sulfate and heparin. Recently, these enzymes have been cloned and expressed in bacteria for application in the chemoenzymatic synthesis of glycosaminoglycan-based drugs. OST activity assays have largely relied on the use of radioisotopic methods using $[35\text{S}]3'$-phosphoadenosine-5'-phosphosulfate and scintillation counting. Herein, we examine alternative assays that are more compatible with a biomanufacturing environment. A high throughput microtiter-based approach is reported that relies on a coupled bienzymic colorimetric assay for heparan sulfate and heparin OSTs acting on polysaccharide substrates using arylsulfotransferase-IV and $p$-nitrophenylsulfate as a sacrificial sulfogroup donor. A second liquid chromatography-mass spectrometric assay, for heparan sulfate and heparin OSTs acting on structurally defined oligosaccharide substrates, is also reported that provides additional information on the number and positions of the transferred sulfo groups within the product. Together, these assays allow quantitative and mechanistic information to be obtained on OSTs that act on heparan sulfate and heparin precursors.

Keywords Enzymes · Mass spectrometry · Bioassays · Sulfotransferases · Coupled assay · Heparin · Heparan sulfate

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Introduction

Glycosaminoglycans (GAGs) are linear polysaccharides that are found ubiquitously in intracellular and extracellular domains [1]. GAGs are divided among four unique families: heparan sulfate, chondroitin sulfate, keratan sulfate, and hyaluronan. These families are differentiated by composition, structure, and the number of sulfo (S) groups that they carry [2]. These differences lead to highly specific roles in human biology, including but not limited to GAG-protein binding [3, 4], structural GAGs [5, 6], cell adhesion and cell-cell interactions [7], and morphogenesis [8].

Heparin is the most highly sulfated member of the heparan sulfate family of GAGs and is a GAG of substantial biological interest, primarily because of its binding to many proteins. One of the most important biological functions of heparin is its capacity to bind to the serine protease inhibitor, antithrombin III (AT) [9] and inhibit the coagulation pathway. In the coagulation cascade, prothrombin is enzymatically cleaved by Factor Xa to form thrombin, which in turn, acts on fibrinogen to form a fibrin clot. AT inhibits this pathway by binding to Factor Xa, preventing the formation of thrombin and ultimately the formation of fibrin. The heparin-AT complex binds to Factor Xa with very high affinity (10^7 M⁻¹ s⁻¹) making this complex a potent inhibitor of the coagulation pathway [10]. Heparin also forms a ternary complex with ATIII and thrombin, inhibiting thrombin activity and preventing blood coagulation [11].

The in vivo biosynthesis of heparin requires enzymatic catalysis resulting in: (1) the formation of the heparosan backbone on the serine-linked tetrasaccharide of the serglycin core protein through the action of polysaccharide synthases; (2) N-deacylation-, N-sulfotransferase (NDST)-1-, and NDST-2-catalyzed N-deacylation and N-sulfonation of the glucosamine (GlcN) residues; (3) the formation of iduronic acid (IdoA) residues from glucuronic acid (GlcA) residues by C₅-epimerase (C₅-epi); (4) the 2-O-sulfonation of IdoA residues by 2-O-sulfotransferase (2-OST); and (5) the 6-O-sulfonation by 6-O-sulfotransferases (6-OSTs) and 3-O-sulfonation by 3-O-sulfotransferase-1 (3-OST-1) of GlcN residues. Commercial heparins are derived from a number of different animal tissues and can vary with regard to sulfonation extent and molecular weight [12].

There is a major concern about the conditions used to isolate and purify many commercial heparins. Heparin production involves steps that occur at the slaughterhouse that are not performed under current good manufacturing practices (cGMP). These steps greatly increase the risk of the introduction of impurities, contamination or adulteration, a clear hazard to the recipients of these heparin-containing products [13]. Such manufacturing pitfalls increase the demand for a safer heparin product produced entirely under cGMP protocols. Our laboratory is investigating a fermentative-chemoenzymatic approach for the preparation of bioengineered heparins.

Under a bioengineering approach to the production of heparin, the heparosan backbone, a repeating →1)-β-d-GlcA(1→4)-β-d-GlcNAc(1→ disaccharide, where GlcA is GlcA and GlcNAc is N-acetylglucosamine, is produced by fermentation of Escherichia coli K5 on glucose and ammonium chloride [14]. Heparosan is then converted to an N-sulfonoheparosan, with a major repeating →1)-β-d-GlcA(1→4)-β-d-GlcNS(1→ (where S is sulfo) disaccharide structure, by enzymatic [15] or by chemical means [16]. Enzymatic conditions rely on treating heparosan with NDST enzymes, which N-deacylates the GlcNAc residue and N-sulfonates the resulting amino group. Chemical conditions begin with harsh basic conditions for N-deacylation of heparosan followed by chemical N-sulfonation using sulfur trioxide trimethylamine complex. The remaining enzymatic steps are believed to be identical to those in vivo: C₅-epimerization of the N-sulfonoheparosan converting most of the β-d-GlcA residues to α-L-iduronic acid (α-L-IdoA) residues, followed by the 2-O-sulfonation of the IdoA residues to IdoA2S, the 6-O-sulfonation of GlcNS to GlcNS6S residues, and finally 3-O-sulfonation of a small percentage of the GlcNS6S and GlcNAc to GlcNS3S6S and GlcNAc3S residues, respectively (Fig. 1).

The primary differences between the in vivo biosynthesis and the plant scale bioengineering of heparin are in the elongation of the heparosan backbone and the N-sulfonation of the...
backbone. In vivo, the heparosan backbone is elongated by KfiA, KfiB, KfiC, and KfiD enzymes in E. coli [17] and in animals by Ext enzymes, in which case the backbone is subsequently N-deacetylated and N-sulfonated enzymatically by NDST enzymes [18]. In bioengineered heparin, the heparosan backbone is formed by E. coli K5 fermentation, then N-deacetylated with sodium hydroxide and N-sulfonated using sulfur trioxide methylamine. Finally, in both the biosynthesis and fermentative chemoenzymatic synthesis of heparin, O-sulfonation is accomplished using O-sulfotransferases (OSTs).

A major challenge to prepare a bioengineered heparin has been the development of assays to readily assess the activity of OSTs [19]. Commonly used radioisotopic assays pose problems for routine use in a quality control laboratory of a manufacturing facility [19]. The most accurate and reliable method of analyzing the bioengineered enzyme activity is nuclear magnetic resonance spectroscopy, which is time-consuming, requiring added steps for product isolation and purification prior to analysis, and requires sophisticated instrumentation, making it potentially very useful [20, 21], but challenging for the routine assessment of enzyme activity.

Sulfonation reactions are also being increasingly recognized as important in the metabolism of drugs, hormones, carcinogens, neurotransmitters, lipids, and peptides [22]. Essential to understanding these metabolic sulfotransferases is the development of robust enzyme activity assays. Sulfotransferase assays have been the subject of a recent review that reported the most commonly used assay to be based on the measurement by scintillation counting of radioactive sulfate incorporated from PAP$^{35}$S [19]. While these assays are widely used and very sensitive, they are also hazardous, expensive, do not provide information on the position and number of incorporated S groups, and have not been widely used to determine kinetic information.

The current study offers two advances in OST assay methodology development. The first is a real-time colorimetric method for the evaluation of heparin OST activity using a coupled enzyme system, and the ability to obtain quantitative information on OST substrate specificities. In this assay, a recycling enzyme, aryl-sulfotransferase IV (AST-IV), takes the 3′-adenosine 5′-phosphate (PAP) product of the OST reaction and recycles it into the 3′-adenosine 5′-phosphosulfate (PAPS) cofactor (Fig. 2) using an additional sulfate donor, p-nitrophenyl sulfate (PNPS). One of the products of this reaction, p-nitrophenol (PNP), is visible and can be measured spectrophotometrically over time.

As this coupled systems offer an indirect measurement, a second method, relying on hydrophilic interaction liquid chromatography (HILIC)-Fourier transform mass spectrometry (FTMS), was developed to assess heparin OST enzyme activity directly based on measuring the products formed using defined substrates of various chain lengths. This lower throughput method is accurate and does not require workup steps usually required by other mass spectrometric methods. The HILIC-FTMS method is also useful for obtaining mechanistic information about the OSTs and exact location and number of the S groups introduced.

**Experimental procedures**

**Materials**

Heparosan was produced through the large-scale fermentation of E. coli K5, a capsular K5 antigen corresponding to a
polysaccharide composed of GlcA and GlcNAc in a 1:1 molar ratio [23]. This product, with a molecular weight of nearly 70 kDa, was chemically de-N-acetylated using NaOH and chemically N-sulfonated using a sulfur trioxide trimethylamine complex (Sigma Aldrich, St. Louis, MO) [12]. The resultant N-sulfoheparosan typically has a molecular weight range of 10–16 kDa with 85 % N-sulfo and 15 % residual N-acetyl groups. PAP and PNPS were purchased from Sigma. PAPS was prepared according to previously published methods [24, 25].

The following enzymes, appropriate expression vectors, and bacterial expression cells were prepared as previously described [26]. These enzymes include rat AST-IV (EC 2.8.2.1) in pET15 expression vector and BL21 expression cells; human C5-epimerase (NCBI, NM_015554.1), mouse 6OST isoform 1 (NCBI, NM_015818.2), and mouse 6OST isoform 3 (NCBI, NM_015820.3) in pMAL-c2x expression vector and Rosetta-gami B expression cells; hamster 2-OST (GenBank no. D88811.1) in pMAL-c2x expression vector and Rosetta-gami B expression cells; mouse 3OST-1 (NCBI, NM_010474.2) in pET28 expression vector and BL21(DE3)RIL expression cells. AST-IV and 3OST-1 possess N-terminal (His)_6 tags. C5-epi, 2-OST, 6OST-1, and 6OST-3 possess N-terminal maltose binding protein tags. A recombinant 6OST-3 prepared in CHO cells was obtained from R&D Systems (Minneapolis, MN) and used where indicated for comparative purposes. Heparin lyases I, II, and III were cloned from the genomic DNA of Flavobacterium heparinum and the expression and purification of the recombinant heparin lyases was conducted in E. coli as previously described [27].

Enzyme preparation, purification, and quantification

Cells on agar plates were selected and grown for 16 h in 5 mL of lysogeny broth media in 14 mL BD Falcon tubes, supplemented with the appropriate antibiotics. The 5 mL culture was then transferred to a baffled 2.8-L Erlenmeyer flask containing 1-L of lysogeny broth media and the appropriate antibiotics. The cultures were incubated at 37 °C and shaken at approx. 180 RPM until the solution optical density at 600 nm reached 0.7–0.9 absorbance units. At this point, the flask was transferred to an incubator shaker at 22 °C and 180 RPM. After 30 min, the first inducer was added to the culture. If necessary, the second inducer was added after an additional 20 min. The flask was then shifted for 20 h.

After incubation was completed, the 1-L solution was centrifuged using a Sorvall centrifuge from Thermo Scientific (Rockland, IL) at 3,500×g for 30 min at 4 °C. The supernatant was discarded and the cell pellet was re-suspended in 20 mL of buffer, containing 25 mM Tris (pH 7.4) and 500 mM NaCl for C5-epimerase, 2-OST, and 6-OST isoforms 1 and 3, or containing 25 mM Tris (pH 7.4), 500 mM NaCl, and 30 mM imidazole for AST-IV and 3-OST isoform 1. The re-suspended solution was sonicated using a Misonix 3000 (Farmingdale, NY) sonicator at 30 W and 50 % cycle (15 s on and 15 s off) for a 3-min total on time. The sonicated solution was then centrifuged at 9400×g for 60 min at 4 °C. The supernatant was retained and the cell pellet was discarded.

The supernatant was passed through a Millipore (Billerica, MA) 0.45 μm sterile filter in preparation for affinity isolation and purification. A 20-mL gravity-flow column was washed three times with 20-mL distilled water, followed by the loading of 3 mL of Ni-NTA resin from GE Healthcare (Piscataway, NJ) or 5 mL of twice with 15 mL of washing buffer, containing 25 mM Tris (pH 7.4) and 500 mM NaCl for amylose, or containing 25 mM Tris (pH 7.4), 500 mM NaCl, and 30 mM imidazole for Ni-NTA. The solutions containing the free enzymes were then passed through the resin, followed by clear of unbound material from the resin with an additional 10 mL of washing buffer. Finally, the bound enzymes were released from the resins using elution buffer, containing...
25 mM Tris (pH 7.4), 500 mM NaCl, and 40 mM maltose for amylose, or containing 25 mM Tris (pH 7.4), 500 mM NaCl, and 300 mM imidazole for Ni-NTA.

Following elution, the enzymes were buffer-exchanged from Tris-buffer containing high concentrations of imidazole or maltose to standard phosphate buffered saline (pH 7.0) by centrifugation in Millipore (Ipswich, MA) Centrifugal Filter Units with a molecular weight cut-off of 3,000 Da. Enzyme solutions were centrifuged at 3,500 × g and re-suspended in phosphate-buffered saline. Solution protein concentrations were determined by the bicinchoninic acid assay, purchased from Thermo Scientific (Rockland, IL), against a bovine serum albumin standard. Protein purity was determined by visualizing by UV spectroscopy at 232 nm. The resultant protein solution was desalted on a BioGel P-2 column. The substrates were then desalted on a BioGel P-10 column and 300 mM imidazole for Ni-NTA.

Preparation of defined substrates for HILIC-MS method

A partial digestion of heparosan polysaccharide was conducted to ~40 % completion over 17 min at room temperature [28, 29]. Heparosan (23 mg at 1 mg/mL) was mixed with 0.28 units (1 mL) of purified heparin lyase III in a 50-mM sodium phosphate (pH 7.6). The total reaction volume was 528 mL. The digestion was stopped by heating at 100 °C for 10 min. The reaction was then repeated at a 600-mg scale of heparosan. The digested polysaccharide was resolved on a BioGel P-10 column, which was eluted with a buffer containing 0.2 M NaCl at a flow rate of 2 mL/h. The fractions were visualized by UV spectroscopy at 232 nm. The resultant oligosaccharides were desalted on a BioGel P-2 column. N-sulfoheparosan (250 mg), containing 100 % GlcNS, was also digested following the same reaction protocol, and the reaction products were also resolved on a BioGel P-10 column and desalted on a BioGel P-2 column. The substrates were then characterized to confirm oligosaccharide structure and purity. Analysis was performed by mass spectrometry and nuclear magnetic resonance spectroscopy. The resulting heparosan and N-sulfoheparosan oligosaccharide substrates ranged from degree of polymerization 2–10.

Preparation of [34S]-PAPS for HILIC-MS method

[34S]PAPS was synthesized enzymatically using a modified method from similar previously described syntheses of [35S]PAPS by substituting Na234SO4 for Na2SO4 (from ISOFLEX USA) [30, 31]. The reaction included 90 mM adenosine triphosphate, 100 mM MgCl2, 1 M LiCl, 0.8 mg/mL pyrophosphatase, 0.8 mg/mL Klyveromyces lactis-expressed aryl-sulfotransferase, 0.8 mg/mL adenosine-5′-phosphosulfate-5′-phosphokinase, and 50 mM Tris–HCl at pH 8.0. The reaction was incubated at 30 °C for 6 h. The [34S]PAPS product was analyzed using high-performance liquid chromatography (HPLC; polyamine II column, YMC America, Inc.) as follows: 100 % water for 10 min, followed with a linear gradient of 0–100 % of 1 M KH2PO4 for 30 min, followed by 100 % 1 M KH2PO4 for 15 min at a flow rate of 1 mL min−1 with detection at 254 nm. Purification of PAPS was achieved on a DEAE-Sepharose fast flow column (GE Healthcare; 1.5 × 60 cm). The diethylaminoethyl column was washed with water, and PAPS was eluted with a gradient of 0–500 mM NaCl at 5.0 mL min−1 for 200 min. Fractions containing PAPS as determined by HPLC on a polyamine II column were pooled and stored at −80 °C. The purity of the [34S]PAPS product was assessed by MS analysis and determined to be 90 %+ with less than 10 % PAP contamination (Fig. S1 in the Electronic supplementary material (ESM)).

Colorimetric method for measuring enzyme activity

OST activity assays were conducted, unless otherwise specified, in transparent, U-bottom, 96-well plates purchased from Greiner Bio-One (Monroe, NC). A typical reaction volume of each well was 250 μL with the following conditions: 125 μL PNPS (10 mM in phosphate-buffered saline, pH 7.0), 25 μL N-sulfoheparosan (1 mg/mL), 25 μL of an OST enzyme (400–800 μg/mL), 25 μL AST-IV (2–3 mg/mL), 25 μL PAPS or PAP (250–500 μM), and 25 μL C5-epi (250–500 μg/mL) or PBS. The concentrations of the substrates and the enzymes were then varied on a case-by-case basis. The 96-well plate was incubated at 37 °C in a temperature-controlled SpectraMax plate reader (Molecular Devices, Sunnyvale, CA). Kinetic plots of PNP formation were generated by sample measurements at 400 nm at even time intervals over at least 30 min. Using the extinction coefficient for PNP at pH 7.0 (at 400 nm, ε = 10,500 M−1 cm−1), output absorbance readings from the plate reader were converted to μM concentrations of PNP and the data were re-plotted in terms of [PNP] as a function of time. Initial enzyme velocities (v0) were calculated as the change in [PNP] over a linear segment of the plot. V1 values, corresponding to varied PAP, PAPS, or enzyme concentrations, were fit to Michaelis-Menten and Eadie-Hofstee functions to yield the kinetic parameters.

Reaction conditions for HILIC method

The activities of OSTs were assessed by incubating 5.0 μg of purified 2-OST with 2.5 μg of N-sulfoheparosan derived decasaccharide and 2 μg PAPS in 22.5 μL of 50 mM MES buffer pH 7.0. The reaction occurred over 24 h with aliquots
removed at 0, 1, 2, 3, 4, 7, and 24 h. Each aliquot was quenched by adding 50 μL of 100 % acetonitrile.

MS analysis

The products were analyzed by HILIC using a 2.0 × 50 mm Luna HILIC column (Phenomenex, Torrance, CA) coupled to an electrospray ionization LTQ-Orbitrap XL FTMS (Thermo Fisher Scientific, San-Jose, CA). The mobile phase A was 5 mM ammonium acetate prepared with HPLC-grade water. Mobile phase B was 5 mM ammonium acetate prepared in 98 % HPLC-grade acetonitrile with 2 % HPLC-grade water. An Agilent 1200 HPLC binary pump was used to deliver the gradient from 10 to 80 % A over 8 min at a flow rate of 250 μL/min after injecting the samples. The optimized MS parameters, used to prevent in-source fragmentation, included a spray voltage of 4.2 kV, a capillary voltage of −40 V, a tube lens voltage of −50 V, a capillary temperature of 275 °C, a sheath flow rate of 30, and an auxiliary gas flow rate of 6. External calibration of mass spectra routinely produced a mass accuracy of better than 3 ppm. All FT mass spectra were acquired at a resolution of 60,000 within a 400- to 2,000-Da mass range.

Results and discussion

Approach

Each OST has two substrates, the S donor PAPS and a phenol or a carbohydrate hydroxyl acceptor. We initially examined the activity of the recycling enzyme, AST-IV, which is critical for transferring the S group from PNPS to PAP to form the PAPS cofactor. Through the action of AST-IV, therefore, the bieznyzic system can be used to assess conversion of PNPS into the visibly detectable PNP. AST-coupled assays were first reported on NodST, an enzyme that catalyzes the formation of 6-O-sulfochitobiose [32–34]. These studies also provided an indirect measurement of NodST’s Michaelis–Menten kinetics. Assaying the heparin OSTs adds several additional levels of complexity. First, these enzymes act on various substrates, ranging from oligomers to polymers, which have multiple non-identical acceptor sites. For example, a polysaccharide substrate with a molecular weight of 10,000 might have ~20 GlcN residues, some substituted with N-acetyl groups and some with N-sulfo groups, and ~20 uronic acid residues, some consisting of GlcA and some IdoA. In addition, the reactivity of these residues might differ whether they are in the center of the polysaccharide chain, at the non-reducing end or at the reducing end. Finally, the sequence context might have a profound effect on whether or not an OST acts and to what extent.

Coupling AST-IV to OST catalysis was then tested with all three classes of site-specific glycan sulfotransferases: 6OST-1 and 6OST-3, 2OST, and 3OST-1. The 6OSTs are structurally the simplest heparin biosynthetic enzymes and are believed to act on unmodified chains of simple structures, i.e., heparosan or N-sulfoheparosan, or slightly more modified disaccharides, i.e., 2-O-sulfonated N-sulfoheparosan, with different reactivities. The 6OST activity assay was coupled to AST-IV to recycle PAPS, maintaining the PAP concentration low to prevent product inhibition [35]. 2OST was examined on N-sulfoheparosan, C5-epi-treated N-sulfoheparosan, and several defined oligosaccharide substrates. Finally, 3OST-1 was studied on a highly modified substrate, heparan sulfate. Each heparin OST assay was first performed on polysaccharide substrates using an AST-IV coupled colorimetric assay to measure activity. This was then followed by the determination of OST activity on defined oligosaccharide substrates followed by HILIC-FTMS analysis. It is important to note that while all of the E. coli-expressed recombinant OSTs used in this study show good specificity and are catalytic, a fairly high enzyme-to-substrate ratio is required. This may be the result of reduced activity due to their truncation (missing their trans-membrane domain), the presence of fusion proteins (with either (His)6- or maltose-binding protein-tags), or the lack of glycosylation (present in the natural OSTs).

AST-IV kinetics

Rapid OST kinetic analysis requires the coupled, bieznyzic recycling activity of AST-IV, which must not be rate limiting. Therefore, we first examined the kinetics of AST-IV using PNPS and PAP as substrates. At concentrations of PNPS and PAP of 5.0 and 0.25 mM, respectively, the rate of PNP release was linear up to 8.5 μg/mL AST-IV (Fig. S2A in the ESM). The enzyme followed conventional Michaelis–Menten kinetics with a $K_m$ for PAP of 25.3 μM with 5 mM PNPS (Fig. S2B in the ESM). This value is similar to that found in the literature [34, 36]. The $V_{max}$ for PAP as substrate was 165.5 nmol min$^{-1}$ mg$^{-1}$ enzyme at 5 mM PNPS, similar to literature values [37].

Assays for 6OST-1 and 6OST-3

Initial experiments revealed that 6OST-1 and 6OST-3 showed no measurable activity on heparosan using the coupled colorimetric assay, consistent with the literature [38], which is unsurprising considering the low percentage of N-acetylated and 6-sulfonated disaccharides present in heparin [16]. However, both enzymes were active on heparan sulfate and N-sulfoheparosan, with activity on the former at least 2-fold higher than that on the latter, indicating that the presence of the 2-O-sulfate and IdoA residues are favorable for 6OST activity (Fig. S3 in the ESM). This reactivity was confirmed
by $^1$H-nuclear magnetic resonance spectroscopy (Fig. S4A in the ESM). Furthermore, disaccharide analysis of the action of 6OST-3 on a partially 2-O-sulfated heparan sulfate revealed that the enzyme strongly favored the N-sulfo and 2-O-sulfo disaccharides over the unsulfated and N-sulfo disaccharides (Fig. S4B in the ESM).

We assessed the rate-limiting step in the bienzymic assay with AST-IV to obtain more quantitative kinetics information on the 6-OSTs. At PAPS and N-sulfoheparosan concentrations of 75 μM and 100 μg/mL, respectively, increasing the 6OST-3 activity up to 80 μg/mL, the highest 6OST-3 concentration tested, resulted in a linear increase in reaction rate (Fig. 3). Hence, under conditions used in this work, AST-IV was not rate limiting. The activity of 6OST-3 on N-sulfoheparosan could be increased by pre-incubating the enzyme with C5-epi (Table 1; Fig. 4), suggesting a preference for the action of this isoform at GlcNS residues in close proximity to IdoA residues.

For polysaccharide substrate, the enhanced reactivity of 6OST-3 due to the C5-epi was not evident (Fig. 5a). Specifically, in the presence of 80 μg/mL 6-OST 3, the $K_M$ and $V_{max}$ values for N-sulfoheparosan were 13.8 μg/mL and 51 nmol/(min mg enzyme), respectively. On the C5-epimerized N-sulfoheparosan as polysaccharide substrate, the $K_M$ and $V_{max}$ values were 14.1 μg/mL and 61.0 nmol/(min mg enzyme), respectively. While the influence of the C5-epi was marginal on the kinetics of 6OST-3 on the polysaccharide substrate, the kinetics with the PAPS substrate was strongly influenced by the C5-epi (Fig. 5b). Specifically, the $K_M$ of PAPS was approximately 4-fold lower in the presence of the C5-epi (8.3 vs.

Table 1: Initial reaction rates for $O$-sulfotransferases on $N$-sulfoheparosan

<table>
<thead>
<tr>
<th>Reaction conditions</th>
<th>Velocity (nmol/min mg OST)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2OST</td>
<td>0.00±0.85</td>
</tr>
<tr>
<td>2OST and C5-Epi</td>
<td>1.63±0.80</td>
</tr>
<tr>
<td>6OST3</td>
<td>45.1±1.04</td>
</tr>
<tr>
<td>6OST3 and C5-Epi</td>
<td>86.4±0.89</td>
</tr>
</tbody>
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33 μM, respectively), while the \( V_{\text{max}} \) values for PAPS were similar at 73.5 or 62.5 nmol \( \text{min}^{-1} \text{mg}^{-1} \) enzyme with or without the C5-epi, respectively.

To understand further the influence of C5-epimerization on 6OST-3 activity, two-substrate Michaelis–Menten kinetic analysis was performed by varying PAPS (5–40 μM) and N-sulfoheparosan (12–100 μg/mL) concentrations, with and without 3 h pre-incubation of 6OST-3 with C5-epimerase. Under conditions where N-sulfoheparosan was preincubated with C5-epi, the primary Lineweaver–Burk plots suggested ternary complex kinetics, represented by coincidental intersection (Fig. 6a, b). The secondary replots against the intercept values (Fig. 6c, d; Table 2) afforded \( K_{\text{M, PAPS}} = 9.8 \) μM, \( K_{\text{M, NSH}} = 37 \) μg/mL, \( k_{\text{cat}} = 0.2 \) s\(^{-1}\), and a \( k_{\text{cat}}/K_{\text{M, PAPS}} = 2.04 \times 10^{4} \) s\(^{-1} \text{M}^{-1}\). Under conditions without C5-epi present, ternary complex mechanism kinetics was also observed (Fig. 7a, b). Secondary replots against the intercept values (Fig. 7c, d; Table 2) afforded \( K_{\text{M, PAPS}} = 39.2 \) μM, \( K_{\text{M, NSH}} = 41 \) μg/mL, \( k_{\text{cat}} = 0.04 \) s\(^{-1}\), and a \( k_{\text{cat}}/K_{\text{M, PAPS}} = 1.02 \times 10^{3} \) s\(^{-1} \text{M}^{-1}\).

Based on the intersection point of the primary plots (Figs. 6a, b and 7a, b), apparent dissociation constants for the PAPS and NSH substrates with 6OST-3 could be calculated. For PAPS, the apparent dissociation constant (\( K_{S, \text{PAPS}} \)) is nearly identical between both epimerized and non-epimerized N-sulfoheparosan (~15–20 μM), whereas the apparent dissociation constant for epimerized and non-epimerized N-sulfoheparosan are distinct (\( K_{S, \text{EPINSH}} \approx 66 \) μg/mL, \( K_{S, \text{NSH}} \approx 3 \) μg/mL). This suggests that 6OST-3 follows a kinetic mechanism wherein an ordered ternary complex is formed, with PAPS first binding to the enzyme followed by the binding of the N-sulfoheparosan. Following the transfer of the S group, the 6-O-sulfonated N-sulfoheparosan is released followed by the PAP product. To verify this order, the binding of 6OST-3 to N-sulfoheparosan was tested in the absence of PAPS using surface plasmon resonance. As shown in Fig. S5 (ESM), 6OST-3 was incapable of binding to N-sulfoheparosan in the absence of PAPS, which confirms the ordered ternary complex mechanism.

Structurally defined N-sulfoheparosan oligosaccharide substrates were employed to investigate further the reactivity and substrate specificity of 6OST-3. These experiments initially used the commercial recombinant mouse enzyme

![Table 2](image)
expressed in Chinese Hamster Ovary (CHO) cells (R&D Systems). The sequence comparison for the commercial 6OST-3 differed slightly from the E. coli-expressed enzyme prepared in our laboratory (Fig. S6 in the ESM). In addition, this enzyme was glycosylated by the CHO cells in which it had been prepared. The CHO cell expressed 6OST-3 showed greater specific activity but identical specificity as the E. coli-expressed 6OST-3 on the substrates tested. In these studies, \[^{34}S\]PAPS and the stable sulfur isotope (Fig. S1 in the ESM), replaced previously used \[^{32}S\]PAPS to improve the sensitivity of the method as the product masses would increase by 2.0159 amu for each S group incorporated [39].

Activity on each individual substrate was assessed by incubating the commercial 6OST-3, or lab prepared 6OST-3, with 50 μM of heparan hexasaccharide, N-sulfoheparosan tetrasaccharide, N-sulfoheparosan hexasaccharide, N-sulfoheparosan octasaccharide, or N-sulfoheparosan decasaccharide, and 120 μM \[^{34}S\]PAPS in 50 mM MES buffer at pH 7.0 for a total reaction volume of 100 μL. Over 24 h, 6-μL aliquots were removed at specific time points. Each aliquot was added to 15 μL of 100 % acetonitrile to quench the reaction. The samples were then analyzed, and the sum of the sulfonated products was plotted. The sulfonation of an N-sulfoheparosan tetrasaccharide over time is presented in Fig. 8. The addition of one S or two S groups was monitored, and the initial rates were converted into molar activity.

A summary of the activities based on varying chain lengths of substrate is presented in Table 3. No activity was observed on the heparan hexasaccharide, consistent with the aforementioned lack of 6OST activity on heparan. The N-sulfo tetrasaccharide was a poor substrate, while the N-sulfo hexasaccharide, N-sulfo octasaccharide, and N-sulfo decasaccharide showed similar activities suggesting a minimum chain length requirement. The crystal structure of 6OST-3 has not been determined, and hence, the binding sites are unknown. However, these data might help to clarify its substrate specificity.

Assays for 2OST

As with 6OSTs, we determined 2OST to be rate limiting in the coupled reaction with AST-IV (data not shown). 2OST showed no measurable activity on either heparan or N-sulfoheparosan. However, addition of C5-epimerase, resulted in...
in the introduction of IdoA residues into N-sulfoheparosan, and formation of measurable activity of 2OST (Fig. S3 in the ESM), as confirmed using both disaccharide compositional analysis (Fig. S7 in the ESM) and nuclear magnetic resonance spectroscopy (Fig. S4A in the ESM). Treatment of N-sulfoheparosan with C_{5}-epimerase and 2OST afforded new peaks in the $^1$H-nuclear magnetic resonance spectroscopy at 3.9, 4.2, 4.3, and 5.2 ppm, corresponding to signals associated with IdoA2S \[12\]. These results are consistent with the reported preference of 2OST for IdoA-containing substrates \[40\].

We proceeded to evaluate the activity of 2OST on an epimerized N-sulfoheparosan substrate. The concentrations of the GAG and OST were increased above those used in the 6OST assay to generate a robust response. Under varied PAPS concentrations maintaining a constant epimerized N-sulfoheparosan concentration of 0.5 mg/mL and a 2-OST concentration of 0.25 mg/mL, the $V_{\text{max}}$ and $K_{M}$ values were calculated to be 35.1 μM and 288 pmol/min mg enzyme, respectively (Fig. 9). These values were similar to those reported in literature \[41\] for 2OST activity on completely de-O-sulfated heparin, a substrate that lack all O-sulfonations

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Specific activity (nmol product/ (min mg protein))</th>
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<tbody>
<tr>
<td>NS tetrasaccharide</td>
<td>0.73</td>
</tr>
<tr>
<td>NS hexasaccharide</td>
<td>11.82</td>
</tr>
<tr>
<td>NS hexasaccharide (Lab Prep. 6OST-3)</td>
<td>10.94</td>
</tr>
<tr>
<td>NS octasaccharide</td>
<td>19.07</td>
</tr>
<tr>
<td>NS decasaccharide</td>
<td>14.01</td>
</tr>
<tr>
<td>Heparosan hexasaccharide</td>
<td>No reaction (&lt;0.01)</td>
</tr>
</tbody>
</table>

Fig. 10 a Time course of 2-O-sulfonation of N-sulfo decasaccharide. The increasing amounts of mono-, di-, and tri-sulfonated products were calculated from the HILIC-MS data. b Time course of N-sulfoheparosan (squares) 2-O-sulfonation from the addition of one sulfo (circles), two sulfo (squares), or three sulfo groups (triangles), total product formed (diamonds), as determined by HILIC-MS

Fig. 9 Michaelis–Menten kinetics (Eadie–Hofstee plot as Inset) of 2OST on completely de-O-sulfonated heparin

Fig. 11 Michaelis–Menten kinetics (Eadie–Hofstee plot as inset) of 3OST-1 on heparan sulfate
but possesses the IdoA residues and N-sulfonations critical for 2OST activity.

Finally, we examined 2OST activity on the structurally defined N-sulfoprocanosanic decasaccharide substrate to confirm 2OST activity on GlcA residues. The N-sulfoprocanosanic decasaccharide substrate showed a peak in its mass spectra at m/z 694.08 ([M–3H]3−, z=−3; MW, 2,085.2886) that disappeared over time with the appearance of a mono-2-O-sulf–N-sulfoprocanosanic decasaccharide product peak at m/z 720.739 ([M–3H]3−, z=−3; MW, 2,165.2454) after 1 h, di-2-O-sulf–N-sulfoprocanosanic decasaccharide product peak at m/z 747.397 ([M–3H]3−, z=−3, MW 22.45.2022) after 1 h, and tri-2-O-sulf–N-sulfoprocanosanic decasaccharide product peak at m/z 774.049 ([M–3H]3−, z=−3; MW, 2,325.159) after 7 h (Fig. 10a). The peak area of each sulfonated product was determined and compared with the peak area of the known concentration of substrate to calculate the concentration of each product as a function of time (Fig. 10b), assuming similar ionization efficiencies. The sum of the sulfonated products was also plotted and its linear region (up to 5 min) was used to estimate an enzyme activity of 0.33 pmol product formed min−1, with a specific activity of 131 pmol product formed min−1 mg−1 protein. It is important to note that loss of 0.1–10 % of S groups can take place through in-source fragmentation increasing with the level of S group substitution. Since liquid chromatography–mass spectrometry (LC-MS) analysis affords different retention times for oligosaccharides with differing levels of sulfation, in-source fragmentation can be clearly distinguished from an analyte-containing oligosaccharides having different sulfation levels.

Assays for 3OST-1

3OST-1 is responsible for installing the AT-pentasaccharide binding site into approximately one third of the heparin chains [11], and thus, recognizes a rare sequence within the precursory substrate structure [42]. 3-OST-1 acts on a polysaccharide containing the sequence: →4)α-d-GlcNY(1→4)α-/α-d-GlcA/α-/IdoA(1→4)α-d-GlcNY6X(1→4)α-d-GlcA(1→4)α-/d-GlcNS6X(1→4)α-/d-IdoA2S(1→4)α-/d-GlcN6S(1→4)α/β-d-GlcA/α-/IdoA2X(1→ (where Y=Ac or S and X=H or S, bold residues correspond to the AT-pentasaccharide-binding site and bold/italicized residue is the one that will be sulfonated at the third position by 3-OST-1). No measurable activity was observed for 3OST-1 acting on heparan or N-sulfoprocanosanic using the coupled colorimetric assay. The more highly modified heparan sulfate polysaccharide, however, is known to act as a substrate for 3OST-1 converting a low AT affinity heparan sulfate into a high affinity chain [11, 42]. A 3OST-1 assay, using coupled AST-IV (Fig. 11), afforded a PAPS K_M for 3OST-1 of 13.5 μM, consistent with previously published literature using radioisotopic methods [43]. In addition, heparin lyase II treatment followed by LC-MS analysis of the resulting high AT affinity heparan sulfates confirmed the formation of a 3-O-sulfo group containing tetrasaccharide consistent with the generation of an AT pentasaccharide-binding site (Fig. S8 in the ESM).

Conclusions

We have outlined two assays for the real-time and near real-time analysis of heparan sulfate sulfotransferase enzyme activity. Under real-time conditions, a colorimetric methodology was used, coupling the OSTs with an AST-IV enzyme to produce a visibly measurable product. This method was also coupled with the C4-epi to gauge the importance of IdoA residues in OST bioactivity. Robust colorimetric assays were run with 6OST-3, to determine detailed kinetic parameters, and products were verified by LC-MS and nuclear magnetic resonance spectroscopy. Variations of PAPS concentration and epimerization of GlcA into IdoA residues had a significant effect on 6OST-3 and 2OST kinetics, but no obvious effect was seen under conditions of varied N-sulfoprocanosanic concentration. In the near real-time methodology, hydrophilic interaction chromatography was used with a defined substrate to measure the overall additions of S groups to the substrate.

References