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Heparinoids activate a protease, secreted by mucosa and tumors, via tethering supplemented by allostery

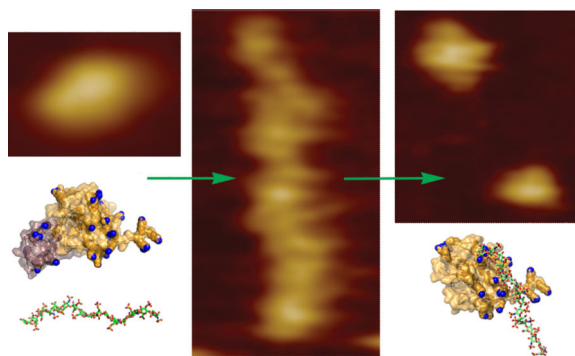
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Abstract



Activation by glycosaminoglycans (GAGs) is an emerging trend among extracellular proteases important in disease. ProMMP-7, the zymogen of a matrix metalloproteinase secreted by mucosal epithelial and tumor cells, is activated at their surfaces by sulfated GAGs, but how? ProMMP-7 is activated in *trans* by representative heparin oligosaccharides in a length-dependent manner, with a large jump in activation at lengths of 16 monosaccharides. Imaging by atomic force microscopy visualized small complexes of proMMP-7 molecules linked by 8-mer lengths of heparinoids and extended assemblies formed with 16-mer lengths of heparin. Complexes of proMMP-7 with polydisperse heparin or heparan sulfate were more diverse. Heparinoids evidently accelerate activation by tethering multiple proMMP-7 molecules together for proteolytic attack among neighbors. Removal of either the prodomain or C-terminal peptide sequence of KRSNSRKK from MMP-7 prevents formation of the long arrays induced by heparin 16-mers or heparan sulfate. The role of the C-terminus in activation assays suggests it contributes to remote, allosteric binding of GAGs. Enhancement of proteolytic velocity of MMP-7 by GAGs indicates them to be effectors of

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ASSOCIATED CONTENT

Supporting Information

Supporting information includes supplementary figures, tables, sources of materials, preparative methods, buffers, and methods details. This material is available free of charge via the Internet at <http://pubs.acs.org>.

V-type allostery. GAGs from proteoglycans appear to assemble proMMP-7 molecules for activation, an event preceding its tumorigenic or anti-bacterial proteolytic activities at cell surfaces.

Keywords

zymogen activation; glycosaminoglycans; protein-carbohydrate interactions; atomic force microscopy; allostery; enzyme kinetics

INTRODUCTION

Zymogens from four major classes of proteases were found to be regulated by sulfated glycosaminoglycans (GAGs) which physiologically radiate from the core proteins of proteoglycans on and outside of animal cells. GAGs accelerate the activation of procathepsins B, L and S (lysosomal cysteine proteases),¹⁻³ the serine protease trypsin,⁴ and aspartic proteases such as β -secretase.^{5, 6} GAGs promote autolytic activation of proMMP-2 and -7,^{7, 8} plus transactivation of proMMP-2 by proteases.⁹⁻¹¹ The zymogens of MMPs comprise a prodomain, a catalytic domain (Figure 1A), and usually a hemopexin-like domain.^{12, 13} GAGs can position the protease to attack substrate, e.g. heparin proteoglycan linking chymase to substrate¹⁴ or GAG chains recruiting MMP-7 to proHB-EGF at surfaces of mucosal epithelial cells in female reproductive organs¹⁵ or to pro- α -defensins to activate bactericidal activity.^{8, 16} GAGs also regulate some active proteases. For example, heparin accelerates antithrombin inhibition of thrombin¹⁷ and chondroitin-4-sulfate (C4S) makes cathepsin K collagenolytic.¹⁸ The work herein asks by what mechanism(s) does GAG-dependent activation of proMMP-7 proceed? Greater understanding should offer perspective on GAG activation of other pathophysiologically important proteases as well.

GAG-stimulated activation of proMMP-7 initiates its role in innate immunity⁸ and very possibly in tumorigenesis.^{15, 19} Heparan sulfate (HS) appears to dock proMMP-7 to epithelial cells.^{20, 21} Proteolytic activity of MMP-7 in mucosal epithelia promotes antibacterial defense in the small intestine,¹⁶ wound closure in the lung,²² tumor cell survival (by decreasing FasL-triggered apoptosis),^{23, 24} and mammary tumor cell proliferation (via release of a domain from the ErbB4 receptor).²⁵ High MMP-7 expression is associated with early stage breast cancer,¹⁹ its metastasis to bone,²⁶ as well as invasion, metastasis and poor outcome of colorectal cancer.²⁷

The activation of proMMPs can be grouped under mechanisms of stepwise proteolysis and allosteric displacement of the prodomain not requiring proteolysis.¹² Stepwise proteolysis can be triggered by destabilizing agents such as proteases and thiol-reactive agents.¹² Allosteric activation without proteolysis has resulted from protein-protein interactions with the pro-forms of proMMP-9, -2, and -3.^{12, 28, 29} Heparin and chondroitin-4,6-sulfate (CSE) activate proMMP-7.^{8, 30} One hypothesized mechanism for this activation is an allosteric bridging of the GAG chain between pro- and catalytic domains to expose the active site.¹³ Such bridging in a 1:1 complex that unmask the active site is the working hypothesis for autoactivation of procathepsin B.¹ It was speculated that heparin might enhance activity of MMP-7 through conformational change.³⁰ Another possibility would be GAGs linking

proMMP-7 molecules, analogous to GAGs of proteoglycans bridging MMP-7 to substrates of pro- α -defensin^{8, 13} or pro-HB-EGF in complex with its ErbB4 receptor.¹⁵ Simultaneous binding of serine protease inhibitors (serpins) and serine proteases to heparin results in inhibition.^{31, 32} Considering such alternative hypotheses and elucidating the mechanism of GAG activation of proMMP-7 is the purpose of this work.

The heparinoids HS and heparin are extended, undulating polymers of disaccharides of uronic acid $\alpha(1-4)$ -linked to *D*-glucosamine with a higher level of sulfation in heparin and more N-acetyl groups in HS.^{33, 34} A tetrasaccharide therein forms one period of the sinusoidal structure (Figure 1B).³⁵ The composition and length of HS especially varies among tissue and cell types.³³ Heparin from bovine and porcine mucosa is widely used in research but polydisperse. Heparin oligosaccharides of defined length have served as a simplifying tool for investigation of GAG-protein interactions and in this study of the activation of proMMP-7. AFM recently imaged carbohydrate-lectin complexes³⁶ and studied forces in an enzyme-GAG association.³⁷ The novel AFM images of GAG-enzyme interactions reveal structural attributes of GAG-induced assemblies with proMMP-7. Heparin fragments appear to activate proMMP-7 primarily by bridging the protease molecules into extended complexes. Formation of these complexes that are efficient at activation requires heparin oligosaccharides of sufficient length, the prodomain, and the C-terminal basic peptide. GAGs then act on mature enzyme to enhance proteolytic velocity, suggesting allosteric transmission across the catalytic domain.

RESULTS

GAGs associating with proMMP-7

The ability of physiological GAGs to interact with proMMP-7 was surveyed by their competition with surface-tethered heparin, using surface plasmon resonance. ProMMP-7 was made stable by the E195A substitution of the general base, which inactivates mature MMP-7 by 1900-fold.³⁸ Heparin, chondroitin sulfate E, and dermatan disulfate (DS) competed well for proMMP-7-E195A and contain two or more sulfate groups per disaccharide repeated in their structures (SI Figures S1, S2). HS, DS, and chondroitin sulfates A and C (averaging one sulfate per disaccharide) and chondroitin sulfate D (with two sulfates per disaccharide) failed to outcompete heparin (SI Figure S2). The lesser affinity of its physiological partner HS for proMMP-7 sufficed for efficient, saturable activation (SI Figure S1B). This suggests that HS may not only recruit proMMP-7 into cell surface complexes,¹⁵ but also accelerate its maturation to the active form.

ProMMP-7 is activated in trans

We asked if GAGs act on individual or pairs of proMMP-7 molecules to induce activation. We exploited oligosaccharides of HS and heparin in assays of proMMP-7 activation for two reasons: (i) to distinguish GAG effects on a single proenzyme from linking of proenzyme molecules and (ii) to introduce homogeneity in size. HS oligosaccharides were available in lengths up to 12 saccharide units (HS dp12). Heparin oligosaccharides are available up to 20 saccharide units (Hep dp20). Heparin oligosaccharides are experimentally well-behaved and may represent physiological partners HS and CSE. Heparin shares similar disaccharide

structure with HS and shares high sulfation with CSE (SI Figure S1C). Incubation of wt proMMP-7 with Hep or HS dp4 to dp12 increased autolytic activation moderately, with appearance of active catalytic domain increasing with oligosaccharide length (Figure 1C,D). The activation was much more complete using Hep dp16, Hep dp20, or low molecular weight heparin. The dramatic boost in activation at lengths of 16 monosaccharides or greater suggests the need for this length for tethering and positioning of the proenzymes together.

We tested the hypothesis of heparin-dependent colocalization promoting activation. E195A-inactivated proMMP-7 is incompetent for the Hep dp20-induced activation that readily proceeds with dilute wt proMMP-7 (Figure 1E, lanes 1–4). Addition of wt proMMP-7 activated part of the proMMP-7-E195A once Hep dp20 was added (Figure 1E, lanes 5–6). Addition of 0.36 equivalents of active wt MMP-7 activated part of proMMP-7-E195A without Hep dp20 and much more with it (Figure 1E, lanes 7–8). Thus, longer heparin chains appear to draw (pro)MMP-7 molecules together for activation to proceed in *trans*.

Hydrodynamics of assembly with heparin fragments

We investigated the average size of proMMP-7 complexes formed in solution as a function of the length of the heparin oligosaccharide, using hydrodynamics. Dynamic light scattering (DLS) established proMMP-7-E195A to be monomeric with estimated MW of 27.6 kDa (and diameter of 2.5 nm; SI Table S1) matching calculated MW of 27.7 kDa. After adding a 10-fold molar excess of Hep dp8, DLS estimated the MW to be 30 kDa and diameter at 2.6 nm, suggesting a single proenzyme in complexes. Addition of Hep dp16 to 5-fold molar excess introduced excessive light scattering dominated by a component appearing to be 6 μm in diameter (SI Table S1), suggesting aggregation, instead of a 2:1 complex hypothesized initially.

ProMMP-7-E195A eluted as a sharp peak from an analytical gel filtration column (Figure 2A). Addition of Hep dp8 to 10-fold molar excess gave rise to a similar but slightly broader peak consistent with DLS evidence of a single proenzyme in complexes. Addition of the Hep dp8 to 20-fold molar excess added a faster migrating peak at fraction 10, suggesting lower affinity association into a higher MW complex. Addition of 4-fold molar excess of Hep dp16 introduced an even larger complex (eluting at fraction 9), while a peak still eluting at fraction 11 likely contained a single proenzyme (Figure 2A). After correcting for retardation in migration of free proMMP-7-E195A in 150 mM NaCl (from 0.67 to 0.81 column volumes), the proMMP-7 with Hep dp8 may have eluted around 2.2-fold the apparent MW of free proMMP-7, consistent with complexes containing two proenzyme molecules. ProMMP-7-E195A with Hep dp16 may have eluted with apparent MW around 4.1-fold that of free proMMP-7.

As a surrogate for apparent MW, rotational correlation times (τ_C of diffusional tumbling) were measured throughout Hep dp8 and dp16 titrations of ^{15}N -labeled proMMP-7-E195A using ^{15}N NMR relaxation (detecting most enzyme states except aggregates). τ_C is proportional to the hydrated volume of a globular protein complex. Addition of 0.6:1 Hep dp8 to proMMP-7-E195A increased τ_C by 1.6-fold to almost 18 ns (Figure 2B). This ratio and the slowed tumbling might suggest formation of assemblies containing possibly two proenzymes per chain of Hep dp8. The drop in τ_C upon addition of the dp8 to 0.8 to 1

equivalents parallels the “template effect” typified by heparin bridging between serpins and serine proteases.^{17, 32, 39} This suggests that the 0.8 to 1.0 equivalents of dp8 competed some proMMP-7-E195A away from some ternary complexes (one dp8 per two proenzymes). Addition of Hep dp16 at 0.4:1 to proMMP-7-E195A slowed tumbling by almost 3-fold to ~31 ns. Both the slowing and the ratio suggest formation of complexes that include cases of two to three (or more) proenzyme molecules bound per Hep dp16. NMR line broadening became severe at 0.5:1 Hep dp16:proenzyme and prohibitive at higher concentrations, consistent with high MW.

Oligomerization and oligosaccharide length

Atomic force microscopy (AFM) enabled us to ask further how do strongly activating complexes with Hep dp16 oligosaccharides differ in structure from weakly activating complexes with dp8? The maximum heights of proMMP-7-E195A appear to peak around 2.5 nm for the free state and complexes with Hep dp16, and just above 2 nm in the Hep dp8 complexes (Figure 3E). These heights are consistent with the diameter of the catalytic domain in the crystal structure (PDB code 1MMQ). ProMMP-7-E195A alone appears as monodisperse monomers, some of which look pear-shaped (Figure 3A). Considering the dimensions of MMP structures containing pro- and catalytic domains, the smaller (sometimes faint) lobe in the AFM-observed pear shape may represent the prodomain and the larger lobe the catalytic domain.

What do apparent volumes from AFM suggest about size of complexes formed? While proMMP-7-E195A is sharply defined in volume, its assemblies with Hep dp8 present have a broad distribution of volumes, and those with Hep dp16 a very broad distribution of volumes. The apparent volumes of the free monomers of proMMP-7-E195A are well-defined around $1.4 \times 10^5 \text{ \AA}^3$, with most between 0.8 and $2.5 \times 10^5 \text{ \AA}^3$ (Figure 3F). In the mixture with Hep dp8 with proMMP-7-E195A, the features are more diffuse with biggest population around 2.0 to $5.5 \times 10^5 \text{ \AA}^3$ (Figure 3F). This being about twice the monomer volume suggests *two* proenzymes to be present in these particles. This is consistent with NMR τ_C evidence for two proenzymes bound per Hep dp8 chain near the 0.6 dp8 per proMMP-7 point in the NMR titration (Figure 2B). AFM images suggest mixtures with HS dp8 are also enriched in apparent tandem dimers, but have a higher proportion of particles containing a single proenzyme and a lower proportion containing multiple proenzyme molecules (Figure 3D,F). Lesser but significant populations with Hep dp8 or HS dp8 appeared around 3.4×10^5 , 4.0×10^5 , and $5.4 \times 10^5 \text{ \AA}^3$ (Figure 3F), suggestive of complexes containing three, four, or five proenzyme molecules, respectively. Less common features with Hep dp8 exhibit an apparent volume of $9.4 \times 10^5 \text{ \AA}^3$ (Figure 3F), suggesting larger complexes containing perhaps eight to ten proMMP-7 molecules.

These AFM data do not account for tip convolution. Deconvolution to minimize the contribution of tip geometry reduced the volume of proMMP-7-E195A and its complexes to around 70% (SI Figure S3); quoted volumes are thus small overestimates.

Striking extended, chain-like arrays with links of diffuse objects formed upon mixing of Hep dp16 with proMMP-7-E195A (Figure 3C). Most of these, extended arrays have apparent volumes of 6 to $30 \times 10^5 \text{ \AA}^3$, though some range up to $60 \times 10^5 \text{ \AA}^3$ (Figure 3F). Such large

features are likely to contain tens of proMMP-7 molecules. Nonetheless, with Hep dp16 the most abundant apparent volumes are near 2.8×10^5 and $4 \times 10^5 \text{ \AA}^3$, suggesting two to four proenzyme molecules in the complexes. Hep dp16 minimizes the number of single enzyme particles present on the mica surface, in contrast to images of dp8-containing mixtures. – This paucity and the extended arrays suggest that Hep dp16 draws smaller complexes together into the extended arrays.

Assemblies with heparinoids

Whether these trends extend to heparin and HS was considered. Bovine heparin or HS, each polydisperse with chain lengths spanning 5 to 50 kDa, was mixed with proMMP-7-E195A and monitored by AFM (Figure 4). In both cases, the features ranged across the size range observed with Hep dp8 to some as large as observed with Hep dp16, but with more variability in morphology. Extended complexes are visible and partly resemble those induced by Hep dp16, both in images with HS and in larger numbers with heparin (Figure 4). Apparent volumes of the AFM features with heparin, HS, Hep dp8 appear, however, to be similar within experimental uncertainties (SI Figure S4). (Numerous smaller particles are present in the polydisperse mixtures, despite the human eye focusing on the larger particles). There is greater spreading of the larger features with HS and heparin. Lower resolution of imaged features seems to accompany longer GAG chains, which might allow more freedom of movement on the surface.

Activation relies on favorable electrostatics

Favorable electrostatic interactions between polyanionic Hep dp16 and the extensive positive charge of proMMP-7 (Figure 1A,B) were tested for importance in activation by adding NaCl up to 1 M in activation assays. 100 mM NaCl decreased activation to about half and 500 mM NaCl to about 10% (SI Figure S5). (Salt had no effect on proteolytic activity of the catalytic domain.)

Prodomain required for assembly

We tested whether the positively charged prodomain (Figure 1A) participates in oligomerization. Without the prodomain, no extended arrays form. Mixing active wt MMP-7 with Hep dp8, Hep dp16, or HS preserved its monodisperse AFM images, heights, and volumes (SI Figure S6), suggesting that the particles mainly contain a single enzyme in each case. Thus, the prodomain appears necessary for heparinoid-induced formation of oligomers.

C-terminal roles in activation and assembly

Since a synthetic C-terminal decapeptide from rat MMP-7 binds heparin,²⁰ we tested whether the corresponding basic peptide from human MMP-7, KRSNSRKK, affects heparin-induced activation of proMMP-7. Deletion of this C-terminus impaired Hep dp16-induced activation, requiring at least 4-fold as much dp16 for as much activation as wt enzyme (Figure 5A). Moreover, synthetic KRSNSRKK peptide competed with Hep dp16-induced activation of wt proMMP-7, as 400 μM peptide nearly abolished it (Figure 5B). The C-terminus must then bind heparin in the activation assay.

A role for the C-terminus in oligomerization was tested by AFM imaging. Hep dp8, dp16 or HS did not enlarge appearance, height or volume of proMMP-7 C particles (Figure 6) Hep dp16 mixed 1:1 with proMMP-7 C was dominated by volumes ($\sim 1 \times 10^5 \text{ \AA}^3$; Figure 6C,F) equaling those of monomeric proMMP-7 (Figure 3A,F). The C-terminus is thus important in oligomerization needed for efficient activation.

GAGs enhance velocity by allostery

We investigated whether GAGs modulate activity of mature MMP-7. HS, DS, and low molecular weight heparin, in a concentration-dependent manner, enhanced MMP-7 catalytic efficiency and velocity of proteolysis of Knight's soluble peptide substrate⁴⁰ of MMPs (SI Table S2). Fitting of progress curves⁴¹ found the enhanced catalytic efficiency (k_{cat}/K_M) mainly results from increased velocity V_{max} or catalytic turnover k_{cat} . The increased velocity and binding outside the active site categorize this as V-type allostery. DS boosted the catalytic efficiency the most of the intact GAGs, i.e by 1.7-fold, along with a 2.1-fold boost of k_{cat} (SI Table S2). HS was the most efficient in needing only 0.1 mg/ml to achieve enhancement of k_{cat} .

We asked whether structural features important in activation of the zymogen are important in enhancing mature MMP-7 activity. Proteolytic velocity increased with length of heparin fragment (SI Table S3). Hep dp16 delivered a 2.8-fold increase in both k_{cat}/K_M and k_{cat} , which was superior to enhancements by Hep dp8, dp12 or low molecular weight heparin (Figure 5C, SI Tables S3, S2). Moreover, enhancement by Hep dp16 saturated at the much lower concentration of 10 μM (Figure 5C).

We checked if the C-terminus participates in this allosteric enhancement. Hep dp16 enhanced the catalytic efficiency and k_{cat} of MMP-7 C by 1.9-fold (Figure 5C), implicating its binding after loss of the C-terminus. MMP-7 C retained only one-third of wild-type catalytic efficiency (Figure 5C). This impairment is surprising because this C-terminus is remote from the active site, outside the fold of this enzyme, and disordered, judging from sharp, random coil NMR peaks. Nonetheless, the C-terminus acts akin to an internal allosteric activator.

DISCUSSION

Trans-activation and assembly via GAG binding

We investigated the question of how proMMP-7 is activated by heparin and HS. The principles may extend to CSE and DS because of their analogous charge complementarity to proMMP-7. Heparin oligosaccharides induce *trans*-activation of proMMP-7 molecules they bridge (Figure 1). The significantly greater activation by heparin chains 16 or more saccharides long suggests side-by-side binding of zymogens, which is corroborated by AFM images (Figures 3, 4). HS and CSE are probable physiological partners for proMMP-7 that are linked to proteoglycans *in vivo*.^{8, 13, 15} Tandem assembly and *trans*-activation of basic proMMP-7 along acidic chains of HS or CSE emanating from proteoglycans can be predicted of mucosal epithelial cell surfaces.

The importance of heparin-binding by the prodomain and basic C-terminus in normal assembly processes is evident from their deletion preventing formation of the arrays induced by Hep dp16 or HS (SI Figure S4, Figure 6). Mixtures of Hep or HS dp8 with proMMP-7 are likely to be enriched in dimers (Figure 3) sandwiched around an 8-saccharide chain bound, as well as 1:1 complexes (Figure 7A). The prodomain and C-terminus of proMMP-7 working together with Hep dp16 to create long tandem arrangements is symbolized in Figure 7B.

A long GAG chain may provide enough freedom for one proMMP-7 to attack the vulnerable sequence linking pro- and catalytic domains of its neighbor on the GAG chain. As activation minimally needs three molecules to interact (two proMMP-7 and GAG chain to link them), the velocity of activation must depend strongly on their concentrations. Extended complexes on longer heparinoid chains (Figures 3C, 4) have locally high concentrations that promote activation. Multiple proenzymes in the oligomers increase the probability that a transiently active proenzyme will contact and hydrolyze a neighboring zymogen.

For covalently intact proMMP-7 to be transiently an activating protease, the pro- and catalytic domains must separate enough to allow a loop of another proMMP-7 to enter its active site. The precedent of prodomain being spatially separated from the active site while covalently joined has been inferred from allosteric activations of proMMP-9, -2, and -3 by protein-protein interactions^{12, 28, 29} and from GAG activation of procathepsin B.¹ ProMMP-9 activated in this fashion was inhibited by TIMP-1, confirming separation of the connected pro- and catalytic domains.²⁸

Allostery

The hypothesis of GAG activation of proMMP-7 being allosteric was deduced from the need for pro- and catalytic domains to separate and for the GAG to bind away from active site.¹³ All GAG constructs tested accelerated proteolytic velocity of MMP-7 (SI Tables S2, S3), establishing them as allosteric activators. The complexes of *catalytic* domain with Hep dp16 or HS containing only one enzyme molecule in over 90% of particles (SI Figure S4) has two implications. First, proteolytic activation of proMMP-7 is likely to free the catalytic domain from GAG-induced oligomers, even while the catalytic domain can continue to associate with GAG chains (Figure 7C). Second, the allosteric enhancement of its proteolytic velocity by the GAG bound is probably transmitted across the catalytic domain.

The need for C-terminus and prodomain in heparinoid-induced assembly and activation (Figures 3, 5, 6, S4) identify them as remote GAG binding sites. The C-terminus must be a key binding site because its deletion seems to diminish the higher affinity component of acceleration by Hep dp16 (Figure 5C). However, the remaining lower affinity component of 1.9-fold acceleration induced by Hep dp16 (Figure 5C and SI Table S4) means that heparin oligosaccharides also bind elsewhere in the catalytic domain to allosterically enhance velocity. Indeed, several other positive charges on the back of the catalytic domain (Figure 1A) were hypothesized to tether rat MMP-7 to the HS of uterine epithelial cells and basement membranes.²⁰ The C-terminus is needed not only for binding heparin but also for full enzyme activity. This suggests a network of communication between C-terminus and active site.

Regulatory bridges of GAGs to proteases

Parallels can be seen between GAG-dependent activations of proMMP-7 and proteases from two other classes. Favorable electrostatics are important in the GAG activations of proMMP-7, procathepsin B, L and S,¹⁻³ β -secretase and procathepsin D.^{5, 6} GAG chains bind *both* pro- and catalytic domains of procathepsin B and proMMP-7. GAG activation of procathepsin D increases with length of heparin oligosaccharide,⁶ quite like *trans*-activation of proMMP-7 (Figure 1). Cathepsin K binding to C4S (CSA) introduces collagenase activity and “beads-on-a-string” oligomeric complexes.¹⁸ However, their regular repetition in crystal lattices⁴² contrasts the irregularity of proMMP-7 upon Hep dp16, heparin, or HS on solution-covered surfaces (Figures 3, 4).

Tethering between active protease and substrate is analogous to tethering between protease molecules. GAG chains were proposed to promote proMMP-2 activation by tethering it to active MMP-2⁷ or thrombin.¹¹ Heparin proteoglycans may draw chymase together with its substrate of thrombin.¹⁴ GAG chains link MMP-7 to pro- α -defensin for its maturation to a bactericidal peptide.⁸ HS proteoglycan positions MMP-7 to process HB-EGF to its mature form to bind its ErbB4 receptor and support cell survival.¹⁵

Joint, ionic binding to heparin, 16 monosaccharides or longer, brings serine proteases close to serpins for inhibition,^{17, 32, 39} similar to 16-mers and longer bridging among proMMP-7 molecules (Figures 1, 3, 4). In contrast to crystallized ternary complexes of serpin-heparin-thrombin,³² multiple copies of proMMP-7 assemble with 16-residue fragments of heparin up to full length heparin and HS (Figures 2, 3, 4). The use of *two* binding sites by proMMP-7 appears to support its repeating bridging by heparin and HS chains.

CONCLUSIONS

The initial hypotheses that GAG activation of proMMP-7 depends on either (i) 1:1 GAG binding with bridging and separation of pro- from catalytic domain¹³ or (ii) formation of ternary complexes with two proMMP-7 bound side-by-side to a GAG chain⁸ (like serpin-heparin-thrombin complexes) must be modified. Heparin-induced activation proceeds mainly in *trans* in assemblies initially containing multiple zymogens, rather than the simple 2:1 complexes expected. (There is precedent for extended complexes of another GAG-dependent protease.⁴²) Heparinoid binding that is highly productive for activation of proMMP-7 bridges it to neighboring proMMP-7 molecules by binding prodomains and C-termini. This tethering together must increase likelihood of proteolytic attack among neighbors. Subsequent to the activation, binding of heparinoid chains to mature MMP-7 (at C-terminus and elsewhere) allosterically enhances its proteolytic velocity. The idea of preventing activation of proMMP-7 as a novel cancer therapeutic strategy with fewer side effects¹³ can now consider intervening in C-terminal and prodomain roles in activation.

METHODS

Surface plasmon resonance

Porcine intestinal heparin (Celsus) was immobilized on a streptavidin chip (GE Healthcare). For competition analysis, 500 nM of proMMP7 pre-mixed with 1000 nM of the GAG in

HBS-P buffer (GE Healthcare) was flowed over the heparin chip at 30 $\mu\text{L}/\text{min}$ at 25°C, followed by 2 min of flow of HBS-P buffer for dissociation, and surface regeneration using 2 M NaCl.

GAG activation of proMMP-7

ProMMP-7 (wt, E195A-inactivated, or with C-terminal KRSNSRKK deletion) was incubated with bovine HS (Sigma), low molecular weight heparin from porcine intestinal mucosa (Sigma), or the heparin oligosaccharides. The reaction mixtures were incubated for 2 or 3 h at 37°C, stopped using LDS sample buffer (Novex), separated on Bis-tris 4–12% SDS-PAGE gels (Novex), and stained with Coomassie Blue. The protein bands were scanned and quantified by Quantity One software (Biorad).

Hydrodynamics

Analytical gel filtration chromatography used an analytical TSK-GEL G3000SW column (Tosoh) at 22°C. The column and samples were equilibrated with gel filtration buffer listed in SI. 50 μM proMMP-7 was mixed with Hep dp8 at 20-fold molar excess or Hep dp16 at 4-fold molar excess and filtered through a 0.22 μm centrifugal filter (Millipore). 80 μL of each mixture was injected, eluted at 0.8 mL/min, and 1 mL fractions collected.

Rotational correlation times τ_C were measured with a Bruker 800 MHz Avance III NMR system from ^{15}N relaxation (see SI), at each point of titrations of proMMP-7-E195A (300 μM) with heparin oligosaccharides.

Atomic force microscopy

ProMMP-7-E195A and truncated forms were prepared at 50 or 100 nM in imaging buffer (10 mM CaCl_2 , pH 7.5; see SI). Mixtures with Hep dp8 or dp16, heparin, HS, or HS dp8 were examined. Each solution was deposited immediately after mixing on a freshly cleaved muscovite mica surface (V-1 grade, Spi Supplies) and incubated for ~5 min. The surface was then rinsed three times with ~100 μL of imaging buffer. AFM images were acquired in imaging buffer in tapping mode using a commercial instrument (Cypher, Asylum Research). SNL tips (sharp nitride levers, SNL-10, Bruker) with measured spring constants of ~0.2 N/m were used. Images were recorded at ~30°C with an estimated tip-sample force < 100 pN, deduced by comparing the free space tapping amplitude (~5 nm) to the imaging set point amplitude. Under such conditions, minimal protein distortion is expected. Image analysis is detailed in SI.

Kinetics of proteolysis

Digestion of soluble Knight's peptide⁴⁰ (FS-6; EMD) was monitored by FRET with excitation at 328 nm and emission at 393 nm. Progress curves of the digestions at 25°C in proteolysis buffer (see SI) were monitored with a BioTek Synergy MX plate reader. k_{cat}/K_m , K_m and k_{cat} were obtained by fitting of a few progress curves as described.⁴¹

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

AFM	atomic force microscopy
CAD	catalytic domain
CSA or C4S	chondroitin-4-sulfate
CSE	chondroitin sulfate E (chondroitin-4,6-sulfate)
dp	degree of polymerization
DS	dermatan sulfate
DLS	dynamic light scattering
FRET	Förster resonance energy transfer
GAG	glycosaminoglycan
HB-EGF	heparin-binding epidermal growth factor
Hep	heparin
HS	heparan sulfate
MMP	matrix metalloproteinase
serpin	serine protease inhibitor
SPR	surface plasmon resonance
wt	wild-type

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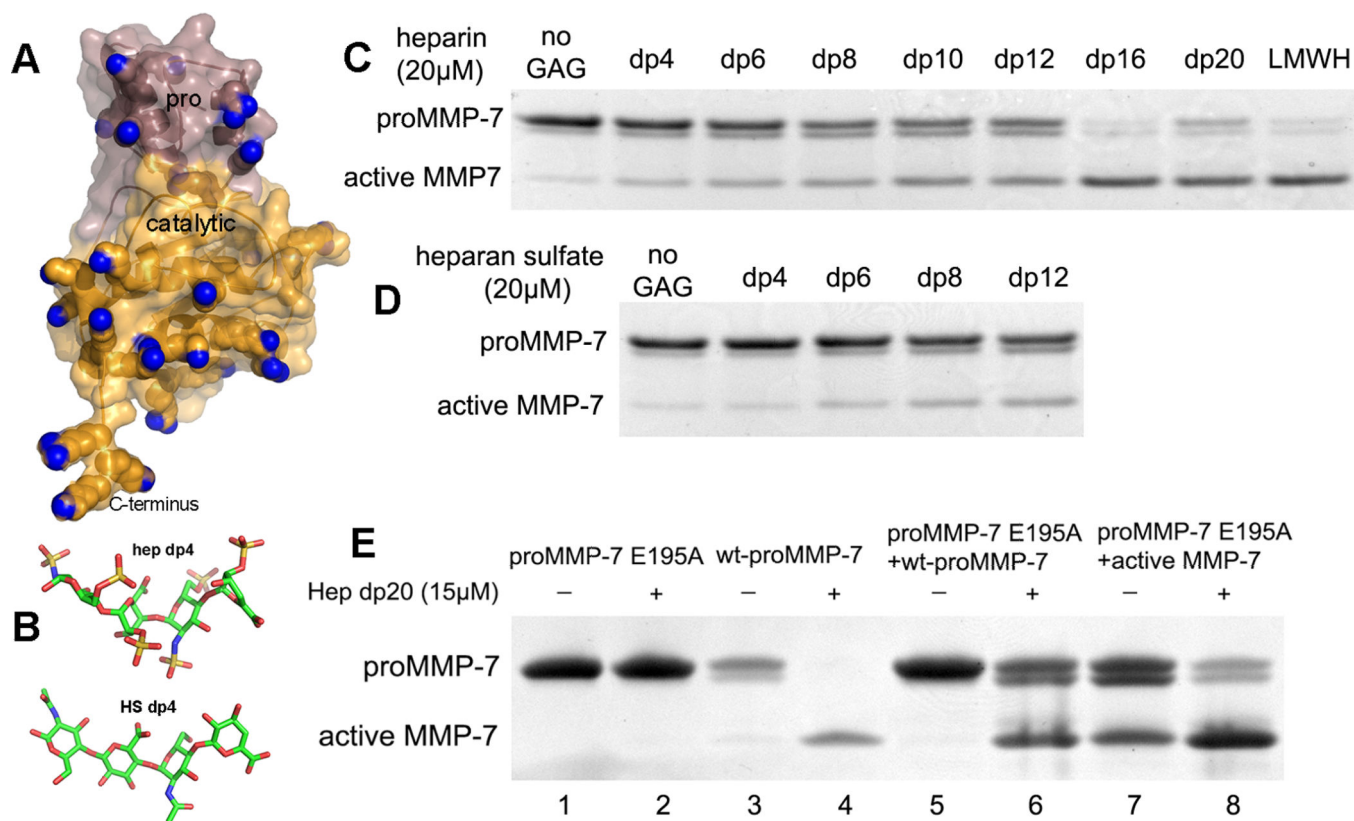


Figure 1. GAGs mediate intermolecular activation of proMMP-7

(A) The positive charges of arginine and lysines are colored blue on the domains in the homology model of proMMP-7.⁴³ (B) Crystallographic coordinates are plotted for tetrasaccharides from heparin and from HS (PDB codes 1BFB, 3E7J). (C, D) Heparin and HS oligosaccharides of defined length accelerate proMMP-7 activation in a length-dependent manner. Wild-type proMMP-7 (5.5 μ M, 27.5 kDa) was incubated with 20 μ M (C) heparin oligosaccharides from four to 20 saccharides units long (dp4 to dp20), low molecular weight heparin (LMWH) (C), or (D) HS oligosaccharides four to 12 saccharide units long at 37°C for 2 h before stopping reactions. (E) In the presence of heparin fragments, proMMP-7 is activated through intermolecular proteolysis. ProMMP-7-E195A (7.3 μ M) was incubated with wt proMMP-7 (1.8 μ M, lanes 5, 6) or active wt-MMP-7 (2.6 μ M, lanes 7, 8) with or without Hep dp20 (15 μ M, *even* lanes) at 37°C for 3 h before stopping the reactions.

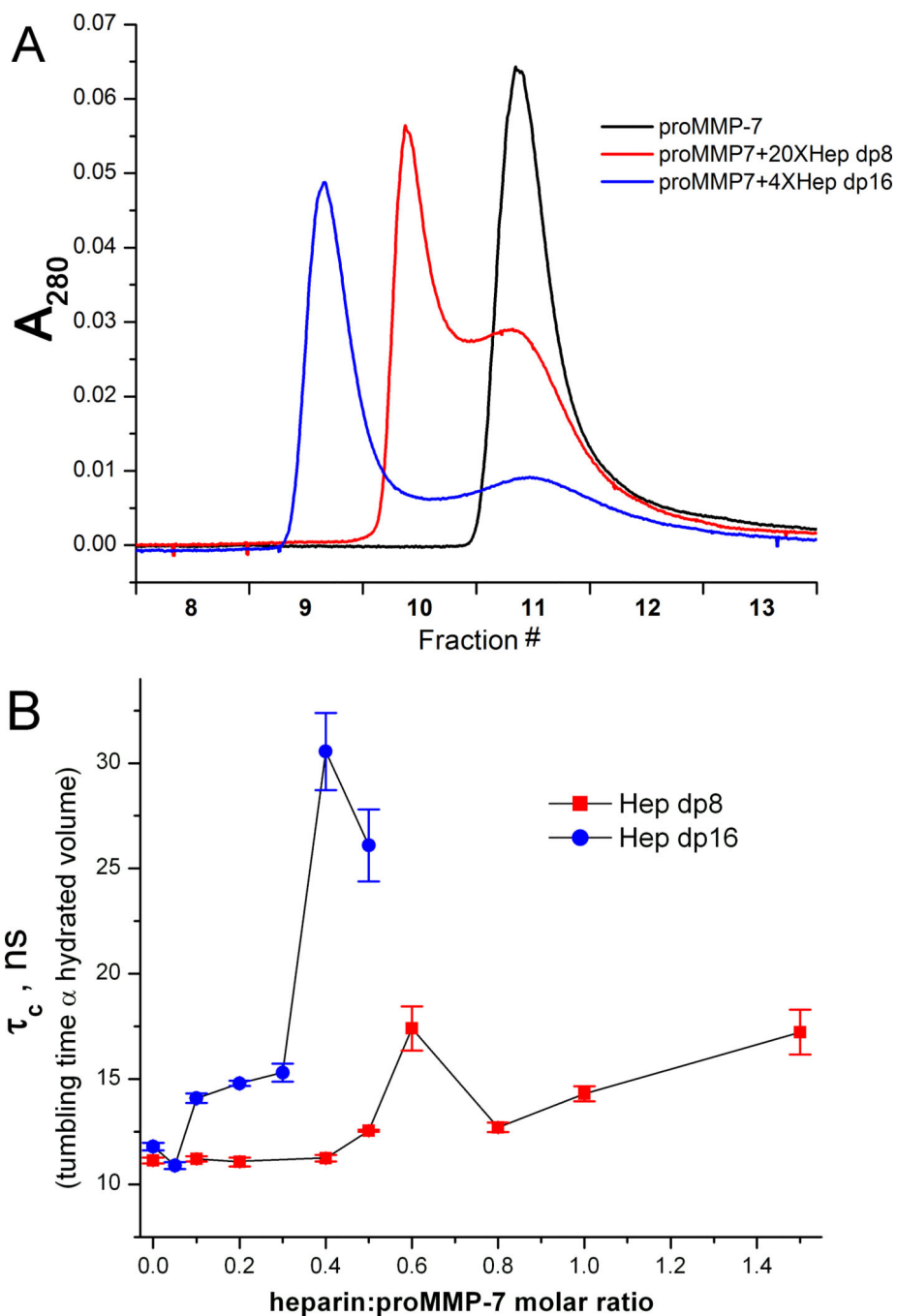


Figure 2. ProMMP-7 and heparin oligosaccharides form complexes containing two or more proenzyme molecules

(A) ProMMP-7-E195A (50 μ M) and its mixtures with Hep dp8 or dp16 oligosaccharides (at molar excesses indicated) were run on an analytical gel permeation column. (B) NMR-detected titrations of proMMP-7-E195A (15 N-labeled, 300 μ M) with Hep dp8 or dp16 were monitored with a proxy for apparent hydrodynamic volume, the rotational correlation time (τ_c) from 15 N NMR relaxation.

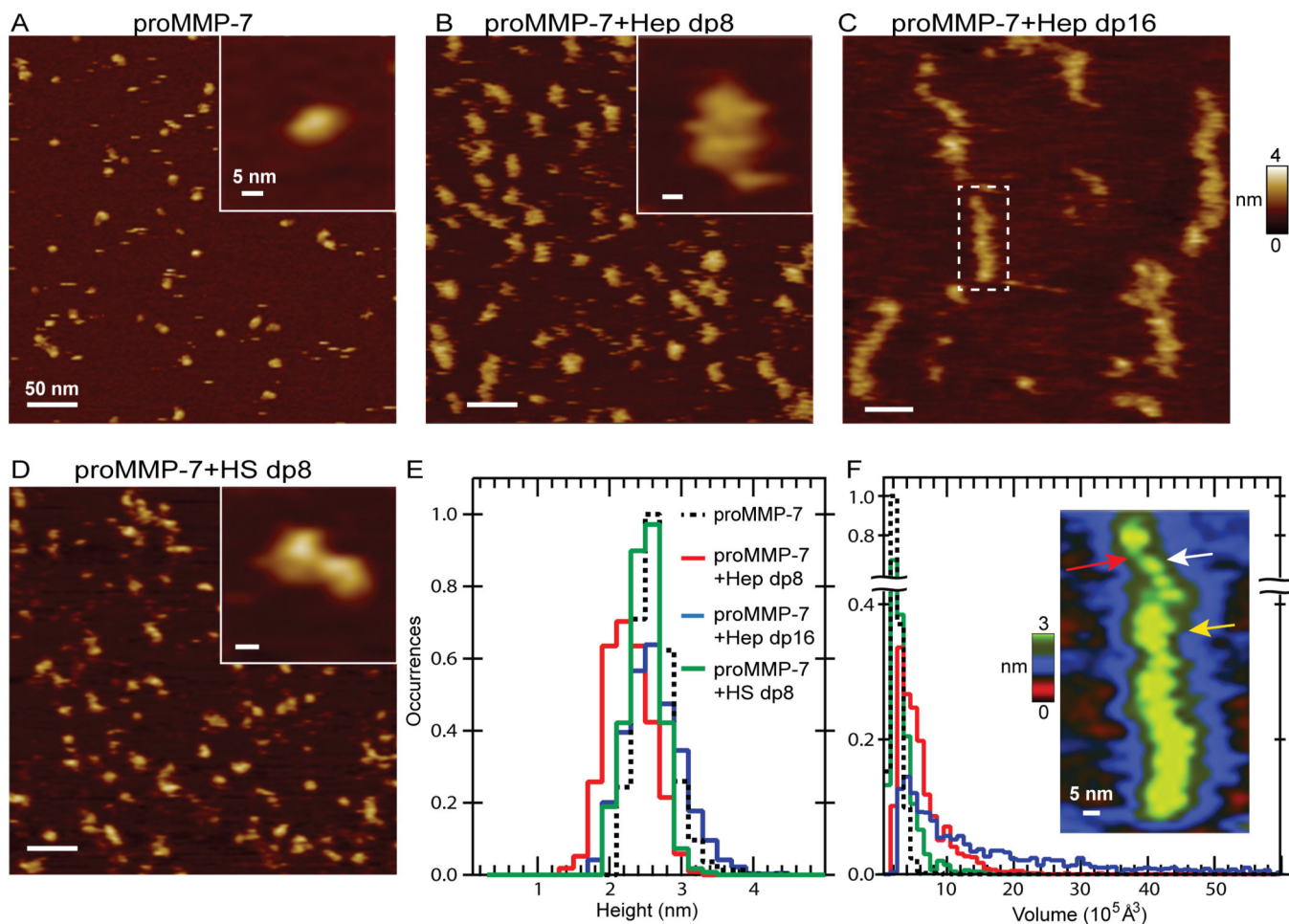


Figure 3. Atomic force microscopy (AFM) reveals tethering of proMMP-7 molecules by heparin oligosaccharides that becomes extensive with lengths of 16 monosaccharide units
 400×400 nm AFM images are shown for (A) proMMP7-E195A (100 nM) only or (B) 50 nM zymogen with 100 nM Hep dp8, (C) 100 nM zymogen with 100 nM Hep dp16, or (D) 50 nM zymogen with 100 nM HS dp8. Insets show enlargements. (E) The maximum heights of proMMP-7-E195A in the free state or with Hep dp8 or dp16 oligosaccharides present are tabulated as histograms with bins of 2 Å. In E and F, the black dashed line represents proMMP-7-E195A only ($N = 903$ features), red the mixture with Hep dp8 ($N = 980$), blue with Hep dp16 ($N = 1379$), and green with HS dp8 ($N = 826$ features). (F) Volumes of the same features are tallied into bins of $1 \times 10^5 \text{ \AA}^3$ in the histogram. The inset of (F) expands the boxed feature from panel C. The red arrow marks an apparent Hep dp16 chain. The white arrow marks a width of one enzyme while the yellow arrow marks a width of two enzymes.

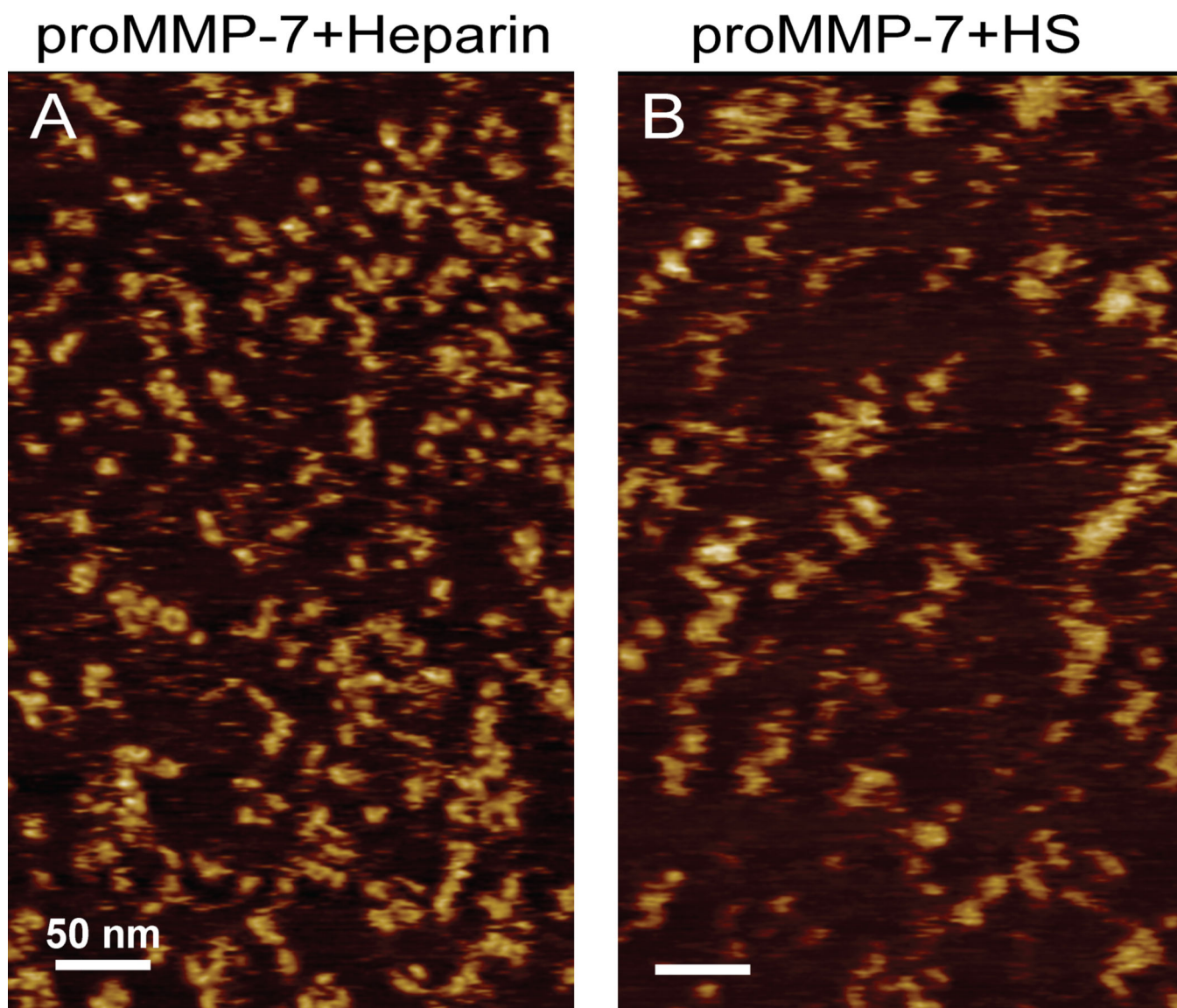


Figure 4. AFM images of proMMP-7-E195A mixtures with heparin (A) and heparan sulfate (B) suggest similarities in the heterogeneous tandem assemblies

300×500 nm images are plotted for mixtures of (A) 100 nM proMMP-7-E195A with 2.5 µg/ml bovine lung heparin or (B) 50 nM proMMP7-E195A with 2.5 µg/ml bovine heparan sulfate. Note that bovine heparin and HS are polydisperse, with chain lengths from 5 to 50 kDa. The vertical scale spans 0 to 4 nm for both panels.

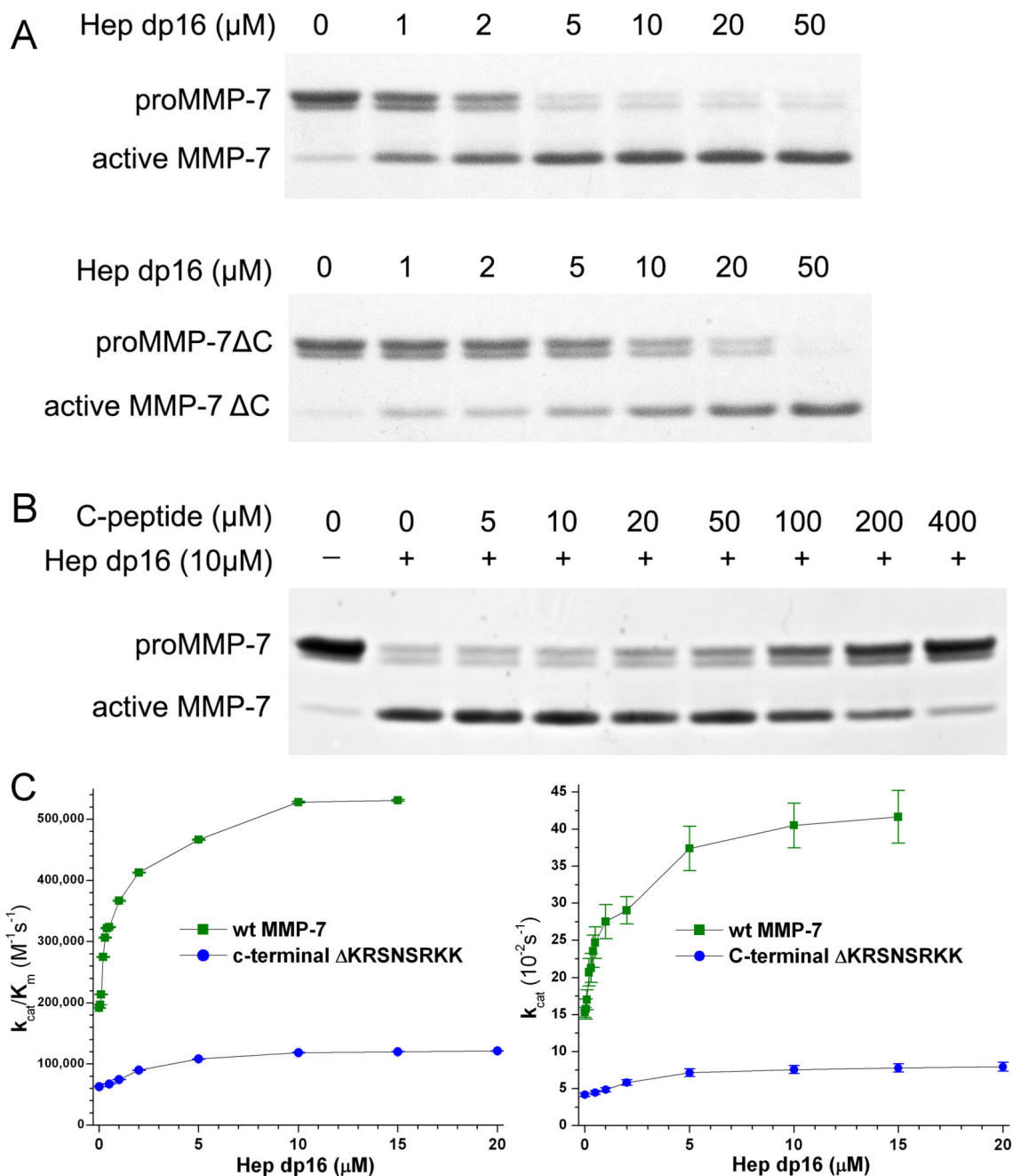


Figure 5. The basic C-terminus is a remote site of allosteric, activating binding by Hep dp16 oligosaccharides

(A) ProMMP-7 (5.5 μM) with wt sequence or deletion of KRSNSRKK was incubated with increasing concentrations of Hep dp16 at 37 $^{\circ}\text{C}$ for 2 h until each reaction was stopped. (B) The synthetic peptide with the KRSNSRKK sequence from the C-terminus interferes in Hep dp16-induced activation of proMMP-7. (C) Hep dp16 effects on catalytic efficiency (k_{cat}/K_M) and proteolytic turnover (k_{cat}) of soluble Knight's peptide were monitored by FRET (25 $^{\circ}\text{C}$, pH 7.5) are compared between active wt-MMP-7 and MMP-7 C.

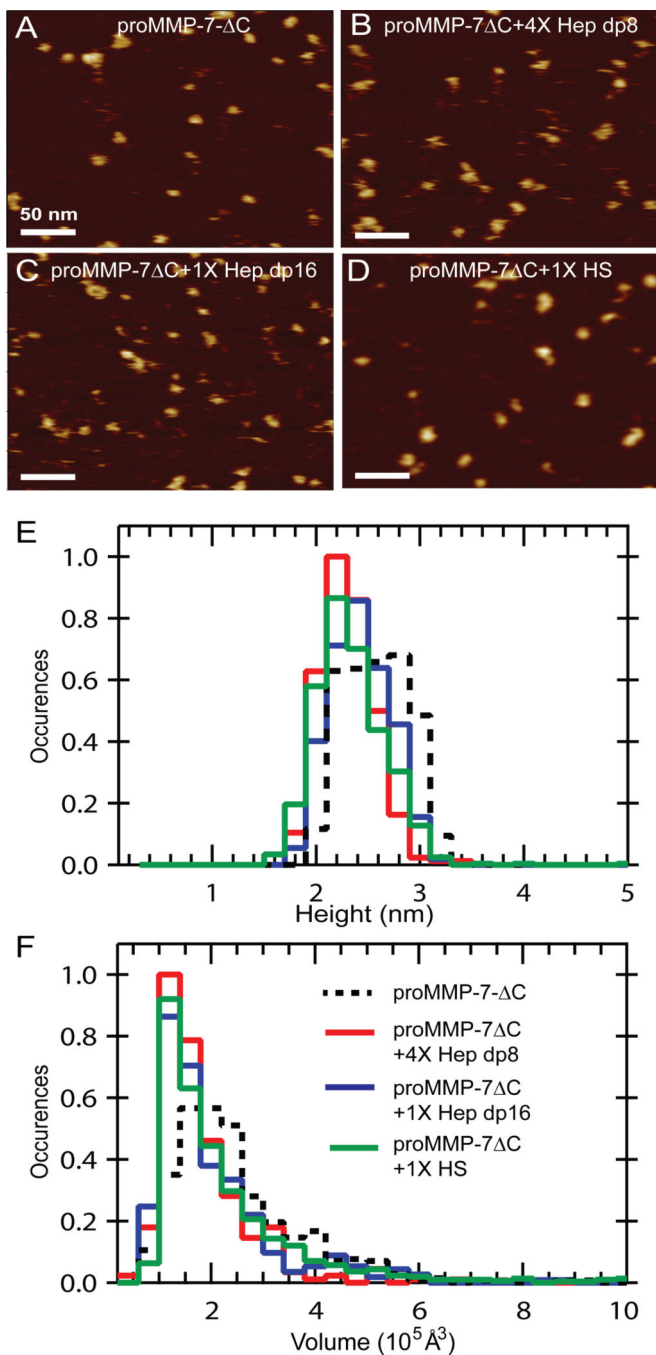


Figure 6. Deletion of the basic C-terminus abrogates formation of extended complexes in the presence of heparin oligosaccharides

400×400 nm AFM images are shown for (A) 50 nM proMMP-7 C alone, (B) 50 nM proMMP-7 C mixed with 200 nM Hep dp8, (C) 50 nM proMMP-7 C mixed with 50 nM Hep dp16, or (D) 100 nM proMMP-7 C mixed with 5 μg/ml HS. The number of features tabulated for the histograms are $N = 456$ for the free state, $N = 284$ for the mixture with Hep dp8, $N = 362$ for the 1:1 mixture with Hep dp16, and $N = 957$ for the mixture with HS.

(B) The height histograms use bins of 2 Å. (C) The volume histograms use bins of 0.4×10^5 Å³.

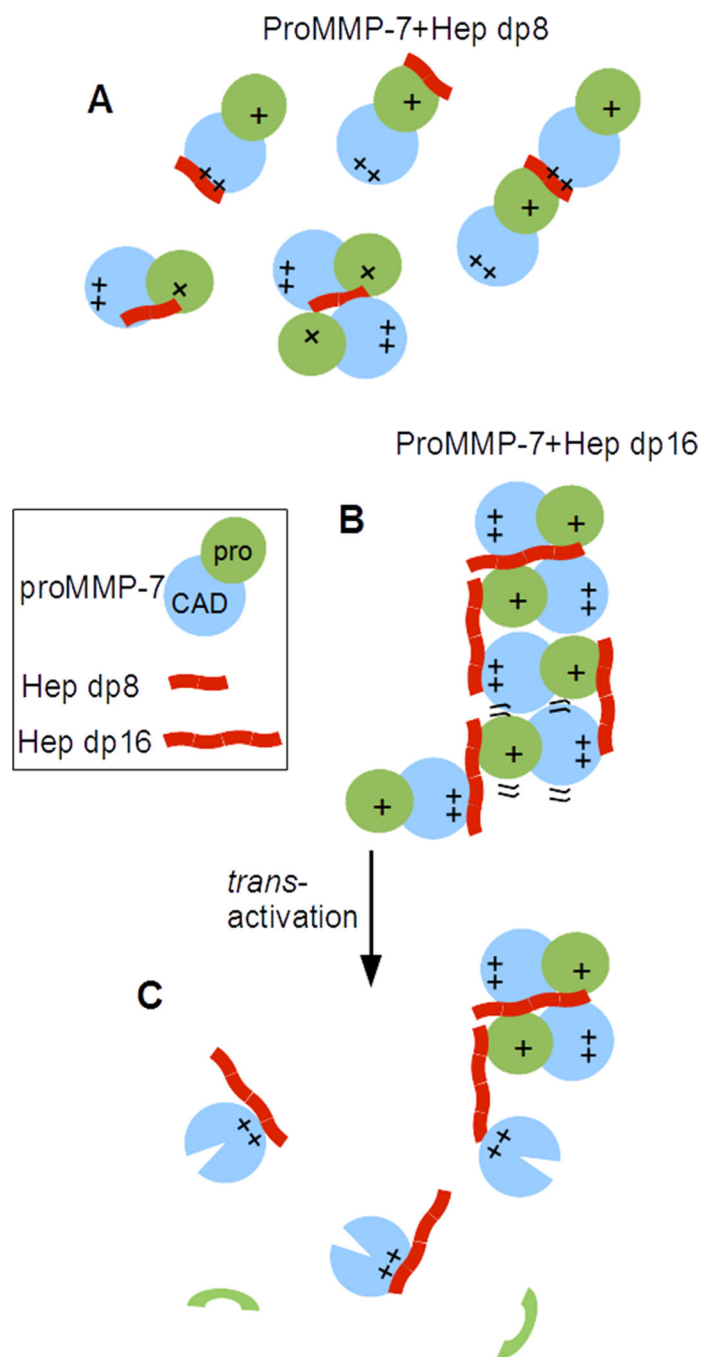


Figure 7. Speculative and simplified models for assembly of proMMP-7 induced by short and long heparin oligosaccharides

(A) 8-monosaccharide lengths (dp8) of heparin are likely to encourage formation of sandwich dimers, in addition to forming 1:1 complexes with proenzymes. (B) 16-monosaccharide lengths (dp16) of heparin trigger formation of extended tandem arrangements that promote proteolysis of neighbors. (C) Removal of the prodomain renders the mature MMP-7 unlikely to remain in the arrays.