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Characterization of low adsorption filter membranes for electrophoresis and electrokinetic sample manipulations in microfluidic paper-based analytical devices

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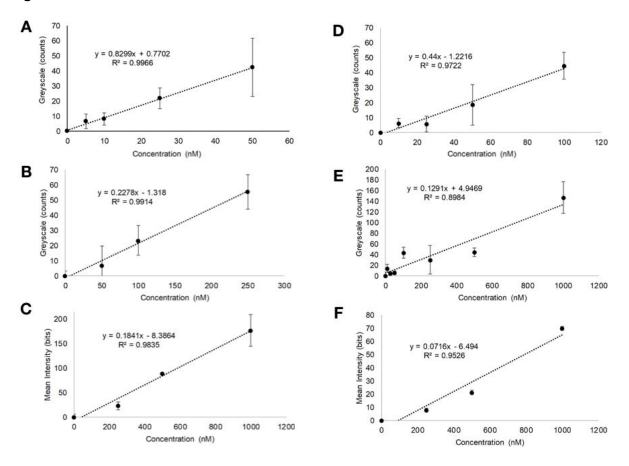
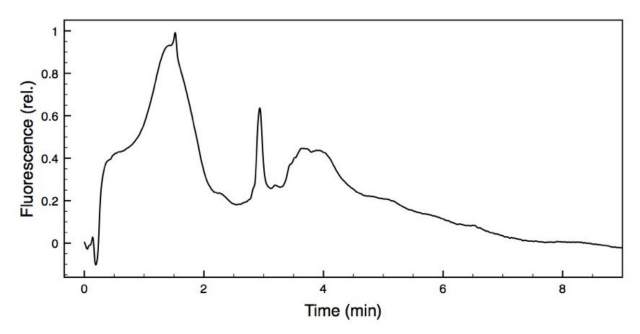
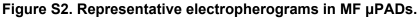


Figure S1. Fluorescence calibration plots.

Blank-subtracted fluorescence calibration plots utilized for detection limit calculations. **A.** Fluorescein on OE66 **B.** Fluorescein on PVDF **C.** Fluorescein on MF **D.** Nile blue on OE66 **E.** Nile blue on PVDF **F.** Nile blue on MF







Electropherograms in both TRIS and borate BGEs were irreproducible and complex, but the representative trace here demonstrates their common appearance: a complex peak profile migrating entirely before 4 minutes. We hypothesize this results from relatively high rates of electroosmotic flow which prevents electrophoretic resolution on these time scales. Further device optimization may improve separation performance and reproducibility.