Endothelial antigen assembly leads to thrombotic complications in heparin-induced thrombocytopenia

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Heparin-induced thrombocytopenia (HIT) is a prothrombotic disorder initiated by antibodies against complexes between human platelet factor 4 (hPF4) and heparin. A better understanding of the events that initiate the prothrombotic state may improve approaches to antithrombotic management. Here, we visualized thrombus formation in an in vivo murine model and an endothelialized microfluidic system that simulate the pathogenesis of HIT. hPF4 released from platelets predominantly bound to peri-injury endothelium and formed HIT antigenic complexes that were dissociated by heparin. In mice expressing both hPF4- and human platelet IgG Fc receptor IIA (FcγRIIA), infusion of the HIT-like monoclonal antibody KKO increased fibrin and platelet deposition at sites of injury, followed immediately by antigen formation on proximate endothelial cells. After a few minutes, HIT antigen was detected within the thrombus itself at the interface between the platelet core and the surrounding shell. We observed similar results in the humanized, endothelialized microfluidic system. hPF4 and KKO selectively bound to photochemically injured endothelium at sites where surface glycolocalyx was reduced. These studies support the concept that the perithrombus endothelium is the predominant site of HIT antigen assembly. This suggests that disrupting antigen formation along the endothelium or protecting the endothelium may provide a therapeutic opportunity to prevent thrombotic complications of HIT, while sparing systemic hemostatic pathways.

Introduction
Heparin-induced thrombocytopenia (HIT) is an iatrogenic disorder initiated by antibodies directed against a complex between a positively charged chemokine, platelet factor 4 (PF4, also known as CXCL4), and heparin (1) that can lead to limb- and life-threatening arterial or venous thrombi (2), which often occur at sites of vascular injury (3). HIT remains a common and serious disorder, even though the use of unfractionated heparin has been restricted to settings such as cardiopulmonary surgery (4, 5), where it remains the anticoagulant of choice (6). Present-day management with intense anticoagulation has not reduced the incidence of death or amputation, provides substantial but incomplete protection against recurrent thromboembolism, and is associated with a risk of bleeding for which antitodes are not available (7). A better understanding of the pathobiology of HIT might lead to novel disease-specific approaches that prevent immune complex–mediated thrombosis, with less reliance on intense systemic anticoagulation.

We have proposed a model to help explain the inordinate risk of thrombosis in patients with HIT compared with the thrombosis risk with other antibody-induced thrombocytopenias (8, 9): PF4 released from activated platelets binds to surface glycosaminoglycan side chains (GAGs) on intravascular and vascular cells. Infused heparin, which has a higher affinity for PF4 than for other GAGs, removes surface-bound PF4, forming circulating antigenic PF4–heparin complexes that fix complement and bind to CD21 on circulating antigen-presenting B cells (10). The antibodies that result bind to PF4-GAG complexes on the surface of platelets (9), monocytes (8), and neutrophils (11), leading to cell activation via Fc receptors (12), which promotes the generation of thrombin and other prothrombotic pathways (13).

Endothelial cells are targeted by HIT antibodies, at least in vitro (14–16), but the mechanism and in vivo relevance are unclear. The endothelium is covered by a complex surface layer termed the glyocalyx, which contains heparan sulfate–rich proteoglycans, glycoproteins, and associated plasma proteins (17, 18) that would be predicted to bind PF4 with greater affinity than the platelet surface. Binding of HIT antibody to cultured endothelial cells induces platelet adhesion and expression of tissue factor (15), but there is little direct evidence that HIT antibodies impair the natural antithrombotic properties of the endothelium in more biologically relevant models.

The spatial and temporal events that initiate thrombus development in HIT are not well delineated but are potentially of considerable import. Studies of platelet degranulation in the...
In this study, we used a previously described passive immunization model of HIT in mice double-transgenic for human PF4 and the human platelet IgG Fc receptor IIA (referred to hereafter as hPF4+/FcγRIIA+) (9, 22) and the cremaster arteriole laser injury model (23) to study in situ thrombus development in HIT. We extended these in vivo findings by using endothelialized microfluidic chambers, which allowed us to introduce localized hematoporphyrin-based photochemical injury to study the contribution of activated endothelium in a humanized system. Our data show that the perithrombus endothelium is the predominant site of HIT antigen formation, surprisingly at sites where the glycocalyx is depleted, followed by lower levels of antigen expression at the interface between the thrombus core and shell. We believe that these studies will serve to shift the focus of future studies on HIT pathogenesis and intervention to factors that regulate antigen assembly and the thrombotic consequences of antibody binding to the endothelium.

Figure 1. Widefield cremaster laser injury in a non-HIT hPF4+ murine model: In situ studies of hPF4 and HIT antigen distribution in thrombi. (A) Representative widefield study of more than 10 cremaster laser injuries in hPF4+ mice, with time 0 indicating the onset of injury. Images from a video of a laser injury; platelets are indicated in red, hPF4 is indicated in green, and the direction of blood flow in the vessel is denoted by blue arrows. Graph shows the accumulation of platelets and hPF4 over the study in relative value units (RVU) compared with time 0. (B) Same as in A, but with green showing binding of KKO to indicate the appearance of the HIT antigen. (C) Representative images from Supplemental Video 1 beginning 5 minutes after injury, the point at which 10^3 U/kg heparin was infused i.v. Platelets are indicated in red and KKO binding in green. The graph indicates that various doses of heparin were infused beginning 5 minutes after the cremaster injury. Percent mean ± 1 SEM for binding of KKO after heparin relative to the 5-minute time point is shown. The dashed line represents no change in KKO binding after heparin infusion compared with the 5-minute heparin time point. Original magnification, ×60.
Results

Distribution of HIT antigen after nonimmune vascular injury in hPF4+ mice. We previously proposed that a key step in the development of HIT was the formation of antigen when positively charged hPF4 binds to negatively charged GAGs found on many cell surfaces (9). This model predicts that HIT antigen will form at all sites of vascular injury, even in the absence of HIT-associated antibodies. We tested this hypothesis in vivo using a cremaster arteriole laser injury model in hPF4+ mice. Following injury, hPF4, detected using a polyclonal anti-hPF4 antibody, and HIT antigen, detected using the HIT-like monoclonal antibody KKO (24), were present almost immediately on widefield microscopy, and the involved area expanded over the ensuing 3 minutes (Figure 1, A and B, respectively). This result demonstrates that hPF4 released as a result of thrombus formation accumulates locally in the form of PF4-containing complexes that can be recognized by HIT antibody in mice expressing hPF4 in their platelets with or without HIT.

We had reported that high doses of heparin attenuated the severity of thrombocytopenia in hPF4+/FcγRIIA+ mice (9). We postulated that this occurred because heparin can dissociate antigenic complexes from platelet surfaces at these concentrations (9), given its higher affinity for hPF4 than for surface GAGs (25). In turn, this generates the circulating hPF4-heparin complexes needed to induce an immune response. (25, 26) Therefore, we asked whether infusing unfractionated or low-molecular-weight heparin into hPF4+ mice would mobilize prebound PF4 and decrease retention of the HIT antigen at the site of thrombus formation. Binding of KKO to the site of thrombus formation decreased proportionally with the dose of either unfractionated heparin (Figure 1C and Supplemental Video 1; supplemental material available online with this article; https://doi.org/10.1172/JCI90958DS1) or low-molecular-weight heparin (Supplemental Figure 1) infused into hPF4+ mice. Importantly, however, even when mice were given 1,000 U/kg unfractionated heparin—a dose approximately 10-fold higher than therapeutic doses (27) — about 25% of the HIT antigen remained, suggesting that the vasculature continues to be a potential target for immune injury. Also of note, mice that did not receive heparin had the highest level of local HIT antigenicity, with an approximate 2-fold increase in HIT antigen compared with mice receiving a dose in the range normally given clinically (100 U/kg) (28).

Study of the prothrombotic state in a murine HIT model. We have previously shown that infusion of KKO or HIT-associated IgG (HIT-IgG) induces thrombocytopenia and a prothrombotic state in hPF4+/FcγRIIA+ mice (8, 9, 22). We recapitulated the prothrombotic state in the cremaster arteriole laser injury model. Injection of KKO into hPF4+/FcγRIIA+ mice enhanced platelet and fibrin
Table 1. Secondary growth of thrombi in hPF4+/FcγRIIA+ mice leading to an occluded vessel

<table>
<thead>
<tr>
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<th>KKO (1 μg/g)</th>
<th>HIT IgG (50 μg/g)</th>
<th>TRA (1 μg/g)</th>
<th>Control IgG (50 μg/g)</th>
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<td>22</td>
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<td>No. of occluded vessels</td>
<td>24</td>
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<tr>
<td>% of occluded vessels</td>
<td>72.7%±6</td>
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Cremaster arterioles were injured by laser, and the accumulation of platelets was assessed. Five minutes later, KKO or TRA or HIT IgG or control IgG was infused i.v. at the indicated dose, and the site of injury was observed. The number of injured and occluded vessels with cessation of blood flow is indicated. A 72.7%±6 of occluded vessels of hPF4+/FcγRIIA+ mice, but not in hPF4+ mice or in hPF4+/FcγRIIA+ mice exposed to the isotype control antibody TRA (24) (Figure 2A).

Patients with HIT may be predisposed to develop thrombi at sites where vessels have been injured by catheters (3). To simulate this sequence in the cremaster model, we induced a thrombus in hPF4+/FcγRIIA+ mice, followed 5 minutes later by infusion of KKO or HIT-IgG. Preexisting thrombi expanded in mice given KKO or HIT-IgG, but not control antibodies (Figure 2B). These data also underestimate the severity of the prothrombotic state, because virtually all the vessels became occluded, limiting further thrombus expansion (Table 1 and Supplemental Video 2). Neither TRA nor normal IgG control infusions led to renewed growth of thrombi or vascular occlusion (Figure 2B and Table 1). Of note, thrombus regrowth began immediately after KKO was infused, but accelerated with time, demonstrating that a preexisting thrombus provides a potent nidus for expansion, leading to vascular occlusion in the setting of HIT antibodies.

Analysis of thrombus development in HIT by confocal microscopy. To begin to understand why thrombus regrowth was enhanced in HIT, we used confocal microscopy to examine the initial steps in thrombus formation following laser-induced arteriole injury in animals preinfused with KKO. Remarkably, the predominant site of KKO binding was the endothelium underlying and surrounding the thrombus, extending both upstream and downstream from the site of thrombosis (Figure 3A). The upstream binding appeared to be related to turbulent blood flow in this area (Supplemental Video 3).

By 5 minutes after injury, KKO began to appear on platelets within the clot, which intensified over time at the boundary between the tightly packed core and surrounding looser shell of the thrombus (21) (enlarged image in Figure 3A and Figure 3B). We posit that this hemispheric zone of antigen was formed by the slow spread of α-granule release outward within the core until reaching the core-shell interface (21). Our studies suggest that more loosely bound platelets on the surface of the thrombus or caught in turbulent flow around the thrombus may make a more immediate and more significant contribution to the detectable PF4 in the thrombus. Moreover, the PF4 released from surface-bound platelets probably adheres better to the peri-injury endothelium, with its glyocalyx, which is rich in high-affinity heparan sulfate and dermatan sulfate (17), rather than to the thrombus platelets, with their lower affinity surface chondroitin sulfate (29, 30). This platelet-bound PF4 is likely swept downstream. The overlap of injured endothelium and accumulation in hPF4+/FcγRIIA+ mice, but not in hPF4+ mice or in hPF4+/FcγRIIA+ mice exposed to the isotype control antibody TRA (24) (Figure 2A).

HIT was induced by adding KKO and hematoporphyrin to infused human whole blood prior to photochemical injury. KKO induced a significant increase in platelet adherence to HUVECs in the injured area compared with that observed with TRA (Figure 5A) and had a similar effect on similarly prepared adult human aortic endothelium (Supplemental Figure 5). Downstream of the injury, in the uninjured endothelium, no difference was noted between KKO and TRA exposure (Figure 5B). KKO also caused an increase of approximately 7-fold in PF4 binding to the injured endothelium compared with binding to the uninjured area, which was accompanied by an increase of approximately 16-fold in KKO binding to the injured endothelium compared with uninjured endothelium (Figure 5C). The injured endothelium showed an
Discussion

HIT differs from other antibody-mediated thrombocytopenias by its propensity for significant thromboembolism (2, 33). We have attributed the prothrombotic state to the concurrent activation of multiple cell types by immune complexes. Platelets are dual-ly activated by immune complexes through FcγRIIA (12) and by increase of approximately 50-fold in P-selectin surface expression (P < 0.0001, Figure 5D) and release of vWF (Supplemental Figure 6) and a decrease of approximately 55% in glycocalyx staining by lectin compared with uninjured endothelium (P < 0.01, Figure 5D). Thus, the increase in binding of PF4 to the endothelium correlated with the loss of glycocalyx and its associated GAGs, as had been seen following other endothelial cell injuries (18, 32). Like the results obtained with infusion of KKO, infusion of HIT-associated IgG (300 μg/ml) along with whole blood led to occlusion of 6 of the 6 channels studied compared with 1 of 6 channels exposed to control IgG (P = 0.02, Table 1).
monocyte-generated thrombin (34). However, to date, the proposed involvement of the endothelium is based entirely on in vitro findings using cultured cells.

To better understand the prothrombotic nature of HIT, we performed video examination of thrombus formation in HIT both in vivo and in vitro. These in vivo cremaster studies involved laser-induced thrombi rather than spontaneous thrombi, but they allowed a detailed temporal analysis of events. We anticipate that the results of future similar studies of spontaneous thrombi in HIT mice would be consistent with our findings. Our in vivo studies showed that released PF4 was associated with formation of the HIT antigen, as identified using the monoclonal antibody KKO and by the binding of HIT-IgG on every thrombus. Infused heparin stripped this bound and antigenic PF4 away from the thrombus, decreasing surface antigenic complexes. These findings suggest that the pathogenic contribution of heparin to the development of HIT is solely the antigenic complexes. These findings suggest that the pathogenic contribution of heparin to the development of HIT is solely the antigenic complexes. 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Proteins that do not have a high affinity for surrounding cell surfaces are probably rapidly washed away, while those like PF4 may adhere to cells within or surrounding the thrombus.

The microfluidic photochemical injury studies, while involving a more diffuse vascular injury than did the cremaster injuries, support the conclusion that the endothelium is the predominant initial harbor for PF4 released from activated platelets and a target for HIT antibodies. While PF4 and KKO bind directly to platelets (9), monocytes (8), and neutrophils (37), we found that binding was more intense on injured endothelial cells. This is surprising, given that the glyocalyx is highly enriched for negatively charged side chains on its constituent proteoglycans, syndecans, and glypicans (17). These are candidate receptors for binding PF4, and one would have expected that with their loss, PF4 binding would have decreased. The loss of the glyocalyx involves multiple glycosidases or sheddases that remove hyaluronic acid residues and many other polyanions in this layer (38). We posit that the enhanced binding to injured glycocalyx may be due to the unveiling of a novel high-affinity binding site in the glyocalyx or the release of novel binding molecules from the injured endothelium that then reside in the remaining glyocalyx.

These studies support a pathogenic scenario, in which binding of PF4 leads to the formation of immune complexes on the endothelium, which may enhance platelet adhesion, enhance thrombin generation, and cause the release of additional PF4, which would sensitize the downstream endothelium and lead to a feed-forward pathway that propagates thrombosis. Such a phenomenon could contribute to the vessel occlusion we observed in the HIT cremaster arteriole model (Figure 2B) and, as observed earlier in HIT, to the presence of long, platelet-rich thrombi, resulting in the alternative name for this disorder of “white clot syndrome” (39). Whether these extended clots are related to a recruitment of undamaged endothelium in HIT needs to be examined. If this model is correct, it will be important to identify as potential therapeutic targets...
the biophysical and biochemical changes in the endothelium that promote antigen and antibody binding. Moreover, these studies suggest that endothelial cell–targeted antiinflammatory and antithrombotic drugs (40, 41) might provide a rational intervention focused on the primary vortex of prothrombotic reactions that lead to thrombotic vascular occlusion.

**Methods**

**Mice and human samples.** Transgenic mice expressing platelet-specific hPF4 (42) (hPF4+) and/or human FcγRIIA expressing its R131 isoform (43) (FcγRIIA+) were studied. All transgenic mice were on a Cxcl4−/− background (44), as murine PF4 is not targeted by HIT-associated antibodies (24). This knockout setting, common to all the mice studied, is not specified hereafter. Genetic alterations were confirmed by the appropriate PCR analyses (42, 43). Mice were studied at 6 to 10 weeks of age. Only male mice were studied for the cremaster vessel injuries; however, we have not noted any prior sex difference in thrombosis in the passive immunization HIT murine model (8).

Human blood (10–25 ml) from healthy volunteers for in vitro studies was drawn by gravity through a 19-gauge butterfly into sodium citrate (Sigma-Aldrich; 0.38% final concentration). Blood samples were stored at room temperature and used within 1 hour of being drawn. Deidentified plasma samples were obtained from patients who had a high pre-test probability of HIT on the basis of their clinical history (45), a positive PF4/heparin ELISA, and a positive serotonin release assay (46). For plasma samples from healthy donors, whole blood was centrifuged (200 g, 15 minutes), followed by centrifugation (2,000 g, 15 minutes) of the resultant platelet-rich plasma.

**Heparins, antibodies, pooled IgGs, and other labeled probes.** Unfractionated porcine heparin (BD) and low-molecular-weight heparin (enoxaparin; Novaplus) were used in this study. KKO, a mouse IgG2bK anti-hPF4/heparin monoclonal antibody, and TRA, a monoclonal IgG isotype control antibody (24), were purified from hybridoma supernatants. F(ab′)2 fragments of the monoclonal anti-mouse CD41 antibody MWReg30 (BD Biosciences) were used to detect murine platelets in the cremaster laser injury model. Anti–fibrin 59D8 monoclonal antibody was provided by Hartmut Weiler of the BloodCenter of Wiscon-

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**Figure 5. Studies of HIT in the endothelialized microfluidic channel photochemical injury system.** (A) Platelet accumulation along HUVECs after photochemical injury perfused for 15 minutes with whole human blood containing either KKO or TRA. Data represent the mean ± 1 SEM. *P < 0.0001, for KKO versus TRA exposure by 2-tailed Student t test. (B) Images demonstrate platelets bound to representative fields of photochemically injured and uninjured areas of the channels shown in A from whole blood containing TRA or KKO. (C) Images of representative uninjured and injured areas and graphs of overall comparative measurements showing enhanced binding in injured versus uninjured areas of PF4 (white) and KKO binding (green) after infusion of whole blood, as in A. The mean relative binding for injured versus uninjured endothelium ± 1 SEM is shown. *P < 0.01 and **P < 0.001, by 2-sided Student’s t test for injured versus uninjured areas, with an expectation of 1 and no change in binding (dashed line). (D) Same as in C, except for P-selectin staining (yellow) and lectin binding (yellow) in injured endothelium versus the downstream uninjured area. Lectin staining of the endothelium was decreased after injury. Original magnification, ×20.
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Annexin V labeled with Alexa Fluor 647 was purchased from Thermo Fisher Scientific. Recombinant FXα labeled with Alexa Fluor 488 was provided by Rodney Camire of the Children’s Hospital of Philadelphia. Mouse anti-human P-selectin antibody (clone CTB201) was purchased from Santa Cruz Biotechnology Inc. Polyclonal rabbit anti-human vWF antibody was purchased from Dako. hPF4 was visualized using polyclonal rabbit anti-hPF4 antibody (Abcam). All antibodies were either labeled using Alexa Fluor antibody-labeling kits according to the manufacturer’s instructions or species-appropriate Alexa Fluor-conjugated secondary antibodies (all from Thermo Fisher Scientific). IgG was isolated from plasma from HIT patients and healthy donors using protein G agarose (Pierce, Thermo Fisher Scientific).

Cremaster laser injury studies. Intravital microscopy was performed as previously described (48). Vascular injury was induced with an SRS NLI100 pulsed nitrogen dye laser (440 nm) focused on the vessel wall through the microscope objective. Arterioles of 20 to 40 μm diameter were selected, and the laser was pulsed until the vessels were perforated and a small number of red blood cells escaped. Antibodies and unfractionated porcine heparin (BD Biosciences) were infused as 100-μl boluses via a catheter placed into the jugular vein. Widefield and confocal microscopy were performed as described previously (20). Data were collected and widefield time-lapsed images of platelet and fibrin accumulation were analyzed using Slidebook 6.0 (Intelligent Imaging Innovations). Confocal Z-stacks were analyzed using Velocity 6.3 (PerkinElmer). hPF4, FcγRIIA+, and hPF4+/ FcγRIIA− mice were studied. We studied 1-10 injuries per mouse during a maximum experimental time of 1 hour.

Endothelialized microfluidic studies. Microfluidic studies were performed using a BioFlux 200 Controller (Fluxion) with an attached heating stage set to 37°C as described previously (34). The BioFlux controller was used in conjunction with a Zeiss Axio Observer Z1 inverted microscope equipped with a motorized stage and an HXP-120 C metal halide illumination source. The microscope and acquisition were controlled using BioFlux Montage software with a MetaMorph-based platform (Molecular Devices). HUVECs and adult human aortic endothelial cells (both from Lonza), at passage 3-4 (5 × 10⁶ cells), were seeded onto fibronectin-coated (50 μg/ml, Sigma-Aldrich) channels of 48-well BioFlux plates (Fluxion), allowed to adhere, then cultured at 37°C under 5% CO₂ in endothelial cell growth media (Lonza) until they reached confluence.

An HT-like state was induced by adding KKO (10 μg/ml) to sodium citrate-anticoagulated whole blood from healthy donors immediately before infusion. Platelets in the whole blood were labeled by incubating the blood with 2 mM calcium AM (Thermo Fisher Scientific) for 20 minutes prior to infusion. To cause photochemical injury without disrupting the endothelial cell lining, hematoporphyin (50 μg/ml final concentration; Sigma-Aldrich) was also added to the whole blood prior to infusion (31). Channels were exposed to blue light using the HXP-120 C light source with 475-nm excitation and 530-nm emission filters, allowing real-time, concurrent visualization of calcein AM-loaded platelets and ROS generation from hematoporphyin. An exposure time of 50 ms at the highest intensity setting on the light source was used for all experiments. The blood was recalcified immediately before infusion with calcium chloride (11 mM final concentration). Infusion into endothelialized channels was done at a rate of 10 dynes/cm². Following injury, channels were washed with PBS and fixed with 2% paraformaldehyde (BD Biosciences) for confocal studies of P-selectin, PF4, and KKO. To measure the glycosyla thickness, HUVECs lining the channels were stained with 1 μg/ml DyLight 488–labeled Lycopersicum esculentum lectin (Vector Laboratories) for 20 minutes prior to fixation (49). Endothelial cell nuclei were identified by staining with 5 μg/ml Hoescht 33342 (Thermo Fisher Scientific) for 20 minutes. Stained channels were imaged with a Zeiss LSM 710 laser scanning confocal microscope. Data were analyzed using Slidebook 6.0 (Intelligent Imaging Innovations) for photochemical injuries or Volocity 6.3 (PerkinElmer) for confocal images acquired after fixation.

Statistics. Differences between 2 groups were compared using a 2-sided Student’s t test or a Mann-Whitney U test. Differences between more than 2 groups were determined by 2-way ANOVA with Sidak’s correction for multiple comparisons. Occlusion studies were compared using a 2-sided Fisher’s exact test. Statistical analyses were performed using Microsoft Excel 2011 and GraphPad Prism 6.0 (GraphPad Software). Differences were considered significant when P values were less than or equal to 0.05.

Study approval. All animal procedures were approved by the IACUC of the Children’s Hospital of Philadelphia and in accordance with NIH guidelines (Guide for the Care and Use of Laboratory Animals. National Academies Press. 2011.) and the Animal Welfare Act. Human blood was collected after signed, informed consent was provided by healthy donors, and approval for studies using human blood was obtained from the Children’s Hospital of Philadelphia Institutional Human Review Board in accordance with Declaration of Helsinki principles.

Author contributions

VH designed and performed most of the experiments, was primary author of the manuscript, and prepared the first draft of the manuscript. JI performed microfluidic studies and helped write the related sections of the manuscript. GMA and SEMck provided important insights into the proposed studies and outcomes interpretation. DBC and LR helped with study design, data analysis, and editing of the manuscript. MP provided overall scientific guidance and helped prepare the manuscript.

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