Impedance Microbiology-on-a-Chip: Microfluidic Bioprocessor for Rapid Detection of Bacterial Metabolism

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Abstract—Detection of a few live bacterial cells in many industrial or clinical samples is a very important technological problem. We have developed a microscale technique for concentrating bacterial cells from a dilute sample, by factors on the order of $10^3$ to $10^7$, and detecting their metabolic activity by purely electrical means. The technique was implemented on a silicon-based microfluidic chip where the cells are concentrated and incubated in a chamber with a volume of 400 pl. Concentration and capture are obtained by the use of dielectrophoresis on the bacterial cells, and metabolism detection is achieved by means of impedance measurements of the medium in which the bacteria are incubated. Performing impedance-based detection at the microscale results in drastically reduced detection times for dilute bacterial samples, thanks to the ability to efficiently concentrate and capture the cells in an extremely small volume. Such concentration eliminates the need to amplify the bacterial population by long culture steps. This detection technique can be used for a wide variety of applications.

Index Terms—impedance microbiology, biochip, bacterial metabolism, bacterial detection

I. INTRODUCTION

DETECTION of a few live bacterial cells in clinical samples, water, pharmaceuticals, food, or cosmetics (and the raw materials used to produce these) is a very important technological problem. Food, cosmetic, and drug manufacturers must screen their raw materials and/or finished products for total bacterial content before being used for manufacturing, or shipped to customers, respectively. In the industries mentioned above, materials that are likely to contain pathogens or other undesired microorganisms are often subjected to various sterilization procedures. In these cases the purpose of screening is to ensure that no live bacteria remain after sterilization, so the detection method should only be sensitive to live cells. Currently, the official food screening procedures established by regulatory agencies (Food and Drug Administration, FDA, and US Department of Agriculture, USDA) are very sensitive, but require a long time to complete (up to seven days) with the work of highly skilled laboratory personnel [1], [2]. By the time a pathogen is identified in a product or its raw materials, the product would probably have been sold and used. The same is true for raw materials used for production of pharmaceuticals, where the current standard for purified water for final products specifies a maximum bacterial content of 100 cfu/ml (colony-forming-units per ml) [3]. The most common sample volume used by the pharmaceutical industry is 100 ml, which means that the screening protocol should be able to detect 10000 cfu total. This is, in fact, the target cell count that we have chosen for testing our detection technique. Few automated methods exist with sensitivity and reliability comparable to those of the manual methods; and the most sensitive automated techniques cannot discern between live and dead cells. To identify live bacteria with high sensitivity the current detection methods rely on a 24 to 48 hour growth step to amplify the population of live cells.

One common automated bacterial detection method is based on the changes in the electrical characteristics of a medium where bacteria are cultured. These changes are produced by the release of ionic metabolites from live cells, measured by electrodes in contact with the medium [4]. The most common embodiment of this technique, commonly called “Impedance Microbiology,” involves monitoring over time the ac impedance of a pair of electrodes immersed in the culture medium, at a single frequency. If the impedance changes beyond a certain threshold, a positive detection is indicated. This technique was first identified more than 100 years ago, and has been used for many years in bacterial growth monitoring, bacterial load control on perishable products, and bacterial detection. There are numerous reports in the literature dealing with its application to rapid bacterial detection [5]–[7]. And equipment for automated impedance monitoring ranging from laboratory, bench-top size up to industrial scale is commercially available from several vendors.

Unfortunately, the detection time of the conventional impedance-based method can be quite long when the concentration of bacterial cells present in the sample is very small [4]. The lower the initial concentration of microorganisms, the longer it takes for the impedance to change by a measurable amount. This limitation can be overcome if those few cells are confined into a very small volume while the impedance is being measured. In this way the effective cell concentration is increased without increasing the number of cells. As we proposed in a previous publication [8], this concept leads to the idea of implementing the impedance-based detection in a microfabricated device, where volumes on the order of nanoliters or picoliters can be easily created. Such application of microfabrication technologies seems especially fitting, since miniaturization of the impedance-based assay has the potential

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to directly reduce the time for detection by a significant factor. A microfabricated device capable of concentrating cells from a dilute sample completely eliminates the need for amplifying the bacterial population by growth. This new technique of “Impedance Microbiology-on-a-Chip” can be used not only in the front end of a system for determining the presence of live microorganisms in a sample, but for many applications in which a bacterial culture needs to be monitored.

In previous publications [8], [9] we described preliminary experiments designed to test the feasibility of the proposed detection method. In this report we present results of tests of the microscale impedance-based technique for real-time detection of bacterial metabolic activity and growth using two microfabricated devices with two types of bacteria: Listeria innocua, a non-pathogenic species of the genus Listeria, and Listeria monocytogenes, a pathogenic species of the same genus. L. innocua and L. monocytogenes are very closely related and are often found together in certain food products [10]. However, the technique presented here is not exclusive to Listeria cells and can be used with any other culturable microorganism, including most yeasts and molds.

II. MICROFLUIDIC DEVICES

A. Device Design

Two silicon-based microfluidic biochips were fabricated for testing the proposed detection scheme. The design and fabrication of the first biochip was described in detail somewhere else [9]. This first biochip has rectangular chambers connected in series by channels, both etched to a depth of 12 µm onto a crystalline silicon substrate. The chambers have volumes between approximately 75 pl and 5.3 nl. Interdigitated platinum electrodes were created at the bottom of the chambers to measure the metabolism of cells injected into the chambers.

The second biochip or bioprocessor was designed to collect and concentrate the cells from a liquid sample, and detect their metabolism and growth as they are incubated inside the chip. The overall design concept is based on a large straight channel (main channel) through which the sample can be flowed at the desired rate while the cells contained in it are deviated by dielectrophoretic (DEP) forces [11] into a small channel that leads into the incubation/measurement chamber, which has a volume of 400 pl. In the incubation chamber the cells are retained and concentrated by DEP, and their metabolism is measured by platinum interdigitated electrodes in direct contact with the liquid in the chamber. In both the main channel and the incubation chamber, the DEP field is generated by interdigitated electrodes buried under a silicon dioxide layer and excited with separate ac voltage sources. A schematic diagram of the deviation and capture process is shown in Fig. 1. The main channel has a cross sectional area large enough to accommodate a maximum flow rate of approximately 2 µl/min while keeping the peak fluid velocity low enough to guarantee that the DEP forces are adequate for deviating all of the cells into the incubation chamber. The DEP excitation voltage is limited to 20 Vpp or less, the cells are deviated at a very shallow angle with respect to the long axis of the channel. The flow rate in the deviation channel is externally controlled between 0 and approximately 10 nl/min, independently of the flow rate in the main channel. To accurately measure the temperature in the chip, a platinum resistive-temperature-detector (RTD) is built into it, made of a meandering platinum resistor. Each biochip contains two identical concentration and detection devices, so that two tests can be performed in parallel, or one of the devices can be used for control or reference measurements while the other receives the sample to be tested.

Inlet

Fluid only

Fluid and cells

DEP deviation electrodes

Outlet

Main

Outlet

Deviation

Fluid only

Fluid and cells

DEP capture electrodes

Fig. 1. Principle of operation of the dielectrophoresis-based deviation and capture of cells in the bioprocessor.

B. Bioprocessor Fabrication and Packaging

The fabrication process of the bioprocessor started with bare 4” silicon wafers, with a (100) surface and a thickness of 500 µm. Silicon dioxide was thermally grown on the wafers and subsequently patterned with conventional photolithography (using Clariant AZ1518 positive photoresist, Clariant Corp., Somerville, New Jersey) followed by etching in buffered hydrofluoric acid (BHF). This oxide layer serves as a hard mask for etching the channels in an anisotropic potassium-hydroxide-based etchant to a nominal depth of 12 µm. After etching the channels, the hard mask was removed by etching in BHF and the wafers were thermally re-oxidized to create a 2000 Å layer of silicon dioxide. Over this oxide a first metal layer, for the DEP electrodes, was deposited by sputtering of aluminum to a thickness of 1000 Å (MRC-903 Sputterer, Materials Research Corporation, Orangeburg, New York) and patterned by lift-off to create the DEP electrodes (conventional lithography using Clariant AZ4620 positive photoresist, which was used for the remainder of the fabrication process). After patterning the aluminum, a 3500 Å thick layer of silicon dioxide was deposited by plasma enhanced chemical vapor deposition (PECVD, STS-310PC, Surface Technology Systems plc, Newport, United Kingdom) to completely isolate the DEP electrodes and prevent electrolysis of the liquid in the channels. Windows were defined in the PECVD oxide layer using conventional lithography and opened by reactive-ion-etching (STS-340, Surface Technology Systems plc) to access the power dissipated into the chip by the DEP excitation signal is enough to raise its temperature beyond 100°C. To ensure a very high deviation efficiency at a DEP excitation voltage of 20 Vpp or less, the cells are deviated at a very shallow angle with respect to the long axis of the channel. The flow rate in the deviation channel is externally controlled between 0 and approximately 10 nl/min, independently of the flow rate in the main channel. To accurately measure the temperature in the chip, a platinum resistive-temperature-detector (RTD) is built into it, made of a meandering platinum resistor. Each biochip contains two identical concentration and detection devices, so that two tests can be performed in parallel, or one of the devices can be used for control or reference measurements while the other receives the sample to be tested.
the bond pads on the first metal layer. Subsequently, the second metal layer, which creates the measurement electrodes and the RTD temperature sensor, was deposited by sputtering 800 Å of platinum over a titanium adhesion layer (Perkin-Elmer Sputterer Model 2400, Perkin-Elmer Inc., Wellesley, Massachusetts) and patterned by lift-off. Subsequently, a 10 000 Å-thick layer of gold over a titanium adhesion layer were deposited by electron-beam evaporation (Varian Inc., Palo Alto, California) and patterned by wet etch to create robust bond-pads. After dicing the wafers, a glass cover was anodically bonded to each die at 400°C with a voltage of 1 000 V for 60 min. The glass cover was made from 4”, 500 μm thick, polished Pyrex™ glass wafers type 7740 (Corning Inc., Corning, New York), which were custom diced and ultrasonically drilled to create holes where input/output tubes were attached. The holes in the glass were aligned to the input/output channels in the die before anodic bonding. Fig. 2 shows a cross-section of the packaged device.

After fabrication, each die was fixed onto a custom-designed printed circuit board carrier that allows it to be easily connected to the equipment that measures the impedance, generates the DEP signals, and measures and controls the temperature. The carrier contains an integrated heater and generates the DEP signals, and measures and controls the connected to the equipment that measures the impedance, designed printed circuit board carrier that allows it to be easily connected to the equipment that measures the impedance, before anodic bonding. Fig. 2 shows a cross-section of the packaged device.

The impedance of the electrodes in the chip was measured with an Agilent 4284A LCR meter (Agilent Technologies Inc., Palo Alto, California) connected to the chip through an Agilent 34970A switching unit fitted with two Agilent 34905A RF multiplexer cards. The multiplexer card allowed the sequential measurement of up to four sets of electrodes with the same LCR meter. All of these instruments were connected to a computer through a GPIB interface. The impedance measurement process and the chip temperature control were automated by custom LabView® (National Instruments Corp., Austin, Texas) virtual instruments. With this setup, it was possible to acquire impedance vs. frequency curves (the complex impedance could be measured at multiple frequencies between 100 Hz and 1 MHz) from up to four pairs of electrodes at fixed time intervals for an indefinite length of time. In the first biochip the impedance of interdigitated electrodes in a 5.27 nl chamber was measured at 52 frequencies logarithmically spaced between 100 Hz and 1 MHz, with a 50 mVpp voltage excitation. In the bioprocessor the impedance of interdigitated electrodes in the incubation chamber was measured at 51 frequencies logarithmically spaced between 100 Hz and 100 kHz, with a 150 mVpp voltage excitation. Sinusoidal and square-wave DEP signals were generated by Agilent 33120A synthesized signal generators.

IV. MATERIALS AND METHODS

The following materials and methods were employed for all the tests performed with the first biochip and the bioprocessor.

A. Bacterial Growth Media

The composition of Luria-Bertani broth (LB), used for bacterial culture, was 10 g/L tryptone, 5 g/L yeast extract (both from Difco Laboratories, Detroit, Michigan), and 3.3 g/L dextrose in de-ionized water. The nominal conductivity of this medium was 2.2 mS/cm. Half-LB (HLB) was prepared by
mixing equal parts of LB and de-ionized water. The nominal conductivity of HLB was 1.1 mS/cm. *Listeria* cells grow in LB to a maximum population on the order of $5 \times 10^8$ cfu/ml after incubation at 37°C for 16 hours. *Listeria* cells grow in HLB to a final population not more than 30% lower than that achieved in LB (data not shown).

**B. Preparation of Bacterial Suspensions**

*L. monocytogenes* v7 and *L. innocua* F4248 cells were grown in LB at 37°C for at least 16 hours. After growth, the cells were washed by repeated centrifugation and resuspension in sterile HLB, LB, or DI water, depending on the experiment. Washing guaranteed that the cells were suspended in a medium completely free of any metabolic by-products. If heat-killed cells were desired, an aliquot from the as-grown cells was placed in a water bath at 80°C for 20 min and then washed in the same way as live cells. When fluorescent cells were needed a 1 ml aliquot from the as-grown live or heat-killed cells was mixed with about 1 µl of the green fluorescent dye DiOC$_6$(3) (3,3′-diethyloxacarboxyamine iodide, Molecular Probes Inc., Eugene, Oregon), diluted to 1 nM in dimethyl-sulphoxide (DMSO). The mixture was incubated at room temperature for 30 min to allow the cells to absorb the dye, and washed in the same way described above. The washed cells were diluted in sterile HLB, LB, or DI water, depending on the experiment, to the approximate concentration desired for injection into the chip. An aliquot from the diluted cells was diluted further and plated on Brain-Heart-Infusion (Difco Laboratories, Detroit, Michigan) agar plates. The plates were incubated at 37°C for 24 hours and the resulting colonies were counted to determine the actual concentration of viable cells in the suspension injected into the chip.

**C. Immunomagnetic-Based Cell Capture**

*L. monocytogenes* v7 cells were captured with anti-*listeria* Dynabeads™ (Dynal Biotech, Norway). *L. monocytogenes* were cultured in LB broth at 30°C for 24 hours, achieving a concentration of approximately $5 \times 10^8$ cfu/ml. Heat-killed cells were obtained by placing an aliquot from the as-grown culture in a water bath at 80°C for 20 min. A 1 ml aliquot from the culture (live or heat-killed) was centrifuged and resuspended in 1 ml phosphate-buffer-saline (PBS), then mixed with 40 µl of the stock bead suspension and incubated at room temperature for 10 min. The beads were collected with a magnet, most of the liquid was removed, and the beads were resuspended in 1 ml PBS-Tween (PBS with 0.05% v/v Tween-20) and incubated at room temperature for 10 min. This was repeated twice resuspending in 1 ml PBS-Tween, and twice again resuspending in 1 ml LB broth.

**D. On-Chip Cell Concentration and Metabolism Detection**

All the experiments were performed with the bioprocessor heated to (37 ± 0.1°C). The cells suspended in de-ionized water were injected at an input flow rate of approximately 1.7 µl/min, with the deviation and capture electrodes excited with a 16 Vpp square signal at 100 kHz. The broad spectrum of a square signal produced slightly higher DEP forces than a pure sinusoid at the same fundamental frequency. During injection, the flow rate in the incubation chamber was manually controlled to be between 4 and 10 nll/min. Flow rate control was performed manually by adjusting the pressures applied to the bioprocessor based on the observed velocity of the cells in the channels. After the whole 40 µl of sample had been flowed through the bioprocessor the flow was stopped, the DEP deviation electrodes were turned off, and HLB was injected at a flow rate of less than 0.5 µl/min. As the water in the chamber was replaced by HLB, the excitation voltage on the capture electrodes was increased to 20 Vpp and the frequency to 3 MHz to maximize the DEP forces acting on the cells. Once the incubation chamber was filled with HLB, the flow was stopped, all the microbore tubes leading into and out of the bioprocessor were pinched to seal them completely, and the capture electrodes were turned off. After pinching the tubes, the impedance measurement process was started and the cells were incubated for a minimum of 12 hours. When a reference measurement was desired, cells were injected only into one of the two devices in a bioprocessor, while the other device received sterile HLB only. The device with sterile media provided a baseline impedance in the absence of any metabolic activity.

**V. RESULTS**

**A. Preliminary Tests of On-Chip Bacterial Growth Detection**

The first biochip was used to perform preliminary tests of the impedance technique with live and heat-killed *Listeria* cells incubated inside the chip, suspended in Luria-Bertani broth (LB). Sterile LB and suspensions of different concentrations of *L. innocua* F4248 cells were injected into the chip and incubated there at 39°C (temperature of the heating platform where the chip was mounted, which was estimated to result in an on-chip temperature of ~37°C) for more than 12 hours. The cells were uniformly distributed throughout the whole microfluidic path in the chip. The circuit model shown in Fig. 4 was fitted to each one of the measured impedance vs. frequency curves (one curve per time point) to obtain a value for each of the model parameters at each time point. In the model, $C_{di}$ accounts for the dielectric properties of all the materials surrounding the electrodes, $R_l$ represents the bulk resistance of the liquid, $R_{we}$ is the resistance of the on-chip wiring, and $Z_1$ and $Z_2$ are interfacial impedances given by the following expression [8], [12], [13]:

$$Z_i = \frac{1}{(j\omega)^{m_i}B_i},$$

where $j = \sqrt{-1}$. This is the simplest model that would properly fit the measured data over the whole frequency range and at all times during incubation. It is speculated that the two different interfacial impedances account for the interactions of the electrodes with two distinct groups of species in the solution.

Fig. 5 shows the change over time in the admittance (reciprocal of the measured impedance) at 100 Hz (Fig. 5A), the conductance $G_i = 1/R_i$ (Fig. 5B), and the interfacial
parameter $B_1$ corresponding to the parameter $n_1$ with the smallest value (Fig. 5C), for three different concentrations of *L. innocua*: $2 \times 10^3$, $1.7 \times 10^6$, and $6.4 \times 10^6$ cfu/ml. These concentrations correspond to approximately 1 cfu, 9 cfu, and 34 cfu, respectively, in the 5.27 nl chamber where the impedance was measured. The sharp initial drop in admittance and $B$ parameter corresponding to $1.7 \times 10^6$ cfu/ml was due to a decrease in the temperature of the chip, from 40°C to 38°C, during the first ten hours of incubation. The conductance of all the samples was corrected for changes in temperature, to estimate its value at 39°C, using a temperature variation coefficient estimated from the small random variations in temperature observed on the same data set. It was not possible to find a suitable temperature variation coefficient to correct the values of the interfacial parameter $B$. With the exception of the temperature-induced drop, the shape of the admittance and conductance vs. time curves matches very well the typical shape of a bacterial growth curve. There is an initial lag phase where the cells are metabolizing but not multiplying; during this period the impedance changes very little because the bacterial population is small. The lag phase is followed by a growth phase where the cells multiply exponentially. Such fast-growing cell population releases large amounts of metabolic by-products that in turn produce large changes in impedance. When the nutrients in the medium are depleted and/or the concentration of metabolic by-products reaches a certain threshold, the cells stop multiplying and their population remains relatively constant or starts to decrease very slowly as cells die of starvation or due to the toxicity of the accumulated metabolic by-products. During this final phase the conductance changes linearly due to the linear accumulation of metabolic by-products released by a constant cell population.

The bi-modal increase in conductance observed in the $2 \times 10^5$ cfu/ml sample during the first eight hours of incubation cannot be easily explained except probably by ionic contamination coming into the chamber. But this unexplained change serves to illustrate a very interesting phenomenon. During the exponential growth phase, the $B$ parameter corresponding to the $n$ parameter with the smallest value displays a very large increase (by up to factor of ten in some cases), followed by a slow decay. The other $B$ parameter does not change significantly during the course of the incubation. In the case of the $2 \times 10^5$ cfu/ml conductance curve, the changes observed during the first eight hours of incubation are very unlikely caused by bacterial growth because the relevant $B$ parameter does not change appreciably during this time. For detection purposes, monitoring the changes in the interfacial $B$ parameters, along with the changes in conductance, can improve the robustness of the method against changes in conductance not related to bacterial growth. It is suspected that some species released by the cells during growth rapidly adsorbs onto the surface of the electrodes and thus changes the interfacial impedance significantly. Such species would slowly degrade and/or desorb from the electrodes causing the slow decay in the $B$ parameter.

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**Fig. 4.** Circuit model used to fit the impedance vs. frequency curves obtained from the biochip.

**Fig. 5.** (a) Relative admittance (reciprocal of the measured impedance) at 100 Hz of *L. innocua* suspensions at three different concentrations. (b) Relative conductance $G_l = 1/R_l$; temperature-corrected to 39°C. (c) Relative value of the interfacial parameter $B$ (see Eq. 1). The sharp initial decrease in the curves corresponding to $1.7 \times 10^6$ cfu/ml was due a sharp decrease in temperature during the initial part of the incubation. Values at $t = 0$ are defined as 100%.
after exponential growth stops.

The use of interfacial parameters for impedance-based detection of bacterial growth had previously been discussed in the literature [6], [7], [14], but always fitting very simple circuit models, composed of standard resistors and capacitors, to the impedance measured at a single frequency. Such approach makes the circuit model completely disconnected from the actual phenomena taking place, especially because an infinite number of circuit models would match the measured impedance at a single frequency. The model presented here is derived from a physical analysis of the simplest electrochemical processes taking place at the interface and is, thus, bound to better represent the phenomena being measured. In addition, fitting the model to impedance values at a large number of frequencies over a large range, makes the measurement more sensitive. Such fitting essentially “concentrates” small changes in the impedance at each one of the frequency points into larger changes in a small number of parameters that might have some physical relevance.

Live and heat-killed *L. monocytogenes* v7 cells from pure suspensions were used to test the metabolism detection process on bacteria captured by magnetic immunobeads. The captured cells plus beads were incubated on-chip in LB broth (biochip heater set to 39°C). The impedance was measured during incubation in the same way as in the previous experiments. The concentration of beads injected into the chip was $1.7 \times 10^7$ ml$^{-1}$. Preliminary tests of capture following the same procedure indicate that most of the beads carry at least one cell (data not shown), but the exact number of cells per bead is not known. Approximately 81 beads with live cells and 87 beads with heat-killed cells were manually counted in the 5.27 nl chamber where impedance was measured during the two experiments performed. Fig. 6 shows the admittance curves resulting from these cell incubations plus the incubation of sterile LB. The curve for live cells shows very clearly the three phases of cell growth, while dead cells and sterile media produce very small changes in admittance, as was expected from the absence of metabolism in the latter two samples.

![Fig. 6. Relative admittance of live and heat-killed *L. monocytogenes* cells captured by immunomagnetic beads. Values at $t = 0$ are defined as 100%.](image)

### B. On-Chip Bacterial Concentration and Growth Detection

To achieve a highly efficient concentration and capture process, the DEP forces in the deviation and capture stages of the bioprocessor have to be strong enough to counteract the drag forces exerted on the cells by the fluid around them. Testing of the concentration process in the bioprocessor was carried out with fluorescently-labelled *L. monocytogenes* cells suspended in LB, HLB, and de-ionized water. The DEP forces on *Listeria* cells suspended in HLB or LB are too weak to achieve an efficient concentration process at reasonable flow rates. With the cells suspended in LB or HLB, the injection time would be on the order of tens of hours. However, *Listeria* cells experience very strong DEP forces when suspended in de-ionized water with a conductivity of $1 \mu$S/cm, at an excitation frequency of 100 kHz. Under these conditions, the DEP forces are directed towards the regions where the electric field gradient is largest (positive DEP), so that the cells collect on the edges of the electrodes. Fig. 7a shows *L. monocytogenes* cells being deviated from the main channel towards the deviation channel. Fig. 7b shows cells concentrated in the incubation chamber in large accumulations over the gaps between the capture electrodes. With the right combination of medium conductivity, flow velocities, and DEP excitation voltage and frequency, the concentration efficiency is virtually 100%.

To test the complete process of cell concentration and metabolism detection, fluorescently-labelled *L. monocytogenes* v7 cells, suspended in 40 µl of sterile de-ionized water at concentrations of $(2.3 \pm 0.2) \times 10^5$, $(6.8 \pm 0.4) \times 10^4$, and $(8.7 \pm 0.8) \times 10^3$ cfu/ml, were injected into the chip at a flow rate of approximately $1.7 \mu$l/min and incubated for at least 12 hours at 37°C. The total number of cells injected was approximately 9200, 27200, and 3480, respectively. During injection of the sample with $(2.3 \pm 0.2) \times 10^5$ cfu/ml, the DEP electrodes were off so that the cells were not concentrated in the incubation chamber, resulting in a probability of approximately 0.09 of finding one cell in the chamber. The other two samples were injected with the DEP electrodes activated, which caused most of the cells in the samples to be collected in the incubation chamber. The actual number of cells collected is not known, but it was visually confirmed that only a very small fraction of the cells escaped the deviation and concentration process. During the switch from water to HLB a more significant fraction of the cells were lost because the DEP force was weakened by the increased medium conductivity coupled to instabilities in the flow rate, but thousands of cells could still be seen collected in the chamber, as shown in Fig. 7b. Fig. 7c shows the change in admittance over time for the three *L. monocytogenes* samples plus sterile HLB. The frequencies chosen for plotting each curve, out of the 51 frequencies that were monitored, were those for which the relative impedance change was the largest. As expected, the sterile media did not exhibit any clear metabolic signal at any frequency. And the bacterial sample injected without the DEP-based concentration system active did not generate a clear metabolic signal corresponding to exponential growth for more than seven hours because the number of cells in the bioprocessor was very low. It is very likely that the linear
increase in admittance observed in this sample, and the slow decrease seen in the sterile media, were due to a drift in the background admittance. Such drift was fairly common in any incubation of sterile media (Fig. 5a). After approximately 7.5 hours, the metabolic signal from the non-concentrated sample is clear and its shape agrees well with the signal observed in previous incubations. On the other hand, in the two samples injected with the concentration system active a very strong metabolic signal was visible during the first hour of incubation.

VI. DISCUSSION

The three experiments discussed above prove very clearly how the DEP-driven concentration system reduces dramatically the length of the detection process for very dilute bacterial suspensions. By collecting a large percentage of the cells from a very dilute sample in the 400 pl detection chamber, the deviation and capture system makes the effectiv cell concentration in the chamber orders of magnitude larger than that in the original sample. In the experiments presented, the concentration factor is between 10^4 and 10^5, which is the ratio of the original sample volume (40 l) to the incubation chamber volume (400 pl) times the fraction of cells that are permanently captured in the chamber, assuming that between 10% and 100% of the cells were captured. The exact concentration factor is difficult to estimate since it is impossible to count the cells remaining in the chamber, but it is almost certain that at least 10% of them remain. Most of the cell losses were caused by instabilities in the flow rate in the detection chamber, which could be eliminated with a better flow control system (using a syringe pump, for example). Even larger concentration factors can be achieved by injecting a larger sample, at the expense of a longer injection time. Increasing the concentration factor would allow the detection of bacteria in even more dilute samples. But even with the current concentration factor it would be possible to detect smaller numbers of bacteria than those tested. Unfortunately, it was not possible to use more dilute cell suspensions with the current setup because they made it impossible to control the flow rates through the device. Flow rate control was performed manually based on the observed velocity of the cells in the channels. Fewer cells in the channels would provide less feedback on the flow velocity.

VII. SUMMARY

We have demonstrated a microfluidic technique for concentrating bacterial cells from a dilute sample, by factors on the order of 10^4 to 10^5, and detecting their metabolic activity by purely electrical means. Tests with dilute L. monocytogenes suspensions demonstrated the ability of this technique to significantly reduce the time needed to detect the presence of the bacteria. We have also explored the use of a physics-based circuit model fitted to impedance measurements at a large number of frequencies, as a means of improving the sensitivity and accuracy of the detection of metabolic activity. The microscale application of impedance-based detection of bacteria coupled with a dielectrophoresis-based cell concentration system has the potential to dramatically decrease the time needed to screen industrial and clinical samples for total bacterial content, which can currently take several days using conventional methods.

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IX. COMPETING INTERESTS STATEMENT
Rashid Bashir and Dallas T. Morisette declare competing financial interests.

REFERENCES

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