

Thrombin-free polymerization leads to pure fibrin(ogen) materials with extended processing capacity

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Abstract

Fibrin is a key protein for various clinical applications such as tissue reconstruction. However, in contrast to type I collagen, fibrin shaping has so far faced major limitations related to the necessity to add thrombin enzyme to fibrinogen precursors to induce fibrin self-assembly. Here we report a thrombin-free gelation pathway of fibrinogen solutions by incubation at 37°C in mild acidic conditions. We unravel the biochemical mechanisms underlying the gelation process and draw comparison between fibrinogen and fibrin at both molecular and supramolecular levels in these conditions. The protocol enables to control the viscosity of fibrin(ogen) solutions, and to induce fibrin(ogen) gel formation by simple 37°C incubation, with a reinforcement effect at neutralization. It facilitates processing of fibrin(ogen) materials, for coating, molding and extrusion, and offers new possibilities such as 3D printing. This approach is further compatible with type I collagen processing and can provide advanced tissue engineering scaffolds with high bioactivity.

Introduction

Fibrin is a gold standard material in tissue engineering^{1,2}. It exhibits high bioactivity thanks to the direct adhesion sites for integrins^{3,4}, which impact cell migration, proliferation and fate^{5,6}. Fibrin also indirectly interacts with cells through binding to extracellular matrix proteins⁷, polysaccharides⁸ and growth factors⁹. It is widely used clinically, mainly as a hemostat, sealant and adhesive. *In vivo*, fibrin is a major actor of haemostasis¹⁰, which is induced by a complex reaction cascade initiated by the adhesion of platelets at the site of injury^{11,12}. A key step is the thrombin mediated conversion of circulating fibrinogen into fibrin, which subsequently polymerizes to form a tight plug¹³.

Fibrinogen is a 340 kDa protein made of three pairs of chains, called α , β and γ that assemble in a specific symmetric shape, forming 3 aligned globules linked together by coiled coil connectors

constituted by the three different chains ¹⁴ (Fig.1.A.1). The central domain, called E, gathers the amino-terminal parts of the 6 chains, linked together by di-sulfide bonds. The distal domains, called D, are made of the carboxy-terminal ends of the β and γ chains. The carboxy-terminal ends of α chains, called α Cs, further extend in a random-coiled manner, with globular domains at their ends. Those domains interact together and with the E domain, and hence do not float freely. During clotting, thrombin cleaves two pairs of fibrinopeptides FpA and FpB from the central E nodule, (Fig.1.A.2 (0)), where only two out of four Fps are drawn for clarity purpose) ¹⁵. This cleavage reveals two sites A and B which bind to sites a and b, respectively located on the γ and β chains in the D nodule of an adjacent molecule. Such interactions promote the polymerization of fibrin monomers in a half staggered manner (Fig.1.A.2 (0')), resulting in gels (Fb_N) (Fig.1.A.3) with a highly fibrillar network (Fig.1.A.4). A certain delay, called clotting time, exists before the formation of a macroscopic clot ¹³. Lateral assembly of protofibrils only occurs at the end of the clotting time, which can last from seconds to hours depending on pH and ionic strength ¹⁶.

Difficulty to shape natural materials into 3D structures remains a major obstacle to mimic cell environment ¹⁷. The necessity to mix fibrinogen and thrombin to trigger self-assembly and the involved delay may account for the scarcity of work on fibrin shaping into 3D structures. Fibrin is mostly used in the form of gels that can be molded ¹⁸, or in addition to other materials ¹⁹. Some attempts have been carried out to produce fibrin microbeads ²⁰ or nanostructures ^{21,22}. So far, only one process has been reported for pure fibrin extrusion ²³, where fibrinogen and thrombin kept in different solutions are mixed together with a Y connector before extrusion. In sharp contrast, collagen, another gold standard material in tissue engineering, has been extensively studied ²⁴. The simplicity of collagen self-assembly, induced by neutralization or 37°C heat activation, makes it favorable for multiple shaping techniques, such as molding or extrusion ²⁵. To investigate if similar processes could be used to trigger fibrin polymerization and thus widen the possibilities in fibrin shaping, we thus focused on fibrinogen and fibrin solutions at acidic pH.

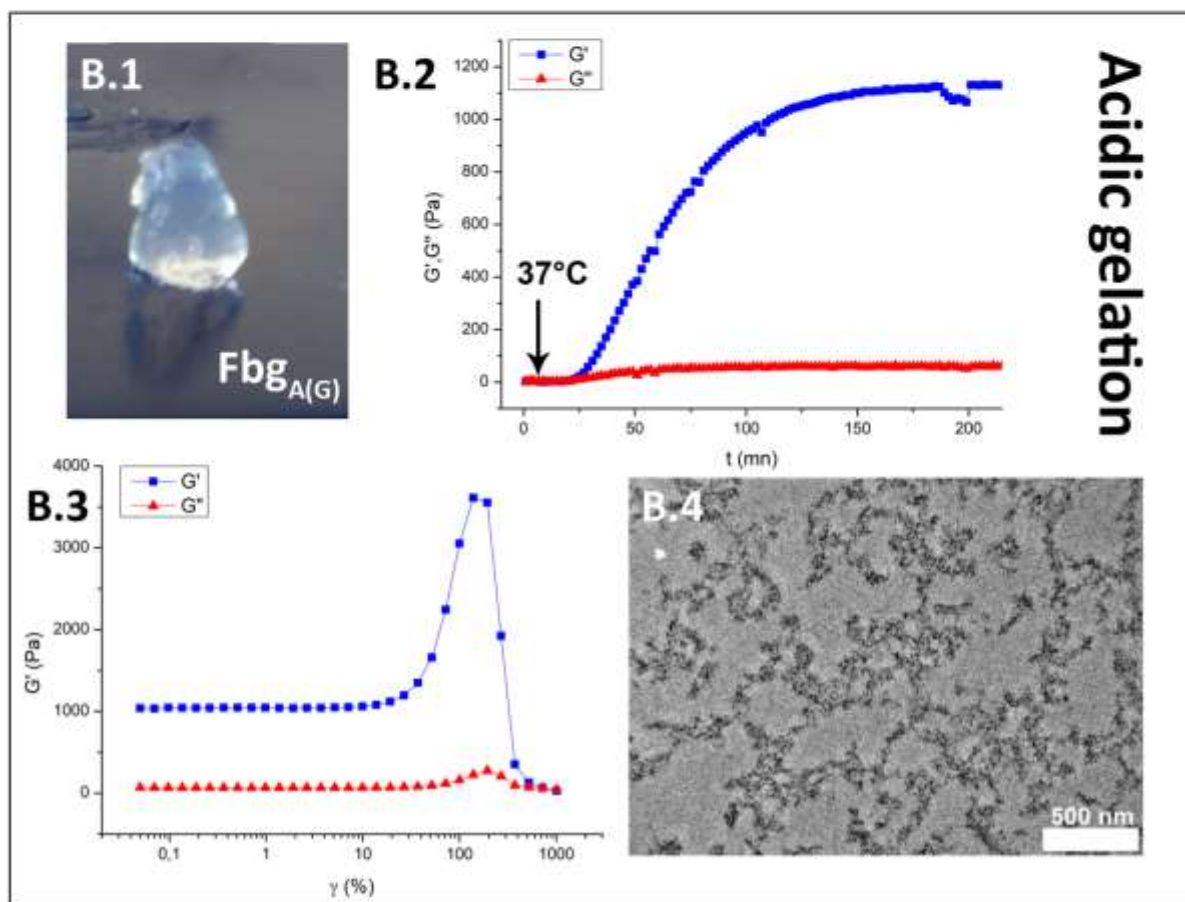
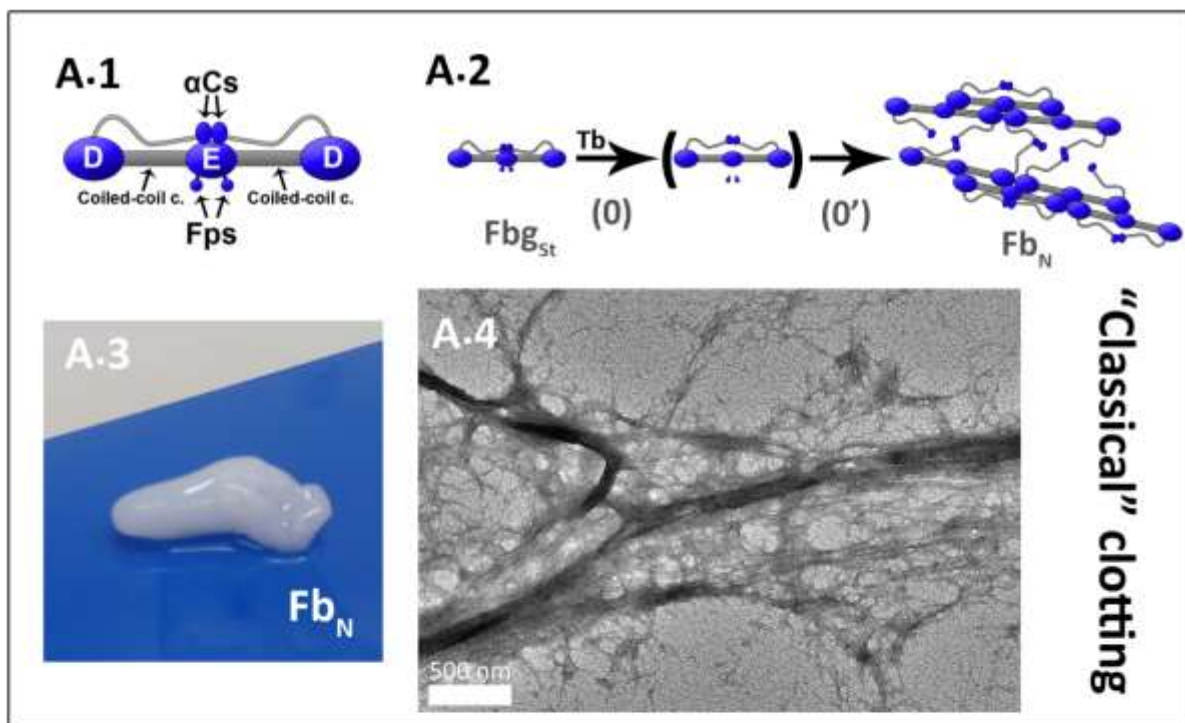


Fig. 1. Two gel formation mechanisms: “classical” neutral thrombin-driven (A) and acidic thrombin-free (B).

(A.1) Schematic structure of fibrinogen, made of two distal D domains and one central E domain, linked by coiled-coil connectors. The E domain bears the two pairs of fibrinopeptides (Fps) A and B. The two carboxy-terminal ends of the α -chains further extend and fold into small globular domains, noted α Cs, which connect to the central E domain. (A.2) “Classical” mechanism where the fibrinopeptides are cleaved (**0**) by thrombin (Tb), converting the fibrinogen proteins (Fbg_{St}) into fibrin monomers that self-assemble (**0'**) in a half-staggered manner to form a fibrin clot (Fb_{N}). (A.3) Image of a fibrin clot. (A.4) Fb_{N} observed by Transmission Electron Microscopy (TEM). (B.1) Image of an acidic gel ($\text{Fbg}_{\text{A(G)}}$) formed by incubation of a 40 mg/mL fibrinogen solution at pH 3.6 (Fbg_{A}) at 37°C. (B.2) Rheological dynamical analysis of gel formation of $\text{Fbg}_{\text{A(G)}}$ initiated by a temperature switch from 20°C to 37°C. (B.3) Mechanical behavior of $\text{Fbg}_{\text{A(G)}}$ under amplitude sweep. (B.4) Structure of an acidic gel $\text{Fbg}_{\text{A(G)}}$ included in araldite and observed by TEM.

Acidic fibrinogen gel formation

Stock solutions of fibrinogen (Fbg_{St}) at 40 mg/mL were acidified to pH 3.6, the highest pH below the isoelectric point (~ 5.5) at which the solution remained stable at ambient temperature (20°C) without precipitating. Without any addition of thrombin these acidic solutions (Fbg_{A}) incubated at 37°C formed within an hour a transparent brittle gel ($\text{Fbg}_{\text{A(G)}}$) (Fig. 1.B.1), while solutions left at ambient temperature set to a gel only after several weeks. Gel formation kinetics was increased at lower pH, with instantaneous gelation at pH 2.5. Formation of fibrinogen transparent gels at acidic pH had already been briefly evoked by Fay & Hendrix²⁶ and Mihalyi²⁷. However no study of this phenomenon has been carried out so far.

Gel formation from acidic solutions at pH 3.6 was dynamically investigated by switching the temperature of geometries from 20°C to 37°C after 5 minutes of rheological properties measurement (Fig. 1.B.2). For approximately 15 minutes G' remained constant, and then rose to reach approximately 1 kPa after three hours, overtaking G'' (factor > 10). During frequency sweep on the newly formed gels $\text{Fbg}_{\text{A(G)}}$, G' increased with frequency, with an inflection point at 1.6 rad/s, while G'' exhibited a reverse bell shape (SI Appendix, Fig. S1). During amplitude sweep, gels exhibited first a plateau with G' around 1 kPa till around 20 % amplitude, then a 3 to 4-fold strain stiffening until 130% strain (Fig. 1.B.3). As a comparison Wufsus *et al.* reported a G' value one order of magnitude higher for “classical” Fb_{N} clots made from 30 mg/mL and 50 mg/mL initial fibrinogen solutions²⁸. Structure-mechanics relationship in “classical” fibrin clots have been

widely investigated but no general picture has been proposed yet. Non-linear elasticity of fibrin clots Fb_N originates from the multi-scale hierarchical structure^{29,30}. The observed behavior in frequency sweep of $Fbg_{A(G)}$ differs from that of classical clots and exhibits a signature reminiscent of rubber plateau in polymer melts with high-molecular-weight entangled chains³¹. However, the strain-stiffening behavior is characteristic of biological gels and in particular of fibrin²⁹. AFM force-unfolding experiments on fibrin proteins suggest that clot elasticity may arise from coiled-coil and γ -nodule unfolding, and from α Cs interactions^{32,33}. Coiled-coil and/or γ nodule unfolding might then account for elasticity and strain stiffening behavior of the acidic gels, while it is unlikely that α Cs are involved as they do not interact at acidic pH³⁴.

Transmission Electron Microscopy (TEM) observations of sections of fibrinogen gels $Fbg_{A(G)}$ included in araldite display a network of fine entities without any specific pattern (Fig. 1.B.4). After chopping a gel in water and depositing the supernatant on a carbon coated grid, some large fiber-like structures can be observed (Fig. 1.B.5). In comparison, a “classical” fibrin clot made by mixing fibrinogen and thrombin at neutral pH exhibits a highly fibrillar network with fibers of various sizes (Fig. 1.A.4).

Underlying Mechanism of acidic gel formation

Acidic fibrinogen gels $Fbg_{A(G)}$ were surprisingly stable to most conditions known to disperse protein-based hydrogels (SI Appendix, Table S1). Change to neutral or basic pH did not dissolve the gels. They were resistant to low and high ionic strength, as well as to high temperature and salt-in solutions. Gels were however soluble in urea. As expected they could also be dissolved in DTT, since the constituting fibrinogen chains are linked together by multiple disulfide bonds. SDS electrophoretic gels of acidic fibrinogen gels dissolved in urea confirmed that no major cleavage was provoked by the acidic conditions and by the thermal treatment at 37°C (SI Appendix, Fig. S2). MALDI-TOF performed on fibrinogen acidic gels $Fbg_{A(G)}$ showed that no fibrinopeptide could be detected after acidic gel formation, while both Fps A and B were visible for fibrin clots Fb_N (SI Appendix, Fig. S3). Thus, the mechanism involved in gel formation at acidic pH is different from the “classical” one ($0+0'$, Fig. 1.A.2).

Circular dichroism measurements on fibrinogen stock Fbg_{st} and acidified solutions Fbg_{A} demonstrate a change in the secondary structure upon acidification, with a decrease in intensity of the 225 nm peak (Fig. 2.A). No further change in acidic solution signature could be observed upon incubation at 37°C, even after formation of a gel $\text{Fbg}_{\text{A(G)}}$. The change appears to be reversible for the solutions Fbg_{A} when raising the pH above the pI (Fig. 2.A). CD spectra of Fbg_{st} have already been reported and a consensus of approximately a third of α -helix content and a third of random structure exists, while the content of the remaining structures fluctuates according to studies³⁵⁻³⁷. In our case the change in structure upon acidification could not be reliably attributed. Similar spectra were however observed in different fibrinogen fragments³⁷⁻³⁹ and partially heat-denatured fibrinogen³⁵, which tends to point out a modification of the helical structure and/or the partial denaturation of the D domain.

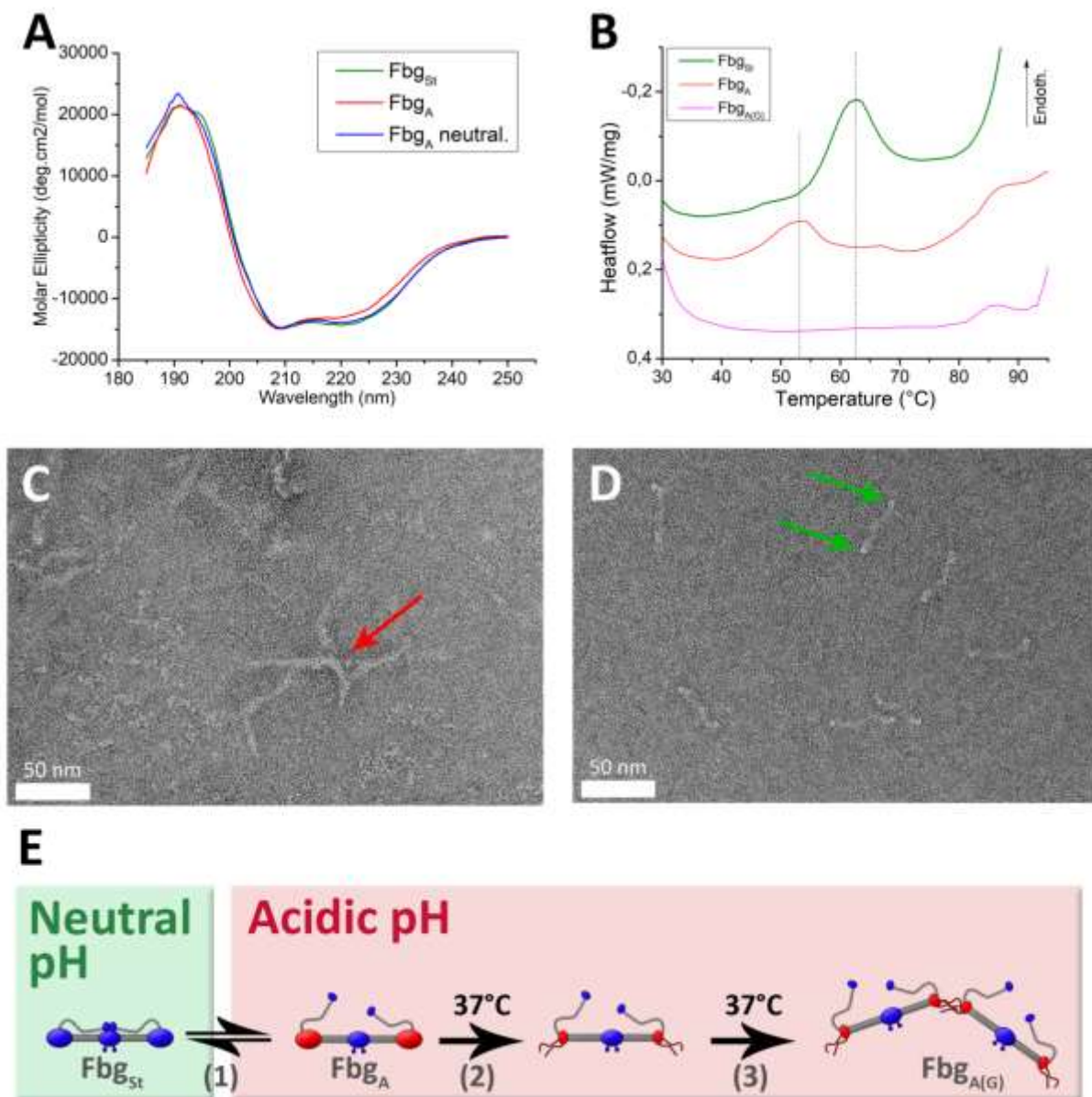


Fig. 2. Mechanism of acidic fibrinogen gel formation. (A) Circular dichroism spectra at 20°C of a Fbg_{st} (green) solution, Fbg_A solution at pH 3.6 with HCl (red), and the same solution brought back to neutral pH by addition of NaOH (blue). (B) Thermograms of Fbg_{st} solution (green), Fbg_A solution pH 3.6 (red), and Fbg_{A(G)} gel pH 3.6. (C & D) Fbg_A and Fbg_{st}, respectively, dried on a carbon coated grid and observed by TEM. (E) Schematic of the mechanism inferred from the obtained results. Upon acidification to pH 3.6 (1) of Fbg_{st}, the D domains of acidic fibrinogen Fbg_A are destabilized in a reversible manner (red). Incubation at 37°C provokes the denaturation (2) of the thermolabile D domains of fibrinogen proteins that subsequently interact by their extremities (3) to give an acidic fibrinogen gel Fbg_{A(G)}.

Differential Scanning Calorimetry of fibrinogen stock Fbg_{St} exhibits a major peak around $63^{\circ}C$ (Fig. 2.B). This low temperature peak of fibrinogen at neutral pH is attributed to the D domain⁴⁰. When acidified, three peaks appear: a major peak around $52^{\circ}C$ and a shoulder around $65^{\circ}C$, corresponding to the thermolabile part of the D domain and to the αCs , respectively⁴⁰, followed by a third peak close to $85^{\circ}C$, attributed to the E domain. The thermolabile D domain is thus destabilized by the acidic condition with a low temperature peak shift of $-10^{\circ}C$, in accordance with previous results⁴⁰. When the acidic solution Fbg_A is first kept at $37^{\circ}C$ for 1 hour before the scan, only the high temperature peak around $85^{\circ}C$ remains (Fig. 2.B). The incubation of acidic solutions at $37^{\circ}C$ for 1 hour thus leads to vanishing of the low temperature peak attributed to the thermolabile D domain⁴⁰. After incubation, the absence of the shoulder at $65^{\circ}C$, corresponding to the αCs and reported to be reversible upon cooling⁴¹, could be due to a too low sensitivity of our setup. Observation of dilute acidic fibrinogen solutions Fbg_A by TEM shows that fibrinogen proteins tend to aggregate preferentially by their extremities (Fig. 2.C), on the opposite to fibrinogen proteins from stock solution Fbg_{St} , which appear dispersed (Fig. 2.D).

Aggregation of fibrinogen in acidic conditions upon incubation in presence of D and E fragments was investigated by dynamic light scattering. Altered aggregation in presence of D fragments was observed, on the opposite to E fragments (Appendix, Fig. S4), which tends to confirm that D domains are involved in aggregation.

As summed up in the sketch (Fig. 2.E), stock fibrinogen Fbg_{St} undergoes upon acidification **(1)** a reversible change in structure that corresponds to a destabilization of its thermolabile D domain (represented by a change of color of the D domain from blue to red) in the acidic fibrinogen Fbg_A . Upon incubation at $37^{\circ}C$ **(2)**, these domains are denatured (represented by the floating carboxyterminal ends of β and γ chains⁴²), promoting interactions between proteins **(3)** and resulting in a gel $Fbg_{A(G)}$.

Effect of neutralization on fibrinogen and fibrin acidic gels

Fibrinogen

The acidic fibrinogen gels $Fbg_{A(G)}$ are not soluble when dipped in a neutral solution and no major change in aspect such as opacification or gel contraction can be observed. The change in rheological properties upon neutralization of acidic gels $Fbg_{A(G)}$ was studied by forming a gel at 37°C in acidic condition and then adding a neutral solution of 100 mM Hepes around the geometries (Fig. 3.A, blue). This resulted in a gel $Fbg_{N(G)}$ with a two-fold increase in storage modulus as well as a different signature in frequency sweep which now increases linearly with the frequency (SI Appendix, Fig. S5). These modified mechanical properties may be related to a change in structure.

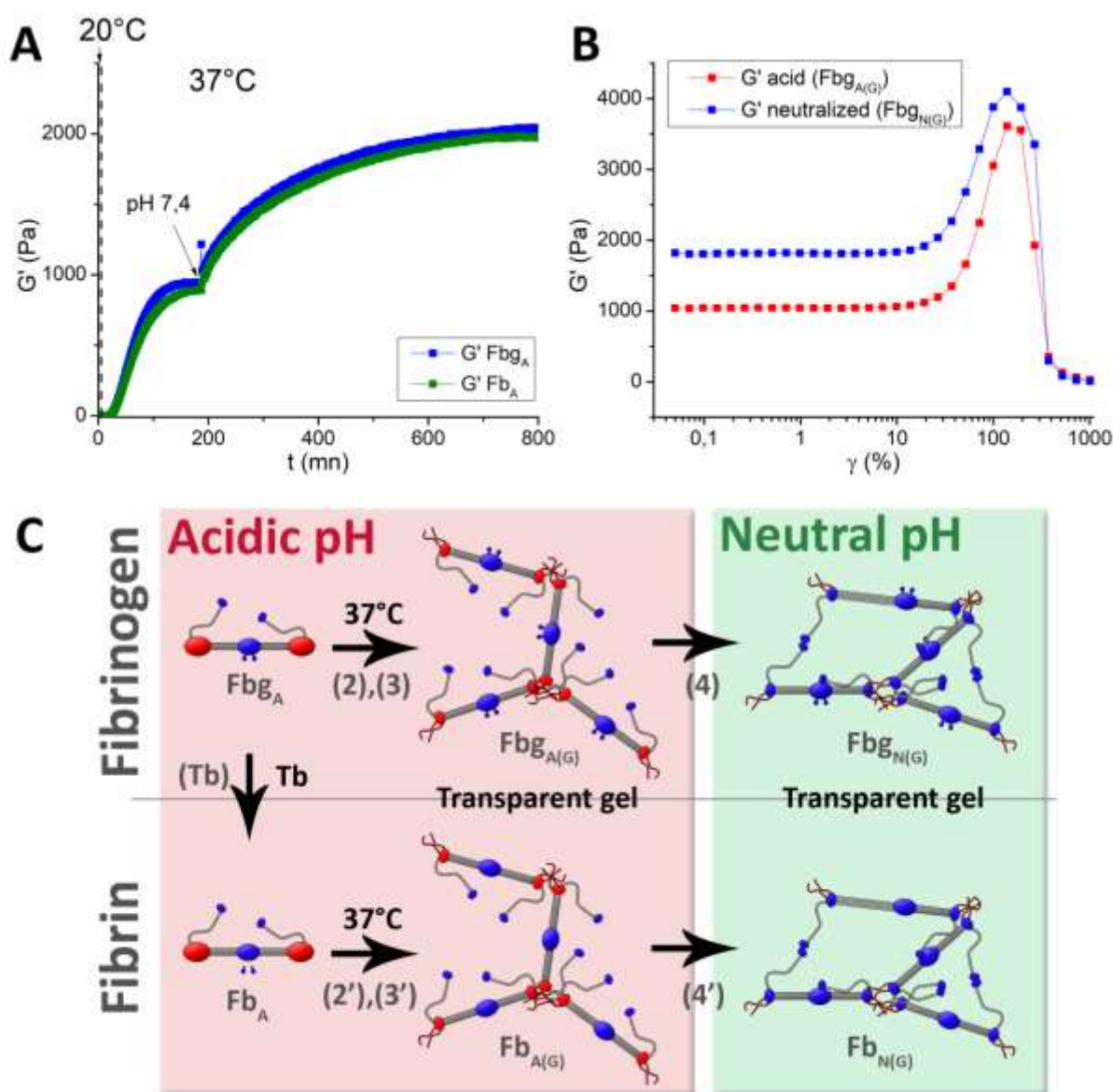


Fig. 3. Effect of neutralization on acidic fibrin and fibrinogen gels. (A) Formation of acidic gels from acidic fibrinogen Fb_{gA} and fibrin Fb_A solutions by incubation at 37°C followed by gel stiffening upon addition of a neutral solution around the geometry. (B) Comparison of mechanical response in amplitude sweep of a neutralized acidic fibrinogen gel (red) compared to a non-neutralized one (blue). (C) Schematic of the underlying mechanisms inferred from the measures. Fibrinogen proteins Fb_{gA} assemble upon incubation (2) forming a gel $Fb_{gA(G)}$ (3) as seen on Fig.2. Upon neutralization, creation of additional interactions inside the gel $Fb_{gN(G)}$ lead to internal constraints and subsequent partial unfolding of some proteins (4). Similar results are obtained in presence of thrombin (Fb_A), both during the gel $Fb_{A(G)}$ formation in acidic conditions at 37°C (2')(3') and upon neutralization (4') to result in a similar gels $Fb_{N(G)}$ as $Fb_{gN(G)}$.

The increase in G' and G'' with frequency is closer to fibrin clot Fb_N signature²⁸. This modification is not surprising as protein global charge is modified secondary to pI crossing. In addition to this, the αCs may form intermolecular interactions. Veklich *et al.* reported that αCs interact together and also with the E domains at neutral pH, while they float freely in acidic conditions. They also demonstrated that αCs cleaved from fibrinogen were dispersed in acidic conditions and polymerized into fibers upon neutralization. It is then likely that the αCs float during gel formation at acidic pH and interact together upon neutralization. This would be facilitated by the forced proximity between proteins in the network. Regarding the amplitude sweep a similar behavior as in the acidic condition was observed (Fig. 3.B), with a G' plateau for low amplitudes, at approximately twice the value, but only a slight increase of peak amplitude between 100 and 200 % strain. Such a phenomenon may be due to a partial initial stretching of the proteins inside the network upon neutralization (Fig. 3.C (4)) because of internal constraints related to electrostatic repulsion and/or αCs interactions in the neutral gels.

Fibrin

Till now, no thrombin was introduced in the solutions, gels being made of fibrinogen proteins (as confirmed by MALDI-TOF results). It was shown in literature that addition of thrombin to fibrinogen in acidic conditions effectively cleaves the Fps, transforming fibrinogen Fb_{gA} into fibrin monomer Fb_A but without inducing clotting⁴³, which becomes instantaneous upon neutralization. The influence of thrombin addition to our system was therefore investigated.

$Fb_{A(G)}$ gels from acidic fibrin solutions Fb_A could be formed by incubation at 37°C , in the same fashion as $Fbg_{A(G)}$, without any impact on kinetics or gel aspect. Same rheological study of acidic gel formation and subsequent neutralization was carried out from fibrin solution Fb_A as for fibrinogen Fbg_A . No significant difference either in formation kinetics or mechanical properties could be observed between $Fb_{A(G)}$ and $Fbg_{A(G)}$, as well as between $Fb_{N(G)}$ and $Fbg_{N(G)}$ (Fig. 3.A). MALDI-TOF performed on a $Fb_{A(G)}$ gel formed by incubation at 37°C confirmed that FpA and FpB are effectively cleaved upon addition of thrombin to acidic fibrinogen (SI Appendix, Fig. S3). Thus, the removal of the Fps by thrombin does not seem to affect the acidic gel formation and neutralization.

Effect of neutralization on fibrinogen and fibrin viscous acidic solutions

Fibrinogen

As already mentioned viscosity increases progressively during incubation at 37°C (Fig. 1.B.2), thus giving the possibility to obtain viscous fibrinogen solutions $Fbg_{A(VS)}$. When neutralized, these solutions form opaque gels $Fbg_{N(VS)}$, stable over time, on the contrary to non-heated solutions Fbg_A which produce precipitates that dissolve over time. This phenomenon was investigated by examining rheological properties. $Fbg_{A(VS)}$ heated prior to measurement were neutralized five minutes after starting the measurement at 25°C by careful addition of a neutral solution of 100 mM Hepes around the geometries, and resulted in gels $Fbg_{N(VS)}$ with G' of a few hundred of Pa after 24 hours (SI Appendix, Fig. S6). Solutions $Fbg_{A(VS)}$ with different initial viscosities were analyzed and the G' of the final gels $Fbg_{N(VS)}$ were plotted as a function of the initial G' of the solutions (Fig. 4.A.1). A sharp increase of final G' as a function of initial G' is observed, followed by a plateau around 350 Pa. Resulting gels $Fbg_{N(VS)}$ exhibit dense regions of aggregated proteins, reminiscent of the structure observed in acidic fibrinogen gels $Fbg_{A(G)}$ as observed by TEM (Fig. 4.A.2). It is likely that the mechanism (5) responsible for formation of the gels $Fbg_{N(VS)}$ upon neutralization of $Fbg_{A(VS)}$ is similar to the mechanism causing acidic gel $Fbg_{A(G)}$ strengthening after raising pH (Fig. 3.C (4)), and was thus hypothesized to involve αCs interaction (Fig. 4.A.3).

Fibrin

The same rheological study was carried out by neutralizing acidic fibrin solutions $Fb_{A(VS)}$ at different initial viscosities. As seen on Fig. 4.B.1, in spite of measures' dispersity, for non-

incubated solutions Fb_A and shortly-incubated solutions $Fb_{A(VS)}$ with low initial storage moduli, the final G' of Fb_N and $Fb_{N(VS)}$ ranged between 500 and 1250 Pa. For longer incubated $Fb_{A(VS)}$ solutions, with higher initial G' , the final value exceeded 1500 Pa, and even 3000 Pa for certain gels. All the fibrin gels Fb_N and $Fb_{N(VS)}$ reached higher storage moduli than pure fibrinogen gels $Fb_{N(VS)}$, independently of the initial solution viscosity. TEM observation of $Fb_{N(VS)}$ gels exhibited a composite structure, with thin fibers connecting aggregate-like islands similar to the ones present in pure fibrinogen gels $Fb_{A(G)}$ and $Fb_{N(VS)}$ (Fig. 4.B.2). Upon neutralization, the non-denatured fibrin proteins self-assemble following the “classical” scheme, bridging the aggregates islands through thin fibers (Fig. 4.B.3). The formation of aggregates by incubation prior to neutralization has a tremendous effect on final gel properties. Indeed, for non- or shortly-incubated solutions Fb_A and $Fb_{A(VS)}$ with low initial G' , the final storage modulus remains below 1.5 kPa, while viscous solutions $Fb_{A(VS)}$ incubated for a longer time all reach higher moduli. The high dispersion of the measures may be explained by the fact that the system is more complex than for fibrinogen, with multiple mechanisms involved.

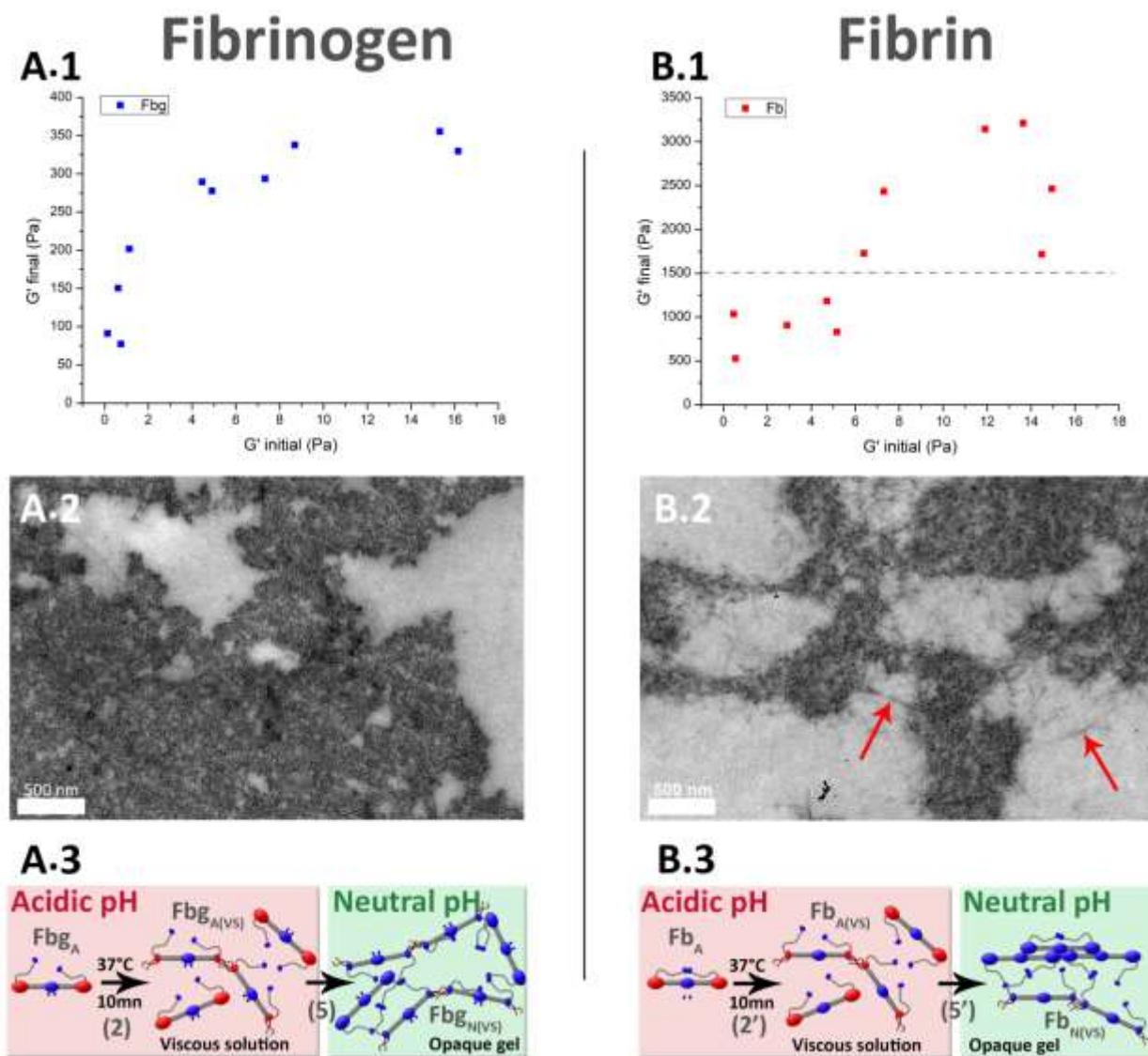


Fig. 4. Effect of neutralization on viscous acidic solutions of fibrinogen Fbg_{A(VS)} (A) and fibrin Fb_{A(VS)} (B) and analysis of the final neutral gels Fbg_{N(VS)} and Fb_{N(VS)}. (A.1 & B.1) Resulting storage moduli (G' final) of respectively Fbg_{N(VS)} and Fb_{N(VS)}, as a function of the initial storage moduli of respectively Fbg_{A(VS)} and Fb_{A(VS)} fixed by different incubation times at 37°C. (A.2 & B.2) Resulting structures, respectively Fbg_{N(VS)} and Fb_{N(VS)}, as observed by TEM. (A.3 & B.3) Schematic sum up of the neutralization (5 & 5') of respectively Fbg_{A(VS)} and Fb_{A(VS)}, where only parts of the proteins are denatured by short incubation times (2 & 2') and form aggregates. For fibrinogen Fbg_{A(VS)}, upon neutralization (5) the strengthening of the network is hypothesized to rely on α Cs interactions promoted by the proximity of proteins. For fibrin solutions Fb_{A(VS)}, upon neutralization (5') the non-denatured fibrin proteins self-assemble following the classical scheme, bridging aggregates formed during incubation at acidic pH (2').

Versatility in acidic solution viscosity and potential applications

The here presented study sheds light on the behavior of fibrinogen and fibrin at acidic pH, and the different steps involved in gelation process are depicted in Fig. 5. Beyond the interest from a biochemical standpoint the investigated phenomena represent a unique opportunity to produce stable materials made of pure fibrin (Fb_N , $Fb_{N(VS)}$, $Fb_{N(G)}$ and $Fb_{A(G)}$) and, more significantly, of pure fibrinogen ($Fbg_{N(VS)}$, $Fbg_{N(G)}$ and $Fbg_{A(G)}$) without any use of chemical or physical crosslinking. Two main processes can be implemented from our work.

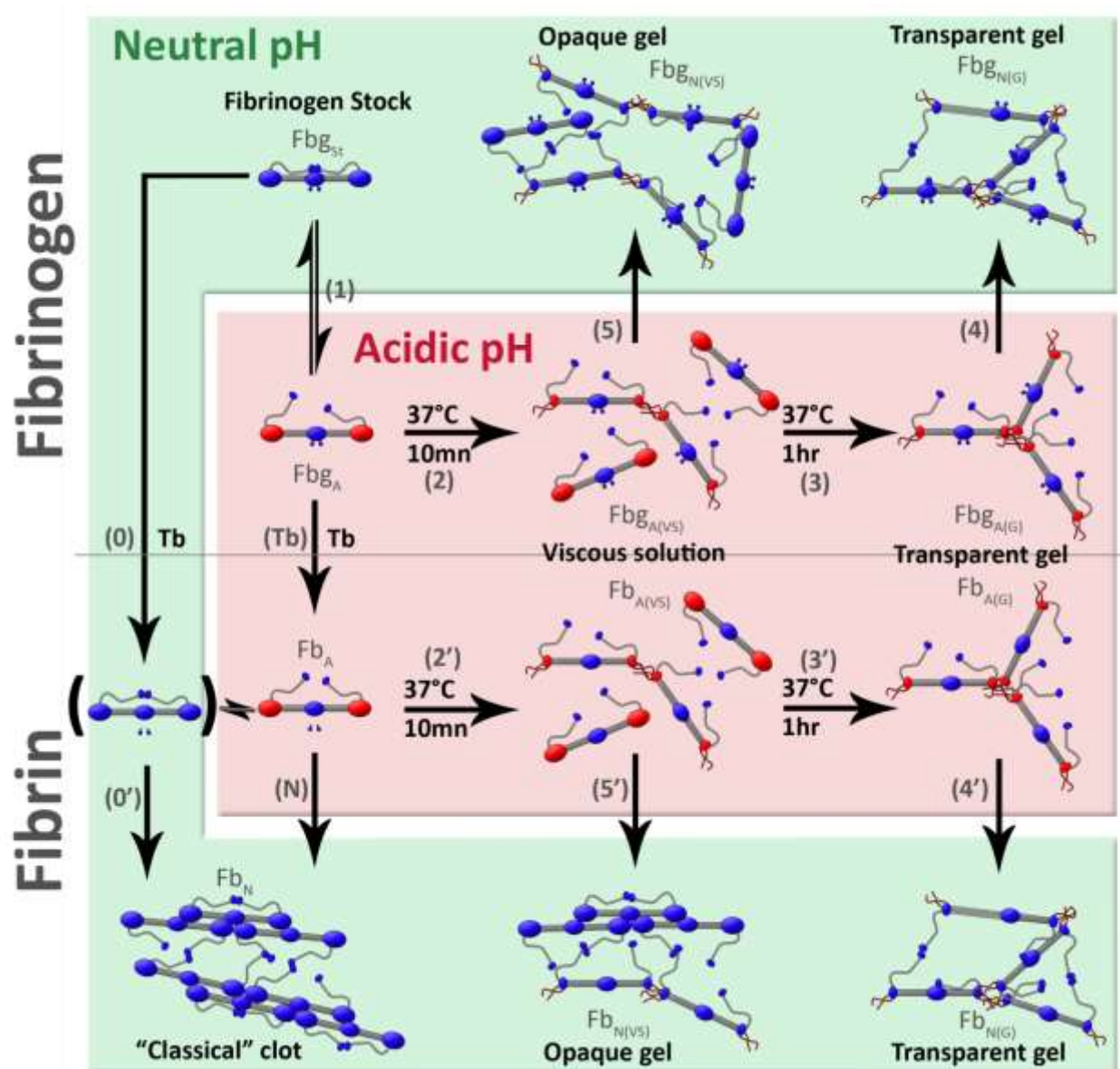


Fig. 5. The different mechanisms of fibrin(ogen) gelation. Classically, fibrin Fb_N is polymerized by the action of thrombin that cleaves Fps from fibrinogen Fbg_{St} (**0**), causing self-assembly in a half staggered manner (**0'**). Upon acidification (**1**), fibrinogen Fbg_{St} undergoes a reversible change in structure which involves the destabilization of the thermolabile D domains and αCs interactions in Fbg_A . Incubation at 37°C denatures this domain, enabling the interactions of “denatured proteins” by their extremities, which causes an increase in viscosity (**2**) of the solution $Fbg_{A(VS)}$ which eventually sets (**3**) to a gel $Fbg_{A(G)}$. Upon neutralization, the formation (**5**) of opaque gels $Fbg_{N(VS)}$ from viscous solutions $Fbg_{A(VS)}$ and the strengthening (**4**) of the transparent gels $Fbg_{A(G)}$ are hypothesized to be partly due to αCs interactions. Addition of thrombin (**Tb**) to freshly acidified fibrinogen solutions Fbg_A causes the cleavage of fibrinopeptides but prevents fibrin Fb_N assembly, which only occurs upon neutralization (N). No effect of the Fps cleavage could be observed on the $Fb_{A(VS)}$ viscosity increase (**2'**) and gel $Fb_{A(G)}$ formation (**3'**) of acidic solutions Fb_A upon incubation, compared to (**2**) and (**3**), as well as after neutralization of $Fb_{A(G)}$ gels (**4'**), compared to (**4**). However, neutralization of viscous fibrin solutions $Fb_{A(VS)}$ (**5'**) produced composites made of acidic aggregates bridged by “classical” fibers, and resulted in stiffer gels $Fb_{N(VS)}$, compared to those obtained (Fb_N) from non-incubated solutions Fb_A after neutralization (N). These phenomena are useful to produce transparent fibrinogen $Fbg_{N(G)}$ or fibrin $Fb_{N(G)}$ gels (**4**)(**4'**) and enable the production of fibrinogen $Fbg_{N(VS)}$ (**5**) and fibrin $Fb_{N(VS)}$ (**5'**) threads, and provide a proof of concept in printing, using incubation of solutions (**2**)(**2'**) to tune the solution viscosity and/or the threads final mechanical properties.

The first is the formation of stable transparent ($Fbg_{A(G)}$ and $Fbg_{N(G)}$) gels by incubation (**2+3**) of acidic fibrinogen solutions. Such homogeneous materials may be valuable for molding or coating processes since they are much easier to process than those obtained by mixing fibrinogen and thrombin at neutral pH (**0+0'**). At low ionic strength, those gels do not contract and are transparent, which represents an obvious advantage for such processes. Fibrin materials ($Fb_{A(G)}$ and $Fb_{N(G)}$) can also be produced the same way (**2'+3'**), with no obvious benefit on structural and mechanical properties as compared to those made of fibrinogen (Fig. 3.A).

The second process is the instantaneous formation of gels ($Fbg_{N(VS)}$, Fb_N , $Fb_{N(VS)}$) from acidic solutions by a simple change from acidic to neutral pH (**5,5'**), on the contrary to the delayed gelation (**0'**) initiated by thrombin (**0**) used in the literature²³. The pH-triggered gelation of stable acidic solutions of fibrin(ogen) enables to transpose multiple technique implemented for collagen to fibrin(ogen). For instance, this makes it possible to produce stable fibrinogen and fibrin threads (Fig. 6.A) by extrusion in neutral buffer, and provides the proof of concept for pure fibrin and

fibrinogen printing as shown on Fig 6.B and SI Appendix, Video S1 where the printing of a honeycomb pattern is displayed.

The ability to tune the viscosity (**2** or **2'**) of the solution is a key asset for processing and allows controlling the mechanical properties of the final gel (Fig. 4.A.1&B.1). Increasing solution viscosity by incubating fibrin(ogen) acidic solution has a paramount effect on thread production by extrusion, as it stabilizes the flow outside the needle. Viscosity of solutions is also a paramount parameter for printing: low viscosity fibrin solutions may be, for instance, interesting for inkjet printing, while viscous solutions are sought for extrusion printing. The acidic pH of the solutions limits the incorporation of cells directly in the inks, on the contrary to classical pathway using thrombin only. Use in bioprinting may then require to print cells and fibrin(ogen) separately. Fibrin(ogen) inks may be used as a support matrix where cells can be printed in between fibrin(ogen) threads, in the fashion of Atala's group¹⁹, or by means of co-extrusion⁴⁴.

Additionally, acidic conditions and extrusion are compatible with collagen solutions, enabling production of composite materials. For instance, fibrin(ogen) solution can be mixed with collagen solution (60 mg/mL, 3 mM HCl, 0.8 mM citrate, pH 3.6) at any rate from 100% fibrin(ogen) to 100% collagen and extruded in the same neutralizing buffer (100 mM Hepes, 2.5% PEG). Co-extrusion can also be performed, with the possibility to tune dynamically thread composition using a double extrusion device (SI Appendix, Fig. S7) and to produce fibrin(ogen) to collagen gradients (Fig. 6.C).

Test were performed to ensure that modification of fibrin(ogen) caused by acidic incubation did not alter fibrin(ogen) biocompatibility and bioactivity. L929 murine fibroblasts, commonly used to test biocompatibility in the ISO standard, were cultured on wells coated with "classical" fibrin Fb_N, acidic fibrinogen and fibrin gelled by incubation Fbg_{N(G)} and Fb_{N(G)}, and viscous acidic fibrinogen and fibrin gelled by neutralization, Fbg_{N(VS)} and Fb_{N(VS)}. As observed on Fig. 6.C, all materials induced a similar increase in metabolic activity from day 1 to day 6, due to the high

proliferation of L929 on the materials till confluence (SI Appendix, Fig. S8). On all materials, cells adopted typical L929 cells morphology, demonstrating that the various processes do not prevent cell adhesion to matrix. Interestingly, at day 1 and 6, a higher metabolic rate is observed ($p < 0.0001$ & $p = 0.0019$ resp.) on non-self-assembled fibrin(ogen), obtained by total incubation $Fb_{N(G)}$ and $Fb_{N(G)}$ or by neutralization of viscous fibrinogen solution $Fb_{N(VS)}$, compared to materials containing self-assembled fibrin, “classical” clot Fb_N and neutralized viscous fibrin solution $Fb_{N(VS)}$.

Further test with primary human fibroblasts seeded on $Fb_{N(VS)}$ fibrinogen and $Fb_{N(VS)}$ fibrin, obtained by extrusion of the corresponding solutions in a neutralizing buffer (100 mM Hepes 2.5% PEG, pH 7.4), was performed to ensure that the process and the difference of structure did not alter cell adhesion. Culture of Normal Human Dermal Fibroblasts on threads exhibited high affinity toward both types of threads, with high cellular densities (Fig. 6.E&F). After three days, both types of threads were covered with a monolayer of cells, without colonization inside the threads, as confirmed using confocal microscopy (SI Appendix, Fig. S9).

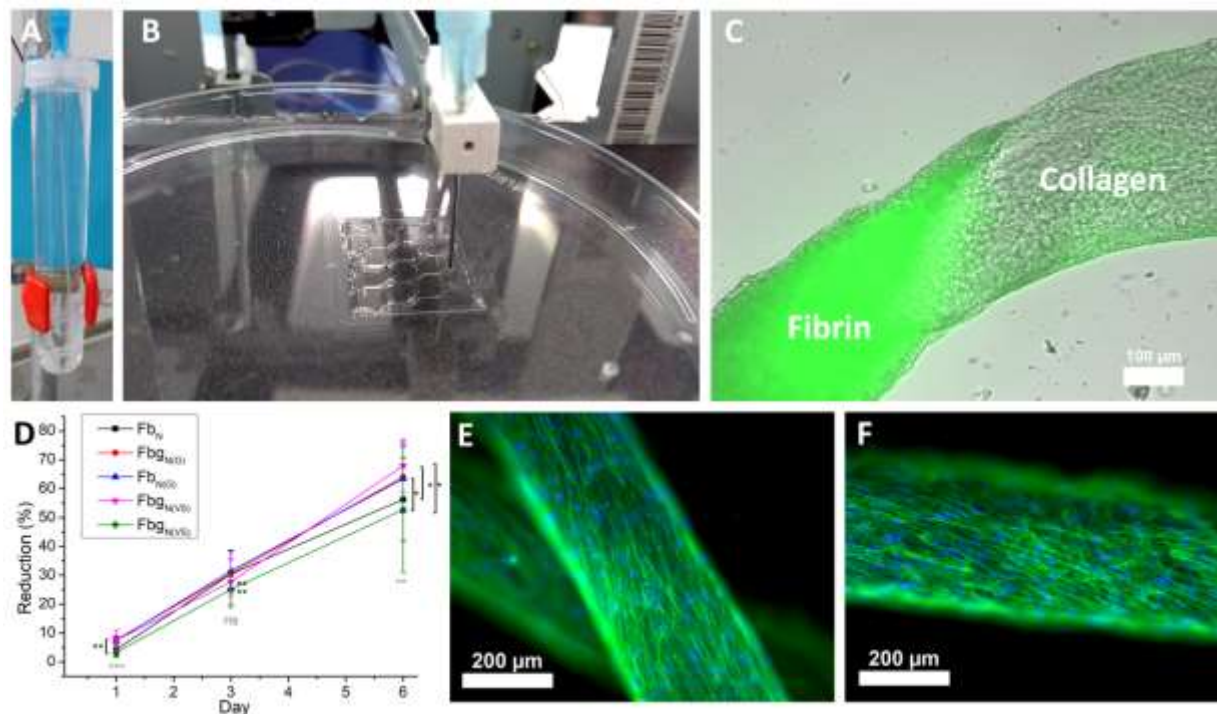


Fig. 6. Use of acidic fibrin and fibrinogen solutions for production of new materials. (A) Image of the extrusion of a viscous fibrinogen solution in 100 mM Hepes 2.5% PEG solution pH 7.4. (B) Printing of a pure viscous fibrin solution on a Petri dish. (C) Pure fibrin to collagen gradient in a continuous thread produced by alternate extrusion of viscous acidic fibrin and collagen solutions (60 mg/mL, 3mM HCl, 0.8mM citrate pH 3.6) in the same conditions (100mM Hepes & 2.5% PEG) observed with a fluorescent microscope. Green: FITC-labelled fibrin. (D) Reduction rate of L929 murine fibroblasts seeded in wells coated with “classical” fibrin Fb_N, acidic fibrinogen and fibrin gelled by incubation Fbg_{N(G)} and Fb_{N(G)}, and viscous acidic fibrinogen and fibrin gelled by neutralization, Fbg_{N(VS)} and Fb_{N(VS)}. (E & F) Normal Human Dermal Fibroblasts 3 days after seeding on respectively a pure fibrinogen Fbg_{N(VS)} and a pure fibrin Fb_{N(VS)} thread. Blue: nuclei (DAPI); green: actin (phalloidin).

Ability to produce fibrinogen materials represents an obvious interest for biomedical applications. Indeed, it spares the need of thrombin, reducing cost and risks as thrombin used clinically is not autologous whereas fibrinogen can be directly extracted from blood patient⁴⁵. On a more academic point of view, ability to produce fibrinogen materials enables the investigation of their effect on cell behavior such as, for instance, the influence on cell phenotype of remaining fibrinopeptides in thrombin-free materials^{46,47} or the presence of self-assembled fibrin.

In summary, our present results widen the possibilities in terms of fibrin processing, and enable production and shape control of fibrinogen materials. Furthermore, the processes are implemented

in similar conditions as those used for collagen, which should enable the development of hybrid scaffolds based on these two proteins. The possibility to tune the biochemical and structural properties of these materials should be an asset in order to induce specific and spatially-controlled interactions with the cells and obtain appropriate cellular response to mimic the complexity of the tissues.

Materials & Methods

Fibrinogen and thrombin solutions

Fibrinogen stock solutions were prepared by dissolving 1 g of human fibrinogen lyophilized from 20 mM citrate pH 7.4 solutions (Merck 341576, >90% clottable proteins) into distilled water as recommended by the manufacturer (final volume of app. 22 mL). The final pH of the stock solutions was 6.6 to 6.8. The concentration (40 ± 2 mg/mL) was measured with the extinction coefficient at 280 nm (1.51 mL/mg) 27. The solutions were aliquoted and kept at -80°C . Lyophilized thrombin from bovine plasma (Sigma) was dissolved in sterile PBS 1X to reach a concentration of 200 U/mL. The solutions were aliquoted and kept at -20°C .

FITC-labeled fibrinogen was prepared by dissolution of 20 mg of FITC CeliteR in 3 mL of 0.13 M carbonate buffer pH 9.2 subsequently mixed with fibrinogen stock solution at 40 mg/mL at a 1:3 ratio and dialysed 48 hours against 2 L 20 mM citrate pH 7.4, aliquoted and stored at -80°C . For extrusion, 10% in volume of FITC-labeled fibrinogen [Fbg-FITC] was added to fibrinogen stock solution, following the same protocol as usual afterwards.

Rheology

Rheology was performed on an MCR 302 rheometer (Anton Paar), equipped with a temperature control of both upper and lower geometries (Peltier hood PTD200). A sanded plate-plate geometry (PP25/S) was used for gel formation study and subsequent gel characterization.

For acidic gel formation study, significant amount of water was poured around the geometry, and the hood lowered to prevent from drying. Measurements were performed at constant amplitude and frequency ($\gamma=0.1$ %, $\omega=1$ rad/s). The temperature was kept at 25°C for 5 mn and then switched to 37°C (zero-gap calibration was performed at 37°C prior to measure). Once gel formed, a frequency sweep ($\omega = 0.1$ to 100 rad/s; $\gamma=1$ %) was performed, followed by an amplitude sweep ($\gamma= 0.05$ to 1000%; $\omega = 1$ rad/s).

Gel neutralization experiments on acidic gels were carried out following the protocols above, except that a solution of 100 mM Hepes pH 7.4 was carefully added around the geometry after 3 hours of measure, once acidic gel formation completed.

For the study of neutralization of viscous solutions a solution of 100mM Hepes, pH 7.4, was carefully added around the geometry, after 5 mn of measure. Temperature was kept constant at 25°C during the whole experiment. Frequency and amplitude sweep were performed as described above.

Transmission electron microscopy

For inclusion in araldite, hydrated samples were crosslinked with paraformaldehyde, glutaraldehyde and osmium tetroxide 4 wt. %. They were subsequently dehydrated using baths with increasing concentrations in ethanol, progressively transferred to propylene oxide and incorporated in araldite resin prior to sectioning.

Observation of solutions was performed by dilution of solutions by a factor 10000 with the corresponding buffer and deposition of a few drops on a charged carbon coated TEM grid. The solution was sucked up after a few seconds with a filter paper on the side of the grid. Gels were chopped on a glass slide and a carbon coated TEM grid was put in contact with the mixture and dried by sucking up the supernatant with a filter paper. A drop of uranyl acetate 0.5% was then added and removed the same way.

The observations were made on a transmission electron microscope FEI Tecnai Spirit G2 operating at 120 kV. Images were recorded on a Gatan CCD camera.

Differential Scanning Calorimetry

Differential Scanning Calorimetry was performed on a DSC Q20 from TA. Small volume (20 μ L) sealed pans were used. As a reference, a similar pan with comparable weight of relevant buffer was used. Solution was kept at 20°C for 5 min for equilibration, followed by a ramp at 10°C/min till 100°C.

Circular Dichroism

Circular Dichroism was performed on a J-810 CD from Jasco, with a temperature control Peltier module (PTC-4235). As the intensity was too high for stock solutions, even with a 0.01 mm cuvette, stock solution was diluted by a factor three with 20 mM citrate solution, pH 7.4 to reach a concentration of approx. 13 mg/mL that was further used for all the experiments. Spectra were smoothed with a Savitsky-Golay filter with a polynomial order of 2 and a smoothing window of 10 points.

Dynamic Light Scattering

Dynamic light scattering was performed on a Malvern Zetameter with solutions 1 mg/mL fibrinogen solutions with or without addition of D or E fragments at a ratio Fbg:fragment = 1:2 in mol. Solutions at pH 7.4 were in 20 mM citrate while acidic solutions were dialysed against 50 mM Glycine buffer and pH adjusted to 3.6.

SDS-PAGE

Acidic fibrinogen gels were made by incubating acidic solutions of fibrinogen at 37°C overnight. They were subsequently dipped into either a 20 mM citrate solution pH 7.4 or a 8 M Urea solution. As for the controls, the same protocol was carried out using a neutral stock solution of fibrinogen and a "classical" clot made by mixing thrombin with the stock solution at neutral pH. After 4 days, the solutions in which the three different forms of fibrin(ogen) had been agitated were sampled and run on 10% SDS-PAGE gels.

Mass spectrometry

Matrix-Assisted Laser Desorption/Ionization (MALDI) coupled with a Time Of Flight (TOF) mass spectrometer was used to study the release of the fibrinopeptides FpA and FpB, respectively 1536.6 and 1552.6 Da in mass ^{48,49}, after acidic fibrinogen and fibrin gel formation. As the fibrinogen stock solution initially contained some fibrinopeptides A, the stock solution was first dialysed 24 hours against 20 mM citrate pH 7.4 with a 12-14 kDa cut-off membrane. Each gel was rinsed with mechanical agitation using 50 % acetonitrile and 50 % of the relevant buffer for each gel (20 mM citrate pH 7.4 for neutral fibrin gel, 20 mM Citrate 60 mM HCl pH 3.6 for acidic fibrin and fibrinogen gels). The rinsing solutions were measured by MALDI-TOF. As a control, a clot formed by mixing thrombin to fibrinogen stock solution at neutral pH was rinsed and both FpA and FpB could be observed at 1536.6 and 1552.6 Da respectively. The attribution of these peaks to FpA and FpB was further confirmed by MS/MS.

Cell culture

Normal Human Dermal Fibroblasts (PromoCell™) at passage 17 were seeded at 420,000 cells/mL on fibrin and fibrinogen threads in low glucose DMEM with 10% fetal bovine serum (GIBCO). After 72 hours culture, threads were rinsed with PBS 1X and fixed with PFA 4 %. Cells were permeabilized with Triton X-100. Fluorescent labeling of the nuclei (DAPI, Invitrogen) and the actin filaments (Alexa FluorR 488 phalloidin, Invitrogen) was performed. Observations of the samples were carried out under fluorescence microscope (Axio Imager D.1, Zeiss) and confocal microscope (Leica SP5 upright Confocal).

Murine fibroblasts L929 at passage 11 were seeded to a final cell density of 2×10^4 cell/cm² in the wells of 48 well-plates coated with the different fibrin(ogen) materials. Fibrinogen solutions were acidified to reach 60 mM HCl and supplemented with 10 mM CaCl₂ final, and 20 U/mL thrombin final for fibrin solutions or PBS 1X for fibrinogen solutions. Classical Fb_N coating was performed using same stock solution supplemented with same concentrations of CaCl₂ and thrombin. Viscous fibrin(ogen) solutions were incubated as for thread production, deposited on wells and neutralized with DMEM, to produce Fbg_{N(VS)} and Fb_{N(VS)}. To produce Fbg_{N(G)} and Fb_{N(G)}, solutions were incubated one hour at 37°C to form a gel and neutralized by adding DMEM. Alamar blue tests were performed on 6 replicates at day 1, 3 and 6 by adding resazurin solution to fresh whole medium to reach a volume of 300 µL at a concentration of 0.01 mg/mL, incubating for 4 hours and measuring fluorescence excitation at 560 nm and emission at 590 nm using a Varioskan Lux reader from Thermo Scientific. Fluorescence imaging was performed by similar labeling as on NHDF cells and observations were done on an EVOS inverted microscope from Thermo Fisher Scientific.

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Author contributions

C. R., T. C. and L. T. conceived the study. C. R., G. M., B. H. and L. T. performed experiments. C. R. analyzed data and developed the figures. C. R., G. M., T. C. and L. T. wrote the manuscript. All authors commented on the manuscript.

Materials & Correspondence

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Competing interests

The authors declare no competing interests.

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Supplementary Information

Thrombin-free polymerization leads to pure fibrin(ogen) materials with extended processing capacity

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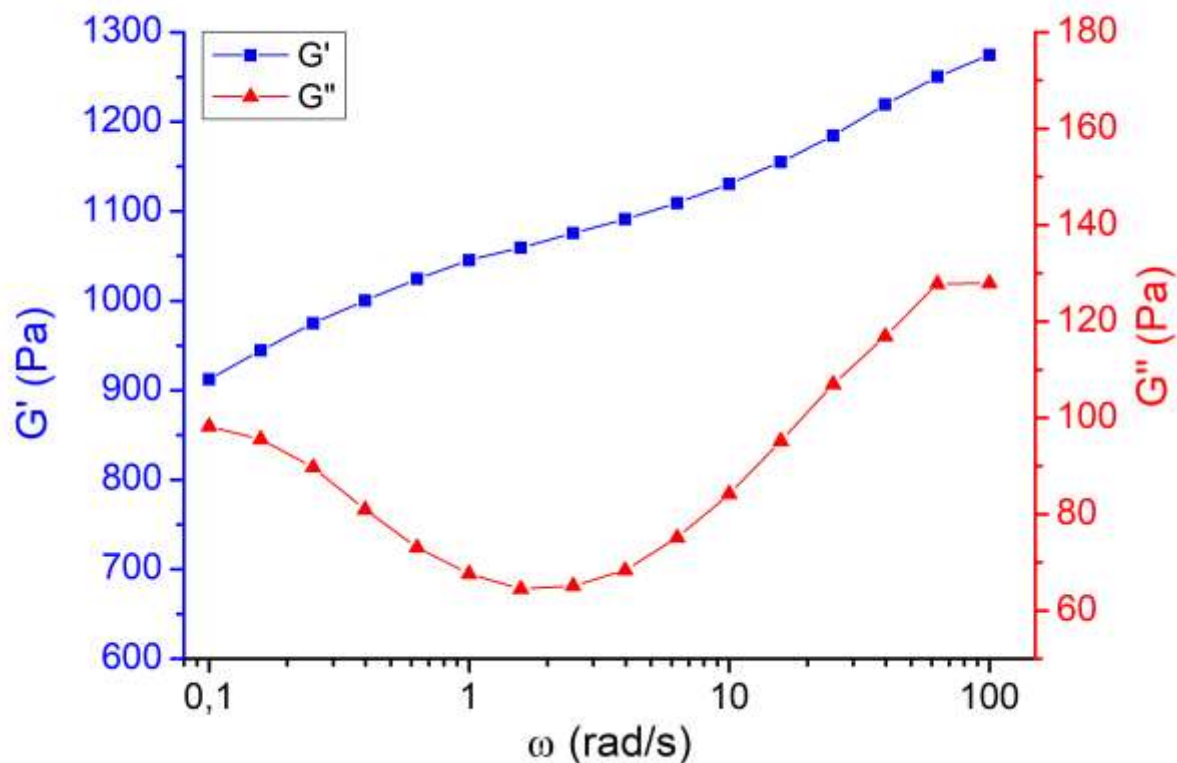


Fig. S1: Mechanical response of an acidic fibrinogen gel $\text{Fbg}_{A(G)}$ in frequency. Rheological study ($\gamma=1\%$) of an acidic fibrinogen gel $\text{Fbg}_{A(G)}$, polymerized prior to measure by 37°C -incubation between the geometries.

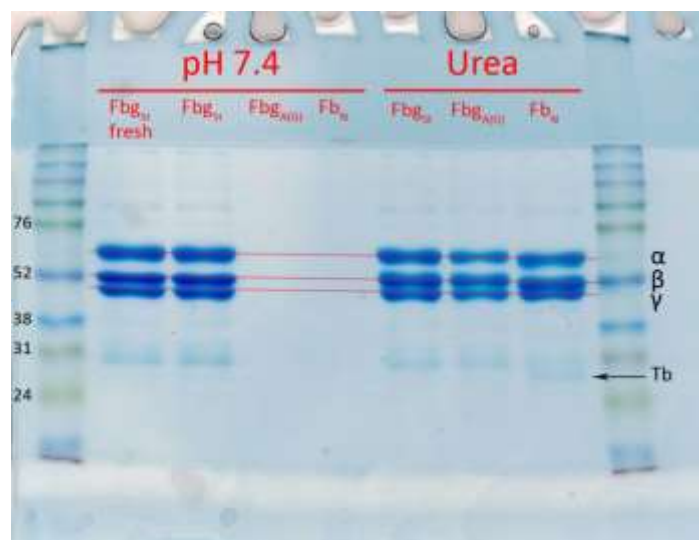


Fig. S2: Solubility tests of acidic fibrinogen gels FbgA(G). Samples were left for 4 days in two different solutions: “pH 7.4” (20 mM citrate) and “Urea” (8 M). Supernatants were then 30ipette and run on SDS PAGE gels to investigate the presence of fibrin(ogen) proteins and possible degradation products. The samples were acid fibrinogen gels FbgA(G), neutral fibrin clots FbN and a stock solution thawed 4 days before FbgSt or the day of the SDS PAGE “FbgSt fresh”. The three main bands correspond to the three chains α , β and γ of the fibrinogen, of respectively 67, 55 and 48 kDa(1). The band around 31 kDa in FbN sample in urea corresponds to thrombin’s B chain.

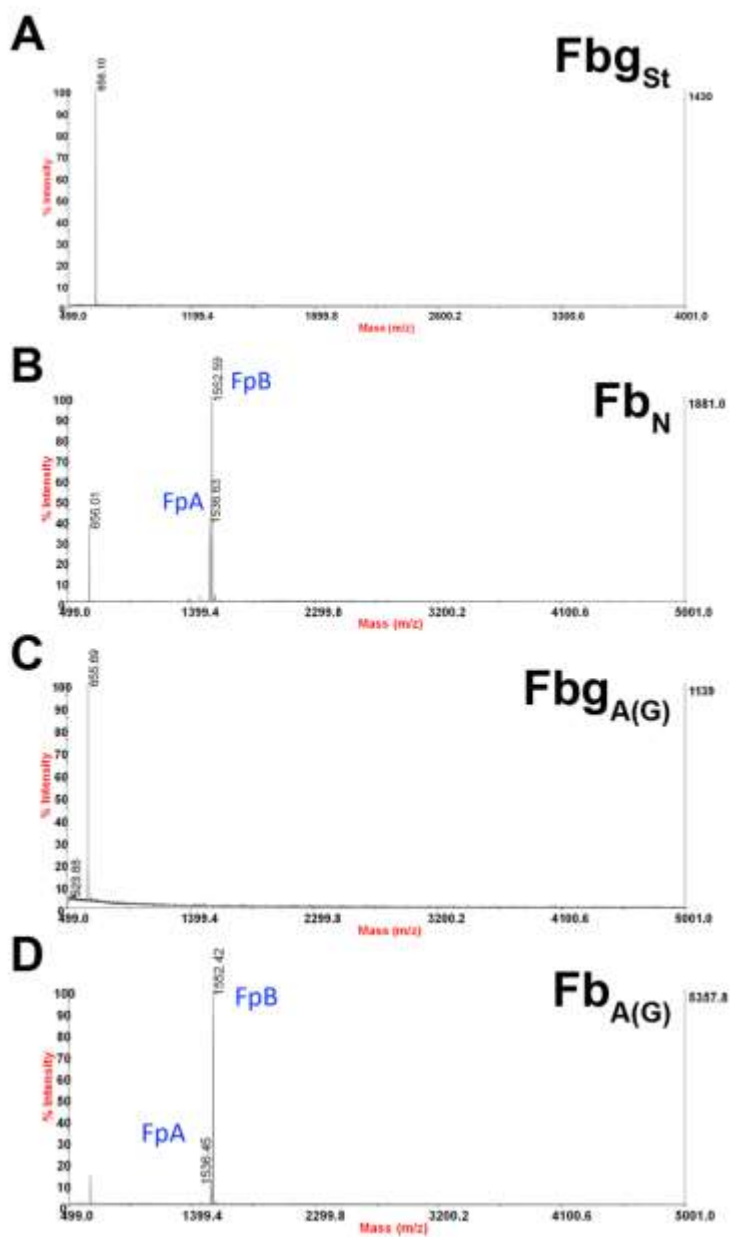


Fig. S3: MALDI TOF of fibrinogen and fibrin in various conditions. MALDI TOF spectra of: (A) fibrinogen stock solution Fbg_{St} (negative control), (B) supernatant after rinsing a fibrin clot Fb_N obtained by mixing thrombin with fibrinogen stock solution (positive control), (C) supernatant after rinsing an acidic fibrinogen gel Fbg_{A(G)} obtained after 1 hour incubation at 37°C, (D) supernatant after rinsing an acidic fibrin gel Fb_{A(G)} obtained after 1 hour incubation at 37°C.

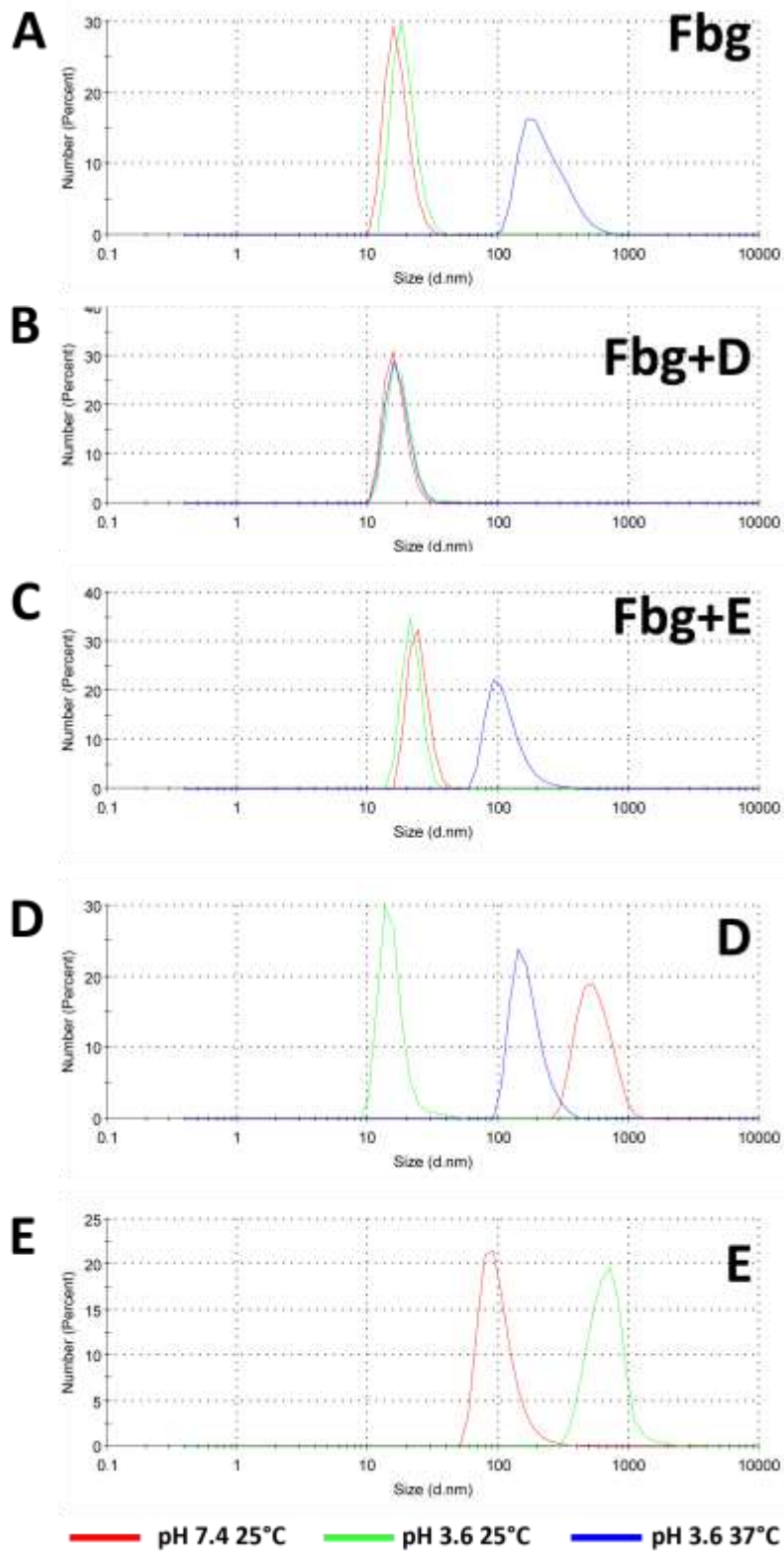


Fig. S4: Acidic aggregation of fibrinogen with or without D or E fragments monitored by Dynamic Light Scattering (DLS). DLS counts (number) of fibrinogen (A), fibrinogen + D fragments (B), fibrinogen + E fragments (C), D fragments (D) and E fragments (E) solutions at pH 7.4 25°C (red), pH 3.6 25°C (green) and 37°C (blue).

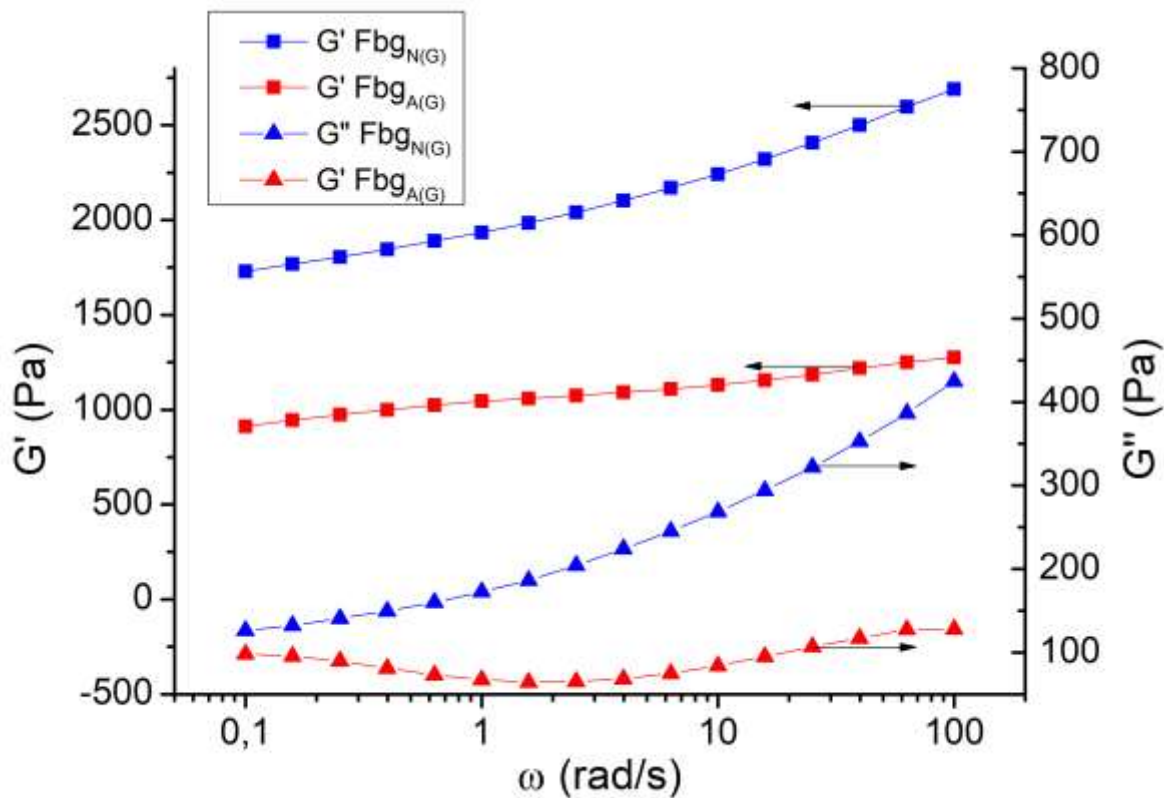


Fig. S5: Difference in frequency between an acidic fibrinogen gel $Fbg_{A(G)}$ and a neutralized fibrinogen gel $Fbg_{N(G)}$. Same rheological study ($\gamma=1\%$) performed on an acidic fibrinogen gel $Fbg_{A(G)}$ (red), polymerized by 37°C-incubation between the geometries prior to measure, and a $Fbg_{A(G)}$ gel subsequently neutralized ($Fbg_{N(G)}$, blue).

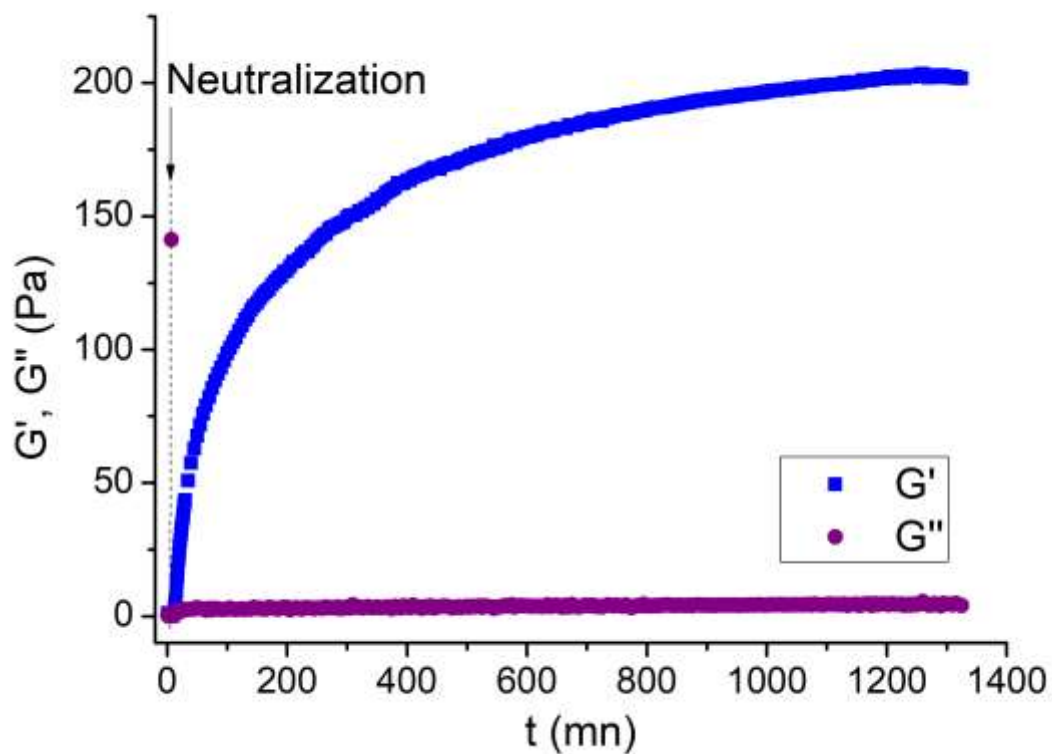


Fig. S6: Neutralization of a viscous acidic fibrinogen solution FbgA(VS). Measurement of the rheological properties of a pre-incubated solution of acidic fibrinogen FbgA(VS) is performed. Five minutes after the start of the measure, a neutral 100 mM Hepes solution is poured around the geometry.

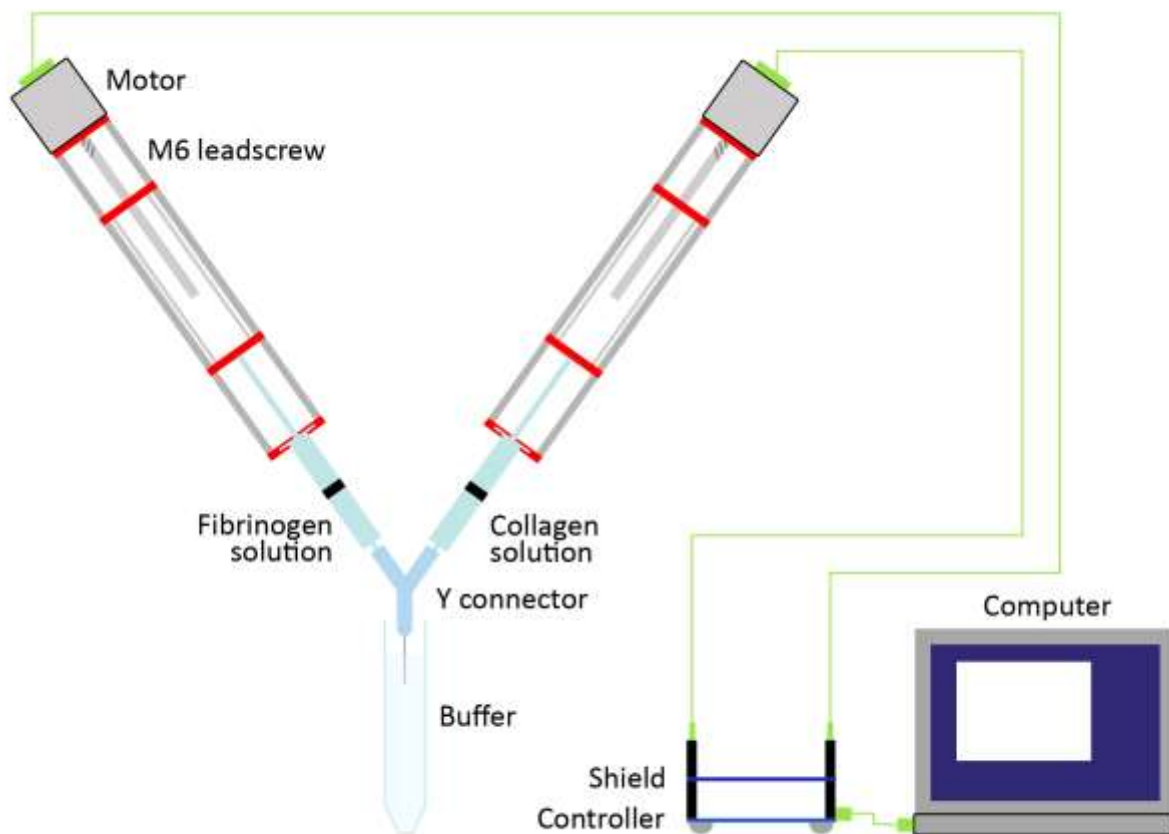


Figure S7 : Schematic of the double extrusion device for gradient production. This lab-made device was designed to dynamically tune fibrin(ogen) and collagen content and produce collagen to fibrin(ogen) gradients thanks to controlled extrusion of both syringes thanks to a controller and to a Y connector to mix both solutions.

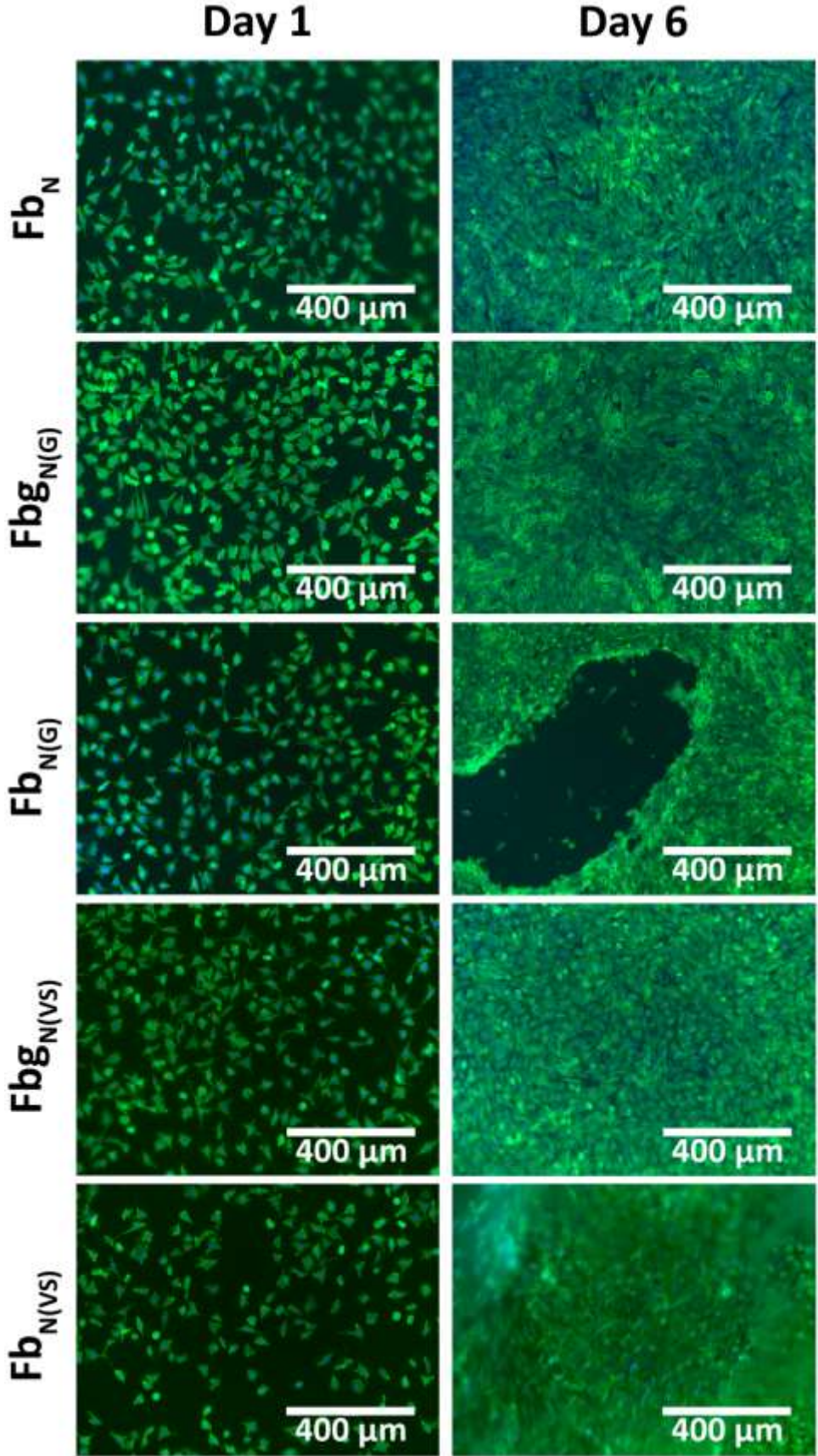


Fig. S8 : Effect of the different fibrin(ogen) materials on murine fibroblasts. Immunofluorescence images of L929 murine fibroblasts seeded at 2×10^4 cells/cm² in wells coated with “classical” fibrin Fb_N, acidic fibrinogen and fibrin gelled by incubation Fbg_{N(G)} and Fb_{N(G)}, and viscous acidic fibrinogen and fibrin gelled by neutralization, Fbg_{N(VS)} and Fb_{N(VS)} and observed at day 1 and 6. Blue: nuclei, Green: actin.

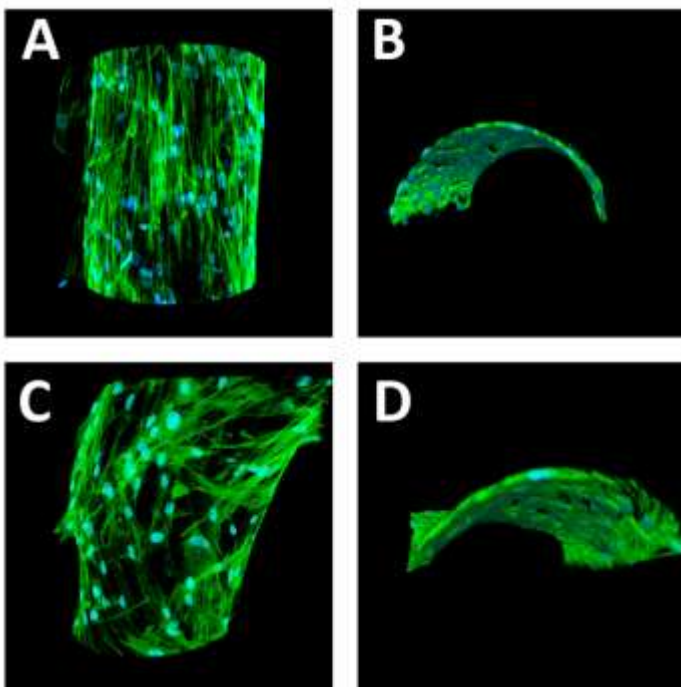


Fig. S9 : Fibrin(ogen) threads colonization by primary human fibroblasts. Confocal immunofluorescence images of NHDFs seeded on fibrin Fb_{N(VS)} (A & B) and fibrinogen Fbg_{N(VS)} (C & D) threads. Blue: nuclei, Green: actin. Dimensions: A&B 394x394x158 μm; C&D 394x394x119 μm.

Parameters	Solutions	Dissolution/Kinetics
Neutral pH	20 mM citrate pH 7.8	No
Basic pH	100 mM Tris pH 10.7	No
Ionic strength	PBS 10X	No
Temperature	100°C, 1h30, 20 mM citrate pH 7.8	No
Weak bonds	8 M Urea	Yes/ 10 to 15 mn
Disulfide bonds	100 mM DTT pH 5/6	Yes/ few days
Dilution	MQ water	No
Salt-in	100 mM KSCN	No
Salt-in	100 mM KI	No
Salt-in	100 mM CaCl ₂	No

Table S1: Solubility of acidic fibrinogen gels FbgA(G) in different solutions.

Video S1: Printing of a pure viscous fibrin solution in a Petri dish.

1. Doolittle RF (2010) Fibrinogen and Fibrin. *Encyclopedia of Life Sciences*, ed John Wiley & Sons, Ltd (John Wiley & Sons, Ltd, Chichester, UK). Available at: <http://doi.wiley.com/10.1002/9780470015902.a0001409.pub2> [Accessed September 16, 2016].