Increasing Binding Efficiency via Reporter Shape and Flux in a Viral Nanoparticle Lateral-Flow Assay

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Supporting Information

INTRODUCTION

Point-of-care (PoC) devices enable medical testing at the site of patient care and are becoming increasingly important. The lateral-flow assay (LFA), one of the most widely used and commercially available PoC formats, is inexpensive, simple, portable, and rapid, and hence is well-suited for testing in settings lacking complex laboratory equipment. In a typical LFA format, analytes suspended in a fluid from a sample are dispensed onto one end of a test strip, typically a porous membrane. Capillary forces drive the fluid to flow along the strip; the analytes, entrained in the flowing fluid, are captured by antibody-coated reporters dried on an upstream conjugate pad. Capture antibodies at the test line recognize and capture the analyte-reporter complexes to produce a visual signal. Although traditional LFAs are widely employed to detect pregnancy and infection, their use elsewhere is often limited by lack of analytical sensitivity and difficulties in quantification. The performance of LFAs can be improved by tuning assay parameters, including fluid velocity, reagent concentration, and reporter particle size. Increasing the analytical sensitivity of LFAs would expand their usefulness, including pathogen monitoring in food and biopharmaceutical manufacturing. Research efforts toward developing sensitive and quantitative LFAs suitable for a broader range of applications have focused on new assay membranes, reporter materials, and read-out methods. In addition to assay materials, the affinity and kinetics of the antibodies used, and readout modalities, the sensitivity of LFAs relies on the transport and binding of analytes and reporters to the sensing surface. Binding effectiveness is measured by the number of binding events, which depends on the balance between the rate at which analytes encounter the sensing surface and the duration of their encounters with receptors there. The rate at which targets encounter the sensing surface, therefore, is as critical as molecular binding kinetics in increasing the number of binding events. In many microfluidic-based sensors, targets are delivered to the sensing surface through convective and diffusive transport processes, for which Péclet (Pe) and Damköhler (Da) numbers can be calculated to identify the mechanisms that dominate transport and binding. Reaction-limited conditions, the binding-reaction rate is much slower than the rate of transport to the surface (i.e., Da ≫ 1), and hence, analytes must encounter the surface for long binding times. Decreasing the flow rate can increase the encounter duration but does not necessarily increase total successful binding, as the rate at which targets encounter the sensing surface...
surface also decreases. To increase LFA sensitivity, there is a need to understand the effects of flow rate on the binding of analytes and reporters.

Most LFA reporters are spherical nanoparticles of gold, latex, carbon, or silver, which are inexpensive and are readily manufactured in a range of sizes.\(^2\) Recent studies of adhesion of biological particles to surfaces, however, suggest that nonspherical nanoparticles may bind more efficiently than spherical nanoparticles.\(^2\) At least two mechanisms have been proposed to explain this increased surface binding efficiency. First, torques on nonspherical particles in shear flows can generate tumbling, rolling, or lateral drifting dynamics\(^2\) that increase the rate of encounter with the surface. Second, a nonspherical particle has greater surface area and lower hydrodynamic drag than a sphere of the same volume.\(^2\) These mechanisms would also be expected to affect binding of nonspherical LFA reporters. Indeed, a recent study showed that gold-coated silica nanorods used as reporters improved the sensitivity of an LFA for proteins by 50-fold compared to spherical gold nanoparticles.\(^3\) Surprisingly, the role of shape in LFA sensitivity has not yet been systematically explored, in part because many synthetic nanorods are very heterogeneous in both size and shape.\(^2\)

In previous work, we developed functionalized M13 filamentous bacteriophage (phage) as an efficient, inexpensive, and biocompatible reporter for very sensitive LFAs.\(^2\) M13 phage are filamentous viral nanoparticles (900 nm long and 6 nm diameter) that infect and replicate within Escherichia coli. Phage produced within E. coli are very uniform in size and shape, and can readily serve as a scaffold on which to attach multiple biorecognition and read-out-signal molecules. Recently, we showed that the flow field within an LFA membrane couples to the phage shape and promotes reorientation for binding, and hypothesized that binding efficiency can be increased by tuning the size and shape of reporters.\(^4\)

In this paper, we show that the binding efficiency in an LFA is modulated by reporter shape and is coupled to flux. We used three filamentous phage of different sizes as model reporters in a simplified LFA. Phage were functionalized on their coat proteins with biotin, a recognition molecule, and AlexaFluor 555 carboxylic acid (Succinimidyl Ester, Life Technologies #A21009) on their coat proteins as previously described.\(^5\) Biotinylated Fluor-phage were detectable by epi-fluorescence microscopy and capturable by NeutrAvidin (Figure 1).

**Biotinylation of Phage**. Asla Biotech generously provided wild type filamentous Pf1 bacteriophage (Pf1) and potato virus M virus-like particles\(^6\) (PVM), and M13 bacteriophage (M13) was from the American Type Culture Collection (#34768). Phage were biotinylated on the primary amines of the major phage coat proteins using EZ-Link Sulfo-NHS-LC-Biotin reagent (ThermoFisher Scientific, #21327), according to the manufacturer’s instructions. We used \(10^7\) molecules of biotin reagent to modify \(\sim 10^9\) phage, and removed excess biotin using a 7K MWCO Zeba spin desalting column (Thermo Scientific #89877).

**AlexaFluor 555 Labeling of Phage** (Fluor-Phage). Biotinylated Pf1, PVM, and M13 phage were further modified with AlexaFluor 555 carboxylic acid (Succinimidyl Ester, Life Technologies #A2-0009) on their coat proteins as previously described.\(^6\) Biotinylated Fluor-phage were detectable by epi-fluorescence microscopy and capturable by NeutrAvidin (Figure 1).

**Confirmation of Fluor-Phage Biotinylation Using Enzyme-Linked Immunosorbent Assay (ELISA)**. A 100 \(\mu\)L portion of biotinylated Fluor-phage or control nonbiotinylated Fluor-phage at concentrations of 0, 10^4, 10^5, and 10^6 pfu mL\(^{-1}\) in 2% (w/v) BSA were added to NeutrAvidin-coated 96-well plates (Pierce Reacti-bind, Thermo Scientific #15128). After 2 h incubation at room temperature, the plate was rinsed four times each with PBST (PBS containing 0.05% (v/v) Tween 20) and PBS, respectively. Next, bound biotinylated Fluor-phage were allowed to react with 100 \(\mu\)L of horseradish peroxidase (HRP)-conjugated StreptAvidin (1:5000 dilution, ThermoFisher Scientific, #N100) for 1 h at room temperature, and then were rinsed. A 50 \(\mu\)L portion of TMB was added to the plate to develop the color for 15 min, and the reaction was terminated by adding 50 \(\mu\)L of 2 N H\(_2\)SO\(_4\). Absorbance was measured at 450 nm in an ELISA reader (Tecan, Infinite M200 PRO).

**Flow Experiments**. Fusion 5 LFA matrix strips (GE Healthcare & Life Sciences #8151-9915), with average pore diameter 8.9 ± 0.1 \(\mu\)m, average pore throat diameter 3.1 ± 0.1 \(\mu\)m, porosity 0.81 ± 0.01, and average fiber diameter \(D = 4 ± 2 \mu\)m; cut into 3 mm × 50 mm × 0.35 mm strips) were modified with 3 \(\mu\)L of 1 mg mL\(^{-1}\) NeutrAvidin (Thermo Scientific, #3100) in 50 mM acetate buffer at pH 4.0 that was spotted at a fixed distance of 1 cm from the downstream end of the strip. After drying in air for 2 h at room temperature, each functionalized strip was placed on a large coverslip (48 mm × 65 mm, Gold Seal), covered with a second, smaller coverslip (22 × 22 mm, Fisher Brand), and placed on the stage of an epi-fluorescence microscope (Leica DMI 3000B) equipped with a 63× oil immersion objective lens (NA 1.4), as shown in Figure S-1. For model LFA...
experiments, biotinylated Fluor-phage were suspended in a background solution containing 0.1% (w/v) hydrolyzed polyacrylamide (HPAM, FLOPAAM 3330, SNF, weight-averaged molecular weight of 8 MDA) and 30% (w/v) glycerol in PBST (PBS containing 0.01% (v/v) Tween 20) to increase the fluid viscosity (and thereby reduce the flow speed) and to nearly match the index of refraction of the fibers in the LFA strip, improving the image quality. Subsequently, 25 μL of 2-fold serially diluted phage at concentrations of \( \frac{1}{5} \times 10^8 \), \( \frac{1}{5} \times 10^7 \), \( \frac{1}{5} \times 10^6 \), \( \frac{1}{5} \times 10^5 \), \( \frac{1}{5} \times 10^4 \), and \( \frac{1}{5} \times 10^3 \) pfu mL\(^{-1}\) was dropped on the upstream end of the strip. Biotinylated Fluor-phage were imaged in capillary-driven transport through the membrane using a sCMOS camera (pco.edge 4.2, 30 frame s\(^{-1}\), 30 ms exposure time, 208.3 μm × 213 μm image field of view). The position of the fluid interface scaled as the square root of time, following Washburn’s equation. Recording began immediately after liquid breakthrough in the field of view (t = 0 s) and ceased when the phage particles exhibited only Brownian (rather than advective) motion, typically after 6 min. Three movies in different membranes were analyzed for each concentration and type of phage.

### Counting of Bound Phage

**Counting of Bound Phage.** In flow experiments, phage bound to fibers over time and accumulated in the images as static particles (Figure S-2a). To locate and count phage bound to fibers over time, we manually analyzed nine 3 s intervals (starting at 20, 60, 100, 140, 180, 220, 260, 300, and 340 s) in each of the three movies for a given phage and concentration and identified groups of bright pixels as bound phage. We set the position of each bound phage as its brightest pixel (Figure S-2b). This position differed by less than a pixel from the centroid position, calculated by weighting each pixel with its intensity; hence, we obtained half-pixel accuracy.

**Preimage Processing for Particle Image Velocimetry (PIV).** We measured the local velocity at different positions in the field of view using PIV. Phage bound to the membrane generated a static background signal that impaired PIV processing. Hence, prior to PIV processing we removed bound phage and other static background in the images. For each 3 s movie we created a background image (Figure S-1c) that contained the minimum intensity at each pixel over time. This background was subtracted from each image in the movie (Figure S-2d).

**PIV Analysis.** To obtain the flow velocity profiles, PIV was performed on the images in 3 s movies of a sample containing M13 phage at a high concentration of \( \frac{1}{5} \times 10^8 \) pfu mL\(^{-1}\). A high phage concentration improved statistical correlations and reduced error across the field of view of the image. After preprocessing, we performed PIV (JPiV software, downloaded from http://www.jipv.vennemann-online.de/) using an interrogation window of 8 pixel × 8 pixel, a vector spacing of 3 pixel × 3 pixel, and a median filter of 3 pixel × 3 pixel. Four sequential images were correlated and averaged to generate PIV output at 20, 60, 100, 140, 180, 220, 260, 300, and 340 s. From the flow vectors obtained from PIV, flow rates and directions were decoupled and represented as separated maps (Figure S-3).

**Determination of Flow Direction on Fiber Surface.** To locate the fibers, a bright-field micrograph of fibers was acquired in the same focal area as each flow movie. Because the intensity and transparency of fibers varied across each image (Figure S-4a, left), we segmented the fiber features from the background manually; fiber areas in the image were painted with black to generate extreme contrast, after which the threshold was applied to the painted image (Figure S4a, center). To obtain the fiber directions, each fiber segment was labeled using connected component labeling. Next, an axis-aligned bonding box was applied to the labeled fibers. The orientation of fibers was calculated (in degrees, relative to the + x-axis) from the length and width of the bounding box (Figure S-4a, right). Finally, the flow direction obtained from PIV was correlated with the fiber orientation. The range of relative angles between flow and fiber varied from 0° (parallel) to 90° (vertical). Four different formulas were used to calculate the relative angle, depending on the angular ranges of the fiber (φ) and flow (θ) directions (Figure S-4b).

### Correlation of Local Flow with Bound Phage

**Correlation of Local Flow with Bound Phage.** The output from the PIV algorithms (Figure S-5a) provided both the magnitude and direction of flow at each location within the image. From this output, the local flow rate (Figure S-5d) and direction (Figure S-5e) were obtained for each position on a fiber (Figure S-5f) at which a phage bound.

### RESULTS AND DISCUSSION

#### Characterization of Biotinylated Fluor-Phage

Three filamentous phage of different dimensions (PVM, length 200 nm and width 13 nm; M13, length 900 nm and width 6 nm; and Pf1, length 2000 nm and width 6 nm; Table 1) were functionalized with biotin and AlexaFluor 555. Fluorescence micrographs confirmed the successful fluor labeling of biotinylated phage (Figure 2). We assayed the biotinylation

#### Table 1. Phage Dimensions and Relative Volume, Surface Area, and Swept Volume

<table>
<thead>
<tr>
<th>Phage</th>
<th>Diameter (nm)</th>
<th>Length (nm)</th>
<th>Relative Volume</th>
<th>Relative Surface Area</th>
<th>Relative Swept Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVM</td>
<td>13</td>
<td>200</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>M13</td>
<td>6</td>
<td>900</td>
<td>0.96</td>
<td>2.1</td>
<td>91</td>
</tr>
<tr>
<td>Pf1</td>
<td>6</td>
<td>2000</td>
<td>2.1</td>
<td>4.6</td>
<td>1000</td>
</tr>
</tbody>
</table>

*Phage were approximated as cylinders of radius r and length l, so that the volume was calculated as \( \pi r^2 l \). The PVM volume was chosen as the reference volume for relative comparisons. The surface area was calculated as \( 2\pi rl \). The PVM surface area was chosen as the reference surface area for relative comparisons. The swept volume, the volume of the sphere swept out when phage rotated in all three dimensions, was calculated as \( 4\pi r^3/3 \). The PVM swept volume was chosen as the reference swept volume for relative comparisons.

**Figure 2. Phage reporter series.** Bacteriophage (a) PVM (length 200 nm and width 13 nm), (b) M13 (length 900 nm and width 6 nm), or (c) Pf1 (length 2000 nm and width 6 nm) were covalently modified with biotin and AlexaFluor 555. Fluorescence micrographs confirmed the successful fluor labeling of biotinylated phage (Figure 2). We assayed the biotinylation of the Fluor-phage via binding on NeutrAvidin-coated plates. Biotinylated Fluor-phage exhibited a positive signal whereas control nonbiotinylated Fluor-M13 gave no significant signal (Figure S-6), confirming that only biotinylated Fluor-phage bound specifically to the NeutrAvidin-coated plate.

#### Comparison of Number of Bound Phage

**Comparison of Number of Bound Phage.** To compare the binding efficiency of phage of different lengths in the Fusion 5 membrane, we counted the number of bound phage as a function of time and concentration (\( \frac{1}{2} \times 10^8 \), \( \frac{1}{4} \times 10^8 \), \( \frac{1}{8} \times 10^8 \), \( \frac{1}{16} \times 10^8 \), and \( \frac{1}{32} \times 10^8 \) pfu mL\(^{-1}\)), as shown in Figure 3a–c for PVM,

**Figure 3.**

- (a) Fluor-PVM
- (b) Fluor-M13
- (c) Fluor-Pf1

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M13, and Pf1 phage. The number of phage bound on the membrane increased with the offered concentration of phage and with time for each phage series. In all experiments, phage predominantly bound early after breakthrough of the fluid. Over half of bound phage bound to fibers within 100 s, before the bulk flow rate had decreased by about 50% (Figure S-7a). After flow ceased, unbound phage exhibited Brownian motion from which their diffusivity could be calculated from the mean-square displacement (Figure S-7b). Because the flow rate was related to phage flux, we counted the flowing phage in the field of view at 20, 60, 100, 140, 180, 220, 260, 300, and 340 s by subtracting the number of bound phage from the total number of phage in the field of view. The number of flowing phage in the membrane was nearly constant over time (Figure S-8a–c), as the relatively large pore diameter (9 µm) did not hinder phage transport. The flux of phage through a fixed field of view decreased over time (Figure S-8d–f). Because the number of flowing phage was constant, the decrease in flux reflected a decrease in the flow velocity (Figure S-7a) and capillary flow rate. The rate at which phage bound to the fiber network, calculated from the instantaneous derivative of the curves in Figure 3, was proportional to phage flux (Figure 4). This result suggested that increasing the chance that phage encounter fibers (by increasing their flux) promoted binding.

The three phage have different widths and lengths (Table 1). Because we varied both the dimensions and concentration of phage, we examined the dependence of binding on several metrics related to phage morphology, including total volume, surface area, and swept volume (per unit volume of solution). To calculate the total volume and surface area of phage, we approximated phage as cylinders of volume \( \pi r^2 l \) and surface area \( 2 \pi rl \), where \( r \) and \( l \) are radius and length of the phage, respectively, and multiplied by the phage concentration. To calculate the total swept volume, we calculated the volume of the sphere that would be created by a phage rotating in all three dimensions around its center of mass (swept volume = \((4/3)\pi r^3\)) and multiplied by the phage concentration. The total number of bound phage from a 25 µL solution is shown as a research article.
function of number, total volume, total surface area, and total swept volume of phage in Figure 5a–d, respectively. Generally, the number of bound phage increased with each metric. At a fixed concentration of $1 \times 10^8$ pfu mL$^{-1}$, for example, the number of bound Pf1 (longest) phage was greater than that of PVM (shortest) phage (Figure 5a). The total volume occupied by phage (Figure 5b) did not collapse all the binding curves, suggesting that binding was not controlled by crowding. Similarly, the total spherical volume swept out by phage as they flowed through the membrane (Figure 5d) did not collapse the binding curves, suggesting that phage did not tumble in flow. Instead, the number of bound phage collapsed for all three phage onto a single curve as a function of the total surface area (Figure 5c). These results indicated that the binding efficiency was related to phage surface area, not volume or swept volume. The surface area of phage increased linearly with the number of capsid proteins (CPs), $\sim$1600 CP for PVM, $\sim$2800 CP for M13, and $\sim$7000 CP for Pf1 (R$^2$ = 0.987), but not length because of the variation in diameters; hence, the number of bound phage also collapsed for all phage onto a single curve as a function of the number of capsid proteins (Figure S-9). This result suggests that a similar fraction of capsid proteins bore biotins or that the biotin density was sufficient for efficient capture. The larger surface area (and hence number of capsid proteins) of the elongated phage increased the probability of interaction between phage and fibers in the membrane, suggesting that binding efficiency should increase with aspect ratio for phage reporters of fixed volume.

Surprisingly, the total swept volume did not correlate with the binding efficiency of phage, implying that phage do not rotate during flow. To test this idea, we used Jeffery orbit theory to evaluate the period of rotation of each phage in a steady linear flow field,$^6$ valid for a ellipsoidal particle in an infinite fluid for which Brownian motion can be neglected. We first confirmed that our system approximately satisfied the two Jeffery orbit conditions: (1) the flow was laminar and steady, and (2) advective transport dominated over diffusivity. Given the open pore network, we estimated the Reynolds number as $Re = \rho DV/\mu$, where $\rho = 1094$ kg m$^{-3}$ was the background solution density, $D$ was the average fiber diameter, $V$ was the phage velocity as a function of time (Figure S-7a), and $\mu = 8.67$ cP was the solution viscosity measured at a fixed shear rate of 42 s$^{-1}$ using an Ubbelohde viscometer. Similarly, we estimated the Péclet number as $Pe = DV/D_\phi$, where $D_\phi = 0.134$ mm$^2$ s$^{-1}$ was the measured translational diffusivity of the Biotin-FlourM13 (Figure S-7b). At all times, $Re \ll 1$ (Table 2), confirming that the flow was laminar and steady, and $Pe \gg 1$, confirming that Brownian motion could be neglected. Finally, phage were rodlike with high aspect ratios and could be approximated as ellipsoids. We therefore applied Jeffery orbit theory to calculate the phage tumbling period, $T = \gamma (r + 1/r)$, where $\gamma$ was the average shear rate (calculated by dividing the flow rate by the pore diameter) and $r$ was the phage aspect ratio. The tumbling period increased with phage length (Table 2).

**Table 2. Péclet and Reynolds Numbers, and Corresponding Tumbling Period of Phage as a Function of Time**

<table>
<thead>
<tr>
<th>time (s)</th>
<th>Reynolds number</th>
<th>Péclet number (for M13 phage)</th>
<th>tumbling period (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>$1.7 \times 10^{-2}$</td>
<td>$1.0 \times 10^{5}$</td>
<td>PVM: 32, 33, Pf1: 72</td>
</tr>
<tr>
<td>100</td>
<td>$8.8 \times 10^{-2}$</td>
<td>$5.0 \times 10^{5}$</td>
<td>PVM: 66, 63, M13: 143</td>
</tr>
<tr>
<td>180</td>
<td>$5.1 \times 10^{-2}$</td>
<td>$3.0 \times 10^{5}$</td>
<td>Pf1: 13, 97, M13: 238</td>
</tr>
<tr>
<td>260</td>
<td>$1.5 \times 10^{-2}$</td>
<td>$9.1 \times 10^{5}$</td>
<td>PVM: 53, 388, Pf1: 575</td>
</tr>
</tbody>
</table>

Effect of Local Flow Conditions on Phage Binding. We hypothesized that the local flow near/on the fiber surfaces controlled binding of phage. Hence we identified the location of fibers in the fluorescence movies by acquiring bright-field images in the same focal area. Next, the flow profile was extracted across the entire fiber area, and the speed and direction of the local flow were obtained at the positions where phage bound. These comparisons allowed us to relate the magnitude and direction of the local flow to the likelihood that phage bound there.

The speed and direction of the local flow at the locations of bound phage varied with time (Figure S-10), reflecting the decrease in capillary flow rate over time. To determine the effect of the local flow speed, we first normalized the flow speed by the difference between the local minimum and maximum speed for each PIV result (at 20, 60, 100, 140, 180, 220, 260, 300, and 340 s after breakthrough) and then calculated the average local flow speed across the entire fiber area. Finally, we identified domains in which the local flow speed was slower and faster than the average local flow speed. Distributions of the number of bound phage as a function of the local flow speed are shown in Figure S-11. Each type of phage (PVM, M13, Pf1) bound in greater numbers in the faster local flow domain than in the slower domain, and more so as the aspect ratio increased (Figure 6). The result that phage were more likely to bind in the faster local flow domain is consistent with the idea that higher phage flux promotes binding, as suggested by the bulk flow studies (Figure 4).

Similarly, we compared the likelihood of binding to the angle at which the local flow encountered the fiber (Figure S-11). Here, we surmised that vorticity generated by flow near the fiber surface might reorient the phage and hence influence their binding interaction. In contrast to the results for local velocity, however, phage did not exhibit a preferred local flow direction for binding (Figure S-12). Instead, the distribution of local flow angles across the bound phage population was constant within the error of the measurements. This omnidirectional binding likely arises from the large number of potential bonding interactions offered by the biotins attached to the phage major coat proteins.$^{30}$ Phage approaching the fiber could be captured on any of the biotins, thereby diminishing the dependence on orientation.

**CONCLUSIONS**

We investigated the effect of two assay parameters (phage flux and size) on the binding efficiency of phage reporters in fibrous glass LFA membranes. Filamentous phage of different lengths and widths were functionalized with biotin and AlexaFluor 555 on their coat proteins. Fusion 5 LFA strips were functionalized with NeutrAvidin. The number of phage bound to the fibers in the membrane was counted as a function of time in movies recorded during capillary flow. The rate at which phage bound correlated with their bulk flux through the membrane. At the
Figure 6. Binding efficiency depends on local flow rate. (a) PIV analysis and the (b) extracted flow speeds. (c) Flow speeds at the positions of the fibers, with (d) positions where individual phage bound overlaid (as ovals). The average local flow rate (vertical dashed lines in parts e–g) was calculated across the entire fiber. Slower and faster flow domains featured local flow rates that were slower and faster, respectively, than the average local flow rate. Phage binding only occurred on the fiber surface. (e–g) Number of bound phage as a function of local velocity for (e) PVM, (f) M13, and (g) Pf1. The orange vertical dashed lines indicate the boundaries between slower and faster local flow domains. For each phage type, a greater number of phage bound in the faster local flow domain than in the slower domain. Insets to parts e–g: Percent of phage binding in slower and faster domains. Error bars indicate the standard deviation measured from three independent experiments in different Fusion 5 membranes.

local (pore) scale, phage were more likely to bind in regions within the membrane with faster local flow (and hence higher local flux). The number of bound phage also increased with phage length and aspect ratio. The capture cross-section controlling the binding efficiency was highly correlated with the phage surface area (but not with total phage volume, concentration, or swept volume) and with the number of capsid proteins, consistent with the idea that increasing the average bulk flux increased the likelihood that phage encountered a fiber.

Our results suggest that increasing the flux of phage increases the binding efficiency of elongated reporters in fibrous membranes. In turn, this suggests that careful tuning of membrane design variables such as the pore diameter and membrane geometry (which control the local and bulk flow rates) could increase binding efficiency and thus the sensitivity of LFAs. Similarly, increasing the surface area of reporters increases their binding efficiency, relative to a spherical particle of the same volume, by increasing the capture cross-section. This result suggests that using reporters of other shapes could be an effective and simple way to increase the analytical sensitivity of this important class of point-of-care diagnostics.

More broadly, our understanding of how reporter flux and size affect capture and binding at the pore scale may prove useful toward optimizing particle capture in a variety of microscale open porous media for applications in on-chip chromatography, virus purification, membrane chromatography, and microfluidic nanocomposite fabrication.

ASSOCIATED CONTENT

Supporting Information

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Figures including experimental schematic, imaging details, and additional results (PDF)

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