Static Mode Microfluidic Cantilevers for Detection of Waterborne Pathogens

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HIGHLIGHTS
- Low-cost manufacturing of microfluidic cantilevers
- Detection of waterborne parasites in a cantilever format
- New manufacturing method utilising laser cutting and polyimide for cantilevers

ABSTRACT
This paper reports on the first demonstration of polymeric microfluidic cantilever sensors. Microcantilever sensors, magnetic beads, and microfluidic technology has been combined to create a polymer based biosensor. Using cheap materials like polyimide, a simple fabrication method has been developed to produce cantilevers with an embedded microfluidic channel. The advantage of this approach is that the addition of a microfluidic channel enables the analysis of smaller volumes and increases the capture efficiency in applications detecting rare analytes. As a proof of principle the system has been applied for the detection of the waterborne protozoan parasite Cryptosporidium, achieving sensitivity comparable to QCM, whereas a previous set-up without the microfluidic channel was unable to detect the parasite.

Keywords: microfluidics, cantilever, Cryptosporidium, detection

Introduction
Cantilever biosensors have demonstrated impressive sensitivity for the detection of nucleic acids, proteins and cells [1-4]. However, in solution, when operated in the resonance mode, viscous damping severely degrades the resolution [5]. Alternatively, cantilevers can be operated in static mode, with surface stress determining the degree of cantilever bending. While this eliminates the problem of viscous damping for measurements in liquid, the challenge then becomes effective delivery of the sample to the cantilever surface. This challenge is especially important in applications where relatively large analyte sample volumes are necessary, e.g. environmental monitoring [6]. In order to address this, immobilisation strategies can be optimised to attempt to maximise capture efficiency of the sensor or external forces can be utilised to enhance delivery [7].
Previously, cantilevers have been embedded within microfluidic systems [8, 9]; and more recently, smaller-scale microfluidics which fits onto the cantilever surface itself is demonstrated. For example, the Manalis group have developed microfluidics upon cantilevers, manufactured from silicon and employed in the resonance mode. This highly successful strategy has lead to the weighing of single cells in fluid [5]. Very few other microfluidic cantilever systems have been reported [10]. However, the materials and fabrication approaches are expensive. Additionally, while the latter work provides an interesting method of weighing individual microorganisms, specificity in pathogen detection is not offered.

*Cryptosporidium* is a protozoan pathogen, which is highly problematic for the water industry due to a low infectious dose [11] and high degree of robustness which enables long survival times in water along with resistance to standard disinfection by chlorination [12]. Several biosensor technologies have been applied to the detection of *Cryptosporidium* as reported in a recent review article [13]. Both quartz crystal microbalance (QCM) [14] and piezoelectric macrocantilever (PEMC) [15] approaches utilised relatively large flow cells and delivery of the sample to the sensor surface was not characterised.

Here we present the low-cost manufacture of polymeric microfluidic cantilevers and demonstrate the effectiveness of this set-up in improving transport to the sensor in both the detection of pathogens and DNA. The approach reported here has the advantage of ensuring effective sample delivery to the surface of the sensor, enabling high capture efficiency, which is useful in the situation of detecting rare pathogens. Miniaturisation of sample delivery in this way limits the throughput of devices, although there is potential to negate this problem through parallelisation or effective sample pre-processing. Previous unpublished work by the authors using microcantilevers without microfluidic channels presented low sensitivity to *Cryptosporidium* oocysts whereas use of the microfluidic channel has enabled a detection limit of $1 \times 10^5$ oocysts/mL. However, the main advantage of the system presented here over previous microfluidic cantilever set-ups is that since the device is made entirely of polyimide it is both cheaper and easier to manufacture.
2. Materials and Methods

2.1 Cantilever manufacture

The sensor was precision fabricated using a photolithography method. Firstly, a sheet of polyimide (7.6 micron thick, 3 inch x 50 inch, VHGLABS Kapton® (Polymide)) was sputter-coated with an adhesive layer of chrome (5 nm) followed by a layer of gold (20 nm) using gold evaporation system (BOC Edwards Auto 500). Secondly, this gold-coated polyimide was attached to a sheet of 20 μm thick positive photoresist (photopolymer dry film resist, ORDYL), and the two sheets were bonded together using pressure applied at 95°C. Thirdly, a mask (fabricated by microlithography) was employed to control the UV exposure (exposure time of 30 seconds) creating patterns of microchannels. Fourthly, the UV exposed sheet was developed (Developer conc. for 4615 dry film Mega Electronics Ltd) for 20 seconds removing the positive photoresist in the exposed areas. These areas define the microfluidic channels. Fifthly, the microchannels were sealed using 25 μm polyimide tape as a top layer. This process is summarised in Figure 1A. Finally, a short pulsed (65 ns) laser of wavelength 532 nm was used to cut the structures into individual microcantilever microfluidic chips, with cantilever dimensions of 1.5 mm in length and 300 μm in width. Each cantilever contained one U shaped microfluidic channel with channel sizes of 60 μm in width, 20 μm in height and total of 3 mm in length (Figure 1B).

2.2 Cantilever Set-Up and Operation

The cantilever set-up developed in this paper includes a rotary valve, microcantilever chip with a microchannel fabricated on top that is connected with tubing to a gravity fed pumping system (1) via the rotary valve (2), laser diode (7), position-sensitive detector (PSD) (8), a magnet (http://uk.rs-online.com/web/p/magnets/6950184/) and microscope with a digital CCD camera (9) (Figure 1C; numbers relate to the labels in Figure 1C). The magnet is located immediately below the cantilever and has a pull force of 3.3kg. The cantilever system is set up on an optical table (4) (Newport Laminar Flow isolator) to reduce vibrations. The system is mounted in a non-transparent box (3) made of PMMA (5mm thickness), with thermal insulated materials (10mm thickness), which reduces the external disturbance from air flow, background light, and temperature variations in the lab[±0.5 degree]. The rotary
valve switch device is computer-controlled via RS-232 and is used to switch between the flow different liquids into the microchannel on the cantilever surface. With gravity pumping of the liquid (1 mL/h), spikes in the results curve can be significantly reduced. The optical resolution of the microscope is 5 μm, which is used to confirm that the laser beam is on the tip of the cantilever. The laser beam, at an angle of 45, reflected by the cantilever is aligned on to a position-sensitive detector (PSD), at a distance of 5cm from the cantilever, and an amplifier is used to amplify the current signal from the PSD and convert into voltage signals. A National Instrument data acquisition card is then used to record data in LabView.

2.3 Detection of Cryptosporidium
Reagents: Viable C. parvum oocysts were purchased from Creative Science Company, Moredun Research Institute. Magnetic beads and goat polyclonal antibody immunoglobulin G (IgG) specific to C. parvum were purchased from Waterborne Inc. Phosphate-buffered saline (PBS) was obtained from Sigma-Aldrich.
Functionalization of cantilever microfluidic biosensor with protein G, antibody IgG and immobilization with C. parvum solution: The sensor was functionalized with protein G solution (20 mg/mL) for 2 hours, IgG solution (20 μg/mL) for another 2 hours [16] and finally exposed to C. parvum solution (between 1x10^5 oocysts/mL and 1x10^7 oocysts/mL in DI water) for 10 mins causing the oocysts’ immobilization on the surface of the sensor. After each step was complete, the sensor was rinsed with PBS solution (10mM, pH 7.4). After immobilization of oocysts, the biosensor was left to stabilize and afterwards it was incubated with magnetic beads solution (Crypto-Grab, Waterborne Inc, 2.5 mg/mL) for 20 minutes. Finally the sensor was rinsed with PBS solution. Every rinsing was performed in order to remove the unbound reagents. The protocol was performed in room temperature. The flow rate for all steps was 1 mL/hr.

3. Results and Discussion
3.1 Cantilever manufacture
Microfluidic channels embedded in silicon cantilevers have previously been manufactured using dry etching. In order to utilise low-cost polyimide materials an alternative fabrication method was required for the production of microfluidic channels. A method using simple lithographic techniques was employed, as described in detail in the materials and methods, and illustrated in Figure 1B.
3.2 Cantilever Characterisation

Following production of the cantilevers, the system was characterised using optical microscopy. Figure 2 shows an optical microscope image of polyimide fabricated cantilevers with embedded microchannels. The width of the cantilever was designed to be 300 μm and the channel is 60 μm wide. Images from several cantilevers were taken, and an average of 5 measurements revealed the channel width was 60 μm ± 3 μm, illustrating that the variability in fabrication was small and that this is therefore a reproducible method. The images illustrate that cantilevers of different lengths can be manufactured using this protocol, though for all subsequent experiments cantilevers of length 1.5 mm were employed.

In the cantilever set-up illustrated in Figure 1C cantilever performance was tested. Flow through the microfluidic channel had no influence upon deflection with the cantilever remaining stable. Various flow rates were trialled and an upper of limit of 1mL/hr was determined. This was limited primarily by the choice to operate using gravity driven flow. While the bonding technique could tolerate higher pressures, and therefore flow rates, pumping of fluids through the channel was observed to result in spikes in the cantilever read-out.

The final performance characterisation involved system calibration with magnetic beads (Figures 1D and 4A). Figure 3A illustrates the schematic of detection employed for the waterborne parasite under investigation. Although detection of whole cells has been demonstrated in cantilever systems, this has typically been in resonance frequency set ups and without the use of specific recognition elements [5][17]; it remains a challenging task in mass-sensitive systems as coupling of the binding event to the system deflection is critical and this is often weak for larger analytes like cells. This is particularly important in the type of assay reported here where analyte binding is a key step. Additionally, factors such as surface stress also contribute to the observed signal. Therefore, the use of magnetic beads was selected to amplify the signal. Figure 1D illustrates the operation and set-up with this detection principle with a magnet located beneath the cantilever holder. To determine that the magnet strength and magnetic bead concentration were appropriate a series of experiments flowing different concentrations of magnetic beads through the system were performed. As seen in Figure 4A, quantitative results were obtained with a series of dilutions indicating that the cantilever read-out was proportional to the magnetic
bead concentration within the channel, thus confirming this approach was suitable for quantitative pathogen detection.

3.3. Pathogen Detection

The microfluidic cantilever system was applied to the detection of the waterborne protozoan pathogen, *Cryptosporidium*. Detection of this pathogen is challenging since it is often present at low concentrations. However, since ingestion of only a few oocysts is sufficient to cause disease it is important to maximise capture efficiency of oocysts within any biosensor system.

Our initial work (unpublished) exploring the potential of cantilever sensors to detect this pathogen were unpromising with the parasite going undetected even at high concentrations. The most likely explanation for this was the sample size and time required for delivery of the pathogen to the surface. Since, an identical set-up was employed during cantilever functionalisation, limitations in delivery of one of the immobilisation reagents and/or the antibody to the surface might also have contributed to the poor detection.

The time allowed for oocyst exposure to the surface was 10 mins. In the set-up without a flow system using 1mL of solution the time was insufficient to result in a high capture efficiency on the cantilever surface. The time taken, t, for a particle to diffuse a distance, d, is given by:

\[ d \sim \sqrt{Dt} \quad \text{Equation 1} \]

where D is the diffusion coefficient (5x10^{-10} cm²/s for oocysts) [6], [18]. This would suggest that oocysts diffuse around 0.002 mm in ten minutes.

However, consideration of diffusion may not be appropriate for oocysts as it has been reported that for micron-sized particles [24], hydrodynamic and gravitational forces are often significant compared to Brownian forces [19]. In the static case, hydrodynamic forces are not relevant and the gravitational force can be determined using the particle free settling velocity, \( U_s \). This is given by:

\[ U_s = \frac{2\Delta \rho g \alpha^2}{9 \mu} \quad \text{Equation 2} \]

where \( \Delta \rho \) (kg/m³) is the particle density (1045.4) minus the density of water (997), g is the acceleration of gravity (9.81 m/s²), \( \alpha \) is the particle radius (2.5 μm for C.
parvum) and \( \mu \) is the water viscosity \((8.91 \times 10^{-4} \text{ kg/ms})\), and is \(0.74 \text{ \(\mu\)m/s}\) for C. parvum. Our calculated figure compares to the slightly lower values of \(0.35\) and \(0.5 \text{ \(\mu\)m/s}\) reported in the literature. Although oocyst travel by sedimentation is around an order of magnitude greater than that of diffusion, and additionally is focused in the direction of the substrate, this is still unlikely to enable efficient delivery of oocysts to the cantilever surface within ten minutes, since using an average of the above values of \(0.53 \text{ \(\mu\)m/s}\), allows for a distance of only \(0.31 \text{ mm}\) to be covered. If a test volume of \(0.1 \text{ mL}\) was utilised it would take days (assuming the volume was solely located on top of the cantilever). However, the non-flow set-up also has the disadvantage that in the flow cell set-up, which is wider, longer and deeper than the cantilever, many oocysts will initially be distributed under or to the sides of the cantilever and therefore be unable to reach the binding surface, especially allowing for sedimentation. Oocysts could not be detected even after \(1 \text{ hr}\).

Within the microfluidic cantilever set-up, both diffusion and settling are still valid methods of oocyst transport to the surface within the channel laminar flow environment. Given the volume of the channel \((0.0036 \text{ \(\mu\)L})\) and the flow rate \((1 \text{ mL/hr})\) it is clear that the transit time within the channel is much less than \(1 \text{s}\). With a channel height of \(20 \text{ \(\mu\)m}\) the maximal distance (in the z direction) to be travelled by an oocyst within this time is \(10 \text{ \(\mu\)m}\) (allowing for the size of the oocyst). It must however be remembered that there is an even distribution of oocysts across the channel height and many will need to travel significantly less than this distance to reach the binding surface. While it is clear that not all oocysts will reach the surface even in the microfluidic cantilever set-up the chances are greatly improved. Increasing the number of encounters with the immobilised antibodies increases the likelihood of a binding event occurring and will therefore increase the capture efficiency of the system.

With the microfluidic cantilever system a series of different Cryptosporidium concentrations \((10^5 \text{ to } 10^7 \text{ oocysts/mL})\) were investigated, with each concentration repeated five times. Following capture of the oocysts, the system was flushed with magnetic beads to amplify the signal. Figure 3B shows representative traces of the experiments, from the oocyst addition stage until the final detection point at which the unbound magnetic beads are removed from the system. One trace for each
concentration is shown along with a reference sample where no Cryptosporidium was added. As the magnetic beads flow through the system little difference is observed between the different samples. However, upon rinsing of the magnetic beads from the system the reference sample returns to zero, whereas for the oocyst samples magnetic beads remain bound to oocysts within the system and can be utilised to determine the Cryptosporidium concentration in the sample. In short, Figure 3B illustrates that quantitative detection of oocysts can occur within the range $10^5$ to $10^7$ oocysts/mL.

The results of all five experiments have been averaged and are presented in Figure 4. The results indicate a linear relationship ($R^2 = 0.96$) confirming detection in the range $10^5$ to $10^7$ oocysts/mL. The upper limit of $10^7$ oocysts/mL was the highest concentration tested in this set-up and could potentially be extended. This is limited by the space for oocyst binding within the microchannel. Interestingly, a calculation of the maximum coverage of the microchannel area revealed that it would be saturated with $\sim 1\times 10^6$ oocysts, using an oocyst diameter of $5\mu$m, a channel area of $18$ mm$^2$ (assuming oocysts only bind to the immobilised antibody and not to other channel surfaces) and assuming a maximum close-packing of 74%. This calculation reveals that although the use of the microchannel improves the capture efficiency, the system still misses some oocysts. By decreasing the flow rate more time would be available for oocysts to bind within the channel, thus increasing the sensitivity. There is thus a trade-off between reaching highly sensitive detection limits and achieving a reasonable throughput/detection time, which is a recurring challenge for biosensor system for waterborne pathogens.

For practical applications, achieving a low limit of detection is the critical parameter. Lower concentrations were found not to yield a measurable response. While the sensitivity of the approach is comparable to the $1\times 10^5$ oocysts/mL detection limit reported for QCM-D detection of this parasite (Poitras 2009), lower concentrations have been determined, by Mutharasan and colleagues (Campbell 2008), with a macrocantilever set-up. However, this operates with a recirculating flow system, which could potentially also increase the capture efficiency of the microcantilever sensor. Additionally, sensitivity could be improved by increasing the magnetic bead concentration or utilising a more powerful magnet.
Conclusions

The results in this paper represent the first example of a microfluidic microcantilever sensor fabricated in polyimide. Using polymer materials to manufacture the system is an advance over previous work, allowing for cheap and easy fabrication, resulting in cheap sensors which can be rapidly produced. A further advantage of this approach relates to the effective sample delivery enabled by confining the sample to a narrow layer above the cantilever surface. Transport of the analyte of interest to the capture region is often the time-limiting step and this design offers a mechanism of effective surface delivery. This is likely to prove advantageous for applications detecting rare analytes as well as in applications where very small samples are to be processed. For larger samples throughput within the microfluidic channels is potential challenge though parallelisation is an option to overcome this possible limitation. Future work could incorporate cantilever sensors on the ends of optical fibres moving towards a miniaturised portable system [20].

Furthermore, this paper has applied the system for the detection of the problematic waterborne protozoan parasite Cryptosporidium, demonstrating sensitivities comparable to existing literature reports and particularly showing greater sensitivity than QCM. Future work will concentrate on the optimisation of the system as well as developments in the immobilisation chemistry and the sample pre-processing to deliver even lower limit of detection, suitable for real-world application of this technology to waterborne pathogen detection.

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References


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Figure 1. Scheme of integrated microfluidic microcantilever sensor. A) Schematic of the fabrication process. B) Layout of the microfluidic channel on the cantilever. C) Cantilever set-up. The numbers refer to: 1) the gravity-fed sample input system; 2) the rotary valve for switching between solutions; 3) insulating box; 4) optical table; 5) cantilever system; 6) sample waste outlet; 7) laser diode; 8) microscope with digital CCD camera; 9) position sensitive diode (PSD). D) Operation of the cantilever sensor with the magnet.
Figure 2. Cantilever characterisation. Optical microscope images of fabricated microchannels on microcantilevers. A) A fabricated array of microfluidic cantilevers all cut from one piece of patterned polyimide. Different channel lengths are demonstrated here to show the fabrication can be easily controlled to produce cantilevers of different lengths. The dimensions illustrated here show that the cantilever is 315 µm wide. B) A close up image of a single cantilever channel of the dimensions utilised in this article. The microchannel structure can be clearly observed. The red scale bar shown in the image is 300 µm.
Figure 3. Cantilever detection of waterborne pathogens. A) Schematic illustrating the functionalisation of the cantilever to detect Cryptosporidium oocysts and the addition of magnetic beads which enables enhancement of the detection signal. B) Detection of oocysts at a range of different concentrations ranging from a control sample of zero to a set of concentrations from $1 \times 10^5$ to $1 \times 10^7$ oocysts. Initially the oocysts solution is passed through the cantilever microchannel and although binding takes place this is insufficient to trigger cantilever bending. After the introduction of the sample a brief rinsing step with PBS is applied. Subsequently, magnetic beads are passed through the channel (at this stage where the beads are incubated in the channel little difference is observed between different oocyst concentrations) and finally the channel is rinsed with buffer removing any unbound beads. In the final stage of the results curve, the measurement of deflection indicates the amount of bound microbeads, and therefore also the concentration of oocysts within the cantilever channel, and it is clear that the biosensor can distinguish between different concentrations of pathogen.
Figure 4. Plot of deflection (nm) against magnetic bead concentration (left graph) and oocyst number (right graph) showing a linear trend in cantilever response against magnetic bead concentration (confirming that the magnetic enhancement of detection is quantitative) and oocyst exposure.