

Title

A naturally derived biomaterials formulation for improved menstrual care

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Summary

Adequately managing menstruation is an important factor for the overall quality of life for women. With a growing discussion of the global need for improved menstrual health and hygiene, it is clear that better management of menstruation can positively influence social, educational, and professional outcomes. Herein, we describe a biopolymer-based formulation that gels blood in a mechanism alternative to coagulation. We first tested several biopolymer mixtures with blood and quantified increases in viscosity, identifying that high-molecular-weight alginate in combination with glycerol could rapidly absorb blood. We then demonstrated that this powder could be deployed as both a traditional menstrual pad filler and as an additive to menstrual cups to reduce leakage and spillage, respectively. Finally, we include an antimicrobial polymer to impair the growth of *Staphylococcus aureus*, a bacterium associated with toxic shock syndrome. Collectively, our work describes a biodegradable formulation derived from renewable resources, that can improve menstrual care.

Introduction

Menstruation is a natural biological process for which cultural stigmas and an absence of societal or infrastructural support can have profoundly negative effects for women and girls. Menstruation can last between three to seven days and occurs at monthly intervals roughly from ages 13.6 to 49.6, with a median of 451 cycles per lifetime.¹ Among several menstruation-associated factors that disproportionately impact women, poor performance of menstrual products is a major factor in absenteeism,² especially in regions where menstruation is considered taboo.³ In low-to-middle income countries (LMIC) as well as low-income areas in the United States,⁴ the inaccessibility or unaffordability of menstrual products can lead to alternatives (newspapers, rags, leaves) that increase the risk of vaginal infection.⁵

Another concern with current menstrual products is the environmental consequences of waste generated by single-use disposable pads and tampons. A woman can use up to 15,000 disposable menstrual products in her lifetime⁶ with most products containing synthetic materials either as supporting components (topsheets and backsheets in pads, or strings in tampons), as packaging or as the absorbent component for some products (*i.e.*, superabsorbent polymers like crosslinked polyacrylic acid (PAA)).⁷ PAA, which is also used in other absorbent products like diapers and absorbent pads, is non-degradable with <1% degrading under landfill conditions.⁸ Because of environmental concerns, reusable menstrual cups have become more popular. Regardless of the type of menstrual product, the challenges associated with menstrual care, such as leakage and spillage during use and management, can lead to shame and distress that leads to withdrawal from daily activities.⁹

To improve the quality of life in women undergoing menstruation, we aimed to develop a biomaterials-based menstrual product that would improve the management of menstrual fluid by reducing its leakage during use, by minimizing its spillage during changing, and by simplifying the process to change/replace menstrual products in the absence of appropriate facilities. In using a biodegradable material to absorb this fluid and increase its resultant viscosity, we hypothesized that we could improve menstrual hygiene and care with broadly familiar products in a more sustainable manner.

In this study, we first characterized an array of polysaccharides for their ability to increase the viscosity of blood. Among these materials, we found that high molecular weight alginate in combination with a glycerol additive could rapidly absorb and gel blood to produce a highly viscous gel that maintained stiffness for long durations. We then incorporated this powder formulation into common menstrual care products. We found that when used as a menstrual pad filler, this formulation substantially reduced blood permeation with increased blood retention and reduced transfer when compared to other common absorptive fillers. We then used this formulation as an additive for menstrual cups and showed that it eliminated blood spillage during *in vitro* menstrual cup removal. Finally, to minimize bacterial growth, we showed that the incorporation of an antimicrobial polymer into this powder formulation inhibits *Staphylococcus aureus* without impairing the formulation's ability to solidify blood. Collectively, our work describes a biodegradable, naturally-derived biomaterial product capable of improving menstrual care.

Results

Polysaccharide solutions mixed with blood can substantially increase viscosity

As a practical assessment of viscosity, we developed an assay in which we measured the time required for an 80 μ L droplet to flow down an acid-cleaned glass test tube (**Figure 1A**). Using increasing concentrations of low molecular weight (LMW) alginate, we confirmed that droplet flow time was dependent on the tube tilt angle (**Supplemental Figure S1**) and had a good correlation with kinematic viscosity as measured by a cannon-Fenske viscometer (**Figure 1B**). Menstrual fluid is a complex mixture of blood, tissue, and mucus that is unable to coagulate¹⁰ due to an abundance of fibrinolytic proteases.¹¹ To mimic human menstrual blood, we used defibrinated porcine blood, which has previously been used as the basis for simulated menstrual fluids.¹² To determine whether polysaccharides can increase the viscosity of blood, we prepared well-mixed equi-volume mixtures using 1 % wt/vol polysaccharide solutions combined with water or blood (0.5 % wt/vol final polysaccharide concentration) and measured their flow time. Several polysaccharides had high flow times when mixed with water or blood (**Figure 1C**), but only kappa-carrageenan, high molecular weight (HMW) alginate, xanthan gum, and iota-carrageenan had significantly greater flow times when mixed with blood compared to water. Alginates in this test were in the sodium form, lacking divalent cations and are hereafter referred to as alginate. When testing LMW alginate samples that would be more sensitive to degradation over time than HMW alginate, we found that flow times had an expected dependence on concentration, which remained consistent over time (**Supplemental Figure S2**). Despite the increased viscosity of well-mixed polysaccharide-blood mixtures, their practical application would depend on their miscibility with menstrual fluid. To test this, we added blood to select polysaccharide solutions without mechanical agitation and found that only alginates could reach homogeneity (**Figure 1D**). Inversion of tubes after the 4 h incubation showed the formation of a cohesive hydrogel when blood was mixed with HMW alginate, but not the other polysaccharides (**Figure 1E**). Alginate is well known for hydrogel formation mediated by divalent cations, as shown with calcium-mediated crosslinking,¹³ a mechanism also identified in carrageenans.¹⁴ Calcium is present in blood¹⁵ and vaginal fluid throughout the menstrual cycle,¹⁶ so it is likely that calcium-mediated crosslinking contributes to the increased viscosity of alginate mixed with blood. To investigate the properties important for flow time when mixed with blood, we tested several alginates with varied molecular weights and ratios of D-mannuronate to L-guluronate (M/G) as well as comparing water, NaEDTA-treated blood, and defibrinated blood. At low viscosities, M/G ratios and use of EDTA as an anticoagulant influenced flow time, but these effects were dampened at high alginate viscosities where polymer length is the dominant factor (**Supplemental Figure S3A-B**). The viscosity of HMW alginate with different blood samples also revealed an important role for plasma in the gelation process. As shown in **Supplemental Figure S3C**, when divalent cations are chelated by EDTA, the resultant alginate-blood mixture has a faster droplet flow time than alginate mixtures with defibrinated blood. Comparable experiments with plasma instead of whole blood showed that plasma from NaEDTA-treated blood flowed faster by an even greater margin than plasma from defibrinated blood. Together, these results indicate that calcium in the plasma is a major driver of gelation with alginate. Additionally, we characterized the molecular properties of the LMW and HMW alginates used. As one would expect, size exclusion chromatography analysis revealed that the molecular weight of LMW alginate was lower than that of HMW alginate, though the difference is only ~2 to 4-fold. Additionally, NMR characterization of M/G ratios revealed 1.24 to 1.81, respectively (**Table S1**). Collectively, these results indicate that the molecular weight of alginate plays an important role in the calcium-mediated gelation of defibrinated blood.

Next, we examined the alginate distribution in static mixtures of alginate solutions with blood by tracking the diffusion of rhodamine-labeled alginate spiked into the initial alginate solutions. We found that from its initial position in the bottom of the tube, alginate diffused to the top region (**Supplemental Figure S4A**). Alginate was confirmed to increase the viscosity of top, middle, and bottom regions by droplet flow test (**Supplemental Figures S4B and S4C**).

Alginate-based powders are highly absorptive for blood

For practical application, we aimed to develop a powder formulation that could absorb and solidify blood. Powders were prepared by solubilization in specific aqueous solutions, dried, mechanically ground, and then sieved by size (**Figure 2A**). To test blood retention, we added 0.5 g of each test powder to the top of a ~50 mm² mesh screen, that was retained within a 25.4 mm inner diameter cylinder (**Figure 2B**). We tested the absorptivity of these powder formulations by adding 5 mL of blood and incubating for 3 min, after which we removed the cylinder to allow unabsorbed blood to bypass the powder and flow through. Initial tests with non-sieved powders showed that alginate in the absence of glycerol (“alginate:glycerol (1:0.0)”) performed similarly to the gold standard superabsorbent polymer, cross-linked polyacrylate (**Figure 2C**). During experiments, we found that blood did not completely permeate the alginate:glycerol (1:0.0) powder, leaving a dry core. We then prepared alginate powders containing several concentrations of glycerol, which has previously been used as a plasticizer in films and increases its hydrophilic character.^{17,18} Of the several alginate-to-glycerol ratios, we found that a 1:0.5 ratio led to maximal blood absorption (**Figure 2C**). For this ratio (AG^{0.5}), particle sizes ranging 0.2-0.5 mm had superior performance (**Figure 2D**), possibly due to the surface/volume ratio that enhances contact between polymer and blood with little powder aggregation. To determine if blood absorption was maintained over time, we incubated powder-blood mixtures for up to 60 min and found that the alginate-glycerol powders rapidly absorbed blood to levels that were maintained for the duration of the experiment, whereas polyacrylate and alginate powders absorbed blood slowly during the first 20 min and then phase-separated by 60 min (**Figure 2E**). We then measured the release of alginate from these gels using a powder mixture spiked with alginate rhodamine, and found that ~13% of alginate was released after 8 h of incubation in blood and PBS (**Supplemental Figure S5**).

Materials properties of alginate-glycerol formulations

For insight into the performance of the alginate-glycerol material, we examined its material properties. Using contact angle measurements, we confirmed the greater hydrophilic nature of alginate-glycerol films (*i.e.*, lower water contact angles) compared to alginate films (*i.e.*, higher contact angles), an effect that does not substantially change over 100 s (**Figure 3A**). Examination of the morphology of these powders by scanning electron microscopy at low magnification shows a generally irregular particle shape for pristine alginate, AG^{0.5} and polyacrylate particles (**Figures 3B, 3C, and 3D**, respectively). Higher magnification shows that alginate has a dense internal structure (**Figure 3B, inset**), similar to what has been observed previously for alginate powders¹⁹ whereas the incorporation of glycerol leads to greater texture and a more porous open structure (**Figure 3C, inset**). By comparison, crosslinked polyacrylate has a smooth surface morphology that is indicative of a crystalline structure previously observed²⁰ (**Figure 3D, inset**). Particle sizes determined from the micrographs revealed that pristine alginate had the smallest projected area and diameter, followed by AG^{0.5} and polyacrylate (**Figure 3E-3F**). We then determined whether the calcium in blood had a similar gelation effect as calcium in buffer by comparing their rheological properties when mixed with alginate-glycerol powder. Using a strain sweep, we determined the linear viscoelastic range and used 1 % for frequency sweep analysis (**Supplemental Figure S6**). As shown in **Figure 3G**, the storage and loss moduli are greater when mixed with blood compared to CaCl₂, which suggests that other components of blood (*e.g.*, cells) play a role in the gelation process. Additionally, the greater magnitude of storage to loss modulus in both conditions supports a stiffer gel-like behavior previously observed with alginate hydrogels.²¹ ICP-MS analysis of the defibrinated porcine blood we used showed an extracellular Ca²⁺ concentration of 2.47 mM, which is similar to what is found in menstrual fluid (2.5 mM²²) and similar to the 2.5 mM used in the calcium buffer. Taken together, these results indicate that the inclusion of glycerol into the alginate powders likely improves its dissolution into blood and that these AG^{0.5} mixtures with blood have comparable rheological properties to mixtures with aqueous CaCl₂ solutions.

Alginate-glycerol powders absorb blood with high retention as menstrual pads

To test the absorptivity of our alginate-glycerol powders, we simulated their application as a menstrual pad, but without topsheets or backsheets to enable the measurement of blood leakage. As

shown in **Figure 4A**, various fillers (nothing, crosslinked polyacrylate, AG^{0.5}, or the commercial cellulose-based filler) were spread as a single layer within a folded sheet of cotton gauze and then taped to the opening of a silicone-based artificial vagina (**Figure 4B**). We used this device to simulate the channeling of menstrual fluid by the vaginal canal onto an irregular but focused area, testing the effect of overloading a small area. We also did not observe significant differences between the absorptive capacity of commercial tampons tested with defibrinated blood or an FDA-recommended Syngina Fluid (**Figure S7**). We tested for blood permeation by rapidly (< 30 s) adding 8 mL of blood to the top of the artificial vagina. This volume within a short timeframe simulates prolonged accumulation (e.g., sleeping or a change in body position) or spontaneous events (e.g., laughing, sneezing, coughing). After 1 h, the pad was removed, and the amount of blood that permeated the pads or was not absorbed into the pad (*i.e.*, retained above) was measured, as shown in **Figure 4C**. Our alginate-glycerol powder and the commercial cellulose filler had similarly high levels of blood absorption and low levels of excluded (*i.e.*, permeated or retained) blood compared to the absence of a filler and the polyacrylate filler (**Figure 4D**). While absorptivity is an important parameter, the transfer of moisture can be a source of discomfort that can lead to rashes. To mimic mild compression and contact that might lead to moisture transfer, we placed blood-absorbed pads on top of 96-well microtiter plates, centrifuged them at low speed, and measured the quantity of blood that was discharged (**Figure 4E**). We found that the commercial filler released nearly all of the absorbed blood, comparable to the empty gauze, whereas the polyacrylate retained an intermediate quantity of blood, and the AG^{0.5} filler retained the maximum amount of blood (**Figure 4F**). This effect can also be observed by gently squeezing these materials (**Supplemental Video S1**). Collectively, these experiments show that the alginate-glycerol powder rapidly absorbs blood with high retention.

Alginate-glycerol powders eliminate spillage when used in menstrual cups

Menstrual cups are silicone cups that are inserted into the vaginal canal to create a seal against the vaginal wall, retaining menstrual fluid until it can be removed and cleaned. These menstrual products are favored over disposable products because they are reusable and have a low likelihood of leakage during use. However, poor placement and/or manipulation during removal can lead to leakage and spillage of fluid. To test our formulation as a sustainable material that can be used in complement with environmentally-friendly menstrual cups, we spread 1.5 g of the AG^{0.5} powder in a 25.4 mm cotton tube (“stockinette”) and coiled it within a menstrual cup (**Figure 4G**). Cups alone or those with stockinettes filled with nothing or AG^{0.5} powder were placed in the Syngina, a device used by the FDA to test the absorptive capacity of menstrual care products²³ (**Figure 4H**). To evaluate the retentive capacity of the menstrual cups, 15 mL of blood was added to the top to simulate a daily average of menstrual fluid.²⁴ After incubation for 4 h at 37°C, five women were blinded to the content of the menstrual cups and asked to remove them with care to minimize spillage. As shown in representative experiments (**Figure 4I**), the alginate-glycerol powder improved blood retention. When examining individual attempts, each woman spilled blood when removing menstrual cups that contained nothing or the empty stockinette, which could be highly variable between attempts. By contrast, spillage was consistently rare when the AG^{0.5} powder was used (equivalent to a few drops when occurring) (**Figure 4J** and **Supplemental Figure S8**). Outside of the Syngina, inversion of these menstrual cups shows that, unlike the empty stockinette, the AG^{0.5} powder improves blood retention (**Supplementary Video S2**), this blood is not readily exuded (**Supplementary Video S3**), and little blood remains in the menstrual cup (**Supplementary Video S4**). The summation of results shows that the use of the stockinette with powder eliminates spillage of blood to levels significantly reduced compared to empty or stockinette conditions (**Figure 4K**).

Suspension of bacterial growth

Highly absorptive menstrual products have been correlated with toxic shock syndrome, where toxigenic *S. aureus* from the vaginal tract is able to proliferate and produce toxins. This is predominantly observed in tampons,²⁵ but is possible with other menstrual hygiene products, including menstrual cups.²⁶ Alginate can potentially act as a nutrient source for some microbes and so including a strategy to inhibit bacterial growth would be important to minimize the risk of toxic shock syndrome.

Because of the significance of the endogenous vaginal microflora,²⁷ we aimed to inhibit bacterial growth within the alginate-glycerol powder without complete eradication in a minimally-leeching formulation because of potential collateral effects to the vaginal microbiota. We used trimethyl chitosan (TMC) because it has antimicrobial activity and will anchor in the alginate via ionic crosslinking and polymer entanglement to minimize leaching. When alginate and TMC were dissolved in solution in the first steps of preparation, we found significant inhibition of bacterial growth when mixed with blood spiked with 5×10^5 cfu/mL (**Figure 5A**) or 5×10^6 cfu/mL of *S. aureus* (**Figure 5B**). This effect was observed after 4 and 8 h of incubation. Mixing alginate-glycerol with TMC in powder form, without co-solubilization, led to less consistent antimicrobial activity, which is likely due to heterogeneity in TMC distribution (**Supplementary Figure S9**). To test whether including TMC altered the alginate-glycerol powder function, we measured the flow of these powders mixed with blood. As shown in **Figure 5C**, the addition of a lower weight fraction of TMC led to slight increases in flow time at low powder concentrations. TMC alone can increase droplet flow times, though not to the same levels as the alginate-glycerol formulations. Interestingly, an equal amount of TMC to alginate-glycerol dramatically reduces droplet flow times, which we expect is due to ionic crosslinking between the polyanionic alginate and polycationic TMC, sequestering alginate from calcium-mediated crosslinking and gel formation. As schematically shown in **Figure 5D**, the inclusion of TMC can impart antimicrobial properties while retaining the ability to solidify blood.

Discussion

Here, we describe the design of a biodegradable, blood-absorbent biomaterial based on a natural polysaccharide that improves the performance of menstrual products by minimizing blood leakage and spillage. First, we developed a screening method to quantify the gelation of blood by mixing several polysaccharides. This strategy identified HMW alginate as an optimal candidate because of the dramatic increase in viscosity and spontaneous miscibility with blood when tested in liquid form. For application as an absorptive powder and to maximize the dissolution of alginate powders in blood, we developed a glycerol-supplemented formulation that both accelerated the stiffening of blood into a gel-like consistency and increased its capacity for blood absorption, an effect that depended on particle size. We then tested the practicality of this material to improve menstrual care by using it as the absorptive component of a menstrual pad and as a complementary component to menstrual cups. In both cases, the use of our alginate-glycerol powder formulation minimized blood leakage and spillage. Finally, to inhibit bacterial growth, we included the quaternary ammonium polysaccharide trimethyl chitosan, which inhibited the growth of *S. aureus*, a bacterium responsible for toxic shock syndrome.

Traditional menstrual hygiene products manage menstrual fluids by their absorption or collection. While the principles can be traced to antiquity, their early modernized designs can be identified in products as early as the tampon patented in the US in 1933,²⁸ the Kotex menstrual (“sanitary pad”) marketed in 1921,²⁹ and the menstrual cup patented in the US in 1937.³⁰ Since their inception, menstrual products have focused on managing menstrual fluid in its liquid state, which faces the same challenges of any other liquid: leakage and spillage/dripping during changing or replacing the menstrual product. Although menstruation is a natural biological process and its healthy progression is not traditionally considered a disease or disorder, its poor management has a major global impact for women.³¹ In many settings, including workplaces in LMIC, the lack of access to facilities with privacy, clean water, and discreet disposal leads to anxiety and stress. This negatively impacts the productivity of women, their ability to remain in the workforce, their income, and their quality of life.³² By increasing the stiffness of blood, producing a more gel-like material, we show greater performance compared to traditional menstrual products.

Alginate has been explored in various biological applications, including wound dressings, hemostatic products, and implantable devices,¹³ but has not been demonstrated as a bulk absorbent material for biological fluids, especially menstrual fluid, which does not coagulate. Notably, in our use of alginate we rely on the blood-derived divalent cations to mediate the ionic cross-linking that leads to

gelation,³³ enabling the gelation of a non-clotting defibrinated blood that mimics the lack of coagulation in menstrual fluid.¹¹ This contrasts with other materials that leverage the presence of clotting factors within blood to mediate coagulation.³⁴ Building on its broad applications in biotechnology, our use of alginate to increase the viscosity of non-coagulating blood to improve menstrual care represents an advancement towards women's health, which is traditionally understudied.

The use of a biodegradable formulation based on renewable resources is an important consideration for menstrual care products. Current commercial products largely utilize non-degradable materials derived from non-renewable sources, such as plastics in the packaging or crosslinked polyacrylate in the absorptive material. Given the biodegradability of alginate,³³ the formulation presented in this work allows for convenient disposal of the solidified blood mixture. The established large-scale manufacturing of alginate from natural sources³³ and its approval for use in several products by the Food and Drug Administration³⁵ also simplifies its safe and scalable implementation. An additional benefit of using a known biopolymer is the option for further functionalization. With toxic shock syndrome as an important concern, we included TMC and showed that at low concentrations, it inhibits the growth of *S. aureus* without impairing the blood-gelation function of alginate, illustrating the potential versatility of alginate-based menstrual products.

While we believe our work represents exploratory but important progress in improving the quality of life for women, there remains several questions that would benefit from further investigation. For example, the large scale processing and manufacturing of alginate-glycerol materials may require additional adjustments that maintain batch-to-batch performance and homogeneity. Additional considerations should monitor several parameters including water content and particle size distribution, especially with regards to blood absorptivity. When considering use of the alginate-glycerol material as an additive to current menstrual products, understanding how powder parameters (i.e., particle size) impact the robustness of function in conjunction with menstrual pads or cups, as well as its compatibility with existing waste streams,³⁶ would be important factors. In LMIC countries where improved menstrual care products would have strong impact, cost will be a major factor. Although a more comprehensive and detailed cost analysis would be necessary, based on the amount of purified, commercially obtained materials used in this study, we estimate the amount per menstrual cup would cost ~\$0.20. To meet regulatory approval for commercialization, several additional tests would be required, including the assessment of cytotoxicity and allergenicity. Finally, to improve our understanding of the performance of menstrual care products, the development of established *in vivo* models would help address potential gaps and limitations in the evaluation of materials.

In conclusion, our work addresses a major factor in the quality of life for women globally. In addition to developing new materials for menstrual care, we hope our work will stimulate greater technological investment and scientific interest in women's health.

Experimental Procedures

Resources Availability

Lead Contact. Requests for further information or data should be directed to the lead contact, Bryan Hsu (bhsu@vt.edu).

Materials Availability. Raw materials are available from the commercial suppliers indicated in this section.

Data and Code Availability. All data reported in this paper are available from the lead contact upon request.

Materials. Low molecular weight (LMW) sodium alginic salt, l-carrageenan, k-carrageenan, pectin, and the super-absorbent cross-linked sodium polyacrylate were purchased from Sigma-Aldrich (USA). High molecular weight (HMW) sodium alginic salt, carboxymethylcellulose, chondroitin sulfate, gelatin type A, trimethyl chitosan (TMC, low molecular weight, ~85% deacetylated, degree of quaternization > 50%, #912700) and sodium hyaluronan were obtained from Thermo Scientific (USA). High-grade alginate samples with different molecular ranges and M/G ratios were ordered from the Promega Company (USA), chitosan was purchased from Polysciences Inc (USA), carboxymethyl chitosan was purchased from Santa Cruz Biotechnology (USA), and xanthan gum was purchased from TCI Chemicals (USA). Alginate rhodamine (high viscosity, 1,000-1,500 cP, AL-512) was purchased from Creative PEGWorks. Aseptically confirmed porcine blood (defibrinated or NaEDTA-supplemented) was purchased from Lampire Inc. (USA). Fuchsin acid-certified was ordered from Neta Scientific (CMX-01606-25G). Aqueous polymer solutions were prepared with ultrapure water (18.2 M Ω cm at 25 °C) from a Milli-Q® IQ-700 ultrapure water purification system (Millipore Sigma, USA).

Size Exclusion Chromatography (SEC). The molecular weight and the polydispersity index (PDI) for the alginates and TMC were determined by SEC. Samples were prepared by dissolving 0.1 – 0.5% sample in 100 mM sodium nitrate buffer and filtering through a 0.45 μ m syringe filter. 20 μ L of the sample was loaded onto a Shodex LB-806M SEC column at a flow rate of 1 mL/min. Molecular weight determination and concentration were measured by multi-angle light scattering (MALS) and refractive index (RI), respectively, via in-line Wyatt detectors. All data was analyzed in Astra 8 software and normalized to commercial dextran standards (Wyatt).

Nuclear magnetic resonance (NMR) spectroscopy. Commercial alginate samples (HMW and LMW) were first depolymerized by partial hydrolysis with HCl and resuspension in 99.9% D₂O (5-10 mg/mL).³⁷ The probe temperature was set to 353 K, followed by the insertion of the NMR tube with the hydrolyzed sample and equilibration for 15 min. Afterward, we set the relaxation delay to 2 s, locked the solvent to D₂O and the number of scans to 128. Additional details for recording the spectra and determining the sample M/G ratio are detailed elsewhere.³⁷

Biopolymer solutions. Polymers weighted in an analytical scale were dissolved at concentrations ranging from 0.5 to 6.0 % wt/vol in ultrapure water, vortexed and heated in a water bath at 50 °C until complete solubilization. For measurements in polymer/blood mixtures, aqueous polymer solutions were mixed with equal volumes of aqueous polymer and blood at room temperature and were homogenized by gently pipetting the mixture up and down. Solutions were left on the bench room temperature for 15 to 30 min to eliminate bubbles before conducting any test.

Blood-absorbent powder formulation. Alginate-based powder formulations were prepared by mixing HMW alginate to glycerol aqueous solution to test various alginate to glycerol solution ratios (**Table 1**). For powder solubilization, the aqueous glycerol solution was spread in a 101.6 mm square weighing boat, and the powder alginate reagent was evenly dispersed on the liquid surface, followed by gentle mixing at the liquid surface that formed a solid mixture. The homogeneous solid mixture was cut into small pieces with a disposable spatula and dried at 50 °C for 48 h, followed by manual grinding with liquid nitrogen to obtain mm-sized particles. The ground mixture was manually sized with a set of sieves with openings that ranged from 0.2 to 1.5 mm (US Standard Sieve Series, USA, A.S.T.M. E11)

to collect particles within different size ranges (<0.2 mm, 0.2 – 0.5 mm, 0.5 – 0.7 mm, 0.7 – 1.0 mm, and 1.0 – 1.5 mm range; for example, particles in the 0.2 – 0.5 mm size range passed through the sieve with 0.5 mm and were retained on top of sieve with 0.2 mm opening size). The processed samples were stored in at 4-8 °C before use to minimize product degradation and contamination.

Test tube flow test. Regular glass tubes were first washed with a commercial detergent and rinsed with DI water five times. These tubes were soaked in hydrochloric acid solution (1.0 M) for 1 h, followed by copious rinsing with DI water and three final rinses with ultrapure water and drying at 70 °C. Polymer aqueous solutions were mixed with equal volumes of ultrapure water or porcine blood (defibrinated or NaEDTA-treated) by pipetting the mixture up and down with low-retention tips (VWR, USA). For tests with blood supernatant, defibrinated or NaEDTA-treated blood was centrifuged at 2,000× g for 15 min at 4 °C, and the supernatant was collected for blood mixing. After complete bubble removal, aqueous or blood polymer mixtures (80 µL) were pipetted ~ 10 mm below the top of the pre-cleaned test tubes positioned at an angle of 45° at room temperature to track the time interval for the droplet to travel 80 mm towards the tube bottom starting (triplicate measurements) (**Figure 1A**). The maximum flow time interval observed was 8 h for mixtures that did not flow.

Flow viscometer measurements. Kinematic viscosity measurements were performed in pre-calibrated Cannon-Manning glass viscometers (viscometer sizes 75, 150, 200, 300, 400, 500, and 600) with pre-mixed aqueous or blood polymer solutions. After sample loading, the viscometer was placed in a water bath at 40 °C and kept for 15 min to reach equilibrium before measuring the mixture flow time.³⁸ The kinematic viscosity was calculated by converting the efflux time in seconds by the viscometer constant (**Table S2**) at 40 °C for measurements in triplicate.

Spontaneous alginate/blood mixing. The HMW alginate solution (1 % wt/vol) was first prepared with 1:100 of alginate rhodamine by mixing the fluorescent polymer into the HMW alginate solution protected from light. The polymer (1 mL) was transferred to a 5 mL conical tube, and 1 mL of blood was carefully added on top of the polymer phase and at room temperature without shaking. Fluorescence was measured from sample aliquots (80 µL) by resuspension in NaEDTA solution to a final concentration of 100 mM NaEDTA, followed by centrifugation at 2,000×g for 15 min at 4 °C. Fluorescence was measured from the supernatant in an Agilent Biotek Cytation plate reader with emission and excitation wavelengths of 520 nm and 583 nm, respectively, in black 96-multiwell plates. The total alginate concentration was determined based on the concentration of fluorescent alginate and the ratio of HMW alginate to alginate rhodamine.

Blood absorption capacity. Powder formulations (PAA, or the formulated alginate-based mixtures with different alginate to glycerol ratios, 0.5 g) were added to a 25.4 mm cross-sectioned acrylic tube (~25.4 mm height) on top of a pre-weighted HDPE screen (46 mesh, ~50 mm²), above a pre-weighted Petri-dish (101.6 mm diameter) (**Figure 2A**). Defibrinated blood (5 mL) was transferred at a rate of 5 mL/min using a syringe pump (Model 300, New Era Pump System, USA) at the top of the powder, and the system was maintained on the bench to measure blood absorption over time (static, room temperature). At the end of the experiment, the cross-sectioned tube was removed, and the contents collected in the Petri dish and retained at the top of the screen were weighed to determine the amount of blood that leaked from and retained in the powder for measurements in triplicate.

External tamponade test (menstrual pads). The powder formulation was first tested as a menstrual pad filler and challenged for blood retention in a silicone-based, anatomically similar vaginal model (Ice Lady-Clear, 247.65 mm×98.43 mm (L×W), Interactive Life Forms, LLC, USA) (**Figure 4A-C**). Powder formulations (1.5 g) were added to a cotton-based gauze (~1.5 g, 50.8 mm × 76.2 mm) for a simple yet practical design of the external tamponade element in a pad format for testing blood retention and loss during the pad transfer. The pads filled with different materials (AG^{0.5}, PAA, or the commercial absorbent element from pads) were taped underneath the vaginal model, and the defibrinated porcine blood (8 mL) was added from an opening at the top using a transfer pipette in a ~30 s interval. After 1 h of incubation at room temperature, blood leakage and retention were determined by weighing the blood that flowed through and the blood retained in the tamponade element. To determine blood release due

to mechanical force, pads were detached from the vaginal model and placed on top of pre-weighted 96 well plates, followed by spinning in a swing bucket centrifuge (centrifuge 5810-R Eppendorf, rotor A-2-DWP) at 1,000 g for 5 min at 25 °C. The pad and the plate were weighted to determine the fractions of blood retained and released in the pad, respectively.

Internal tamponade test (menstrual cups). The alginate powder formulation (AG^{0.5}) was also assessed as a filler in the menstrual cups as a method to minimize spillage and blood loss during cup removal. This test was conducted with an in-house designed Syngina (synthetic vagina) adapted from the FDA-approved methodology for testing tampons absorbency,²³ using a bottle with a larger opening (~35.9 mm inner diameter) and a circulating water-heating system kept at 37 °C (**Figure 4G-H**). Powder formulations (~1.5 g) were dispersed in a 25.4 mm tube-shaped stockinette (~1.0 g, 152.4 mm length) and the tube was arranged in a helical shape inside of a regular menstrual cup (CVS Health, cup size A, maximum working volume 15 mL, medical-grade silicon, #722537). The menstrual cup was placed inside the Syngina, and defibrinated blood (15 mL) was transferred to the cup with a transfer pipette (~5 mL/min) from the top opening. After the incubation period (1 h at 37 °C), the menstrual cup was removed using a pre-weighted white superabsorbent glove with a pre-weighted white absorbent pad underneath the experimental setup, and both the glove and the pads were weighted and imaged to determine the blood messiness during menstrual cup removal. A blinded, randomized test was conducted in duplicate for each sample (empty cup, filled with the empty stockinette, and filled with the stockinette containing the alginate powder) by different female coworkers from Hsu Lab.

Absorptivity test of commercial tampons. The tampon absorptivity test was conducted with pre-weighted regular-size tampons (o.b.[®] Original[™] regular tampons, Playtex Manufacturing Inc.) placed in the center of the Syngina at 37 °C. The tested fluid was pumped from the top of the Syngina at 50 mL/h using a syringe pump (Model 300, New Era Pump System, USA) and stopped when the tampon was visually saturated and the fluid started leaking from the tampon.²³ The tampon was weighed again on an analytical scale, and the amount of fluid absorbed was determined by the weight difference before and after fluid absorption.

Alginate release test. To study the release of alginate from the polymer/blood mixture, the blood-absorbent powder was prepared with an aqueous glycerol solution containing alginate rhodamine (0.2 mg/mL), leading to a mixture with 1:100 of alginate rhodamine to HMW alginate. The powder mixture was processed as described above (see Blood-absorbent powder synthesis section), protected from light. The processed powder was mixed with defibrinated blood (1 g powder per 10 mL of blood), and 300 µL of gel was transferred to the bottom of a 24-multiwell plate. Phosphate buffered saline (PBS) or defibrinated porcine blood (2 mL) was added on top of the gel, and the plate was incubated at 37 °C on a rocker with orbital shaking (50 RPM) for 24 h. Sample aliquots (80 µL) were collected and processed as described in the Spontaneous alginate/blood mixing methodology section. The total amount of alginate released was based on the alginate rhodamine signal, the ratio of alginate rhodamine to alginate, and the initial concentration of alginate in the gel.

Antimicrobial assay. The AG^{0.5} formulation was supplemented with the antimicrobial agent TMC (0.2 to 20 mg per 20 mg of alginate) to investigate the antimicrobial performance of blood-absorbent powder against *S. aureus*. The homogenous formulation was prepared by adding TMC during the alginate solubilization in the glycerol aqueous solution, followed by snap freezing at -80 °C and lyophilization for at least 24 h. After drying, the powder was crushed and used for the antimicrobial test. The heterogeneous formulation was prepared by vigorously mixing the pre-processed AG^{0.5} formulation with TMC prior to the antimicrobial test. For the bacterial inoculum preparation, *S. aureus* was streaked from the frozen bacterial stock in a Luria Broth (LB) agar plate and incubated at 37 °C overnight. One single colony was used to inoculate 5 mL of LB broth in a test tube, followed by overnight incubation at 37 °C and 200 rpm. The overnight cell culture was back-diluted to an OD₆₀₀ of 0.125 (~10⁸ cfu/mL) and spiked in aseptically-tested defibrinated porcine blood to a concentration of 5×10⁵ and 5×10⁶ cfu/mL of *S. aureus*. The TMC-supplemented powder (20 mg) was transferred to the bottom of 24-well plates, followed by the addition of 1 mL of blood with different bacterial loads (tests performed, at least, in triplicate for each powder formulation), and incubation at 37 °C and 200 rpm in an incubator with orbital

shaking. To test bacterial inactivation, blood mixture aliquots (100 μL) were collected after 4 h and 8 h, serially diluted in LB broth and plated in LB agar plates. After overnight incubation at 37 °C, we determined the number of colony-forming units and converted them to the final bacterial concentration in the gelled blood mixture. The antimicrobial performance was determined as the fold-change (FC) in the bacterial concentration of polymer-containing samples compared to samples with no polymer added.

Morphological analysis. For the morphological analysis through Scanning Electron Microscopy (SEM), powder samples of pristine alginate, superabsorbent, and AG^{0.5} were placed over the copper tape. The samples were then coated with 10 nm Pt/Ir coating using the sputter for a few min and probed using JEOL IT500 (JEOL) with an accelerating voltage of 5 kV, and magnifications of 50X and 500X. The minimum and maximum particle dimensions, as well as the particle area, were calculated from the 50X images using the ImageJ software.

Contact angle measurements. The surface hydrophobicity of the pristine alginate and AG^{0.5} powder was investigated through water contact angle by using an optical tensiometer (Biolin Scientific Theta Flow). A thin film of alginate and alginate glycerol powder was prepared over the glass slide. The film was prepared with a 1% solution of alginate and AG^{0.5} in water, which was then poured over the glass slide and dried in the oven at 50 °C. The sessile drop mode of the OneAttension software was used for the static contact angle measurement. A drop of water (5 μL) was placed on the thin layer of the pristine alginate and AG^{0.5} film using the automatic dispenser and imaged with a 5 MP resolution against monochromatic light for 100 s in the equipment software. The angle at the left and right sides of the droplet was automatically determined by the software, and their average was used as the contact angle of the sample (measurements in triplicate).

Rheology. The strength of the AG^{0.5} gels with CaCl₂ and defibrinated porcine blood was assessed through the oscillatory rheological technique using an Anton Paar MCR302 rheometer (Anton Paar GmbH). The strain and frequency sweep were conducted with a 50 mm parallel plate geometry of the rheometer. The gels were prepared by mixing 1 g of AG^{0.5} powder with 10 mL of blood or CaCl₂ (2.50 mM) solution and leaving them on the bench to remove bubbles. The porcine-defibrinated blood used presents 2.47 mM of Ca²⁺ measured by the ICP-MS method. The strain sweep was performed to determine the optimum strain value from the linear viscoelastic region for each gel at 37 °C. The storage (G') and loss (G'') modulus were recorded as a function of frequency between 0.1 to 100 Hz.

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Author's Contributions

Conceptualization, B.B.H.; Investigation, B.B.H., R.A.B., H.K., J.M., L.G., C.C.; Resources, B.H.H.; Writing – Original Draft, B.H.H., R.A.B., H.K.; Writing – Review & Editing, B.H.H., R.A.B., H.K.; Funding Acquisition, B.B.H.

Declaration of Interests

R.B. and B.B.H. are inventors on a pending patent application related to the materials described in this paper.

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Figures and scheme titles and legends

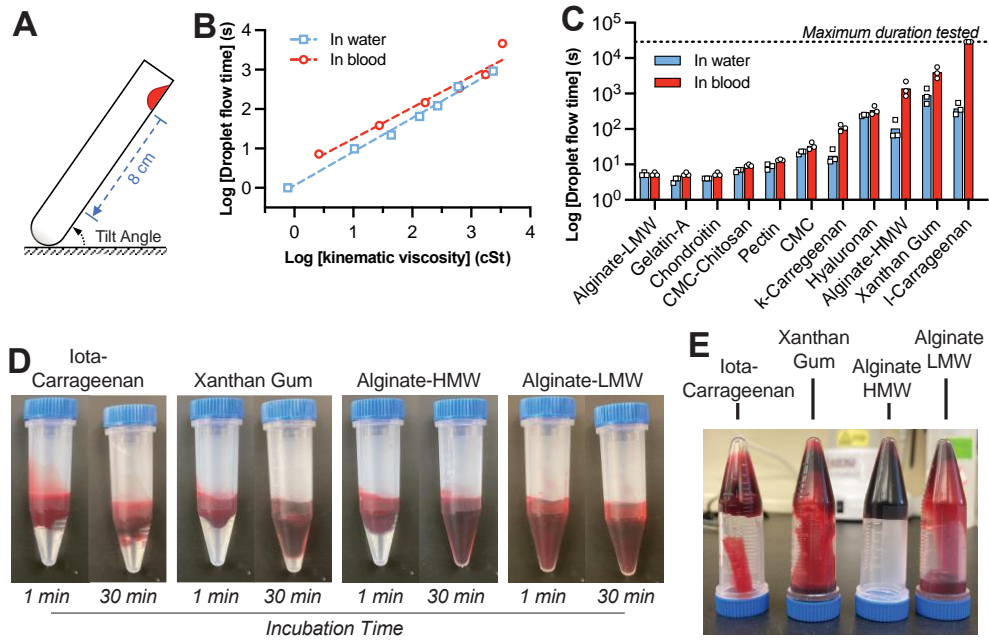


Figure 1. Biopolymers can increase the viscosity of well-mixed solutions of water and defibrinated blood. (A) Setup of the droplet flow test. (B) Droplet flow time scales proportionally with the kinematic viscosity of the alginate solution. (C) The screen of flow times for biopolymer solutions well mixed with water or defibrinated blood to a final concentration of 0.5 % wt/vol. Statistical comparisons were performed by two-way ANOVA with a Bonferroni multiple comparisons test. ****, $p < 0.0001$ (D) Highly viscous biopolymer solutions can remain phase separated from blood if not well mixed, which (E) affects their ability to solidify blood. Bars in (D) and (E) correspond to 20 mm.

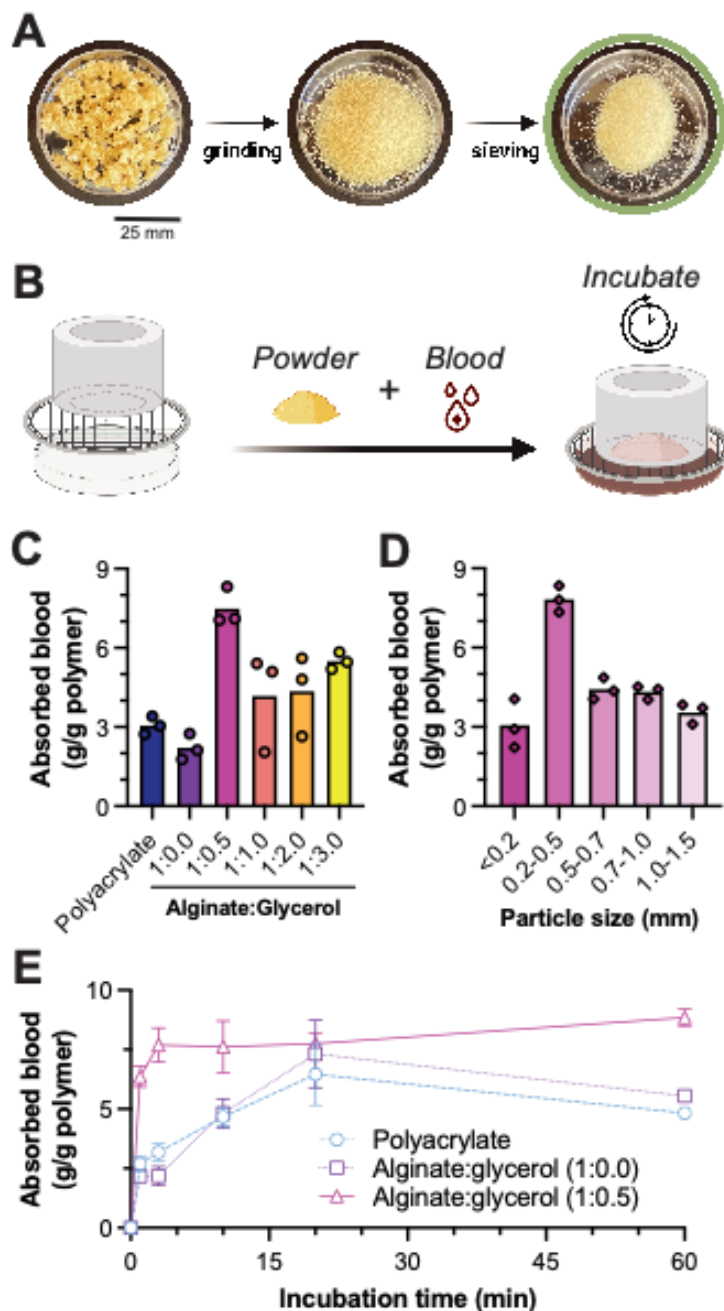


Figure 2. Alginde-based formulations can be tuned to promote maximum blood absorption capacity. (A) Images of the processed alginde-glycerol formulation after drying, grinding, and sieving with particle sizes ranging from 0.2 to 0.5 mm. Bar corresponds to 25 mm. (B) Schematic representation of the setup used for testing powder blood retention. (C) Blood retention for alginde formulations presenting different alginde to glycerol ratios (particle size of 0.2-0.5 mm) after 3 min of incubation. (D) Impact of the particle size on blood retention for AG^{0.5} after 15 min incubation. (E) Time series experiment to determine blood retention for super-absorbent PAA, high-molecular-weight alginde reagent, and the AG^{0.5} (particle size 0.2-0.5 mm).

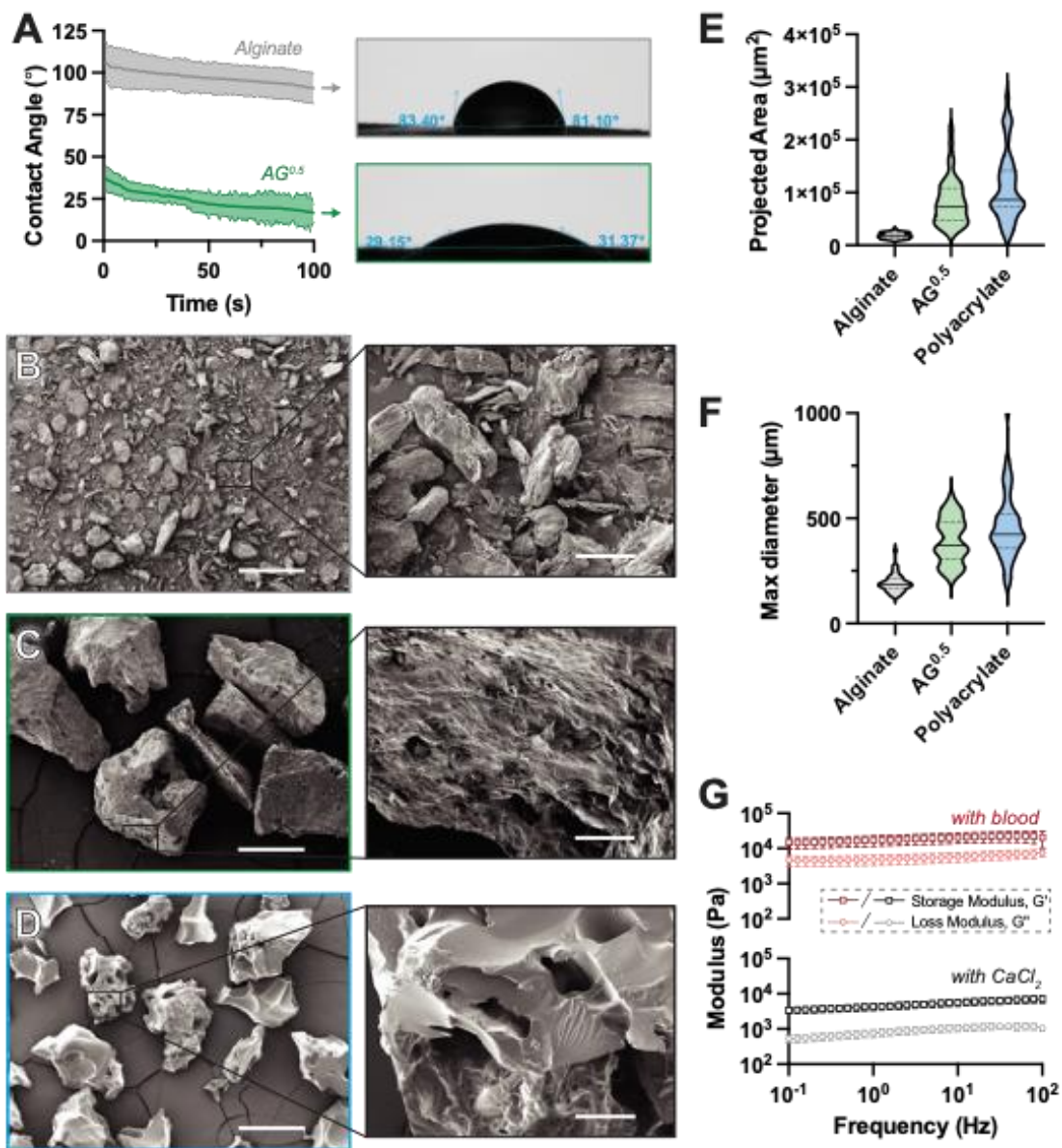


Figure 3. Materials properties of alginate formulations. (A) Water contact angle measurements of alginate or $AG^{0.5}$ films over time with representative images. (B) SEM micrographs of alginate, (C) $AG^{0.5}$, and (D) cross-linked polyacrylate particles. Scale bars represent $500\ \mu\text{m}$ for the low magnification figures or $50\ \mu\text{m}$ for the higher magnification inset figures. (E) Projected particle area and (F) particle diameter were calculated from the micrographs using ImageJ. (G) Frequency sweep of $0.1\ \text{g/mL}$ of pristine alginate or $AG^{0.5}$ powders mixed with blood or calcium chloride solutions at a final concentration of $2.5\ \text{mM}$.

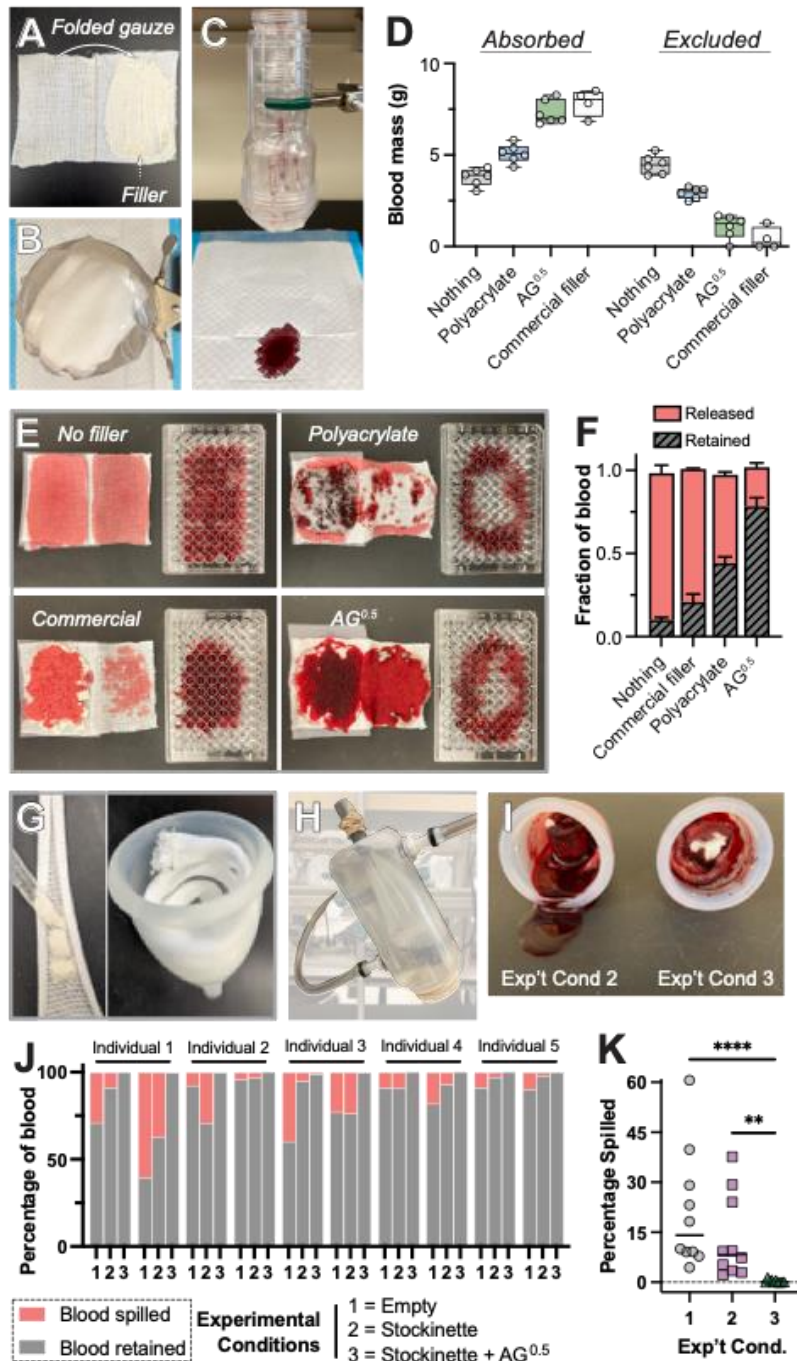


Figure 4. Application of alginate-glycerol formulations for menstrual care. (A) Scheme pad assembly and (B) attachment to a silicone-based artificial vagina. (C) Representative experiment showing the result of blood application to the device and permeation through a pad. (D) Blood retained within the pad was measured, as well as blood that permeated the pad or remained unabsorbed but pooling above the pad. (E) Representative results of testing for blood retention in the used pads and (F) the measured results. (G) For testing alginate-glycerol formulations to solidify blood in menstrual cups, powders were spread in the stockinette and then coiled into the cups. (H) Cups were inserted into a Syngina, received blood, and incubated for 4 h at 37°C. (I) Representative images show the impact of alginate-glycerol formulations on blood consistency with menstrual cups. (J) After individuals removed the incubated menstrual cups, the fraction of blood spilled was measured. (K) Compiled results show significantly less blood is spilled when a stockinette containing the alginate-glycerol formulation is used. Statistical analysis was performed by Kruskal-Wallis test with Dunn's multiple comparisons test. **, $p < 0.005$; ****, $p < 0.0001$.

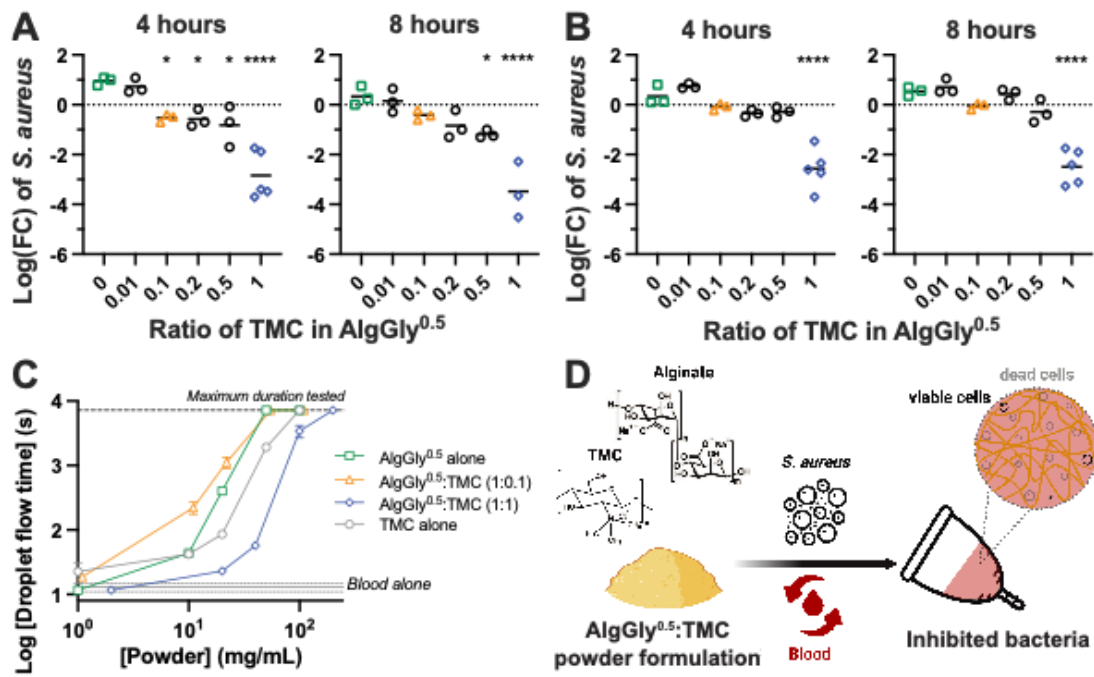


Figure 5. Antimicrobial activity of alginate-glycerol formulations processed with TMC. Formulations containing 20 mg of AG^{0.5} and varying amounts of TMC were mixed with blood inoculated with (A) 5×10^5 cfu/mL or (B) 5×10^6 cfu/mL of *S. aureus* and incubated for 4 and 8 h at 37°C. (C) Flow tests of these powders mixed with blood. (D) A schematic representation of powder function when mixed with blood and *S. aureus*. Statistical analysis was performed by one-way ANOVA compared to a TMC ratio of 0 with a Dunnett post-test. ****, $p < 0.0001$.

Tables and table titles and legends

Table 1. Alginate-based formulations for the production of blood-absorbent powder.

Alginate: Glycerol (w/w)	Alginate (g)	Glycerol (g)	Water (mL)
1:0 (AG ⁰)	2.5	0	10
1:0.5 (AG ^{0.5})	2.5	1.25	10
1:1 (AG ¹)	2.5	2.5	10
1:2 (AG ²)	2.5	5.0	10
1:3 (AG ³)	2.5	7.5	10

Supplemental Video Titles and Legends

Video S1. Alginate powder enhances blood retention in menstrual pads. Gentle squeezing of ~1.5 g of commercial menstrual pad filler or alginate-based powder soaked with 5 mL of defibrinated porcine blood.

Video S2. Pouring of blood from menstrual cups containing various materials. 10 mL of defibrinated porcine blood was incubated in menstrual cups with various fillers for 1 h (1= cup alone; 2=empty stockinette; 3= stockinette containing ~1.5 g of the alginate formulation). After tilting, to demonstrate the fluidity of blood contained within the menstrual cups, the cups were fully inverted to demonstrate blood retention.

Video S3. Retention of blood within various materials. 10 mL of defibrinated porcine blood was incubated in menstrual cups with various fillers for 1 h (1= cup alone; 2=empty stockinette; 3= stockinette containing ~1.5 g of the alginate formulation). Materials were examined to demonstrate blood retention during removal.

Video S4. Remaining blood within menstrual cups after material removal. After removal of absorbent element (see Video S3), cups were completely inverted to show the blood remaining in the cups.