Extracellular Histones Inhibit Fibrinolysis through Noncovalent and Covalent Interactions with Fibrin

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Abstract
Histones released into circulation as neutrophil extracellular traps are causally implicated in the pathogenesis of arterial, venous, and microvascular thrombosis by promoting coagulation and enhancing clot stability. Histones induce structural changes in fibrin rendering it stronger and resistant to fibrinolysis. The current study extends these observations by defining the antifibrinolytic mechanisms of histones in purified, plasma, and whole blood systems. Although histones stimulated plasminogen activation in solution, they inhibited plasmin as competitive substrates. Protection of fibrin from plasmin digestion is enhanced by covalent incorporation of histones into fibrin, catalyzed by activated transglutaminase, coagulation factor FXIII (FXIIIa). All histone subtypes (H1, H2A, H2B, H3, and H4) were crosslinked to fibrin. A distinct, noncovalent mechanism explains histone-accelerated lateral aggregation of fibrin protofibrils, resulting in thicker fibers with higher mass-to-length ratios and in turn hampered fibrinolysis. However, histones were less effective at delaying fibrinolysis in the absence of FXIIIa activity. Therapeutic doses of low-molecular-weight heparin (LMWH) prevented covalent but not noncovalent histone–fibrin interactions and neutralized the effects of histones on fibrinolysis. This suggests an additional antithrombotic mechanism for LMWH beyond anticoagulation. In conclusion, for the first time we report that histones are crosslinked to fibrin by FXIIIa and promote fibrinolytic resistance which can be overcome by FXIIIa inhibitors and histone-binding heparinoids. These findings provide a rationale for targeting the FXIII–histone–fibrin axis to destabilize fibrin and prevent potentially thrombotic fibrin networks.

Keywords
► fibrin(ogen)
► fibrinolysis
► histones
► neutrophil extracellular traps
► FXIII

Introduction
Histones are released from damaged, dying, or activated cells during infection, inflammation, and trauma. A major source of extracellular histones are neutrophils, which eject webs of decondensed chromatin as neutrophil extracellular traps (NETs). NETs trap and kill microbial pathogens, either directly or by acting as a scaffold for the localized activation of coagulation, a process termed “immunothrombosis.” However, NETs have been described as a double-edged sword, as their overproduction or inadequate clearance can be harmful to the host and contribute to pathological microvascular thrombosis in sepsis and disseminated intravascular coagulation. NETs are also implicated in the development of large vessel occlusions, and their presence in circulation is associated with poor prognosis in cardiovascular and cerebrovascular diseases. The mechanisms by which NETs promote thrombosis are unclear but are likely to be multifactorial and involve recruitment and activation of soluble clotting factors, platelets, and leukocytes. NETs have been identified in patient thrombi and...
intercalate into fibrin networks, making clots more stable and resistant to degradation. As the major protein component of NETs, histones are causally implicated in the formation of arterial and venous thrombi. However, most studies have focused on the procoagulant activities of histones, with fewer studies examining their antifibrinolytic mechanisms.

Fibrinolysis requires two key steps: the generation of plasmin from the inactive zymogen plasminogen, catalyzed by tissue type-plasminogen activator (tPA), followed by the subsequent digestion of fibrin by plasmin. These events are influenced by the fibrin structure and network architecture, which is in turn determined by the conditions at the time of fibrin formation. In blood, fibrinogen circulates in an environment rich in macromolecules and noncovalently binds a multitude of plasma proteins and polymers which can influence fibrin polymerization, structure, and susceptibility to fibrinolysis. While it is generally accepted that thicker fibrin fibers lyse more quickly than thinner fibers on a macroscopic scale, the relationship between fibrin structure and fibrinolysis is complex and varies depending on the fibrinogen-interacting protein or polymer and their electrostatic charge. Negatively charged DNA promotes the formation of densely packed networks of thick fibers less susceptible to plasmin digestion, whereas polyphosphate, another anionic polymer, gives rise to heterogeneous clot structure and attenuated plasmin generation. It has been known for some time that histones, which are rich in positively charged lysine and arginine residues, interact with fibrinogen and increase resistance to fibrinolysis. However, the exact mechanisms behind these effects have not been fully elucidated. Histones interfere with the lateral organization of protofibrils, resulting in a fibrin network that is more resistant to tPA-mediated fibrinolysis, despite being composed of thicker fibers. It has been suggested that histone-induced structural alterations in fibrin mediate lytic resistance, as histones neither inhibited the caseinolytic activity of plasmin nor acted as plasmin substrates. Protamine, another strongly cationic protein, also induces thicker fibrin fibers, but with increased susceptibility to tPA lysis. These observations suggest that structural alterations in fibrin do not fully explain the antifibrinolytic effects of histones.

Another important determinant of the mechanical and biochemical stability of fibrin is factor XIII (FXIII) which is bound to fibrinogen in circulation. FXIII is activated by thrombin in the presence of calcium ions to produce the active transglutaminase, FXIIIa. FXIIIa introduces covalent isopeptide bonds (or crosslinks) between glutamine and lysine residues in the α and γ chains of fibrin monomers within individual fibers. This creates γ – γ dimers and high molecular weight α – α and γ – α polymers, which increases clot stiffness and mediates red blood cell retention in clots. FXIIIa also covalently crosslinks other plasma proteins to fibrin, including antifibrinolytic molecules such as α2-antiplasmin, plasminogen activator inhibitor 2 (PAI-2), and thrombin activatable fibrinolysis inhibitor (TAFI). This allows FXIIIa to inhibit fibrinolysis, which is particularly apparent under flow and clot contraction conditions. Together, these activities make FXIIIa a critical determinant of fibrin network stability and thrombus composition and size, and a target in thrombotic disease.

We previously found that histones colocalize with fibrin in patient thrombi and make clots more resistant to fibrinolysis. In the current study, we investigated the mechanisms behind these observations and show that histones competitively inhibit plasmin to delay fibrinolysis. Furthermore, the antifibrinolytic effects of histones are enhanced by covalent crosslinking to fibrin, catalyzed by FXIIIa. Our results suggest blocking histone–fibrin crosslinking with FXIIIa inhibitors or histone-binding heparinoids is an effective strategy to destabilize clots containing histones. These findings are relevant for the prevention of thrombotic disease associated with histone release.

**Methods**

**Reagents**

FXIII (code 02/170), tPA (code 98/714), and freeze-dried control plasma (code 06/158) were from NIBSC (South Mimms, United Kingdom). Thrombin was from Diagnostic Reagents Ltd. (Thame, United Kingdom). Antifibrinogen antibodies (A0080) were from Dako (Glostrup, Denmark). Anti-histone H3 (ab1791) and anticitrullinated histone H3 (ab5103) were from Abcam (Cambridge, United Kingdom). Anti-histone antibodies (A00186) were from Genescript (New Jersey, United States). Mixed histones (calf thymus, IIA) and recombinant peptidyl arginine deiminase 4 (PAD4) were from Sigma (Poole, United Kingdom). Individual recombinant human histones expressed with an N-terminal His6-tag were from Amsbio (Oxford, United Kingdom). Human fibrinogen (plasminogen-depleted) was from Merck (Watford, United Kingdom). FXIII-depleted plasma and glu-plasminogen were from Hyphen Biomed (Neuville-sur-Oise, France). T101 was from Zedira (Darmstadt, Germany) and prepared in HEPES-buffered saline (HBS; 10 mM HEPES pH 7.4, 150 mM NaCl). Chromogenic substrates for plasmin (S-2251) and tPA (S-2288) were from Chromogenix (Milan, Italy). Low-molecular-weight heparin (LMWH; Dalteparin/Fragmin) was from Pfizer (Kent, United Kingdom).

**Solution Chromogenic Assays**

For studying plasmin/tPA inhibition by histones, rates of hydrolysis of chromogenic substrates (S-2251 or S-2288 for plasmin and tPA, respectively) were obtained over a range of substrate and histone concentrations using online apps. Kinetic parameters were obtained by fitting plots of plasmin velocity as a function of S-2251 concentration to Michaelis–Menten kinetics in GraphPad Prism (GraphPad software version 8.1.1, La Jolla, California, United States). Data were also plotted as double-reciprocal (Lineweaver–Burk) plots to identify inhibitory mechanisms. For solution plasminogen activation assays, histones were incubated with glu-plasminogen (250 nM) and tPA (10 nM) in reaction buffer (10 mM Tris pH 7.7, 100 mM NaCl, 0.01% tween-20), before addition of S-2251 (0.6 mM, final concentrations). Rates of plasminogen activation were measured by hydrolysis of S-2251 and calculated from plots of absorbance versus time squared using online apps.
Crosslinking Analysis

For crosslinking analysis in purified systems, fibrinogen (8.5 μM) was mixed with mixed histones and clotted for 2 hours at 37°C with thrombin (5 nM) in HBS + 5 mM CaCl₂. Insoluble fibrin was isolated by centrifugation (7,000 g/10 minutes) and resuspended in clot solubilization buffer (0.2 mM Tris-HCl pH 8, 8 M urea, 4% SDS, 40 mM DTT) for 1 hour at 37°C and analyzed by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and Coomassie staining or western blotting. Time-course analysis of crosslinking, reactions were stopped at the indicated time points with clot solubilization buffer and analyzed as above.

Crosslinking assays with individual recombinant histones expressed with N-terminal polyhistidine (His₆) tags were performed as above and allowed all histone subtypes to be identified with a single anti-hiss₆ tag antibody. Citrullinated histones were prepared by incubating mixed histones with recombinant PAD4 as previously described.¹⁵ and used in crosslinking reactions as above.

For crosslinking analysis in plasma, sixfold diluted plasma (normal or FXIII-deficient) was clotted (7.5 mM CaCl₂, 2 nM thrombin, final concentrations) in the presence of histones or T101. After 2 hours, clots were dissolved in solubilization buffer (60°C, 1 hour) and analyzed by western blot. For FXIII-reconstituted plasma, purified FXIII was added to FXIII-deficient plasma prior to clotting.

Whole blood was clotted (11.25 mM CaCl₂, 5 nM thrombin, final concentrations) in the presence of histones and/or T101 (200 μM) for 2 hours at room temperature. Clots were collected by centrifugation (300 g/15 minutes) and homogenized in cell lysis buffer (10 mM HEPES pH 7.4, 150 mM NaCl, 1 mM EDTA, 1% Triton-X 100) using a Dounce homogenizer. Insoluble fibrin was isolated by centrifugation (7,000 g, 15 minutes, 4°C) and the homogenization process repeated twice. The fibrin pellet was dissolved in clot solubilization buffer for 2 hours at 60°C and analyzed by SDS-PAGE and Coomassie staining or western blotting.

Turbidimetric Analysis of Fibrin Clotting and Structure

Fibrinogen (8.5 μM) was clotted with thrombin (5 nM) in the presence of increasing histones with or without CaCl₂ (5 mM) and/or T101 (200 μM). Clotting was monitored for 2 hours in a Spectramax M5 plate reader (Molecular Devices, Berkshire, United Kingdom) at 405 nm at 37°C. Times to 50% clotting were determined using online apps (examples are provided in - Supplementary Fig. S4A, available in the online version). Optical density values (τₕ) were transformed to turbidity values using τ = 1 - e⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻ Sağlık = (A – A₀)/A_max, where A_max is the maximal and A₀ the final absorbance after complete lysis). In parallel plates, clots were dissolved in clot solubilization buffer at different time points following application of plasmin. Samples were analyzed by SDS-PAGE and Coomassie staining and band intensities quantitated using ImageJ software (ImageJ, National Institute of Health, United States).

Statistical Methods

Means and standard deviations were calculated using GraphPad Prism. The effects of histones on clotting time, lateral aggregation, and fibrin mass/length were analyzed by one-way ANOVA (analysis of variance) with Dunnett’s or Tukey’s multiple comparison testing in GraphPad Prism.

Results

Histones Inhibit Plasmin to Delay Fibrinolysis

The effect of mixed histones, consisting of the four core histone subtypes (H2A, H2B, H3, and H4) plus the H1 linker histone, on the activity of plasmin and tPA is shown in - Fig. 1A. Plasmin hydrolysis of the chromogenic substrate S-2251 was inhibited by histones, with an IC₅₀ of 11.5 μg/mL. By contrast, histones had little effect on the activity of tPA on its substrate S-2288 (-Fig. 1A). Inhibition of plasmin was competitive; mixed histones increased the K_m of plasmin for S-2251 up to sevenfold, with little effect on the V_max (-Fig. 1B). Completed
Fig. 1 Histones inhibit plasmin to delay fibrinolysis in a purified system. (A) tPA or plasmin (5 nM) was incubated with S-2288 or S-2251, respectively, and the indicated concentration of mixed histones. Initial rates were calculated from plots of absorbance (405 nm) versus time and expressed relative to the condition without added histone. Points represent means ± SEM from duplicate wells. (B) Plasmin activity was measured with increasing amounts of S-2251 in the presence of the indicated histones. The inset shows a Lineweaver–Burk plot to illustrate competitive inhibition of plasmin by histones. (C) Completed reactions from (A) for 240 μg/mL histones were analyzed by SDS-PAGE and Coomassie staining. H1 histone, core complex (H2A, H2B, H3, and H4), and digested histones are annotated. The control sample represents histones without tPA or plasmin incubation. (D) Plasmin was added to the surface of fibrin clots prepared by mixing fibrinogen, thrombin, and Ca²⁺ with the indicated concentration of histone. Fibrinolysis was monitored by absorbance (405 nm) and turbidity values normalized as described in the Materials and Methods section. Lysis curves represent means of three experiments. (E) Fibrin clots with or without histones (240 μg/mL) were incubated with plasmin and solubilized at the indicated time points for reducing SDS-PAGE and Coomassie staining. The positions of fibrin(ogen) α, β, and γ chains and crosslinked variants (γ – γ dimer and α(γ)-polymers), together with digestion products of γ – γ dimer (γ′–γ′) and β chain (β′), are annotated. Bands corresponding to the β chain (F), γ – γ dimer (G), and α(γ) polymers (H) were quantitated and expressed relative to their starting intensities. Data represent mean ± SEM (n = 3). ***p < 0.001 (I). Rates of plasminogen activation by tPA in the presence of increasing histones were measured by hydrolysis of S-2251 and calculated from plots of absorbance versus time squared. Points shown are means ± SEM from duplicate wells. (J) Internal clot lysis assays catalyzed by tPA were performed by clotting purified fibrinogen with thrombin in the presence of increasing histones, Ca²⁺, plasminogen, and tPA. Inhibition of fibrinolysis by histones is expressed as extension to 50% lysis times. Errors bars are ± SEM from duplicate wells. SEM, standard error of mean; tPA, tissue type-plasminogen activator.
Histones Are Crosslinked to Fibrin by FXIIIa

Histones are rich in lysine residues which could potentially serve as amine donors in transglutaminase reactions. This prompted us to investigate whether histones could be covalently incorporated into fibrin by FXIIIa.

Commercial fibrinogen, which has contaminating FXIII activity (see Supplementary Fig. S1, available in the online version), was clotted with thrombin in the presence of increasing amounts of mixed histone (30–120 µg/mL) under crosslinking (Ca\(^{2+}\)) or noncrosslinking conditions (Ca\(^{2+}\) or +T101, a FXIIIa inhibitor). Fibrin was isolated from the clot milieu by centrifugation and solubilized for analysis. Noncovalently and covalently bound histones were detected by western blot with antibodies specific for histone H3, which we used as a representative member of the core histone complex previously identified in patient thrombi.

Histone H3 co-purified as a noncovalently bound monomer (~14 kDa) with noncrosslinked fibrin (~Ca\(^{2+}\) or +T101 in Fig. 2A). Under crosslinking conditions (~Ca\(^{2+}\)), which is required for dissociation of the inhibitory FXIII-B subunits from the catalytic FXIII-A subunits, the intensity of the histone H3 band was reduced and replaced by high molecular weight histone H3 immunoreactivity, beginning just above the position of the γ–γ dimer (~62 kDa). This indicated histone H3 had been crosslinked to fibrin, resulting in a heterogeneous mixture of histone–fibrin crosslinked polymers. High molecular weight histone–fibrin crosslinking was not apparent when Ca\(^{2+}\) was omitted or T101 was included, indicating the involvement of FXIIIa.

A time-course analysis of histone–fibrin and fibrin–fibrin crosslinking is shown in Fig. 2B. Distinct bands of histone H3 (~70 kDa and ~100 kDa) could be detected approximately 5 to 10 minutes after initiation of clotting, which roughly correlates to the molecular masses of the α-chain of fibrin and crosslinked γ–γ dimer. Interestingly, higher molecular weight histone H3 immunoreactivity (>98 kDa) was detected only after γ–γ dimer formation was complete (120 minutes; Fig. 2B). It is possible that histone H3 immunoreactivity could originate, at least in part, from histones crosslinked to one another, which subsequently co-purified with fibrin. A band approximately 30 kDa in size was apparent in Fig. 2B, which is a potential crosslinked histone dimer. However, the distinctive banding pattern of histone H3 early in the time course suggests that histones are conjugated to crosslinked fibrin.

We extended our crosslinking analysis by testing individual recombinant histones. Results in Supplementary Fig. S2 (available in the online version) show that all histone subtypes (H1, H2A, H2B, H3, and H4) could be crosslinked to fibrin in a FXIIIa-dependent manner. Bands corresponding to histone dimers were apparent for histones H2A and H3, but these were also observed in the presence of T101, and so the exact nature of these histone oligomers and whether they are crosslinked forms require further study.

Histones in NETs may be citrullinated by PAD4 during their release from the cell, potentially altering histone interactions. To determine whether citrullinated histones could still bind fibrin, we repeated our crosslinking analysis with citrullinated histones, generated in vitro by incubating histones with recombinant PAD4. Similar to unmodified histones, citrullinated histone H3 co-purified with fibrin in noncovalently bound and high molecular weight crosslinked forms, with crosslinking being dependent on FXIIIa (Supplementary Fig. S3, available in the online version).

To show that histones could be crosslinked to fibrin in more complex and physiologically relevant environments, plasma was clotted in the presence of histones and T101. With clotting (i.e., recalcification) crosslinked fibrin (γ–γ dimers and high molecular weight polymers) was detected, but not in the presence of T101, demonstrating effective FXIIIa inhibition (Fig. 2C). High molecular weight histone H3 was again detected in clots, but not when FXIIIa was inhibited. Supporting FXIIIa involvement, fibrin–fibrin and fibrin–histone crosslinking was not detected in clots from FXIII-deficient plasma, but was restored by the addition of purified FXIII to normal plasma levels (Fig. 2D).

Analysis of fibrin isolated from whole blood clots, clotted in the presence of mixed histones, is presented in Fig. 2E. Inclusion of T101 resulted in loss of γ–γ dimers and α, polymers and the reappearance of α and γ chains, demonstrating effective FXIIIa inhibition in blood (Fig. 2E). The histone H3 western blot shows co-purification of histone H3 (~14 kDa) and high molecular weight histone H3 immunoreactivity, which was prevented by FXIIIa inhibition. Inhibition of crosslinking with T101 coincided with increased intensity of noncovalently bound histone H3 monomer, consistent with histones binding fibrin without being crosslinked.

Histones Accelerate Fibrin Polymerization and Enhance Clot Structure Independent of Crosslinking to Fibrin

Binding of histones to fibrin(ogen) alters fibrin polymerization and fibrin network structure. To determine whether these effects are dependent on histone–fibrin crosslinking, we clotted
purified fibrinogen with thrombin in the presence of histones, with or without Ca$^{2+}$ and/or FXIIIa inhibitor (T101). Raw clotting curves are presented in Fig. 3(A–C) illustrating that histones shortened clotting times (time to 50% maximum absorbance, Fig. 3D) and increased rates of lateral aggregation of protofibrils, defined by the slope of the turbidimetric curve (Fig. 3E). These effects were unaffected by the omission of Ca$^{2+}$ (Fig. 3B and red bars in Fig. 3D, E) or the presence of T101 (Fig. 3C and green bars in Fig. 3D, E).

Histones increased the maximum optical density of fibrin clots (Fig. 3A–C), which reflects the mass-to-length ratio of fibrin fibers. Mass-to-length measurements were calculated from turbidity (see the Materials and Methods section and Supplementary Fig. S4 [available in the online version]) and the results are presented in Fig. 3F. Histones increased fibrin fiber mass-to-length ratios in a dose-dependent manner, reaching a maximum of approximately 2.2-fold over the concentration range studied, indicative of thicker fibrin fibers,
Fig. 3 Histones accelerate fibrin polymerization and enhance clot structure independent of crosslinking to fibrin. (A–C) Purified fibrinogen was clotted by thrombin in the presence of the indicated amounts of histone in buffer containing Ca^{2+} (A), without Ca^{2+} (B), or containing T101 (200 μM) plus Ca^{2+} (C). Fibrin formation was monitored by absorbance (405 nm). Curves are average of triplicate measurements. (D) Time to 50% clotting (half-maximum absorbance), (E) rates of lateral aggregation of prototibrils, and (F) fibrin fiber mass-to-length ratios calculated from turbidity measurements from A–C (see the Materials and Methods section and Supplementary Fig. S4 [available in the online version] for details). Rates in (E) are relative to clots without Ca^{2+} and histones. Error bars in D–F represent 95% confidence intervals of the mean from three measurements. Statistical analysis was performed using one-way ANOVA and is relative to conditions without histones. *p < 0.05, **p < 0.001. ANOVA, analysis of variance.

The Antifibrinolytic Potency of Histones Is Enhanced by Crosslinking to Fibrin
Clot lysis profiles examining the effect of histones and FXIII on tPA-mediated fibrinolysis in normal, FXIII-deficient, and FXIII-reconstituted plasmas are shown in Fig. 4A–C, and the calculated extension to 50% clot lysis times in Fig. 4D. Consistent with our results in purified systems (Fig. 1J), histones delayed fibrinolysis and extended clot lysis times in normal plasma (Fig. 4A, D). However, histones were less effective at delaying fibrinolysis in FXIII-deficient plasma, causing only a modest extension to lysis times (Fig. 4B, D). Reconstitution of FXIII activity by the addition of purified FXIII to FXIII-deficient plasma restored the antifibrinolytic potency of histones to that measured in normal plasma (Fig. 4C, D).

The relationship between FXIII and histones was explored in whole blood using ROTEM, which measures viscoelastically clot strength over time. To simulate internal fibrinolysis, blood was spiked with tPA prior to clotting, which resulted in complete lysis after approximately 35 minutes (presented in the TEMograms in Fig. 4E). The importance of FXIIIa for clot strength and stability was apparent with inclusion of T101, which significantly reduced clot strength and lysis times (Fig. 4E–G). Addition of histones stabilized clots, and 60 and 120 μg/mL histones significantly delayed lysis (Fig. 4F). Histones were less effective at stabilizing clots in the presence of T101, but still delayed clot lysis, particularly at 120 μg/mL (Fig. 4F). Histones also caused small but significant increases in clot strength with or without FXIIIa inhibition (Fig. 4G).

Low-Molecular-Weight Heparin Prevents Histone–fibrin Crosslinking and Improves Lysis of Clots Containing Histones
Cationic histones have high affinity for heparin, which carries a strong negative charge. This led us to investigate whether LMWH could interfere with histone–fibrin crosslinking and their inhibitory effects on fibrinolysis.

Therapeutic doses of LMWH prevented histone–fibrin crosslinking in clots formed from purified fibrinogen, without affecting fibrin–fibrin crosslinking (Fig. 5A). LMWH at 0.5 IU/mL prevented crosslinking of 60 μg/mL histone and a higher dose of 1 IU/mL inhibited crosslinking of 120 μg/mL histone. Prevention of histone–fibrin crosslinking by LMWH coincided with increased noncovalently bound histone (~14 kDa) and LMWH-accelerated lysis of clots containing histones in a purified clot lysis system (Fig. 5B–D). Histones increased the maximum turbidity of the clot lysis curves (Fig. 5B, E), as previously noted (Fig. 3), but not in the presence of LMWH (Fig. 5C, E). On the contrary, histones...
reversed the increased clot turbidity caused by LMWH up to a concentration of 120 μg/mL, after which turbidity began to increase (Fig. 5E).

**Discussion**

A growing body of evidence implicates histones and NETs in the pathophysiology of thrombosis by promoting coagulation and inhibiting fibrinolysis.6–8 In this study, we show that histones stabilize fibrin through several mechanisms, summarized in Fig. 6. By acting as competitive plasmin substrates, histones protect fibrin from degradation, which is enhanced by covalent crosslinking of histones to fibrin, catalyzed by transglutaminase FXIIIa. Accordingly, fibrinolysis is enhanced by blocking histone–fibrin crosslinking with FXIIIa inhibitors (Fig. 4) or LMWH (Fig. 5). Noncovalently bound histones accelerate lateral aggregation of protofibrils, resulting in thicker fibrin fibers with higher mass-to-length ratios (Fig. 3). In the absence of FXIIIa activity, histones have a more modest effect on fibrinolysis but still enhance clot strength (Fig. 4). Together, these results show that by interacting noncovalently and covalently with fibrin, histones inhibit fibrinolysis and enhance clot stability.

Histone–fibrinogen interactions have been studied via several approaches including turbidimetric, proteomic, and biophysical methods.13–15,17 However, the exact histone–fibrin crosslinking inhibits fibrinolysis.
binding site(s) on fibrin(ogen) have yet to be determined. It is possible that electrostatic forces mediate the binding of positively charged histones to fibrinogen, which is predicted to contain uneven regions of positive and negative charges distributed on the D- and E-domains. When mixed, solutions of histones and fibrinogen form gels that can be studied turbidimetrically and we recently showed that fibrinogen gelation and accelerated clotting are dependent on histone positive charge. In the current work, citrullinated histones, which have reduced positive charge due to conversion of arginine to citrulline, retained their ability to bind fibrin (Supplementary Fig. S3, available in the online version). This suggests histone–fibrin interactions might be more than simply electrostatic, and hydrophobic interactions may contribute. We also found that histones increased the rate of fibrin protofibril aggregation and enhanced clot structure without being crosslinked to fibrin (Fig. 3). Consistent with this, previous electron microscopy and small-angle X-ray scattering experiments coupled with rheological studies showed that histones enhanced fibrin structure and mechanical stability in the absence of FXIIa activity. Taken together, our results suggest that noncovalent interactions between histones and fibrin(ogen) alter the structural and mechanical properties of fibrin.

Plasmin is a broad specificity serine protease with a preference for peptide bonds adjacent to arginine and lysine residues, making histones good candidate targets. We found that all histone subtypes could be digested by plasmin and competed with fibrinogen clotted with thrombin in the presence of plasminogen, tPA, Ca^{2+}, and the indicated histone without (B) or with 1 IU/mL LMWH (C). (D) Extension to 50% clot lysis times by histones calculated from clot lysis curves in B and C. (E) Maximum turbidity of clot lysis curves in B and C. Error bars in D and E represent ± SEM from duplicate measurements. SEM, standard error of mean.
plasminogen has been previously explored in the context of plasminogen's nonfibrinolytic activities in inflammation and cell migration. All histone subtypes bind plasminogen, with histones H1, H3, and H4 having $K_D$ values in the nanomolar range. Analogous to cell surface plasminogen receptors, it is possible that histones provide a surface for colocalization of reactants through histone–lysine plasminogen–kringle interactions. However, histones competitively inhibit plasmin to delay fibrinolysis, being digested in the process, and protect fibrin from degradation. Crosslinked histones are more effective at inhibiting fibrinolysis than noncovalently bound histones and increase the biochemical stability of fibrin. Blocking histone–fibrin crosslinking with histone-binding heparinoids, or FXIIIa inhibitors, improves lysis of clots containing histones. This suggests targeting the FXIII–histone–fibrin axis could be effective in destabilizing clots containing histones to prevent thrombosis. tPA, tissue type-plasminogen activator.

Although our results have been generated in vitro, they have implications for the pathophysiology and treatment of thrombotic disease. By promoting fibrinolytic resistance, histone–fibrin crosslinking protects fibrin from elimination, potentially contributing to a thrombotic tendency and resistance to pharmacological therapies. Histones are usually undetectable in circulation, but levels can rise as high as 230 μg/mL, for example following trauma, in line with the concentration range used in this study. In plasma from sepsis patients, histone H3 was measured at concentrations up to 60 μg/mL and total histone levels above 75 μg/mL were associated with poor outcome. Increased levels of circulating histones are also associated with poor prognosis in stroke, myocardial infarction, and venous thromboembolism, and have been identified in patient thrombi. However, it is unclear how circulating levels of histones in prothrombotic conditions correspond to levels found in patient thrombi.

A major source of extracellular histones are neutrophils, which release NETs in response to diverse infectious and noninfectious stimuli. However, other cell types release histones through NETosis-independent mechanisms including necrosis and apoptosis. Caution is required when studying NET components such as histones and DNA in isolation, which may not approximate to their activity in NETs. Isolated NETs have shown inconsistent ability to enhance thrombin generation in the absence of platelets, even though histones possess in vitro procoagulant activity. The ability of histones to activate coagulation may depend on whether they are in a free, octameric, or DNA-bound (nucleosome) state. It is also possible that NET-associated granular enzymes, such as elastase and myeloperoxidase, in addition to histone modifications such as proteolysis and citrullination, modify the interactions between NETs and the coagulation and fibrinolytic systems. Citrullinated histones are often used as "NET-specific" markers, and citrullinated histone H3 has been identified in arterial, venous, and...
microvascular thrombi in humans and animals.\textsuperscript{6,8} Although histone citrullination weakens the interaction with fibrinogen,\textsuperscript{15} our results show that it does not prevent binding or FXIIa-mediated crosslinking of histone H3 to fibrin (\textsuperscript{[online only]}. Supplementary Fig. S3).

Histones and DNA exert antifibrinolytic effects, either alone or in combination,\textsuperscript{11,17} but the extent to which histones remain bound to DNA in circulation is unclear. Endogenous DNases promote the degradation of NETs, which would release histones, and their inadequate clearance is associated with thrombosis in animal models.\textsuperscript{38} The presence of DNA and histones in patient thrombi is clinically relevant to clot stability, as the ex vivo lysis of clots retrieved from stroke patients is improved with a combination of DNase and tPA.\textsuperscript{39} However, DNase is not always effective as it only removes the DNA component of NETs, leaving behind histones which mediate tissue injury, initiate thrombosis, and stabilize clots.\textsuperscript{40–42} Our results emphasize that although fibrin is the primary target of thrombolytic therapy, other factors need to be considered for effective fibrinolysis. Fibrin formation and lysis are influenced by numerous fibrin (ogen)-binding proteins and polymers in addition to histones\textsuperscript{17,24} and DNA\textsuperscript{3,11} such as Von Willebrand Factor\textsuperscript{43} and polyphosphate.\textsuperscript{12} It is likely that fibrin exists in a complex and heterogeneous network in vivo providing multiple targets for novel therapies.

Targeting FXIIa to prevent thrombosis has a potential advantage over other anticoagulants in that thrombin generation and fibrin formation can still take place, allowing normal hemostasis but reducing potentially thrombotic fibrin networks.\textsuperscript{23} Recently developed small-molecule inhibitors of FXIIa have shown promise in animal models, reducing clot weight without prolonging bleeding in models of venous stasis.\textsuperscript{44} Our results show that histones can be added to the list of FXIIa substrates that inhibit fibrinolysis, which includes α2-antiplasmin, TAFI, and PAI-2.\textsuperscript{19} In these cases, crosslinking is proposed to localize antifibrinolytic activity to the fibrin surface and contribute to fibrin stability, and we find that crosslinked histones have a similar effect. It is possible that blocking histone–fibrin crosslinking is a potential antithrombotic benefit of FXIIa inhibition, which would be relevant for thrombotic disease characterized by histone release.

Heparinoids are the mainstay of anticoagulant treatment and suppress thrombus formation by inhibiting thrombin generation. Histones bind negatively charged heparin and its derivatives with high affinity,\textsuperscript{32} which may be of therapeutic benefit. Heparinoids neutralize the damaging effects of histones in a range of settings, including toxicity in sepsis,\textsuperscript{45,46} thrombosis,\textsuperscript{5} thrombocytopenia,\textsuperscript{47} and platelet activation.\textsuperscript{48} Heparinoids also bind and dismantle NETs, possibly by displacing DNA.\textsuperscript{5,17} LMWH accelerated lysis of clots containing histones (\textsuperscript{[Fig. 5B–D]}, which correlated with reduced histone–fibrin crosslinking and increased levels of noncovalently bound histones (\textsuperscript{[Fig. 5A]). This indicates that LMWH does not prevent binding of histones to fibrin, but rather the crosslinking reaction itself. Increased amounts of noncovalently bound histones were also observed when histone–fibrin crosslinking was prevented by FXIIa inhibition (\textsuperscript{[Fig. 2]). Presumably under these conditions, histones are bound to fibrin at their crosslinking sites but not covalently attached. However, unlike the situation with FXIIa inhibition (\textsuperscript{[Fig. 3}), noncovalently bound histones did not increase fibrin clot turbidity in the presence of LMWH (\textsuperscript{[Fig. 5}). The relationship between histones, LMWH, and clot structure is likely to be complex, as LMWH can directly alter fibrin structure, increase clot strength, and delay fibrinolysis.\textsuperscript{39} In our system, LMWH increased clot turbidity, which was reversed by histones (\textsuperscript{[Fig. 5E]), suggesting that LMWH and histones neutralize each other’s effects on fibrin structure. Profibrinolytic activities have previously been described for heparin and its derivatives in plasma systems, mainly through reduced TAFI activation, which usually attenuates fibrinolysis.\textsuperscript{50} Preventing histone–fibrin crosslinking and destabilizing clots containing histones are potentially novel antithrombotic and profibrinolytic mechanisms for LMWH beyond anticoagulation, which warrant further investigation.

**What is known about this topic?**

- Histones and DNA in the form of neutrophil extracellular traps (NETs) are released into the circulation as part of the innate immune response.
- While beneficial in the fight against infection, NETs contribute to thrombosis by promoting coagulation and stabilizing clots, making thrombi resistant to pharmacological thrombolysis.
- Histones enhance the mechanical and lytic stability of fibrin.

**What does this paper add?**

- Histones inhibit plasmin as competitive substrates.
- Histones are noncovalently and covalently associated with fibrin in purified, plasma, and whole blood model systems.
- Covalent crosslinking of histones to fibrin, catalyzed by activated transglutaminase FXIIa, enhances the ability of histones to inhibit fibrinolysis.
- FXIIa blockers or histone-binding heparinoids inhibit histone–fibrin crosslinking and improve fibrinolysis, suggesting that the FXIII–histone–fibrin axis can be targeted to destabilize clots to prevent the formation of potentially thrombotic fibrin networks.

**Conflict of Interest**

None declared.

**References**


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Histone–Fibrin Crosslinking Inhibits Fibrinolysis  Locke, Longstaff

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