Combinatorial Polyacrylamide Hydrogels for Preventing Biofouling on Implantable Biosensors

Unfortunately, continuous monitoring has remained challenging due to poor sensor signal and limited lifetimes of sensors. After implantation in the body of a patient, biofouling—the nonspecific adsorption of biomolecules—often occludes these devices and contributes to a reduction in their performance and lifetime. For devices that contact blood, the adhesion and aggregation of platelets from flowing blood significantly shorten functional device lifetimes.\(^{[4,5]}\) Once these devices foul, high-risk and costly invasive surgeries are often required to replace them, greatly increasing patient and hospital burden.

To address these challenges and bridge the gap between devices and long-last implants, devices are often coated with a soft material to mitigate biofouling. “Active” coatings such as drug-eluting\(^{[6]}\) or lubricant-infused polymer materials\(^{[7]}\) can be used to reduce adhesion and activation of platelets. While these approaches show promise in some applications, they offer only short-term benefits on account of the eventual depletion of the active molecules from the coating material,\(^{[8–10]}\) leading to major bleeding in patients near the end of the functional device lifetime.\(^{[11]}\) In contrast, “passive” coating strategies that modify the chemistry at the interface of the devices and bodily fluids are scalable, tunable, and inexpensive. The “gold-standard” materials used for passive anti-biofouling coatings are poly(ethylene glycol) (PEG)\(^{[12]}\) and its derivatives,\(^{[13–15]}\) which form a tight hydration layer through hydrogen bonding with water that is hypothesized to contribute to anti-biofouling properties.\(^{[16]}\) Nonetheless, PEG has several critical drawbacks, including degradation through hydrolysis and auto-oxidation leading to the production of reactive oxygen species\(^{[17–20]}\) and reduced antifouling performance over time.\(^{[21]}\) In recent years, many alternatives to PEG have been developed,\(^{[22]}\) including zwitterionic polymers,\(^{[23,24]}\) fluoro- and acrylate-based polymers,\(^{[25–27]}\) and polyglycidols.\(^{[28]}\) Zwitterionic materials in particular have shown promise as exceptional anti-biofouling coatings, though these materials may still face reduced stability in long-term applications due to the presence of hydrolysis-prone ester bonds.\(^{[29–31]}\) There is thus a critical need to develop materials that achieve superior stability and anti-biofouling performance relative to existing materials. Novel approaches that make use of high-performance assays to rigorously test the function of materials are needed to advance the discovery of such next-generation anti-biofouling materials and enable the development of coatings that improve the performance of medical devices.

In this study, we used a high-throughput screening approach to evaluate biofouling of a library of 172 combinatorial copolymer hydrogel materials assembled from 11 distinct acrylamide-based monomers (Figure 1). Our results demonstrate that a subset of these materials exhibit remarkable anti-biofouling function compared to existing gold-standard coatings. These polycrylamide hydrogels exhibit tunable mechanical properties, and the simplicity of their synthesis makes them highly adaptable for biosensor coatings. While combinatorial screening approaches have been used previously for materials discovery efforts\(^{[32–34]}\) aimed at mitigating marine or bacterial fouling,\(^{[35–38]}\) the foreign-body response,\(^{[34,39]}\) or cell adhesion,\(^{[40,41]}\) we screen for the prevention of platelet adhesion, which is a critical initial process for biofouling in blood.

We have rigorously assessed the antifouling behavior of our library of copolymer hydrogels using a high concentration of serum proteins and platelet-rich plasma over prolonged timeframes. Moreover, we have used machine-learning techniques to elucidate the molecular features of these combinatorial copolymer hydrogels that give rise to their observed biofouling performance. We show that our top-performing copolymer hydrogel is superior to current gold-standard materials in terms of enhancing the performance and lifetime of electrochemical biosensors in vitro and in vivo in multiple rodent models.

2. Results

2.1. Design of a Combinatorial Library of Polycrylamide Hydrogels for Anti-Biofouling

Here we first sought to investigate the ability of a large library of chemically distinct polycrylamide copolymer hydrogels to mitigate blood protein and platelet adhesion (Figure 1a). Several acrylamide-derived monomers have been widely used in biological settings (e.g., SDS-PAGE gels) and in biomedical applications (e.g., contact lenses, filters, and drug delivery materials) and have previously demonstrated potential for use as inert materials in the human body.\(^{[42–44]}\) To maximize the scope of our evaluation of polycrylamide materials, we selected 11 commercially available acrylamide-derived monomers and fabricated a library of 172 polycrylamide copolymer hydrogels comprising unique binary combinatorial mixtures (100:0, 75:25, 50:50, 25:75) of these monomers formulated at 20 wt% monomer (Figure 1b,c; Tables S1 and S2, Supporting Information) to generate an extensive combinatorial library of polycrylamide-based hydrogel formulations.

Each of the hydrogels was prepared by photopolymerization of prepolymer solutions using lithium phenyl-2,4,6-trimethylbenzoylphosphinate (LAP) as a radical photoinitiator and a simple LED (\(\lambda = 350\) nm) light source. During synthesis, four copolymer formulations turned opaque on account of insolubility of the polymer chains in the aqueous media and were thus omitted from further evaluation. We synthesized our hydrogels at a monomer content of 20 wt% to generate a library of materials with stiffness values mimicking those of human vein or artery tissues (Figure 1d).\(^{[45]}\) As all monomers evaluated were acrylamide-derived, the reactivity ratios for copolymerization of each pair of monomers \((r_1r_2 = 1)\) were such that each copolymer hydrogel was expected to comprise a statistical incorporation of the two monomers with the most distinct molecular weights: acrylamide (A; \(M_w = 71.08\) g mol\(^{-1}\)) and [tris(hydroxymethyl) methyl]acrylamide (G; \(M_w = 175.18\) g mol\(^{-1}\)). We selected these two formulations for more in-depth mechanical characterization in order to validate that the various members of our hydrogel library exhibited similar mechanical properties.
Figure 1. Polyacrylamide hydrogels as anti-biofouling device coatings. a) Coating substrates with anti-biofouling hydrogels can mitigate platelet adhesion and aggregation. b) Monomers used for combinatorial hydrogel synthesis: acrylamide (A), dimethylacrylamide (B), diethylacrylamide (C), (3-methoxypropyl)acrylamide (D), hydroxymethylacrylamide (E), hydroxyethylacrylamide (F), [tris(hydroxymethyl)methyl]acrylamide (G), acryloylmorpholine (H), (acrylaminopropyl)trimethylammonium (I), 2-acrylamido-2-methyl-propane sulfonic acid (J), N-isopropylacrylamide (K), poly(ethylene glycol) methacrylate (L), 2-methacycloxyethyl phosphorylcholine (M), [2-(methacycloxy)ethyl]dimethyl(3-sulfopropyl)ammonium (N), 2-hydroxyethyl methacrylate (HEMA; O). c) Photopolymerization (λ = 350 nm) of polyacrylamide hydrogels using lithium phenyl, poly(tetrafluoroethylene) (PTFE), poly(vinyl chloride) (PVC), poly(ethylene terephthalate) (PET), parylene C, and poly(methyl methacrylate) (PMMA). In this study, polyacrylamide copolymers were fabricated with elastic moduli similar to human arteries. e) Oscillatory shear rheology of hydrogels formed with low (A100; top) and high (G100; bottom) molecular weight monomers indicates consistency in shear moduli amongst the hydrogels, regardless of composition.
(Figure 1e). These studies indicated that even these highly distinct hydrogels exhibited similar shear storage and loss modulus values, and elastic modulus values similar to human veins and arteries.

2.2. Protein Adsorption and Platelet-Resistant Hydrogels

To rapidly evaluate the extent of biofouling on these hydrogels in a parallel fashion, we developed a high-throughput assay to screen platelet adhesion following incubation in protein serum. While the exact mechanism of biofouling has yet to be precisely elucidated, it is widely hypothesized that such fouling is initiated with the non-specific adsorption of proteins from the biological milieu. This undesirable binding of proteins to the surface of a biomaterial or device leads to the onset of platelet adhesion and aggregation that eventually causes thrombosis. While the majority of anti-biofouling assays appearing in the literature have focused on evaluating adsorption of model proteins at low concentrations (e.g., 1 mg mL⁻¹ bovine serum albumin or 1 mg mL⁻¹ fibrinogen) on materials of interest, these studies do not provide a realistic representation of the complex milieu of biomolecules in blood. Other studies have used more relevant mixtures of biological proteins (e.g., fibrinogen, bovine serum albumin, and lysozyme), or diluted or undiluted serum or plasma to evaluate biofouling of new materials, but only after short exposure timeframes ranging from 10 to 25 min. The ability to repel whole blood has also been studied; however, in these studies blood was applied to the biomaterial of interest for only a matter of seconds. Consequently, previous studies have not provided a sufficiently realistic representation of the complex engineering challenge imposed on anti-biofouling coatings in the body on account of low protein concentrations, insufficiently complex biological milieu, and short assay times.

To identify anti-biofouling materials from our copolymer hydrogel library with translational potential, we therefore sought to subject these materials to severe fouling conditions for prolonged periods of time that would nevertheless provide a simple and high-throughput readout of fouling activity. We chose to use platelet counting to enable a straightforward and realistic metric for anti-biofouling performance of our copolymer hydrogels, as platelet adhesion causes occlusion and leads to thrombosis. In these assays, each hydrogel was first incubated in 50% fetal bovine serum (FBS) for 24 h at 37 °C to introduce nonspecific protein adsorption (Figure 2a) prior to incubation in rat platelet-rich plasma (PRP) for 1 h at 37 °C to introduce platelets that could be counted in an automated fashion. In the blood biofouling process, the adhesion of platelets recruits and instigates resulting platelet aggregation, thus leading to the likelihood of large platelet clumps and high variability of platelet counts. We chose a 1 h incubation as our terminal timepoint to minimizing the variation seen with extremely high platelet counts (Figure 2b). While we anticipate subtle species-dependent effects on platelet adhesion on account of the use of both bovine serum and rat platelet-rich plasma, we assume that adsorption of rat proteins during the second incubation step sufficiently fosters rat platelet adhesion. Although we could not discern clear trends in the behaviors of the polyacrylamide copolymers based on their composition, hydrophobic monomers such as N-isopropylacrylamide (K100) or diethylacrylamide (C100) yielded high mean platelet counts (28 600 ± 11 000 and 9870 ± 13 platelets cm⁻², respectively). In contrast, a 50:50 copolymer of hydroxyethylacrylamide (F) and diethylacrylamide (C), denoted F50–C50, yielded the lowest platelet counts (82.0 ± 15 platelets cm⁻²) of all of the materials tested, significantly outperforming PEG (L: 4810 ± 520 platelets cm⁻²; p < 0.0001), 2-methacrylicoxyethyl phosphorylcholine (M: 1330 ± 210 platelets cm⁻²; p = 0.0001), poly(2-hydroxyethyl methacrylate) (PHEMA; O: 9560 ± 650 platelets cm⁻²; p < 0.0001), and [(methacryloyl)oxyethyl]-dimethyl(3-sulfo propyl)ammonium zwitterion (N; 15 900 ± 4200 platelets cm⁻²; p = 0.0007) hydrogel formulations (mean ± standard error, unpaired t-test) (Figure 2c and Figure S1, Supporting Information). This approach to high-throughput screening of fouling with blood proteins and platelets therefore enabled identification of polyacrylamide-based copolymer hydrogel formulations with superior anti-biofouling properties in comparison to gold-standard polymers. While protein adsorption has previously been investigated as a metric of anti-biofouling performance, we did not see trends correlating IgG and fibrinogen adsorption with platelet adhesion (Figure S2, Supporting Information). This observation may indicate that other proteins and/or specific conformations of adsorbed proteins (rather than quantity alone) may play key roles in platelet adhesion. As protein layers evolve unpredictably over time and culture conditions of single proteins is insufficient and poorly defined for the growth of cells, such as embryonic stem cells and induced pluripotent stem cells, protein adsorption may offer limited utility as a metric correlating to platelet adhesion.

We then performed scanning electron microscopy (SEM) imaging of treated hydrogel samples following lyophilization to observe platelet morphology. On the PEG hydrogel sample, aggregates of platelets were observed throughout the surface of the hydrogel, although their morphology was generally round, indicating that the platelets were not likely activated. In contrast, polyzwit hydrogel samples exhibited negligible platelet adhesion, and leads to thrombosis. To aid in our understanding of the molecular features giving rise to the observed anti-biofouling properties of all of the materials we tested, we analyzed the physicochemical properties of each of the monomers and resulting hydrogels and the contribution of these features to fouling tendencies. In this study to identify such features, we accounted for the composition of the hydrogels, such as the monomers that comprised ~20 wt%, and features of these monomers, including inherent features.
physical and chemical properties, and also of the reactivity ratios, which affect the distribution of the monomers on the surface of the hydrogel. In addition, the use of platelet counts allowed for inputs that can link polymer behaviors to anti-biofouling performance, in contrast to protein adsorption assays that may not yield predictive models.\textsuperscript{[46]}

The long-standing hypothesis regarding the antifouling mechanism of the gold-standard polymers is the presence of a tight hydration layer between the material surface and water that forms a physical and energetic barrier to protein adsorption, determined often by the physicochemical properties of the materials and the surface packing.\textsuperscript{[46]} These properties include surface wettability and hydrophilicity, electrical neutrality, H-bonding, and prevalence of highly hydrated chemical groups. It has also been suggested that surface packing has a role in non-fouling properties, strongly correlated with the ability to form a hydration layer near the surface. Molecular simulations have also suggested that anti-biofouling properties may arise from steric repulsion
of proteins and polymer surfaces—that the density of grafted polymer chains simply causes resistance to protein adhesion and obstructs their adsorption to the surface.\(^{[56,59-63]}\) Numerous other computational and molecular dynamics studies have shown that entropic contributions play a role in preventing biofouling.\(^{[62,63]}\) wherein reduction in ligand mobility would lead to an entropic penalty against protein adsorption.\(^{[64]}\) Penna et al. have identified the importance of ligand mobility and interfacial dynamics, highlighting a balance of hydration properties and a dynamic ligand layer that greatly affects entropic barriers associated with antifouling efficacy.\(^{[65]}\)

Curiously, the compositions of the top-performing hydrogel formulations that yielded the lowest degree of platelet adhesion did not exhibit any intuitive trends, and so we utilized machine-learning analysis to identify mechanistic factors underpinning antifouling activity. We first classified the copolymer hydrogels as “low platelet count” or “high platelet count” with a random forest model, which offers robustness to overfitting and the capability to evaluate relevance of data features via mean decrease of accuracy (MDA) (see the Experimental Section for technical details). We initially generated a simple feature set (Feature Set A) reflective of the polymer formulation, whereby each polymer was represented using one-hot encoding for the alphabet of monomers weighted by the molar ratios of the monomers in the respective copolymer hydrogels. Feature relevance analysis in combination with class-enrichment analysis (Figure 3a and Figure S4a, Supporting Information) pointed to dimethylacrylamide (B), diethylacrylamide (C), and [acrylamido(propyl)trimethylammonium (I) being associated with heavy biofouling. In contrast, acrylamide (A), (3-methoxypropyl) acrylamide (D), and acryloylmorpholine (H) were prominent monomers in those hydrogel formulations exhibiting excellent anti-biofouling performance. As none of these monomers were in the top anti-biofouling hydrogel formulations, we determined that combinatorial formulations, rather than homopolymer formulations, are crucial in development of robust anti-biofouling coatings. This consideration motivated two approaches of weighing both the molecular features of each specific monomer and the ratio of each monomer present in a specific formulation. In this feature set, we increased resolution of the molecular features by generating vectors of physicochemical descriptors for the monomers, “inserting” these vectors into one-hot monomer encoding, and weighting by the molar ratios of the monomers in the corresponding copolymer hydrogels (Feature Set B). The\(^{[66]}\) score (the overall measure of a model’s accuracy) indicated that these feature sets were suitable in data analysis (Table S3,

![Figure 3](advancedmat.de)
Supporting Information). Descriptors, in general, are numerical values that capture various characteristics of the molecules, both simple and complex. For example, a feature can be the number of proton donors, or the complexity measure of the molecular graph. The set of descriptors computed for each monomer included 19 values split between physicochemical descriptors (LabuteASA, TSFA, MolLogP, MolMR), molecular graph descriptors (BalabanJ, BertzCT, Chi0, HallKierAlpha), hydrogen-bonding descriptors (LipinskiHBA, LipinskiHBD), and 3D shape descriptors (FracCSP3, InertialShapeFactor, PM11, PM12, PM13, RadGyration, SpherocityIndex, Asphericity, Eccentricity), all as implemented in the rdkit library[66] (Table S4, Supporting Information).

Inspection of the feature analysis for the monomers weighed by their molar ratios helped to pinpoint relevant characteristics of the individual monomers. These features could then be expressed as descriptors (Figure 3b; Table S4 and Figure S5, Supporting Information). The top three features are from the physicochemical descriptor (MolLogP and LabuteASA) and 3D shape family (InertialShapeFactor). Overall, the most relevant features were found to be connectivity and shape descriptors: (i) InertialShapeFactor describes the mass distribution in the molecule (by three principal moments of inertia), (ii) MolLogP is an estimate of the octanol/water partition coefficient and captures the hydrophilicity/hydrophobicity of the molecule (an energetic characteristic), and (iii) LabuteASA describes the approximate surface area of the molecule. Surface area of the molecules plays a role in anti-biofouling performance because all monomers have a surface area that is accessible to other molecules (e.g., the solvent), and higher values of LabuteASA indicate that more surface is accessible (Figure 3c).

In a second approach, we produced a simplified version of the descriptor-based feature set by summing up descriptor vectors of the monomers weighted by their molar ratios in corresponding copolymer hydrogels (Feature Set C), and our results indicated the importance of the 3D shape descriptor features (Figure S4b, Supporting Information). Expanded analysis of the heterogeneity of the platelet count distribution also indicated the importance of a steric mode in the observed fouling (Figure S6, Supporting Information). Overall, it appears that the relevant molecular features predominantly characterize properties associated with the steric aspect of the surface formation and the geometry of packing of the monomers.

2.4. Mechanical and Physical Properties of the Top-Performing Hydrogel

The mismatch in mechanical properties between many implant materials and native tissues can lead to adverse events such as immune rejection, fibrotic responses, trauma, and thrombosis.[67] Thus, it is crucial to match the mechanical properties of the biomaterial with surrounding tissue.[68−70] Hydrogels in particular are widely tunable, and have mechanical similarity to biological tissue—in particular the human artery and vein—making them useful for implanted biosensors[71] (Figure 1d,e and Figure S7, Supporting Information). While other coating methods may incorporate polymers via either grafting-to or grafting-from approaches, these coatings are often only nanometers thick and have poor mechanical integrity, making them ill-suited to serve as a soft interface between implanted devices and the body.

We first assessed the mechanical properties of our top-performing hydrogel formulation and control hydrogels comprising PEG (20 wt%), poly(2-methacryloyloxyethyl phosphorylcholine) (PMPC) (65 wt%), poly(2-methacryloyloxyethyl phosphorylcholine) (PSBMA) (65 wt%), and poly(2-hydroxethyl methacrylate) (PHEMA) (~35 wt%) using rheology (Figures S7 and S8, Supporting Information). These controls were synthesized using standard methods as described by literature to match the formulation as “gold-standard materials.”[23,72] We found the shear rheological properties were comparable among each material at these formulations—at 1% strain, G′(F50−C50) = 270 ± 4.9 kPa, G′(PEG) = 66.7 ± 28.5 kPa, G′(PMPC) = 32.2 ± 9.4 kPa, G′(PCBMA) = 39.5 ± 12.8 kPa, and G′(PHEMA) = 41.8 ± 9.2 kPa. Moreover, we used fluorescence recovery after photobleaching (FRAP) spectroscopy to evaluate the mesh size of each of these hydrogel materials, which were found to be similar to each other and well below the size of albumin[73] (Figure S7f, Supporting Information). We also used interferometry-based nanoidentation, which enables examination of the mechanics of these materials under wet conditions (Figure S8a,c, Supporting Information), as well as tensile testing (Figure S8b,c, Supporting Information) to characterize the elastic modulus of our top-performing hydrogel formulation. While soft materials commonly used in the medical industry exhibit Young’s moduli values several orders of magnitude greater than that of human arteries, our F50−C50 hydrogel coating exhibits mechanical properties that are similar to human arteries,[74] reducing the risk of chronic inflammation and potential failure of the implant.[75] Due to the stability of polyacrylamide-based polymers, our lead candidate coating will inherently resist degradation and persist longer under biological conditions over control materials such as PEG. Under a variety of expedited degradation conditions, we observed improved stability of polyacrylamide-based hydrogels in comparison to PEG, PHEMA and zwitterionic polymer hydrogels, suggesting that these materials may be subject to biodegradation over time in the body, inherently limiting their anti-biofouling performance over extended timeframes (Figure S9, Supporting Information).

2.5. Hydrogel Coating of Electrochemical Biosensors Extends Sensor Lifetime

In order to assess the usefulness of our new copolymer hydrogel material, we tested its anti-biofouling performance in protecting electrochemical biosensors in blood. Such biosensors generally offer a promising strategy for continuously detecting analytes in vivo, but their lifetimes are limited by biofouling.[58,76] The degree of biofouling can be quantified using cyclic voltammetry (CV) to measure the anodic peak current. By applying a range of voltages through an electrode, electron transfer and current intensity may be monitored through the anodic peak. Changes in this peak can be used to assess interaction between the electrode surface and (FeCN)₄³⁺ ions in solution and the lifetime and quality of the signal. A larger reduction in signal indicates barriers to the electron transfer
We prepared devices comprising a gold (Au) working electrode (WE), a silver/silver chloride (Ag/AgCl) reference electrode (RE), and a platinum (Pt) counter electrode (CE) (Figure 4a,b) with PEG and F50–C50 hydrogel coatings to assess the ability of the hydrogel coating to extend device lifetime. We first performed CV measurements in Na₂EDTA-treated human whole blood spiked with \(15 \times 10^{-3}\) M of ferrocyanide ions, \((\text{FeCN})_6^{3-}\), to establish a baseline (Figure 4c). Then, devices were incubated in ferrocyanide-free human whole blood mixed with \(50 \times 10^{-3}\) M CaCl₂ to initiate expedited blood coagulation and to activate blood platelets (Figure S10, Supporting Information). After 2 h, probes were placed in the same process and blocking of ions from diffusing toward the electrode surface due to biofouling.

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ferrocyanide-spiked blood sample to record CV measurements (Figure 4d–f and Figure S11, Supporting Information). As the addition of CaCl₂ rapidly expedites blood clotting, 2 h was ample time to observe significant differences in probe performance without full clotting of the blood medium.[86] While anodic peak current was measured, the slight appearance of other peaks was observed with all probes, indicating a change in diffusion rate. The shift in potential often occurs due to the change of Cl⁻ concentration in blood from the addition of CaCl₂, as these concentrations affect the electroactive window.[80,81] Considerable peak reduction was observed for the PEG-coated probe (50.8 ± 16%), whereas the F50–C50 hydrogel coating preserved current intensity (76.6 ± 21%) in these assays (Figure 4g). Uncoated, bare devices were also evaluated as controls (44.3 ± 9%) and did not significantly differ from PEG-coated probes (paired t-test, p = 0.931) (Figures S11 and S12, Supporting Information).

Given that our F50–C50 copolymer hydrogel clearly outperformed PEG in a blood contacting setting in vitro, we proceeded to test the performance of these two materials in vivo. To assess these materials, we selected an area of implantation that would have direct blood contact. We first sought to evaluate the fouling of sensing probes implanted into the femoral vein of a rat for 5 d (Figure S13a, Supporting Information). This implantation site provided the best model for direct blood contact while the implantation time was the longest duration in which we were able to ensure device retrieval. Beyond 5 d, we experienced sutures failures and devices were lost. These device bundles comprised two WEs (one PEG-coated and one F50–C50-coated) and one RE (Figure S13b,c, Supporting Information), and exhibited a diameter of ≈500 µm after the hydrogel coating. CV was performed in a (FeCN₃⁺)ₙ-spiked blood sample in vitro to determine the anodic current intensity before and after in vivo incubation (a fresh Pt probe was included in the measurements for comparison). After 5 d, the sensor was recovered from the femoral vein (Figure S13d,e, Supporting Information) and the hydrogel coatings visibly remained intact on the electrode surface (Figures S13f and S14, Supporting Information). We observed significant signal degradation from bare (81 ± 27% reduction) or PEG-coated (83 ± 23% reduction) probes at the end-point of the study (Figures S15 and S16, Supporting Information). Indeed, two-thirds of these probes were completely nonfunctional after 5 d implanted in vivo. In contrast, we could obtain clear signals with notably less signal degradation (24 ± 36% reduction) from all probes coated with our F50–C50 hydrogel (Figures S15 and S16, Supporting Information). The results of this study indicated that F50–C50-coated probes performed much more reliably than bare and PEG-coated probes.

2.6. Hydrogel Coating Enables Real-Time Measurement In Vivo via DNA Aptamer Biosensors

Based on the in vivo pilot study described above, we evaluated these coatings on real-time biosensors that use structure-switching aptamers to continuously measure specific analytes in vivo.[82–85] Briefly, we engineer the aptamers to undergo reversible structure-switching upon target binding, and we

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Figure 5. F50–C50 hydrogel coating of a DNA aptamer biosensor enables continuous real-time monitoring of kanamycin in vivo. a) DNA aptamers that bind kanamycin with high specificity were functionalized onto the tip of Au probes. b) The probe was inserted into the femoral vein of a rat through a catheter. Kanamycin was infused into the other femoral vein. c) The kanamycin signal response should exhibit fast kinetics and dose-dependence. d) Image of the aptamer probe device. e) Representative graph of real-time in vivo sensing of kanamycin after administration of varying concentrations of the drug by injection into the contralateral femoral vein. Filled arrows represent a 200 µL injection of 200 × 10⁻³ M kanamycin and the empty arrows indicate a 100 µL injection of 200 × 10⁻³ M kanamycin. The shaded red box represents peak intensity of initial baseline signal. f) Signal intensities as indicated by peak heights were measured and normalized to baseline signal of each probe. Deviation from the baseline signal between PEG- and F50–C50-coated sensors showed the ability of the F50–C50 hydrogel to maintain higher signal. Data represent mean ± standard deviation, n = 4, unpaired t-test.
conjugate electroactive reporters (e.g., methylene blue, MB) at the distal end of the aptamer (Figure 5a). We continuously measure the electron transfer between the reporter and the electrode using CV, which enables us to monitor the concentration of the analyte in real-time in a reagentless manner in real-time. We and other groups have used such sensors to continuously measure the systemic concentrations of small-molecule drugs in vivo, and even control their levels using closed-loop feedback control, but biofouling of the surfaces of these devices limits their signal quality, and ultimately, their lifetime. We therefore sought to test whether our top-performing hydrogel coating could improve the performance of aptamer-functionalized electrodes in vivo. As a model system we used aminoglycoside aptamers that bind kanamycin, which is a widely used small-molecule antibiotic drug (MW = 484.5 Da; diameter = 1 nm). To reduce the diffusion time of the drug to the probe surface, we reduced the thickness of the hydrogel coating by dip-coating the electrodes with precursor solutions for F50–C50 and PEG hydrogels and immediately cross-linked by photopolymerization (Figure S17, Supporting Information). The mesh size of the resulting hydrogel coating was determined to be 2.33 ± 0.07 nm using FRAP spectroscopy, thus allowing free transport of kanamycin. We performed three experiments with this system. We initially characterized platelet adhesion on probes following incubation in Na$_2$EDTA-treated human whole blood for three days at room temperature. We observed significantly less platelet adhesion on F50–C50-coated probes than PEG-coated or bare probes (Figure S18, Supporting Information). Next, aptamer probes were inserted into a circulating blood flow system for 6 and 12 d (Figure S19a, Supporting Information). At the end of each assay, the probes were analyzed by SEM, which showed that the F50–C50 hydrogel coatings had noticeably less platelet adhesion after both time-points than control probes (Figure S19b, Supporting Information).

Finally, to test the functional efficacy of these hydrogel coatings in vivo to monitor the analyte kanamycin in real-time, we tested our aptamer devices in the femoral vein of a rat as described above (Figure 5b). Kanamycin (0.1 or 0.2 mL at 200 × 10$^{-3}$ μg) was administered intravenously into the contralateral femoral vein at different time-points while the rat was under anesthesia. Due to the required duration of the rat under continuous anesthesia, we were limited inherently by the assay time, to roughly 3.5 h. Because of this assay limitation, we chose to dip coat the aptamer probes with hydrogel coating, which allowed for a thinner coating and thus reduced diffusion time of analyte to aptamer surface, enabling more rapid response times and increasing number of measurements in this real-time test. Aptamer activity was measured using square-wave voltammetry (SWV) at 400 and 60 Hz, and we used the kinetic differential measurement technique to compensate for drift as described previously (Figure 5c,d). After establishing equilibrium over the course of 60–120 min after implantation, we successfully recorded the real-time drug concentration in vivo for the first time with F50–C50 coated devices (Figure 5e and Figure S20, Supporting Information). Signal peaks in response to kanamycin administration were determined at various time points over the course of roughly 200 min following incubation, and normalized to initial peak intensity, due to sensor-to-sensor variations. In these assays, PEG-coated sensors had an average deviation from original signal of 64.2 ± 16.1%, while the enhanced anti-biofouling capabilities of F50–C50 coatings resulted in significantly less deviation, 33.8 ± 4.9%, from the original signal (Figure 5f) (mean ± standard deviation; n = 4, data was analyzed via unpaired t-test where p = 0.01).

3. Discussion

In this work, we developed a library of 172 polyacrylamide copolymer hydrogels—the most extensive library of this type generated to date—to discover anti-biofouling coatings for biomedical devices that outperform current gold-standard materials. Machine-learning analysis of high-throughput, parallel platelet fouling assays suggested that the relevant mechanisms underlying the observed anti-biofouling properties of these materials arise from steric and inertial shape properties of the monomers used in these hydrogels. We also demonstrated that the top-performing formulation exhibits mechanical properties resembling those of human arteries. Finally, we validated the utility of this hydrogel formulation for improving the performance of biosensor devices both in vitro and in vivo.

To the best of our knowledge, only a small subset of polyacrylamide polymers have been explored previously as candidates for anti-biofouling coatings, including those derived from acrylamide, hydroxyethylacrylamide, (3-methoxypropyl)acrylamide, and $N$-isopropylacrylamide. We sought to explore a wide variety of commercially available acrylamide monomers, as well as combinations of these monomers, on account of the ability to screen a large amount of chemical space with materials that have been demonstrated to be amendable to adhesion onto a variety of surfaces. While our platelet screening assay employed the use of both bovine serum and rat platelet-rich plasma, which may consequently lead to subtle species-dependent effects of protein adsorption and platelet adhesion, this method allowed us to screen our large polyacrylamide library in a high-throughput manner. Though we did not evaluate in depth the adhesion of our hydrogel coatings to the device surfaces, these materials are amendable to a variety of adhesion chemistries. Polyacrylamide-based polymers are highly stable over long time-frames under physiological conditions, thereby enabling their use with medical devices requiring long functional lifetimes. While PEG and zwitterionic polymers are considered gold-standard materials for antifouling coatings, our platelet screening assay identified numerous polyacrylamide copolymer hydrogel formulations exhibiting superior anti-biofouling performance, highlighting the promise of polyacrylamide copolymer hydrogels in these applications. Based on the successes of these polyacrylamide-based copolymer hydrogels, we believe that evaluation of copolymer hydrogels based on PEG and zwitterionic polymers will be important and may also yield next-generation coatings with superior anti-biofouling performance.

Our selected formulation for the L100 PEG hydrogel (PEG diacrylate Mn 780 and PEG acrylate 480) was chosen so as to design a PEG hydrogel that did not meaningfully differ in stiffness from the other hydrogels in our library at the same
polymer content. As other PEG monomers have also been used historically for anti-biofouling studies, namely, PEG diacrylates and PEG dimethacrylates of various molecular weights, we evaluated three alternative formulations: (i) PEGDMA740 (15 wt%) plus PEGMA550 (5 wt%), (ii) PEGDA10k (20 wt%), and (iii) PEGDMA10k (20 wt%). Each of these formulations comprised 20 wt% solids and 1wt% photo-initiator. We found that these formulations exhibited shear storage modulus values ranging over two orders of magnitude (Figure S21, Supporting Information) and resembling values determined for the library of polycrylamide-based hydrogels (Figure 1e). Further, we found that these alternative PEG hydrogels exhibited higher IgG and fibrinogen adsorption than the primary L100 PEG control (Figure S22a, Supporting Information), while platelet adhesion was similar for all formulations evaluated (Figure S22b, Supporting Information). For this reason, we were confident in our use of L100 as a representative PEG formulation in all of our subsequent experiments.

The anti-biofouling behavior of the various copolymer formulations was evaluated using machine-learning algorithms with several different feature sets to elucidate the molecular features that give rise to the observed behavior. The features that were common amongst the top performing hydrogel formulations (Feature Set B) were heavily represented by connectivity and shape descriptors. Coupled with other studies,[57,64,65] indicating that ligand mobility makes an entropic contribution to anti fouling performance, this work provides critical insight into the molecular mechanisms underpinning anti-biofouling behaviors and potentially enabling the rational design of next generation coatings. Recently, Rostam et al. developed a (meth) acrylate and (meth)acrylamide polymer library to modulate the foreign body response to implanted materials and made use of machine-learning approaches to analyze important chemistry descriptors driving macrophage phenotype and implant fouling. While this study made use of arrays of neat polymers instead of hydrogels, steric (and electronic factors) were also attributed to feature importance.[14]

We sought to eliminate surface topography as a contributor to differences in biological performance in the PAAm library by polymerizing the hydrogels between glass slides. Although SEM images revealed that the various hydrogel formulations evaluated exhibit different levels of activation of platelets (Figure S3, Supporting Information), the differences in hydrogel surface topography that were apparent in SEM are attributable to the sample processing, wherein hydrogels were lyophilized prior to imaging. To evaluate the true topology of the hydrated hydrogel surfaces, we performed optical microscopy on a subset of materials, including our top-performing polyacrylamide formulation, homopolymer hydrogels that comprise the top formulation, a subset of various polyacrylamide hydrogels across the library, and control materials. Optical microscopy images (Figure S23a–g, Supporting Information) revealed that all of the materials evaluated are relatively flat, with ≪3 µm variation in their height profile, and have negligible surface roughness (Table S5, Supporting Information). As the topography and surface roughness of the hydrogels are similar across all materials evaluated, we believe surface topography is not an important variable within the hydrogel library.

When the top-performing polyacrylamide copolymer hydrogel was used as a coating on an electrochemical biosensor device, we observed little reduction in signal in vitro in the presence of whole blood. The mechanism behind this improved signal could be attributed to two critical factors that are enhanced by the hydrogel. First, the surface of a bare electrode is exposed to a broad spectrum of molecules, leading to broad peaks, while the pores of the hydrogel coating exclude molecules too large to diffuse through the matrix, reducing peak widths observed with hydrogel-coated probes. Second, the intensity of measured peaks is maintained throughout the duration of the assay with F50–C50 hydrogel-coated probes, suggesting that this formulation precludes biofouling[58,96] while allowing molecules of interest to diffuse through the hydrogel matrix and bind reversibly to the surface to produce signal. In contrast, PEG-coated probes lose their ability to sense molecules over time, despite having a comparable mesh size to the F50–C50 hydrogel coating, suggesting that free diffusion of molecules through the coating is impaired by fouling. Indeed, other studies have suggested PEG coatings may not be suitable for use on biosensors of this type as they can hinder electron transfer.[97]

Through in vivo implantation studies, we observed that F50–C50-coated probes performed much more reliably than bare and PEG-coated probes. Indeed, only one probe from each of the control groups showed any response after five days implanted in the femoral vein of a rat on account of fouling in vivo. In contrast, all three F50–C50-coated probes exhibited clear signals with higher anodic currents (Figure S15, Supporting Information). These studies suggest that the majority of signal loss likely occurs in the first few hours of blood exposure and changes relatively little out to 5 d following implantation. Similarly, PEG-coated DNA aptamer sensors showed twice as much deviation from their original signal relative to F50–C50-coated sensors within the first two hours after implantation in the femoral vein of a rat (Figure 5). In this timeframe alone, we notice significant improvement in current peaks with our polyacrylamide-based material. While future work will certainly be needed to evaluate the function of F50–C50-coated sensors over longer durations relevant for certain applications (e.g., catheter-bound sensors), these studies suggest that performance of F50–C50-coated sensors is rather stable through the first 5 d, thereby exhibiting promise for maintaining sensor function over longer time-frames. Overall, these data show that combinatorial screening of copolymer hydrogels offers an exciting avenue for the discovery of anti-biofouling coatings.

Although it may not be intuitively obvious from a materials design point of view, we demonstrate that certain copolymer formulations can deliver considerably better anti-biofouling performance in comparison to gold-standard materials than homopolymers. Considering the success of these polycrylamide-based copolymer materials, it will be important to similarly evaluate the anti-biofouling performance of copolymer hydrogels comprising PEG and zwitterionic polymers. These observations highlight the value of coupling chemical design intuition with high-throughput screening approaches for discovery of new materials. We have demonstrated the applicability of these new hydrogel coatings to the surface of electrodes using facile synthetic approaches that are amenable to
scalable manufacturing, although future studies must be conducted to optimize the adhesion strength of these hydrogels to device surfaces for longer-term biomedical use[88,90] and to enhance the durability of the hydrogel materials.[100] With further improvements, we believe this strategy of using combinations of acrylamide-based monomers may enable the discovery of anti-biofouling materials enabling long-term implantation of biosensor devices to continuously monitor chronic biomedical conditions.

4. Experimental Section

Materials and Reagents: All reagents were purchased from Sigma-Aldrich and used as received, unless later specified. Au, Pt, and Ag wires were purchased from A-M Systems. EDTA-treated human whole blood for flowing in vitro measurements was purchased from BioIVT. Kanamycin monosulfate was ordered in USP grade from Gold BioTechnology.

Hydrogel Preparation: Prepolymer formulations containing 20 wt% acrylamide monomer, 1 wt% LAP as photoinitiator, and 1 wt% methoxybisacrylamide were mixed and pipetted between two glass slides separated by a silicone spacer (0.25 mm ± 0.05 mm). Gels were crosslinked in a Luzchem photoreactor system with 8 W bulbs and an intensity of 2–40 W m⁻² (LZC-4, hv = 350 nm, 15 min). Due to swelling of polyacrylamides in water, they were placed in 1× PBS for at least 24 h before being punched with a 6 mm biopsy punch. PMPC, PSBMA, and PHEMA hydrogels were prepared as described previously.[101,102] Briefly, poly(ethylene glycol) hydrogels were prepared with monomeric solutions in 1 M NaCl with 4% N,N'-methylenebisacrylamide (MBAm) cross-linker. A stock solution of 15% sodium metabsulfite and 40% ammonium persulfate was added to the prepolymer solution at ~1 wt% polymerization and polymerization was initiated at 60 °C. PHEMA hydrogels were prepared in a solution of ethanol:glycol:water (1:5:1.5:5) with tetraethylenglycol dimethacrylate as a cross-linker. A stock solution of 40% ammonium persulfate and 0.5 wt% TEMED was added to the prepolymer solution at 1.5 wt% polymerization and polymerization was initiated at 60 °C. PEG hydrogels were prepared similarly with a prepolymer solution of 15 wt% PEG diacrylate (Mₙ = 780)[102–104] and 5 wt% PEG acrylate (Mₙ = 480). Prior work has shown that PEG molecular weight (Mₙ > 10k) affects anti fouling performance when PEG is presented as brush polymers tethered to a surface. In addition, when molecular weight of PEG exceeds 40 kDa, issues arise such as potential immunogenicity and accumulation in tissues.[105] Here, hydrogels were used to create a fully crosslinked network and, to our knowledge, have not seen effects of PEG molecular weight. Additional PEG hydrogels were synthesized with 20 wt% PEG10k diacylate and 1 wt% LAP (PEGDMA10k), 20 wt% PEG10k dimethacrylate and 1 wt% LAP (PEGDMA10k) and 15 wt% PEG740 dimethacrylate, 5 wt% PEG 550 methacrylate and 1 wt% LAP (PEGMA740_PEGMA550). Prepolymer solutions were mixed and pipetted between two glass slides separated by a silicone spacer (0.25 mm ± 0.05 mm). Gels were crosslinked in a Luzchem photoreactor system with 8 W bulbs and an intensity of 25–40 W m⁻² (LZC-4, hv = 350 nm, 15 min). Hydrogels were placed in 1× PBS for at least 24 h before being punched with a 6 mm biopsy punch.

Hydrogel Synthesis Modifications: All 2-acrylamido-2-methyl-propane sulfonic acid (AMPSam) formulations were made with slightly basic solution of 2:1:1 m NaOH:water. [Tris hydroxymethyl]methylacylamide (THMAM) formulations were made with 50:50 dimethylformamide:water as well as 100% N-isopropylacrylamide (NIpAm), 75% diethylacrylamide (DEAm) [25% NIpAm, and 25% hydroxyethylacrylamide (HMAm)/75% NIpAm. Acrylamidopropyltrimethyl-ammonium (APTAc)-based formulations were used with 2–3× MBAm concentration to achieve comparable mechanical properties.

Platelet Adhesion Test: Fresh whole rat blood was mixed in a 10:1 volume ratio of an acid citrate dextrose (ACD) anticoagulant buffer (containing 2.13% free citrate ion, BD Vacutainer Specialty Venous Blood Collection Tubes) for the preparation of PRP. PRP was obtained via centrifugation at 600×g for 10 min at 10 °C. The platelets were counted using a Countess II Automated Cell Counter (ThermoFisher Scientific) and diluted to 2.5 × 10⁵ platelets ml⁻¹ in 1× PBS. 6 mm punches of hydrogels were placed in ultralow adhesion 96-well plates and incubated for 24 h at 37 °C. Gels were UV-sterilized for 5 min prior to incubation with platelets. 100 µL of PRP was pipetted on top of the hydrogels. The plates were placed on a rotary shaker for 1 h at room temperature. Platelets were rinsed once with 1× PBS and fixed with 4% paraformaldehyde (PFA). Cells were imaged with an EVOS XL Core Imaging System microscope (Life Technologies).

Platelet Detection in Images: Platelets in the images are small round objects ~3–4 μm in diameter. Platelet images are of different color and can include noise such as gel chunks or dust. The noise typically exhibits sizes considerably smaller or larger than platelets. The platelets in the images were detected using a difference-of-Gaussian approach by blurring images using Gaussian kernels of a range of standard deviations in increasing order. Stacks of the differential images between two successively blurred images form a cube, and where blobs are local maxima of intensity. Blobs of noisy objects are avoided by tuning the range of standard deviations used in the process.

IgG and Fibrinogen Adhesion: Hydrogel discs (n = 3 for each sample) were punched into a low-adhesion clear 96-well plate. To measure IgG adsorption to the surface of the hydrogels, 100 μL of 2 μg mL⁻¹ Alexa Fluor 488-AffinityPure Rabbit Anti-Bovine IgG (Jackson ImmunoResearch) was added to each well; the plate was then sealed and placed on a shaker plate overnight. For fibrinogen adhesion, 100 μL of 5 μg mL⁻¹ fibrinogen from human plasma conjugated to AF488 (Invitrogen) was added to each well; the plate was then sealed and placed on a shaker plate for 1 h. Samples were then washed 5× with PBS, after which the gels were transferred to a black 96-well plate and fluorescence readings were taken (BioTek Synergy H1 microplate reader).

Fluorescence Recovery after Photobleaching: Hydrogel samples were loaded with 0.5 wt% FITC–dextran (4 kDa). An Inverted Zeiss LSM 780 laser scanning confocal microscope (Germany) with a Plan-Apochromat 20×/0.8 M27 objective lens was used for FRAP analysis. A pixel dwell time of 1.58 μs was used. Samples were photobleached with 405- and 488-nm argon lasers set to 100% intensity. The samples were placed in a sterile 0.16–0.19 mm thick glass bottom dish (MatTek). ZEN lite (Zeiss) software was used for all FRAP tests. To avoid any extra noise, the high voltage was limited to 700 V. Different tests (n = 5) were made at different locations of the sample. For each test, ten control prebleaching images were captured per second, and then the spot was bleached with a pixel dwell time of 177.32 μs. 390 post-bleaching frames were recorded per second to form the recovery exponential curve. The pixel size was set to be 0.83 μm. The diffusion coefficient was calculated as described in the literature:[106] $D = \frac{\lambda}{6\pi\eta R d}$ where $\eta = 0.88$ for circular beams, $\omega$ is the radius of the bleached ROI (12.5 μm), and $t_{1/2}$ is the half-time of the recovery. To estimate the mesh size ($\xi$) of the hydrogels, the obstruction theory of Amsden et al. was used:[207] $D = D_0 \exp \left( -\frac{\pi r_{f}^2 + r_{b}^2}{\xi + 2r_{f}} \right)$ (1) where $D$ is diffusivity of the solute in the hydrogel, $D_0$ is the diffusivity of the solute within the liquid in the hydrogel (saline-sodium citrate buffer), which is assumed to be the same as in pure water, $r_{f}$ is the radius of the solute (3.51 nm for FITC–dextran 4 kDa), $r_{b}$ is the radius of the polymer chains within the hydrogel network (estimated as 0.65 nm[108] for F50–500 hydrogel, 0.51 nm for PEG400, 2.00 nm for pHEMA[110] 1.00 nm for PMPC[111–113] and 0.9 nm for PCBMMA[114]). The diffusivity ($D_0$) of a solute in a pure liquid was determined with the Stokes–Einstein equation to be 69.91 μm² s⁻¹ using a solution viscosity of $\eta = 0.89 \times 10^{-3}$ Pa s.

Scanning Electron Microscopy: SEM images were acquired with an FEI Magellan 400 XHR microscope with a beam voltage of 1 kV. The sample was lyophilized prior to imaging, pressed onto silicon paint and sputter-coated with Au:Pd (60:40) before imaging. When imaging hydrogels and
adherent platelets, samples were fixed with 4% glutaraldehyde (CFA) for 5 min, then dehydrated with 10%, 30%, 50%, 70%, 90%, 100% ethanol solutions for 10 min each. Samples were then lyophilized, gently pressed onto silver paint and sputter-coated before imaging.

**Performance Classification:** The relevant performance range of the antifouling polymer is platelet count under 100 units. This requirement naturally sets up the data analytic task as classification. From this point, antifouling candidates are considered as members of one of two performance classes. The positive class includes polymers with median platelet count below or equal to 100, and the negative class includes polymers with median platelet above 100 counts. Out of the characterized polymers, there were 76 members in the positive class and 91 members in the negative class. The slight class imbalance was considered irrelevant for practical purposes, and can be mitigated using standard techniques such as over- and undersampling. The data were organized into three different feature sets (A, B, C). Feature Set A was reflective of the molar ratios of the monomers in the respective polymer. Feature Set B increased granularity by generating vectors of physical-chemical descriptors to the monomers and weighting these by the molar ratios of the monomer in the polymers. A simplified version was generated in Feature Set C, by summing up descriptor vectors of the monomers weighted by their molar ratios in the corresponding polymers. These classifiers (Table S3, Supporting Information) showed highest performance with Feature Sets A and B.

**Machine-Learning Data Analysis:** Classification was performed using a random forest (RF) model, as implemented in scikit-learn 0.20.2. As a model, RF offers robustness to overfitting and the capability to evaluate relevance of data features. RF model parameters were selected via cross-validation grid-search; optimized parameters included the number of estimators, maximum estimator depth, and maximum number of features used in the splitting procedure. Three-fold cross-validation was performed over the training set, which included 75% of the available observations. The model produced as a result of the cross-validated grid search was tested on the withheld test set that included 25% of the available observations. MDA was selected to evaluate feature relevance, which captures the drop in accuracy of the trained model under random permutation of the feature whose relevance is being evaluated with the rest of the features. Higher positive values of MDA are indicative of a larger drop in accuracy in the permutation test and point to higher impact of the respective feature on the model predictions. MDA values close to zero are indicative of minimal impact of the feature permutation on accuracy, and help identify irrelevant features. Negative MDA values point at features whose permutation increases model accuracy. Such features do not contribute meaningfully to the model performance. Model performance on the test set was evaluated using standard measures of accuracy in classification tasks, such as recall, accuracy, and f1-score.

**Expanded Analysis of Platelet Homogeneity:** The coverage profile for each formulation was generated in the following manner: (i) partition the image of the polymer surface into square bins, (ii) count the platelet population in each bin, (iii) generate a population histogram for all images obtained for a given formulation, and (iv) normalize the histogram to the total number of population bins obtained for a given formulation. There are several general types of population histograms that one might expect to encounter (Figure S4a–d, Supporting Information). It was recognized that the difference between homogeneous and heterogenous coverage modes is quantitative, not qualitative. In both cases, histograms correspond to distributions that have tails; the thickness of the tails distinguishes the two cases, where the tails fall off faster in the heterogenous cases than in homogeneous cases.

Using normalized population histograms as features, anomaly detection analysis was performed with the Isolation Forest algorithm. Each histogram received an anomaly score; negative values indicate outliers, and positive values indicate inliers. The notions of inliers and outliers are reflective of the structure of the data set and do not have absolute meaning. The anomaly scores were visualized by reducing dimensionality of the data set to two dimensions via t-SNE manifold embedding and encoding the anomaly score of the samples as the color (Figure S4i, Supporting Information). Outlier analysis identifies histograms corresponding to homogenous coverage as the most normal. The change of the anomaly score toward the most abnormal system is associated with increased heterogeneity of the coverage (Figure S4j, Supporting Information). Overall, there were 121 systems that bore signs of increasing heterogeneity of coverage, suggesting that the steric mode of fouling plays a meaningful role.

**Rheological Characterization:** Oscillatory rheology measurements were performed with a TA Instruments AR-G2 rheometer. Amplitude sweeps were conducted at a frequency of 10 rad s\(^{-1}\) from 0.1% to 100%. Frequency sweeps were conducted at 0.1% strain from 0.1 to 100 rad s\(^{-1}\). All tests were performed at 25 °C using an 8 mm parallel plate geometry.

**Tensile Test:** Tensile strength measurements were performed with an Instron series 5560A with 100 N load cell. Tensile tests were conducted at 0.2 mm s\(^{-1}\) at room temperature.

**Nanoindentation Test:** Young’s modulus measurements of the hydrogels were performed using a Piuma Nanoindenter (Optics11) with a probe from the same manufacturer with a stiffness of 38.8 N m\(^{-1}\) and a tip radius of 27.0 µm. Calibration was conducted as on glass under wet conditions as per the manufacturer’s instructions. Each sample was immersed in a sterile saline buffer solution (Opti-free – Replenish; Alcon) before their moindentation experiments. The indentation depth was fixed to be <1 µm to avoid bottom effects. At least eight force curves were used to determine the local Young’s modulus of each sample, using the Optics11 Nanoindenter V2.0.27 software. The results are shown as mean ± standard error of the mean. A Hertzian contact model parameter was used for the fit of the curves assuming that the Poisson’s ratio of the samples is \(\nu = 0.5\).

**Fabrication of Electrochemical Biosensors for In Vivo Assessment:** The WE was prepared with 75-µm diameter gold wire (PFA-coated). Ag/AgCl wires were prepared by incubating silver electrodes in bleach solution, rinsed vigorously with water, and dried. Pt wire was bundled with the Au electrode after bundling with Ag/AgCl wire to prevent shorting. The tip of the device was cut with a razor blade to expose bare gold and rebleached. No surface roughening was applied. To prevent wires from touching the side of the Eppendorf tube walls, they were propped in the middle of the tube. A hole was drilled in the cap of a 0.3 mL Eppendorf tube to fit 0.04 in. Tygon tubing. One inch of tubing was inserted into the drilled hole, and we applied epoxy and cured with UV light (365 nm) for 20 s. Wires were then inserted through the Tygon tubing into the tube.

**Hydrogel Coatings on Electrochemical Biosensors for In Vitro Assays:** 1 µL of prepolymer solution was pipetted into a 100 µL pipette tip. The probes were inserted and polymerized for 5 s, which was photocured at 365 nm. The pipette tip was removed, and the hydrogel coating was visible by eye. The hydrogel was then photocured for an additional 30 s.

**In Vitro Assays with Electrochemical Devices:** Bare, PEG-coated, and F50–C50-coated probes were incubated in EDTA-treated human whole blood at room temperature with 250 µL of 15 × 10\(^{-10}\) m (FeCN\(_4\))\(^{3-}\) to record baseline CV data. This step was limited to 5 min to ensure a steady-state readout while minimizing biofouling. Probes were rinsed in 1× SSC buffer for 5 min to thoroughly remove any intrinsic fouling and transferred to 250 µL whole blood, followed by addition of CaCl\(_2\) to a final concentration of 50 × 10\(^{-3}\) M. After 2 h, probes were transferred to the original tube used for baseline measurement. The device was incubated for 5 min to allow the diffusion of (FeCN\(_4\))\(^{3-}\) throughout the hydrogel. CV scanning was performed on the Au wire (potential range: –0.7 to 0.8 V, step size: 1 mV, scan rate: 0.1 V s\(^{-1}\)).

**Fabrication of Electrochemical Biosensors for In Vivo Assessment:** Bare gold wire was cut into ~10 cm wires. The tip (~1 mm) was bundled with black medical heat shrink tubing twice (ID 0.006 in. × 0.001, Nordson Medical 103-0325). Another two layers of heat shrink were applied ~1–2 mm away, allowing exposure of ~1–2 mm of bare gold. Ag/AgCl wires were bundled likewise to prevent shorting. Two gold wires (WE) and one Ag/AgCl wire (RE) were bundled at the tip with clear shrink wrap (ID 0.02 in. × 0.001, Nordson Medical 103-0249) and at the middle of the probe to provide mechanical strength when pushing through the catheter.
Hydrogel Coatings on Electrochemical Biosensors for In Vivo Assays: For each device, one Au wire was left bare and one was coated with hydrogel. The prepolymer solution of hydrogel was pipetted into a short segment of tubing from a 26G catheter with inner diameter of 0.6 mm. The device was pushed through the tubing and aligned with one of the two bare gold wires. UV light (365 nm) was applied for 5 seconds and the tubing was gently removed. The hydrogel coating was visible on the entirety of one Au probe only, and UV light was applied for 25 more seconds to ensure complete photopolymerization. Probes were stored in SSC buffer and placed in 200 µL of EDTA-treated rat blood (BioIVT) with 15 × 10^3 m (FeCN)_n^4− for 1 h. Probes were incubated for 5 min to allow diffusion of (FeCN)_n^4−. CV scanning was performed on the Au wire (potential range: −0.7 to 0.8 V; step size: 1 µV; scan rate: 0.1 V s⁻¹).

In Vivo Assays with Electrochemical Devices: Live animal studies were performed with male Sprague–Dawley rats under Stanford APLAC protocol number 33226. All rats used in this work were purchased from Charles River Laboratories at a weight of 350–475 g. The rats were anesthetized using isoflurane gas (2.5%) during implantation procedures and monitored continuously. After a small incision to expose the femoral vein, a 24G catheter was inserted into the vein. The needle of the catheter was removed, and the catheter was clamped. The devices were inserted through the catheter and pushed far into the vein. The catheter was clamped, and the device was secured in place, after which the incision was sutured and the animal monitored during recovery and the duration of the 5 d. At the end of the experiments, animals were anesthetized using isoflurane gas and the devices were removed for immediate measurement in blood used for baseline sample. Rats were then euthanized by exsanguination while under general anesthesia.

Aptamer Device Fabrication and Functionalization: The kanamycin aptamer probes were synthesized by Biosearch Technologies with the following sequence: 5′-H5-C(2CH2)_4-GGCACTTGGTTAGTATACACTCC-MB-3′. Probes were titrated at the 3′ end with a 6-carbon linker for self-assembly onto the WE, and conjugated with a methylene blue (MB) redox label at the 3′ end with a 7-carbon linker for charge transfer measurements. The modified DNAs were purified through dual HPLC by the supplier. Upon receipt, the construct was dissolved and diluted to 100 × 10^−6 m using UltraPure water (Thermo Fisher Scientific) and frozen at −20 °C in individual aliquots of 1 µL until use. The WE was prepared with 8 cm pure Au wire (75 µm diameter) insulated with heat-shrinkable tubing (Nordson Medical, 103-0235) to define the aptamer immobilization surface. The exposed gold wire has a length of 1–2 mm with an overall surface area of 0.25–0.5 mm². No surface roughening was applied. Before immobilizing the aptamer probes, the wire was rinsed with acetone, ethanol, and deionized water in a sonicator sequentially, followed by electrochemical cleaning. CV scanning on the gold wire was performed in 500 and 50 × 10^−3 m sulfuric acid solutions (potential range: −0.4 to 1.5 V; step size: 1 mV; scan rate: 0.1 V s⁻¹), each with three scans. An aliquot of the DNA construct was thawed and reduced for 40 min with 2 µL 100 × 10^−3 m tris(2-carboxyethyl)phosphine at room temperature in the dark. The reduced DNA construct was diluted to 1 × 10^−6 m with deionized water, and a freshly cleaned gold electrode was immersed for 1 h at room temperature in dark. Next, the sensor was rinsed with deionized water for 3 min, followed by immersion in 6 × 10^−3 m 6-mercapto-1-hexanol in 1× SSC buffer for 2 h at room temperature in the dark to passivate the remaining gold surface and remove nonspecifically adsorbed DNA. The sensor was rinsed with deionized water for another 3 min and stored in 1× SSC buffer at 4 °C for 12 h before application of the hydrogel.

Electrochemical measurements were conducted using an EmStat (Mater. Mater.) for determination of mass degradation. A custom peak-fitting script was created to fit the SWV measurements with a Gaussian curve on a hyperbolic baseline. Peak currents were then normalized to a baseline peak current to generate the signal gain. All reported gains were obtained via KDM, with the difference divided by the average of gains from the 400 and 60 Hz signals.

Hydrogel Coatings on Aptamer Devices: Acrylamide monomers were purified through a basic alumina column. The hydrogel was applied to the sensing gold wire through capillary force, in which the sensor was dipped into the hydrogel prepolymer solution for 5 s and removed immediately. The hydrogel was then photosensitized by applying 366 nm light for 30 s. No reduction in the MB peak current was observed after UV application. An Ag/AgCl reference electrode (75-µm-diameter Ag wire chlorinated in bleach overnight) was attached to the hydrogel-coated gold wire using heat shrinkable tubing. The final device was placed in 1× SSC buffer at 4 °C overnight before use.

In Vitro Assays of Aptamer Devices in Stationary and Flowing Blood: Aptamer devices were incubated in EDTA-treated human whole blood (Bio-IVT) or inserted via a catheter into a closed-loop flowing blood setup (Masterflex L/S Easy-Load II EW-77200-80 pump head at 1.5 mL min⁻¹) for 6 or 12 d at room temperature (Figure S19, Supporting Information). Probes were then immersed in 2.5% CFA for 15 min and rinsed with PBS. Probes were then freeze-dried, lyophilized, and coated with Au/Pd for SEM imaging.

In Vivo Assay of Aptamer Devices: Live animal studies were performed using Sprague–Dawley rats under Stanford APLAC protocol number 33226. Male rats were purchased from Charles River Laboratories at a weight of 300–350 g. The rats were anesthetized using isoflurane gas inhalation (2.5%) during the entirety of the experiment and drug infusions and monitored continuously. A small incision was made in both legs of the rat to expose both femoral veins. A 20G catheter was inserted into the left femoral vein for sensor probe insertion, and a 22G catheter was inserted into the right femoral vein for drug infusion. 0.1–0.3 mL of heparin (1000 U mL⁻¹, SAGENT Pharmaceuticals) were immediately infused through the catheter to prevent blood clotting. The needle of the 20G catheter was removed, catheter clipped, and the sensor was inserted deep into the vein, secured in place with a surgical suture, and allowed to equilibrate for 1–2 h. A sequence of bolus injection was performed manually using a 5 mL syringe. In each administration of kanamycin through the sensor-free 22G catheter, either 100 or 200 µL of 200 × 10^−3 m kanamycin in PBS buffer was injected. At the end of the experiments, animals were euthanized by exsanguination while under general anesthesia.

Degradation Assay for Hydrogels: Solutions of 12 µM HCl, 12 µM NaOH, 30% peroxide, and 1 mg mL⁻¹ Amano Lipase PS were prepared (Sigma-Aldrich). Hydrogel discs were placed into one of each of these solutions at 50 °C for up to 72 h. At specified time intervals, swollen hydrogel discs were removed from the solution and placed into deionized water for 24 h to remove salts. Samples were then freeze-dried and lyophilized before mass was recorded. Weights were normalized to mass at time 0 to determine mass degradation.

Optical Microscopy and Roughness of Hydrogels: Differential interference contrast (DIC) microscopy (10×) of the gel surfaces was performed with a Keyence VK-X250/260K 3D laser scanning confocal microscope. A 408 nm violet laser was used and image analysis was performed with VK Viewer and MultiFileAnalyzer software from Keyence. Surface roughness measurements used to quantify surface roughness measurements were defined and parameterized by ISO 25178 and measuring height profiles across the x–y plane of the hydrogel samples.

Statistical Analysis: All values of significance were determined using a one-way ANOVA or Student’s t-test with Prism GraphPad 8.4 software. As we noted drastic differences in clotting times between batches of blood, a paired t-test was used to determine significance between PEG-coated and F50–C50-coated probes in Figure 4. In Figure S5, an unpaired t-test was performed to compare performance of PEG-coated and F50–C50-coated sensors as there was one only group per leg of each animal (no pairing). In Figures S17 and S22 (Supporting Information), a one-way ANOVA test was used as there were multiple groups being compared.

Ethical Statement: All animal procedures were performed according to Stanford APLAC approved protocols.
Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest
D.C., J.-C.C., E.A., H.T.S., and E.A.A. are listed as authors on a provisional patent application describing the technology reported in this manuscript.

Author Contributions

Data Availability Statement
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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