Fucosylated Chondroitin Sulfate 9–18 Oligomers Exhibit Molecular Size-Independent Antithrombotic Activity while Circulating in the Blood

Lufeng Yan, Danli Wang, Yanlei Yu, Fuming Zhang, Xingqian Ye, Robert J. Linhardt,* and Shiguo Chen*

ABSTRACT: Fucosylated chondroitin sulfate (FCS) oligosaccharides extracted from sea cucumber and depolymerized exhibit potent anticoagulant activity. Knowledge of the antithrombotic activity of different size oligosaccharides and their fucose (Fuc) branch sulfation pattern should promote their development for clinical applications. We prepared highly purified FCS trisaccharide repeating units from hexasaccharide (6-mer) to octadecasaccharide (18-mer), including those with 2,4-disulfated and 3,4-disulfated Fuc branches. All 10 oligosaccharides were identified by their nuclear magnetic resonance structures and ESI-FTMS spectroscopy. In vitro anticoagulant activities and surface plasmon resonance binding tests indicated those of larger molecular sizes and 2,4-disulfated Fuc branches showed stronger anticoagulant effects with respect to anti-FXase activity, as well as stronger binding to FIXa among various clotting proteins. However, both types of FCS 9–18-mer exhibited molecular size-independent potent antithrombotic activity in vivo at the same dose. In addition, both types of the FCS 6-mer exhibited favorable antithrombotic activity in vivo, although they showed weak anticoagulant activity in vitro. Combining absorption and metabolism studies, we conclude that FCS 9–18 oligomers could remain in the circulation to interact with various clotting proteins to prevent thrombus formation, and appreciable quantities of these oligomers could be excreted through the kidneys. All FCS 9–18 oligomers also resulted in no bleeding, hypotension, or platelet aggregation risk during blood circulation. Thus, FCS 9–18 oligomers with 2,4-disulfated or 3,4-disulfated Fuc branches exhibit potent and safe antithrombotic activity needed for clinical applications.

With the general improvement of human living standards and the aging of the population, thrombotic disease has become a serious human health concern over the past few decades. Heparin and low-molecular weight heparins (LMWHs) are in widespread use as anticoagulants for the treatment of thrombotic diseases, especially during surgical situations. However, heparin and LMWHs always have the risk of serious bleeding during therapy and cannot be orally administered. Thus, researchers have continued to search for other effective and safer anticoagulant drugs.

Fucosylated chondroitin sulfate (FCS), isolated from sea cucumber, has outstanding in vitro anticoagulant and antithrombotic activities and might also achieve in vivo anticoagulant and antithrombotic effects on oral administration after appropriate modification. FCS oligosaccharides of a particular molecular size range can show potent in vivo anticoagulant and antithrombotic activities by selective inhibition of the intrinsic factor Xase (FXase), avoiding side effects such as factor XII (FXII) activation, platelet aggregation, and increased bleeding risk.

While the chemical synthesis of FCS oligosaccharides has been possible, it is much easier and more cost-effective to prepare FCS oligosaccharides derived from polysaccharides. The structural differences of FCS polysaccharides from various sea cucumbers mainly result from differences in the sulfation pattern of Fuc branches. Obtaining effective FCS oligosaccharides with a different sulfation pattern of Fuc branches could expand the clinical application of these new anticoagulant agents.

Many chemical or physical methods have been reported for preparing FCS oligosaccharides, such as acid-catalyzed hydrolysis, γ-Co irradiation, and free-radical depolymerization.
However, these methods will all require high-purity FCS polysaccharides as a starting material to avoid fragments of a second sea cucumber polysaccharide, fucoidan. FCS oligosaccharide preparation methods generally rely on anion-exchange chromatography for polysaccharide purification and gel permeation chromatography for oligosaccharide fractionation, but both methods are labor-intensive and costly and, thus, are inappropriate for large-scale preparation.

We recently established a method for the rapid preparation of purified FCS oligosaccharides from dry sea cucumber, Pearsonothuria graeffei. The resulting FCS oligosaccharides showed potent anticoagulant and antithrombotic activities by selectively inhibiting the intrinsic coagulation pathway while avoiding side effects such as bleeding and hypotension. The chemical process, N-deacetylation–deaminative cleavage, used by this method is the same as that currently relied on in the preparation of LMWHs from heparin. The molecular size of the resulting FCS oligosaccharides prepared by this method was optimized for 6–18 oligomers, resulting in easy production quality control. Prior to the clinical application of FCS 6–18 oligomers, the effects of the molecular size and sulfation pattern of the Fuc branches on anticoagulant and antithrombotic activity and side effects need to be resolved. In addition, because there are no mammalian enzymes that can degrade FCS oligomers because of their Fuc branches, the clearance of FCS oligomers after subcutaneous injection needs to be explored.

Figure 1. HPGPC profiles of (a) depolymerized FCS-Ib, (b) depolymerized FCS-Pg, (c) separated FCS-Ib oligomers, and (d) separated FCS-Pg oligomers, analyzed using a Superdex Peptide 10/300 GL column (10 mm × 300 mm) by a refractive index detector. Samples were size-fractionated using 0.2 M NaCl as the eluent at a flow rate of 0.4 mL/min.

RESULTS AND DISCUSSION

Efficient Separation of FCS Oligomers. Native FCS polysaccharides, FCS-Ib and FCS-Pg, were isolated and purified from Ib and Pg, respectively. Their structures, including mainly trisaccharide repeating units and unique sulfation on Fuc branches, were previously established. We next used a partial N-deacetylation–deaminative cleavage method to selectively depolymerize the glycosidic bond of β-GalNAc4S6S-β1,4-o-GlcA on the native FCS polysaccharides with no obvious sulfation or Fuc branch loss.

The HPGPC profiles of partially depolymerized FCS-Ib and partially depolymerized FCS-Pg are shown in panels a and b of Figure 1, respectively. This treatment afforded a mixture of homogeneous oligosaccharides with trisaccharide repeating units, which were identified as the 3-mer, 6-mer, 9-mer, 12-mer, 15-mer, and 18-mer. However, chromatographic peaks in depolymerized FCS-Pg, especially the 6-mer and 9-mer, were broad and showed some peak splitting, suggesting the partial vacancy of the Fuc branch from original polysaccharides as observed in our previous study. Thus, efficient gel permeation chromatography column (1.0 cm × 120 cm) packed with Superdex 30 prep grade gel filtration resin was used to separate the mixture of depolymerized FCS. Combined with analysis using a Superdex Peptide 10/300 GL column, highly purified FCS oligomers, ranging from the 6-mer to 18-mer, were obtained from both sea cucumbers (panels c and d of Figure 1, respectively). The entire preparation process for FCS oligomers is summarized in Scheme 1.

Structural Characterization of FCS Oligomers. We characterized the structure of these highly purified FCS oligomers based on the published native FCS structure. In the nuclear magnetic resonance (NMR) spectra of the FCS-Ib 6-mer (Figure 2a,c), the H-1 proton of each residue could be assigned to the peaks at 5.68, 5.56, 4.52, 4.53, 4.57, and 4.6.
3.69/3.81 ppm for middle Fuc, terminal Fuc, nonreducing terminal GlcA, reducing terminal GlcA, GalNAc, and 2,5-anhydro-α-D-talitol (anTal-ol), respectively. The C-1 signals at 97.2, 97.3, 104.2, 102.4, 100.6, and 61.4 ppm, respectively, were also similarly assigned. Compared with the unsulfated monosaccharide, the downfield shift of protons caused by sulfation indicated that Fuc is sulfated at the C-2 and C-4 positions, and GalNAc and anTal-ol are both sulfated at the C-4 and C-6 positions. In addition, ESI-FTMS analysis of the FCS-Ib 6-mer revealed an ion at m/z 411.74 for [M – 4H]+ and m/z 549.32 for [M – 3H]3−, confirming a molecular formula of C38H41O21N1S8 (Figure 2f). Thus, the structure of the FCS-Pg 6-mer was mainly established as ε-Fuc3,4diS-α1,3-D-GlcA-β1,3-α-GalNAc4,6diS-β1,4-[ε-Fuc3,4diS-α1,3]-D-GlcA-β1,3-α-anTal-ol4,6diS. Using the same method, we confirmed the structure of FCS-Ib 9–18 oligomers mainly as ε-Fuc2,4diS-α1,3-α-GlcA-β1,3-α-GalNAc4,6diS-β1,4-ε-Fuc2,4diS-α1,3-α-GlcA-β1,3-α-anTal-ol4,6diS and the structure of FCS-Pg 9–18 oligomers mainly as ε-Fuc3,4diS-...
Figure 2. continued
Figure 2. continued
α1,3-β-D-GlcA-β1,3-[α-GalNAc4,6dis-β1,4-[β-Fuc3,4dis-α1,3-]
β-D-GlcA-β1,3-]n-D-anTal-ol4,6dis (n = 2−5) (Figures S5−S8
and S10). All of the 1H and 13C chemical shifts of FCS
oligomers are listed in Tables S1 and S2, although it looks like
there is some contamination from high- and low-molecular
weight oligomers in fractions like the FCS-Ib 9-mer, FCS-
Ib 12-mer, and FCS-Ib 15-mer as reflected by uneven HPGPC
curves (Figure 1c). From the NMR and MS spectra, it could be
confirmed that all of these oligomer fractions are of high purity
with little surrounding contamination and responsible for
further structure−activity study.

FCS Oligomers Exhibit Effective Anticoagulant
Activities by Selectively Inhibiting FXase. The anticoagu-
lant activities of FCS oligomers were evaluated using the
activated partial thromboplastin time (APTT) and thrombin
time (TT) of plasma clotting assays (Table 1) to determine
their ability to inhibit blood clotting through the intrinsic or
common pathways of the coagulation cascade. Both FCS-Ib
oligomers and FCS-Pg oligomers showed intrinsic anticoagu-
lation activity as only prolonging APTT. Their molecular size
had a significant impact on this activity for both types of FCS
oligomers. The concentration of the FCS 18-mer required to
double the APTT was similar to that of LMWHs. With the
decrease in molecular size, the concentration required to
double the APTT was increased. The concentration of the FCS
9-mer required to double the APTT was ∼3-fold higher than
that of the FCS 18-mer. The FCS 6-mer showed weak intrinsic
anticoagulant activity as reflected by APTT. For the same
oligomer size, the FCS-Ib oligomer showed intrinsic
anticoagulant activity slightly higher than that of the FCS-Pg
oligomer as reflected by APTT. This result was consistent with
the previous study that showed that native FCS-Ib showed
stronger intrinsic anticoagulant activity than native FCS-Pg.
We next compared the effects of the FCS oligomers on coagulation factors with a cofactor, including anti-FIIa/AT, anti-FXa/AT, and anti-FXase to further investigate the mechanism of their action on the intrinsic clotting pathway. The results are shown in Figure 3 and Table 1. All of the activities of these oligomers were concentration-dependent,

Table 1. Anticoagulant Properties of FCS Oligomers and Their Effects on Coagulation Factors with a Cofactor

<table>
<thead>
<tr>
<th>compd</th>
<th>APTT&lt;sup&gt;a&lt;/sup&gt;</th>
<th>TT&lt;sup&gt;a&lt;/sup&gt;</th>
<th>anti-FIIa/AT</th>
<th>anti-FXa/AT</th>
<th>anti-FXase</th>
</tr>
</thead>
<tbody>
<tr>
<td>FCS-Ib 18-mer</td>
<td>46.5</td>
<td>54.7</td>
<td>&gt;128</td>
<td>&lt;14.6</td>
<td>324</td>
</tr>
<tr>
<td>FCS-Ib 15-mer</td>
<td>62.7</td>
<td>40.5</td>
<td>&gt;128</td>
<td>&lt;14.6</td>
<td>&gt;1000</td>
</tr>
<tr>
<td>FCS-Ib 12-mer</td>
<td>97.7</td>
<td>26.0</td>
<td>&gt;256</td>
<td>&lt;7.30</td>
<td>&gt;1000</td>
</tr>
<tr>
<td>FCS-Ib 9-mer</td>
<td>160</td>
<td>15.8</td>
<td>&gt;256</td>
<td>&lt;7.30</td>
<td>&lt;1000</td>
</tr>
<tr>
<td>FCS-Ib 6-mer</td>
<td>&gt;256</td>
<td>~7.00</td>
<td>&gt;512</td>
<td>&lt;3.65</td>
<td>&gt;1000</td>
</tr>
<tr>
<td>FCS-Pg 18-mer</td>
<td>61.4</td>
<td>41.4</td>
<td>&gt;128</td>
<td>&lt;14.6</td>
<td>480</td>
</tr>
<tr>
<td>FCS-Pg 15-mer</td>
<td>89.2</td>
<td>28.5</td>
<td>&gt;128</td>
<td>&lt;14.6</td>
<td>&gt;1000</td>
</tr>
<tr>
<td>FCS-Pg 12-mer</td>
<td>129</td>
<td>19.7</td>
<td>&gt;256</td>
<td>&lt;7.30</td>
<td>&gt;1000</td>
</tr>
<tr>
<td>FCS-Pg 9-mer</td>
<td>190</td>
<td>13.4</td>
<td>&gt;256</td>
<td>&lt;7.30</td>
<td>&gt;1000</td>
</tr>
<tr>
<td>FCS-Pg 6-mer</td>
<td>&gt;256</td>
<td>~7.00</td>
<td>&gt;512</td>
<td>&lt;3.65</td>
<td>&gt;1000</td>
</tr>
<tr>
<td>UFH</td>
<td>12.0</td>
<td>212</td>
<td>8.81</td>
<td>212</td>
<td>0.50</td>
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<tr>
<td>LMWH</td>
<td>36.5</td>
<td>69.6</td>
<td>33.1</td>
<td>56.4</td>
<td>1.53</td>
</tr>
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</table>

<sup>a</sup>The APTT and TT of FCS-Ib oligomers were cited from our previous publication. <sup>b</sup>Concentration required to double the APTT/TT of human plasma (APTT/TT doubling). <sup>c</sup>The activity was expressed as international units per milligram (IU/mg) using a parallel standard curve based on the international heparin standard (212 IU/mg). <sup>d</sup>IC<sub>50</sub> value, the concentration required to inhibit 50% of protease activity.

Figure 3. Effects of FCS-Ib oligomers on (a) FIIa and (c) FXa activities in the presence of antithrombin (AT) and (e) intrinsic FXase activity. Effects of FCS-Pg oligomers on (b) FIIa and (d) FXa activities in the presence of AT and (f) intrinsic FXase activity. The results are expressed as means ± the standard deviation (n = 3). See Table 1 for IC<sub>50</sub> values.
First, we investigated the interactions of coagulation factors or cofactors measured by biolayer interferometry (BLI). The binding affinity of UFH with FIIa ($K_D = 1.84 \times 10^{-7}$ M) and UFH with FIIa ($K_D = 1.62 \times 10^{-8}$ M) were very similar. The binding affinities of FCS-Ib with FIXa ($K_D = 1.14 \times 10^{-7}$ M), FCS-Pg with FIXa ($K_D = 8.22 \times 10^{-8}$ M), and UFH with FIXa ($K_D = 1.84 \times 10^{-7}$ M) were also very similar. However, the binding affinity of UFH for AT ($K_D = 5.39 \times 10^{-8}$ M) was ~5-fold stronger than the binding affinity of FCS-Ib for AT ($K_D = 2.76 \times 10^{-7}$ M) and ~20-fold stronger than the binding affinity of FCS-Pg for AT ($K_D = 1.13 \times 10^{-8}$ M). All of these results were consistent with previous reports on binding affinities of a depolymerized FCS ($M_w \sim 4 \text{ kDa}$) with coagulation factors or cofactors measured by biolayer interferometry (BLI). In addition, these results suggested that the anticoagulant activities of native FCSs and FCS oligosaccharides were mainly mediated through direct binding to various coagulation factors.

The binding affinity of the FCS oligomer with the coagulation factor or AT was next examined by competitive binding experiments. For example, FIIa (100 nM) was preincubated with gradient concentrations of FCS-Ib and FCS-Pg prior to interacting with immobilized FCS-Ib to investigate the binding affinity of the FCS-Ib 18-mer with FIIa. If the FCS-Ib 18-mer interacted with FIIa, the amount of FIIa binding to immobilized FCS-Ib would be reduced, resulting in a weaker resonance unit (RU) signal. This method indirectly measures the binding affinity of the FCS-Ib 18-mer for FIIa and has been successfully used in studying the interactions between various proteins and polysaccharides. As shown in Figure S12a, with an increase in the concentrations of the FCS-Ib 18-mer in solution, the level of binding between FIIa and immobilized FCS-Ib was gradually reduced. The maximal binding affinity of FIIa with immobilized FCS-Ib was ~1015 RU when there was no FCS-Ib 18-mer in the solution. The maximal binding affinity of FIIa for immobilized FCS-Ib was reduced to ~132 RU when there was 25 μM FCS-Ib 18-mer in the solution, indicating that the free percentage of FIIa was only 13%.

**Table 2. Binding Affinities of the FCS Oligomer with a Coagulation Factor or Cofactor by Competitive Binding Experiment**

<table>
<thead>
<tr>
<th>compd</th>
<th>FIIa (100 nM)</th>
<th>FIXa (500 nM)</th>
<th>AT (500 nM)</th>
<th>FIXa (500 nM)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>μM</td>
<td>pg/mL</td>
<td>μM</td>
<td>pg/mL</td>
</tr>
<tr>
<td>FCS-Ib</td>
<td>3.65</td>
<td>18.4</td>
<td>1.58 $\times 10^{-7}$</td>
<td>4.95</td>
</tr>
<tr>
<td>18-mer</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FCS-Ib</td>
<td>7.7</td>
<td>25.7</td>
<td>3.33 $\times 10^{-6}$</td>
<td>10.2</td>
</tr>
<tr>
<td>12-mer</td>
<td>8.2</td>
<td>20.5</td>
<td>3.55 $\times 10^{-6}$</td>
<td>20.8</td>
</tr>
<tr>
<td>9-mer</td>
<td>16.8</td>
<td>27.7</td>
<td>7.27 $\times 10^{-6}$</td>
<td>57.5</td>
</tr>
<tr>
<td>FCS-Pg</td>
<td>5.2</td>
<td>26.2</td>
<td>1.78 $\times 10^{-6}$</td>
<td>14</td>
</tr>
<tr>
<td>18-mer</td>
<td>7.2</td>
<td>30.1</td>
<td>2.46 $\times 10^{-6}$</td>
<td>20.8</td>
</tr>
<tr>
<td>FCS-Pg</td>
<td>8.8</td>
<td>29.4</td>
<td>3.01 $\times 10^{-6}$</td>
<td>22</td>
</tr>
<tr>
<td>12-mer</td>
<td>12.5</td>
<td>31.2</td>
<td>4.28 $\times 10^{-6}$</td>
<td>39.5</td>
</tr>
<tr>
<td>FCS-Pg</td>
<td>23.5</td>
<td>38.8</td>
<td>8.04 $\times 10^{-6}$</td>
<td>82.8</td>
</tr>
<tr>
<td>6-mer</td>
<td></td>
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</table>

*The detailed SPR competition sensorgrams are shown in Figures S12–S15. I_Co value of the FCS oligomer molar concentration required to reach 50% of binding competition. $K_D$ is the binding affinity of the FCS oligomer for the coagulation factor or cofactor, and it is calculated by the Cheng–Prusoff formula $K_D = I_Co/[1 + (I_Co/K)])$, where $C$ is the concentration of the coagulation factor or cofactor and $K_D$ is the binding affinity of native FCS for the coagulation factor or cofactor.*
Competitive binding experiments confirmed the interaction between the FCS-Ib 18-mer and FIIa. Once the binding sites of FIIa were occupied by the FCS-Ib 18-mer, they could not be replaced with immobilized FCS-Ib, indicating that the FCS-Ib 18-mer shows fairly strong binding affinity with FIIa. According to competitive binding experiments with various concentrations of the FCS-Ib 18-mer, the IC$_{50}$ value required to reach 50% of binding competition was obtained at ~3.85 μM. We then used the Cheng–Prusoff formula $K_d = IC_{50} / (1 + C/K_d)$ to calculate the binding affinity of the FCS-Ib 18-mer for FIIa ($K_d$) to be 1.67 × 10$^{-7}$ M, which is approximately 1/40th of FCS-Ib–FIIa binding affinity ($K_d = 4.33 × 10^{-5}$ M).

We determined the binding affinities of other FCS oligomers for coagulation factors or AT by using FCS oligomer concentrations required to reach 50% of binding competition (Table 2). In all competitive binding tests, the IC$_{50}$ molar values for the FCS oligomers increased with a decrease in molecular size. The binding affinity of the FCS oligomer for the coagulation factor or AT also decreased with a decrease in molecular size. Both types of oligomers from the 9-mer to the 6-mer showed an even more pronounced decrease in binding affinity with a coagulation factor or AT. Among the coagulation factors and AT, FIXa showed higher binding affinities for FCS oligomers than FIIa and AT, explaining the selective anti-FXase activities of FCS 9–18 oligomers. For the same molecular size, the FCS-Ib oligomers showed some stronger binding affinity with coagulation factors and AT, indicating that FCS oligomers with 2,4-disulfated Fuc branches have stronger binding affinity for these proteins than those with 3,4-disulfated Fuc branches. When the IC$_{50}$ molar values were turned into IC$_{50}$ mass values, it indicated that all FCS-Ib 9–18 oligomers or all FCS-Pg 9–18 oligomers showed similar inhibitory effects on coagulation factors, especially on FIXa.

**FCS 9–18 Oligomers Exhibit Molecular Size-Independent Potent Antithrombotic Activity In Vivo.** We next investigated and compared the in vivo antithrombotic activity of both types of FCS 6-mer to 18-mer through a rabbit brain thromboplastin-induced venous thrombosis model. FCS oligomers or the LMWH (enoxaparin, positive control) were administered dorsally and subcutaneously to male Sprague-Dawley rats before inducing the venous thrombosis. As shown in Figure 4, both types of FCS 9-mer to 18-mer exhibited similarly strong inhibition of venous thrombus formation with an inhibition rate of 92–95% at a dose of 10 mg/kg. This effect was close to that of LMWHs at a dose of 4 mg/kg. Additionally, FCS 9–18 oligomers exhibited similarly strong inhibition of venous thrombus formation with an inhibition rate of 55–65% at a dose of 5 mg/kg. Although these FCS 9–18 oligomers showed anticoagulant activities and binding affinity for various clotting factors commensurate with their molecular size and the sulfation pattern on Fuc branches in vitro, all of these FCS oligomers showed similar antithrombotic activities in vivo at two doses. This indicated that inhibiting thrombogenesis is much more complex than inhibiting a single anticoagulant factor. From the result of IC$_{50}$ mass values in Table 2, it showed that all FCS-Ib 9–18 oligomers at a concentration of 4.5–7.1 μg/mL could inhibit half of 500 nM FIXa, and all FCS-Pg 9–18 oligomers at a concentration of 8.8–12.4 μg/mL could inhibit half of 500 nM FIXa. Thus, the same concentration of FCS 9–18 oligomers has an inhibition ability similar to that of FIXa. Although both FCS-Ib 6-mer and FCS-Pg 6-mer exhibited much weaker anticoagulant activities, they could still interact with a variety of coagulation factors, which caused thrombosis inhibition rates of 66% and 62% at a dose of 10 mg/kg, respectively. Overall, the antithrombotic properties of these FCS oligomers can be attributed to their interaction with various clotting factors, particularly FIXa, finally inhibiting the generation of venous thrombus.

**FCS Oligomers Are Rapidly Absorbed after Subcutaneous Administration.** We tracked the absorption and metabolism of two types of fluorescently labeled FCS oligomers mainly including 6–18 oligomers after subcutaneous administration to rats to determine how long FCS oligomers serve as anticoagulants and antithrombotic agents in the body. First, fluorescein isothiocyanate (FITC) was used to label the two types of FCS oligomers as shown in Figure S16. Ultraviolet (UV) scanning spectra indicated that these FITC-labeled FCS oligomers showed UV absorbance at ~490 nm (Figure S17). HPGPc profiles by refractive index detection indicate that FITC-labeled FCS oligomers show almost the same molecular size distributions as the original FCS oligomers (Figure S18a,b). HPGPc profiles of FITC-labeled FCS oligomers by the UV detector at 490 nm were similar to these obtained by refractive index detection, indicating that FITC labeling was uniform for different oligomers (Figure S18a,b).
Fluorescence scanning spectra of the two types of FITC-labeled FCS oligomers both show maximum excitation wavelengths (Ex) at \( \sim 450 \) nm and maximum emission wavelengths (Em) at \( \sim 510 \) nm, similar to those of FITC (Figure S19).

After successful FITC labeling, the two types of FITC-labeled FCS oligomers were subcutaneously administered to rats to determine their absorption and metabolism. Both FCS-Ib FITC-labeled and FCS-Pg FITC-labeled oligomers showed similar absorption–metabolism curves in the blood (Figure 5).

FITC-labeled FCS oligomers rapidly appear in the blood at 0.5 h after subcutaneous administration. After \( \sim 1 \) h, the concentration of FITC-labeled FCS oligomers in the blood reached its maximum before gradually decreasing. Within the next hour, the concentration of FITC-labeled FCS oligomers in blood is reduced to half of its peak value. Five hours after injection, the concentration of FITC-labeled FCS oligomers in blood was almost undetectable. From the absorption–metabolism curves 1–5 h after administration, we could calculate that the half-lives (\( t_{1/2} \)) of FCS-Ib FITC-labeled and FCS-Pg FITC-labeled oligomers are 1.00 and 1.02 h, respectively.

Figure 5. Absorption and metabolism of FCS oligomers after subcutaneous administration in rats. Concentration monitoring of (a) FCS-Ib FITC-labeled oligomers and (c) FCS-Pg FITC-labeled oligomers in blood. The half-lives (\( t_{1/2} \)) of FCS-Ib FITC-labeled and FCS-Pg FITC-labeled oligomers in the blood 1–5 h after administration were 1.00 and 1.02 h, respectively. HPGPC profiles of the urine samples at different periods after subcutaneous administration of (b) FCS-Ib FITC-labeled oligomers and (d) FCS-Pg FITC-labeled oligomers. The control was FITC-labeled FCS oligomers (1 mg/mL), and the blank was the urine before injection. Samples were analyzed using a Superdex Peptide 10/300 GL column (10 mm x 300 mm) by a UV detector at 490 nm while using 0.2 M NaCl as the eluent at a flow rate of 0.4 mL/min.

Figure 6. Bleeding effect of (a) FCS-Ib oligomers and (b) FCS-Pg oligomers in vivo. The LMWH (enoxaparin, positive control) or FCS oligomers were administered dorsally and subcutaneously to male Kunming mice. Blood loss was determined by measuring the hemoglobin present in the water using a spectrophotometric method. The results are expressed as microliters of blood loss (mean ± the standard deviation; \( n = 6; **P < 0.01 \) vs control).

FITC-labeled FCS oligomers rapidly appear in the blood at 0.5 h after subcutaneous administration. After \( \sim 1 \) h, the concentration of FITC-labeled FCS oligomers in the blood reached its maximum before gradually decreasing. Within the next hour, the concentration of FITC-labeled FCS oligomers in blood is reduced to half of its peak value. Five hours after injection, the concentration of FITC-labeled FCS oligomers in blood was almost undetectable. From the absorption–metabolism curves 1–5 h after administration, we could calculate that the half-lives (\( t_{1/2} \)) of FCS-Ib FITC-labeled and
FCS-Pg FITC-labeled oligomers in the blood were 1.00 and 1.02 h, respectively. These results are consistent with previous reports that a depolymerized FCS ($M_w \sim 12$ kDa) was nearly cleared in 6 h after intravenous administration in rats.

In these experiments, we collected urine samples at different times. The HPGPC profiles of these samples using UV detection at 490 nm are shown in panels b and d of Figure 5. The FITC-labeled FCS oligomers in urine show nearly the same molecular size distribution as the administered FITC-labeled FCS oligomers, indicating their reliable stability in the circulatory system. Moreover, the concentration changes of FITC-labeled FCS oligomers in urine reflected the levels observed in the blood. We could conclude that the appreciable quantity of both types of FCS 6-mer to 18-mer in the blood could be excreted through the kidneys without a degradative metabolism.

**FCS Oligomers Exhibit No Bleeding Risk In Vivo.** Our previous studies showed that subcutaneous administration of a mixture of the FCS-Pg 6-mer to 18-mer did not cause bleeding in vivo. Both types of 9-mer to 18-mer were investigated independently for their bleeding risk to provide a more accurate assessment on this side effect. The LMWH (enoxaparin, positive control) or FCS oligomers were administered dorsally and subcutaneously into male Kunming mice before cutting 5 mm from the tip of the tail. Blood loss was determined by measuring the hemoglobin present in the water using a spectrophotometric method. The results are expressed as microliters of blood loss as shown in Figure 6. Compared with the blood loss of the control group, both types of the FCS 9-mer to 18-mer did not result in obvious blood loss at doses of 120 mg/kg ($P > 0.05$). In contrast, the LMWH caused obvious blood loss at doses of 40 mg/kg ($P < 0.01$). These results suggest that these oligomers effectively result in no bleeding risk in vivo.

**Table 3. Relation between Molecular Information and the Influence on Platelet Aggregation of Different FCS-Ib Compounds**

<table>
<thead>
<tr>
<th>compd</th>
<th>$M_n$ ($10^5$ g/mol)</th>
<th>polydispersity ($M_n$/$M_w$)</th>
<th>Mark–Houwink–Sakurada $\alpha$</th>
<th>intrinsic viscosity $\eta$ (mL/g)</th>
<th>maximum aggregation (%)$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>native FCS-Ib</td>
<td>70.09 ± 0.862</td>
<td>1.442 ± 0.046</td>
<td>0.538</td>
<td>78.54 ± 0.911</td>
<td>41.6</td>
</tr>
<tr>
<td>FCS-Ib D1</td>
<td>13.09 ± 0.256</td>
<td>1.609 ± 0.118</td>
<td>0.970</td>
<td>28.04 ± 0.175</td>
<td>23.1</td>
</tr>
<tr>
<td>FCS-Ib D2</td>
<td>9.994 ± 0.192</td>
<td>1.518 ± 0.091</td>
<td>0.605</td>
<td>14.89 ± 0.084</td>
<td>19.8</td>
</tr>
<tr>
<td>FCS-Ib D3</td>
<td>8.828 ± 0.428</td>
<td>1.556 ± 0.182</td>
<td>0.530</td>
<td>11.58 ± 0.015</td>
<td>15.0</td>
</tr>
<tr>
<td>FCS-Ib D4</td>
<td>7.350 ± 0.122</td>
<td>1.630 ± 0.077</td>
<td>0.493</td>
<td>10.32 ± 0.015</td>
<td>6.49</td>
</tr>
<tr>
<td>FCS-Ib D5</td>
<td>6.668 ± 0.120</td>
<td>1.671 ± 0.057</td>
<td>0.465</td>
<td>9.420 ± 0.091</td>
<td>6.13</td>
</tr>
<tr>
<td>FCS-Ib D6</td>
<td>6.022 ± 0.502</td>
<td>1.723 ± 0.232</td>
<td>0.432</td>
<td>8.714 ± 0.019</td>
<td>5.21</td>
</tr>
<tr>
<td>FCS-Ib D7 (18-mer)</td>
<td>5.398 ± 0.136</td>
<td>1.795 ± 0.097</td>
<td>0.443</td>
<td>7.850 ± 0.021</td>
<td>3.06</td>
</tr>
<tr>
<td>FCS-Ib D8 (15-mer)</td>
<td>4.550 ± 0.165</td>
<td>1.641 ± 0.087</td>
<td>0.465</td>
<td>6.912 ± 0.049</td>
<td>5.50</td>
</tr>
<tr>
<td>FCS-Ib D9 (12-mer)</td>
<td>3.546 ± 0.128</td>
<td>1.753 ± 0.109</td>
<td>0.439</td>
<td>6.269 ± 0.024</td>
<td>4.19</td>
</tr>
</tbody>
</table>

$^a$The maximum aggregation values of platelets caused by different FCS-Ib compounds are cited from our previous publication.14

![Figure 7. Effects of (a) FCS-Ib, (b) FCS-Pg, (c) the FCS-Ib 18-mer, and (d) the FCS-Pg 18-mer administered by intravenous injection on heart rate (●, BPM) and mean arterial blood pressure (■, mmHg) of male Sprague-Dawley rats. After a 10 min adaptation period, the rats were intravenously administered at 10 mg/kg of native FCS or FCS 18-mer as an aqueous solution, and then the systolic, diastolic blood pressures and the heart rate were continuously monitored. The results are expressed as means ± the standard deviation ($n = 5$).](https://dx.doi.org/10.1021/acschembio.0c00439)
anticoagulant and antithrombotic effects and may be safer than LMWHs for clinical application in preventing bleeding.

**FCS Oligomers Exhibit No Hypotension or Platelet Aggregation Risk In Vivo.** Molecular size determines whether an FCS oligomer can activate human FXII leading to dangerous hypotension and a reduced heart rate.\(^{12,25}\) We investigated the in vivo hypotension risk of both types of the FCS 18-mer and compared these to native FCS (Figure 7). Intravenous tail administration of FCS-Ib or FCS-Pg polysaccharides to rats immediately caused hypotension and a reduced heart rate, which were mostly caused by activating FXII.\(^{10}\) After a continuous decrease for ~2 min, the blood pressure and heart rate level out at a value well below normal. In contrast, the administration of the FCS-Ib 18-mer or FCS-Pg 18-mer showed no obvious effect on blood pressure or heart rate in rats. This result was also mostly caused by the inactivation on FXII by the FCS-Ib 18-mer or FCS-Pg 18-mer with a sufficiently small molecular size.\(^ {12}\) We hypothesize that the smaller molecular size of the FCS 9-mer to 15-mer oligomers was insufficient to cause hypotension and a reduced heart rate.

The molecular size also determines whether an FCS oligomer can cause platelet aggregation.\(^ {13-14}\) Solutions of a series of FCS-Ib compounds were studied to determine their association with platelet aggregation (Table 3). FCS-Ib compounds of different molecular sizes exhibited entirely different chain conformations and intrinsic viscosities. The chain conformation characteristic constant, Mark–Houwink–Sakurada \(\alpha\) values of 0–0.5, reflects a rigid sphere in an ideal solvent. Values of 0.5–0.8 correspond to a random coil in a good solvent, and values of 0.8–2.0 correspond to a rigid or rodlike configuration (stiff chain).\(^ {39}\) The native FCS-Ib polysaccharide showed a random coil with a high intrinsic viscosity of 78.5 mL/g in solution and easily caused platelet aggregation (maximum aggregation of 42%). After depolymerization, some \(M_n\) compounds (FCS-Ib D1–D3) still showed a random coil or a stiff chain in solution, but their intrinsic viscosities were significantly decreased along with their decreased \(M_n\). FCS-Ib D1–D3 caused a much lower aggregation maximum for platelets. Even lower-\(M_n\) compounds (FCS-Ib D4–D9) became rigid spheres in solution, and their intrinsic viscosity continued to decrease with a decreased \(M_n\). FCS-Ib D4–D9 did not result in platelet aggregation. On the basis of these results, we could conclude that the platelet aggregation caused by large FCS compounds is highly related with their chain conformation and intrinsic viscosity in solution. Both types of FCS 9-mer to 18-mer are sufficiently small with a rigid sphere conformation and a low intrinsic viscosity in solution to avoid platelet aggregation.

**SUMMARY AND CONCLUSIONS**

In this study, we prepared and purified the FCS 6-mer to 18-mer from two types of sea cucumbers, Ib and Pg. FCS oligomer structural analysis, performed by NMR and ES-FTMS spectroscopy, confirmed that they were all comprised of trisaccharide repeating units of the structure \(-\text{Fuc}C2\text{3,4dis}-\alpha l3-0-\text{GlcA}-\beta l3-1\text{-[\-GalNAc4,6dis}-\beta l4-1\text{-Fuc}C2\text{3,4dis}-\alpha l3-1\text{-GlcA}-\beta l3-1\text{-[\-anTal-e4,6dis}]- (n = 1–5)\), and the only distinction of the oligomers prepared from different sea cucumbers was their Fuc branch sulfation pattern. We next compared their anticoagulant activities in vitro, including APTT, TT, anti-FIIa/AT, anti-FXa/AT, and anti-FXase. The 9-mer to 18-mer of both types of FCS oligomers mainly showed effects on the intrinsic clotting pathway through inhibition of the FXase. SPR competitive experiments verified that these FCS oligomers exhibit the greatest binding affinity for FIXa among various clotting proteins. Both the blood coagulation assays and SPR measurements indicated that the anticoagulant activities of these FCS oligomers vary with the molecular size and Fuc branch sulfation pattern. However, all sizes of FCS 9–18 oligomers with 2,4-disulfated or 3,4-disulfated Fuc branches exhibited similar in vivo antithrombotic effects. Both types of FCS 6-mers also showed favorable antithrombotic activity in vivo, although they showed weak anticoagulant in vitro.

In conclusion, we proposed the mechanism for the use of FCS oligomers for inhibiting thrombosis in blood vessels. On the basis of the blood coagulation assays and SPR measurements, FCS oligomers in the blood can interact with multiple clotting proteins, including FIXa, FXa, and FIIa to inhibit the coagulation pathway.\(^ {30}\) The most significant is the interaction with FIXa to inhibit FXase, preventing the generation of FXa. Meanwhile, these FCS oligomers could also interact with FXa to prevent the generation of FIIa and interact weakly with FIIa to prevent the generation of fibrin. In addition, at the same mass dose of FCS oligomers, although the binding affinities of the smaller oligomers are relatively weak, their molar concentration is relatively higher, giving these more opportunity to interact with clotting factors once they enter the blood circulation. Thus, these FCS oligomers achieve a similar in vivo antithrombotic effect.

By FITC labeling, we found that both types of FCS 6-mer to 18-mer were rapidly absorbed into the blood after subcutaneous administration in rats and appreciable quantities of them were cleared through the kidneys in ~5 h. Evaluation of the side effect indicated that none of the FCS 9–18 oligomers caused bleeding or hypotension in rats. Furthermore, FCS 9–18 oligomers were sufficiently small to behave as rigid spheres with low intrinsic viscosity in solution, avoiding platelet aggregation. Thus, they should be suitable for the large-scale preparation for clinical evaluation in antithrombotic applications.

### METHODS

**Materials.** The native FCS-Pg polysaccharide was isolated and purified from the sea cucumber Pg, as previously described.\(^ {11}\) The depolymerized FCS-Pg polysaccharide was accomplished through partial N-deacetylation–deaminative cleavage as previously described.\(^ {13}\) The polysaccharide was fractionated by gel filtration on a Superdex 200 prep grade column (2.6 cm × 120 cm, GE Healthcare Life Sciences) with 0.3 M NH\(_4\)HCO\(_3\), at a flow rate of 0.3 mL/min. The native FCS-Ib and different sizes of FCS-Ib oligomers were obtained in the same way as in our previous publication.\(^ {14}\) Finally, we obtained ~2 g of FCS oligomers mainly including 6–18 oligomers from 1 kg of each type of dry sea cucumber with optimized purification degradation. After separation of mixed oligomers by GPC, each size of highly purified oligomer was obtained in amounts of 50–100 mg. Unfractionated heparin (UFH) was obtained from Sigma (St. Louis, MO). The LMWH (enoxaparin, 0.4 mL × 4000 AU/mL) was obtained from Sanofi-Aventis. APTT assay kits, TT assay kits, a calcium chloride solution (0.02 M), and standard human plasma were obtained from Siemens Healthcare Diagnostics. Biophen Heparin Anti-FIIa kits, Biophen Heparin Anti-FXa kits, and a chromogenic assay kit for measuring FVIII:C in concentrates were obtained from Hyphen Biomed. Human coagulation FVIII was obtained from China Biologic Products, Inc. (Shandong, China). Human thrombin (FIIa), purified human factor Xa (Fxa), human antithrombin (AT), and purified human factor IXa.
The results are expressed as international units per kg of body weight. Male Kunming mice (body weights of 18 g) and Male Sprague-Dawley rats (body weights of 300 g) were randomly segregated into 20 groups of eight animals each. Measurement of Anticoagulant Activity.

**The APTT and TT assays were determined with a coagulometer (RAC-120) using APTT and TT reagents and standard human plasma as previously described.**

**The results are expressed as international units per milligram using a parallel standard curve based on the international heparin standard (212 IU/mg).**

**Antithrombin (anti-FIIa) and antifactor Xa (anti-FXa) activities** were measured in the presence of antithrombin (AT) and samples and inhibition of intrinsic factor Xase (factor IXa−factor VIIIa complex) in the presence of samples were carried out in a 96-well microtiter plate as previously described. The absorbance change rate was proportional to the FIIa and FXa activity occurring in the incubation mixtures. The experimental results were expressed as the percent of control (n = 3). IC50 values were obtained by fitting the data to a noncompetitive inhibition model for the samples according to Sheehan and Walke. Studies of the Interaction between GAGs and Coagulation Factors or a Coafactor. The interactions of GAGs (including FCS-18, FCS-Pg, and UFP) with coagulation factors or a cofactor (FIIa, FXa, AT, and FXa) were measured using surface plasma resonance (SPR) on a BIACore 3000 system while referring to similar publications. Briefly, the biotinylated GAG was immobilized to an SA chip based on the manufacturer’s protocol. Successful immobilization of GAG was confirmed by the observation of a >200 resonance unit (RU) increase in the sensor chip. The commercial coagulation factor or cofactor was resuspended in HBS-EP buffer. Different dilutions of the coagulation factor or cofactor were injected at a flow rate of 30 μL/min. At the end of the injection, the same buffer was passed over the sensor surface to facilitate dissociation. After dissociation, the sensor surface was regenerated by injecting 30 μL of 2 M NaCl. The response was monitored as a function of time (sensorgram) at 25 °C. Competition of Coagulation Factors or Cofactor Binding to Immobilized Native FCSs by FCS Oligomers. Competitive SPR experiments, using different sizes of FCS oligomers, were performed to compare the ability to bind to FIIa, FXa, FIXa, or AT. A coagulation factor or cofactor was specifically preincubated with graded concentrations of the FCS oligomer and then injected over the native FCS chip at a flow rate of 30 μL/min. After each injection and association, dissociation and regeneration were performed as described above.

Inhibition of Venous Thrombosis. Antithrombotic activity in rats was assessed using rabbit brain thromboplastin as the thrombogenic stimulus. Male Sprague-Dawley rats (body weights of 250–300 g) were randomly segregated into 20 groups of eight animals each. The control group, the LMWH group, and the FCS oligomer groups were administered dorsally and subcutaneously at a dose of 1 mL of 0.86% NaCl/kg of body weight, 4 mg of LMWH/kg of body weight, and 5 or 10 mg of FCS oligomers/kg of body weight, respectively. After 60 min, rats were anesthetized with an intramuscular injection of 100 mg of ketamine/kg of body weight and 16 mg of xylazine/kg of body weight, the inferior vena cava and its branches were isolated, and the branch of inferior vena cava under the left renal NMR spectrum was ligated. A volume of 1 mL of 2% tissue thromboplastin/kg of body weight was injected from the femoral vein. After 20 s, stasis was established by ligating the edge of the left renal vein. After a 20 min stasis, the cavity was then reopened, the ligated segment was opened longitudinally, and the thrombus formed was removed, rinsed, dried for 24 h at 50 °C, and then weighed. The results are reported as means with the standard deviations. Absorption and Metabolism of FCS Oligomers. Male Sprague-Dawley rats (body weights of 250–300 g) were randomly segregated into two groups of five animals each to measure the absorption and metabolism of FCS oligomers. Two groups were administered dorsally and subcutaneously with 10 mg of FCS-18 FITC-labeled oligomers and FCS-Pg FITC-labeled oligomers per kilogram of body weight, respectively. After 0.5, 1.0, 1.5, 2.0, 2.5, 3.0, 4.0, and 5.0 h, 0.3–0.5 mL of blood was collected from the tails and mixed with one-ninth of the volume of a sodium citrate solution (0.109 mol/L). After centrifugation at 3000g for 5 min, plasma was detected as FCS-oligomers−FITC content with a fluorescence microplate (BioTek) from a standard curve based on Ex at 450 nm and Em at 510 nm. Meanwhile, each group’s urine at periods of 0–2.0, 2–3.5, and 3.5–5.0 h was separately collected, combined, and filtered (0.22 μm filter membrane, Millipore) for detecting FCS-FITC-labeled oligomers by HPLC with a UV detector at 490 nm. Bleeding Effects. Blood loss was determined by measuring the hemoglobin present in the water using a spectrophotometric method. Male Kunming mice (body weights of 18–22 g) were randomly segregated into 10 groups of six animals each. The control group, the LMWH group, and the FCS oligomer groups were administered dorsally and subcutaneously with 5 mL of 0.86% NaCl/kg of body weight, 40 mg of LMWH/kg of body weight, and 120 mg of FCS oligomers/kg of body weight, respectively. After 60 min, the tails of the mice were cut 5 mm from the tip and immersed in 40 mL of distilled water for 90 min at 37 °C while being stirred. The volume of blood was determined from a standard curve based on absorbance at 540 nm. Arterial Blood Pressure and Heart Rate. Male Sprague-Dawley rats (body weights of 250–300 g) were randomly segregated into four groups of five animals each to measure the arterial pressure and heart rate. Rats were anesthetized with a combination of xylazine and ketamine as described previously, and a P10 catheter was inserted into both their right femoral vein and artery. The arterial catheter was connected to a pressure transducer (MP150, BIOPAC Systems Inc.) coupled to an acquisition system (AcqKnowledge, BIOPAC Systems Inc.). After a 10 min adaptation period, the rats were intravenously administered 10 mg of native FCS or FCS 18-mer/kg as an aqueous solution, and then the systolic and diastolic blood pressures and heart rate were continuously monitored. At the end of the observation period, the animals were euthanized using KClI (10 mg/kg).

Molecular Information about FCS Components. Molecular information about FCS compounds, including weight-average molecular weight (Mw), polydispersity (Mw/Mn), and chain conformation (Mark−Houwink−Sakurada α), was obtained by size exclusion chromatography equipped with a multiangle laser light scattering system with a refractive index detector (SEC-MALLS-RI, Wyatt Technology). SEC columns (SB-806 HQ and SB-804 HQ, 7.8 mm × 300 mm, Shodex) were protected by a OHpak SB-G guard column (Shodex), and SEC was performed at 25 °C (g) with a sample loading rate of 0.7 mL/min (ViscoStar III, Wyatt Technology) and connected to the SEC-MALLS-RI system. The value of dn/dc (specific refractive index increment) was estimated to be 0.138 mL/g/0.2 M NaCl containing 0.02% Na2SO4 (pH 7.0) was used as the mobile phase at a flow rate of 0.5 mL/min.
0.5 mL/min. The sample was dissolved directly in the mobile phase (2–5 mg/mL) and filtered through 0.22 mm filter membrane (Millipore). The injection volume was 50 mL and run for 100 min. Data acquisition and calculations were performed with ASTRA, version 7.1.2 (Wyatt Technology).

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acschembio.0c00439.

Additional data about the FCS oligomers’ NMR spectra with 1H and 13C chemical shifts, FCS oligomers' high-resolution FTMS spectromaps, SPR sensorgrams with a fitted curve and kinetic constants, SPR competition sensorgrams with IC_{50} curves, and preparation and characterization of FCS oligomers-FITC (PDF)

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### Notes

The authors declare no competing financial interest.

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