Fabrication of a microfluidic flow-through immunoassay for simultaneous detection of multiple proteins Nicole Y. Morgan, Alyssa C. Henry, T.M. Phillips, T.J. Pohida, P.D. Smith (NIH); M. Gaitan, L. Locascio (NIST)



s indicate direction of flow for two stages

Tracking fluid flow

Fluid flow in the prototype device is monitored using

sulforhodamine B in 20 mM phosphate buffer, pH 7.4

Abstract

We have developed a chip-based microfluidic device for multianalyte immunoaffinity capture and detection of proteins. The immediate motivation is an epidemiological study of the immune response to the Human Papilloma Virus (HPV), for which the simultaneous isolation and detection of multiple proteins from a large number of microliter samples of cervical secretions is required. Using the microfabrication facilities at NIST, we are able to make micrometer-scale glass-encapsulated microfluidic systems with any desired two-dimensional configuration. The prototype devices consist of a long glass-encapsulated channel, 50µm x 15µm x 30cm, with a serpentine pattern. Side ports are used for electroosmotic loading of different biotinylated antibodies into each segment of the channel. These antibodies hind to streptavidin that has been covalently bonded to the channel walls via an imine linkage. The robust attachment of the antibodies allows them to be used for multiple sample runs; after each run, the antibody-antigen interaction can be disrupted by an acidic buffer gradient, leaving the tethered antibodies ready for reuse. After the antibodies have been immobilized, the sample under analysis flows through the entir device. Electrical control of the sample flow permits adjustment of the residence time in each segment in order to optimize binding. The channel device architecture has several advantages over existing array technology: the proteins are detected by single-point capture, and much smaller sample volumes can be used

Overview and objectives

Why microfluidics?

Smaller - System volumes less than a microliter are possible allowing detection of smaller amounts of analyte. Faster - Diffusion-mediated processes happen more quickly in

- a smaller system. Multiplexing measurements is more easily accomplished.
- Cheaper With smaller systems, use of expensive reagents (e.g., capture ligands) is greatly reduced.

Develop microfluidic expertise in-house

Focus on specific applications that require extensive ation or integration into a measurement system

- Initial goal develop microfluidic device for immunoaffinity analysis of small samples
- Simultaneous detection of multiple analytes using specific binding interactions With a system volume of 1uL, detection of 10pg/mL concentration

→ sub-picogram detection capability Integrate with optical detection instrumentation

Device operation - schematic



dissociating agent (e.g., acidic buffer)

Interagency collaboration

This area is ideally suited for a collaboration between researchers at NIST and NIH. Microfluidics is an active area of research at NIST Gaithersburg. Device fabrication typically employs techniques and equipment first developed for the semiconductor industry. These facilities, and the technical expertise to operate them, already exist just up the road from NIH.



Top: wet etch

hench and resist spinners w cleanroom facility at Bottom: aligner for NIST, being furbished now photolit

Initial strategy

- Initial devices are made from silicon/glass - system and fabrication techniques are well-characterized chemistry for biomedical applications better understood. easier to incorporate circuit elements in future applications
- Control flow with both pressure and voltage pressure-driven flow (vacuum) for initial chemistry electrokinetic flow of sample through device permits better control, less dilution of sample plug during measurement
- Attach antibodies with avidin-biotin chemistry: - antibodies strongly linked to surface so devices can be reused well-established and widely used chemistry
- Detect bound molecules with LIF inside channels - laser-induced fluorescence of tagged proteins after singlenoint canture
 - -detecting proteins bound in channels gives the most signal, but potentially higher background.

Prototype device



Fluid access to ports is through silicon wafer Strips of PDMS confine aqueous buffers to individal ports





Forming the channels: Oxide laver serves as a mask for silicon etch Anisotropic etch: TMAH etches preferentially perpendicular to the <100> Si planes, leading to a trapezoidal cross-sectio Typical channel dimens 50 um wide, 15 um deep, 30 cm long,



Coating channel with uniform glass layer After etching, grow a thick layer of thermal silicon dioxide. Chemistry for functionalizing SiO₃ surfaces is well-established Anodic bonding to glass wafers has been successful with oxide lavers as thick as 650 nm.

Estimated breakdown voltage ~ 500 V dry. Measured breakdown voltage > 400V (substrate positive) when channel is filled with aqueous buffer (e.g., 20mM nhoenhate)



Sealing the channel: Permanently attached with anodic bonding: sandwich heated to 400 C, voltage of 1200 V applied for 30 minutes. Glass used: optically flat Corning 7740 borosilicate glass (Pyrex)





Releasing captured antigens interactions between the antigen and the antibody can be disrupted by ing with a mildly acidic buffer. The surface-bound antibodies can then be used again for the next sample Fluorescence images of a silicon/glass microchannel with immobilized antibody through two cycles of exposure and rele of the complement

Reusing the capture antibodies

Reuse of capture antibodies Silicon microchannels terminated in anti-human IgG were exposed to human IgG, rinsed with acidic buffer, exposed again to human IgG, and rinsed again. The approximate protei concentration used was 1 nM

Towards quantitative detection

Controlling fluid flow

The flow of the sample through the serpentine channel

nattern is achieved with electroosmotic flow:

Measured fluorescence varies

Silicon microchannels terminated in

anti-human IgG were exposed to increasing concentrations of humar

Current limit of detection ~ 10 pM

ed charge on the interior

walls of the microchannels

buffer to the sidewalls.

attract counterions from the

When voltage is applied along the channel, the electric field pushes these counterions. The moving ions drag nearby solvent molecules along move, causing bulk flow in the microhomoal

the microchannel

By changing the voltages applied to the different fluid reservoirs at different times, a small plug of

fluid can be injected into the device.

(1.3 ng/ml) using a microscope with halogen lamp and CCD camera.

with analyte concentration:

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of sample injection

Fluorescent plug shows some signs of diffusive broadening as well as photobleaching, as it moves through the device.

After injection, the fluorescent plug can be driven through the serpentine channel. Monitoring the flow of a biological sample without photobleaching the fluorophore-tagged proteins, should be possible using a longer-wavelength tracer. Flow through the 30-cm device takes approximately 30 minutes

Current status

Device fabrication

Made prototype devices with glass-encapsulated channels and lithographic back-side ports Arrays of straight channels permit rapid parallel tests of attachment chemistry.

Flow control:

Have shown independent control of flow in different channels. Have demonstrated both pressure-driven and electro-osmotic flow in these channels. Electrical control of flow permits detailed control of injected

- sample plug. EOF mobilities: bare Si0₂ μ_{EOF} = 3.5x10⁻⁴ cm²/V-s. aminated channel μ_{EOF} = -1.7x10⁻⁴ cm²/V-s.

Attachment chemistry:

Robust tethering of antibodies to microchannel surfaces Molecular recognition of complementary antigens with minimal non-specific interactions Demonstrated reuse of immobilized capture antibodies

Future directions

Short-term goals:

Quantitative characterization of functionalized surfaces Further optimize quantitation of analyte capture Begin reliability/reusability testing. Integration into whole-chip detection system

Long-term possibilities:

Incorporate additional on-chip functionality, such as detection hardware or labeling chemistry Explore the use of plastics for biomedical applications of Other clinical and research applications

Connecting to the channels Lithographic patterning of the through-holes allows smallvolume ports with good control of both geometry and surface quality Requires the canability to align the back and front side patterns with better than 1µm precision Reflected light Transmitted light Pictures are glass-side up. Estimated port volume: < 6 pL in anti-human IgG were exposed to mouse IgG, rinsed, and then exposed to human IgG. Protein For pressure-driven flow, use Nanoports (Upchurch) or vacuum For electrokinetic flow, build PDMS reservoirs on top of silicon for concentration was approximately 1 nM. negative or atmospheric pressure).





fluid flow