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Label-free detection of *Escherichia coli* based on thermal transport through surface imprinted polymers.

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Keywords: Surface imprinted polymers, heat-transfer method (HTM), *Escherichia coli*, *Staphylococcus aureus*, point-of-care bacterial testing.

Abstract

This work focuses on the development of a label-free biomimetic sensor for the specific and selective detection of bacteria. The platform relies on the rebinding of bacteria to synthetic cell receptors, made by surface imprinting of polyurethane-coated aluminum chips. The heat-transfer resistance (R_{th}) of these so-called surface imprinted polymers (SIPs) was analyzed in

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3 time using the heat-transfer method (HTM). Rebinding of target bacteria to the synthetic
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5 receptor led to a measurable increase in thermal resistance at the solid-liquid interface.
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8 *Escherichia coli* and *Staphylococcus aureus* were used as model organisms for several proof-
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10 of-principle experiments, demonstrating the potential of the proposed platform for point-of-
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12 care bacterial testing. The results of these experiments indicate that the sensor is able to
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14 selectively detect bacterial rebinding to the SIP surface, distinguishing between dead and
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16 living *E. coli* cells on the one hand and between Gram-positive and Gram-negative bacteria on
17
18 the other hand (*E. coli* and *S. aureus*). In addition, the sensor was capable of quantifying the
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20 number of bacteria in a given sample, enabling detection at relatively low concentrations (10^4
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22 CFU mL⁻¹ range). As a first proof-of-application, the sensor was exposed to a mixed bacterial
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24 solution containing only a small amount (1%) of the target bacteria. The sample was able to
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26 detect this trace amount by using a simple gradual enrichment strategy.
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3 Bacterial pathogens pose a major threat for public health care, causing a wide variety of
4 conditions ranging from common illnesses to mortal infections. Adequate bacterial testing is
5 therefore of major importance in various fields, including medicine (hospital-acquired
6 infection, HAI), environmental and food safety (water-, air- and food-borne bacteria) and
7 even counter-terrorism (Anthrax infection).¹ Food-borne pathogens cause an estimated 5000
8 deaths annually in the US alone,² while HAI's have shown to affect more than 1 in 20 people
9 that are hospitalized in Florida.³ The containment of bacterial infection is further complicated
10 by the increasing number of multidrug-resistant bacteria, causing millions of deaths every
11 year.⁴

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25 Conventional diagnostic techniques are typically very slow and costly with a limited
26 specificity and sensitivity. Samples are collected, labeled and analyzed in the lab by trained
27 users using immunosorbent techniques such as ELISA.⁵ More recently, sensitive molecular
28 techniques have been developed that are able to selectively detect bacteria in a faster manner.
29 These techniques make use of genetic screening, PCR or real-time PCR and are a huge
30 improvement in comparison to the more classical techniques but they still require sample
31 preparation and expensive equipment that needs to be used in a lab environment by skilled
32 personnel, limiting their use in point-of-care applications.⁶⁻⁸ Therefore, a reliable test that can
33 be used "on-the-bench" could lead to a more accurate and faster diagnosis, thereby improving
34 the prognosis for the patient.

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50 Biosensors offer an elegant alternative to these expensive techniques as they are typically fast,
51 cost-effective and label-free and therefore more suitable for point-of-care applications.
52 Various biosensor platforms have been developed for bacterial identification. These devices
53 employ numerous transducer mechanisms including impedance spectroscopy,⁹⁻¹¹ optical
54 detection,¹²⁻¹⁴ and microgravimetry.¹⁵⁻¹⁷ Although these devices are very sensitive and
55 selective and overcome the problem associated with more expensive, complicated techniques,
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3 there are some drawbacks associated with their use. Biosensors rely on biological receptors
4 such as antibodies,^{18, 19} bacteriophages²⁰ or aptamers²¹ to detect their target. Although these
5 receptors display a high affinity for their target they can be disadvantageous in terms of
6 stability and the complex nature of their synthesis procedure.
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13 The use of synthetic receptors might overcome the problems associated with the use of natural
14 receptors in sensor applications. Synthetic receptors made by molecular imprinting mimic the
15 target sensitivity and selectivity of an enzyme²² but are more stable, reusable, easier and
16 cheaper to produce and have an unlimited shelf-life.²³ Over the past two decades, the concept
17 of molecular imprinting has been extended towards surface imprinting of thin polymer layers
18 for macromolecular templates using various imprinting approaches.²⁴⁻²⁸ Surface-imprinted
19 polymers (SIPs) have been incorporated into numerous biomimetic sensor applications.
20 Platforms based on optical techniques such as ELISA, confocal and fluorescence microscopy
21 have proven to be very selective and extremely sensitive.²⁹⁻³³ However, since these techniques
22 rely on expensive equipment, operated by experienced users in a lab environment and are
23 difficult to miniaturize their applicability for point-of-care testing is limited. Therefore, label-
24 free, low-cost alternatives have been developed based on classical biosensor read-out
25 platforms based on microgravimetric,³⁴⁻³⁶ electrochemical,³⁷⁻³⁹ and optical detection
26 principles.⁴⁰⁻⁴²
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48 In this article, the authors present a new platform for bacterial detection based on the heat-
49 transfer method (HTM). This read-out technique has proven to be a versatile tool for
50 biosensing⁴³ and was combined with SIPs in a biomimetic assay for the detection of cancer
51 cells.⁴⁴⁻⁴⁶ Due to its relatively simple and low-cost nature, HTM offers several benefits over
52 classical biosensing techniques. Impedance spectroscopy, quartz crystal microbalances and
53 surface plasmon resonance *f.e.* require some degree of temperature control, in addition to the
54 sensing hardware, to function optimally. HTM on the other hand, requires little
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3 instrumentation as it functions as a sensing and temperature control platform at the same time
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5 which limits the cost price of the device. Additionally, the data interpretation is relatively
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7 straightforward and the device can be easily scaled down in terms of point-of-care sensing.
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9 Furthermore, HTM does not rely on the electrical conductivity or piezoelectric properties of
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11 the platform material, any solid material can be used as platform provided that it does not
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13 inhibit the heat flow through the SIP layer.
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18 In this paper, polyurethane-coated aluminum chips were imprinted with *Escherichia coli* and
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20 *Staphylococcus aureus* by the so-called micro-contact printing approach that was previously
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22 developed by Dickert *et al.*^{27, 47} Rebinding of bacteria to the SIP lead to a measurable increase
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24 in thermal resistance, detected by HTM. In order to establish a proof-of-principle, it was
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26 assessed whether or not the sensor was able to distinguish between living and dead bacteria
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28 and could selectively detect *E. coli* and *S. aureus*. Additionally, a limit of detection (LoD) was
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30 determined and a first proof-of-application was demonstrated, detecting a trace amount of *E.*
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32 *coli* (1 %) in the presence of an excess of *S. aureus*.
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38 **Experimental Methods**

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41 **Bacterial culturing and sample preparation** The characterized strains of *Escherichia coli*
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43 (ATCC® 8739™) and *Staphylococcus aureus* (ATCC® 6538™) were obtained from DSM-Z.
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45 20 ml nutrient broth (NB, x929.1 ROTH) and Caso broth (TSB, x938.1 ROTH) were
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47 inoculated with a single colony of *E. coli* and *S. aureus* respectively and allowed to grow
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49 overnight at 37°C while shaking. Prior to imprinting, 1ml of the overnight culture was diluted
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51 in 20 ml of the respective broth, and allowed to grow at 37°C for 3 hours or until OD600 of 1
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53 was obtained. Afterwards, the cells were harvested by centrifugation and the pellets were
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55 washed one time with PBS, and resuspended in phosphate buffered saline to achieve the
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57 desired concentration.
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6 **Preparation of bacteria-imprinted polyurethane layers** Polyurethane layers were
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8 synthesized by dissolving 122 mg of 4,4'-diisocyanatodiphenylmethane, 222 mg of bisphenol
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10 A, and 25 mg of phloroglucinol in 500 μL of anhydrous tetrahydrofuran (THF). All reagents
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12 had a purity of at least 99.9% and were used as received (Sigma-Aldrich N.V., Diegem,
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14 Belgium). The mixture was polymerized up to the gelling point at 65°C for 200 minutes while
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16 gently stirring. The solution was diluted in anhydrous THF in a 1:5 ratio. Polyurethane layers
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18 with an average thickness of $1.2 \pm 0.1 \mu\text{m}$, as measured with a profilometer (Dektak3ST,
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20 Sloan Instruments Corporation, Santa Barbara, CA) were created by spincoating the solution
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22 during 60 s at 2000 rpm onto 1 cm^2 aluminum substrates.
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27 In parallel, bacteria-covered homemade polydimethylsiloxane (PDMS) stamps were formed
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29 by applying 400 μL of a bacterial suspension in PBS to the stamp. The bacteria were allowed
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31 to sediment to the surface of the stamp for 10 minutes and the excess fluid was removed by
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33 spin coating the stamp at 3000 rpm for 60 s to create a dense monolayer of bacteria on the
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35 stamp surface. PDMS stamps were made using the Sylgard 184 silicone elastomer kit
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37 (Malvom N.V., Schelle, Belgium). The bacteria-covered stamps were gently pressed onto the
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39 semi-cured polyurethane layer to ensure full contact and the bacteria were allowed to sink into
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41 the layer passively due to the weight of the stamp. Finally, the polymer was cured for 18
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43 hours at 65°C under inert atmosphere after which the stamp was removed from the surface.
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Template bacteria were washed off with a 70 % ethanol in MilliQ and PBS, leaving behind
selective binding cavities on the surface, creating SIPs for *E. Coli* and *S. aureus*.

55 **Sensor setup & measuring methodology** The sensor setup and its use in cell-binding assays
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57 has been described earlier.³⁴⁻³⁷ The proportional-integral-derivative (PID) settings (P = 1, I =
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59 8, D = 0) used were optimized in a previous study.³⁸ The system is allowed to stabilize in PBS

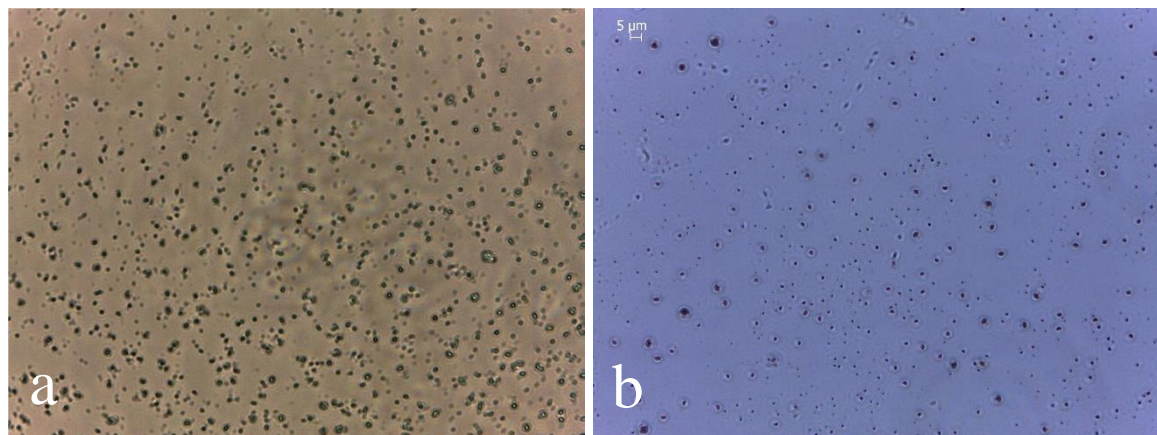
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3 buffer (pH = 7.4) at the beginning of each experiment. Bacteria are introduced to the system
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5 by injecting 3 mL of a bacterial solution (1×10^7 CFU mL⁻¹ in PBS) at a controlled flow rate
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7 of 2.5 mL min⁻¹. The system is left to stabilize after which the system is flushed with PBS at a
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9 flow rate of 0.25 mL min⁻¹ for 12 minutes (total volume 3 mL) to remove any unbound
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11 bacteria from the SIP layer. The HTM setup monitors the thermal resistance (R_{th}) at the solid-
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13 liquid interface at a rate of one measurement per second.
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20 **Imprint characterization** Microscopic imaging of the cell-imprinted polyurethane surfaces
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22 was performed with a Leica DM750 optical microscope (Leica Microsystems, Diegem,
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24 Belgium. All SIPs were imaged at magnifications 640x and 1000x. ImageJ 1.44P (National
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26 Institute of Health, Bethesda, MA, USA) was used to determine the number of cell imprints
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28 per area unit on microscopic images of the SIPs. The average surface coverage of cell
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30 imprints on the polyurethane layer was calculated based on cell imprint counts of three
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32 different samples for each type of SIP and five locations on each sample. An Agilent 5500
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34 AFM system was used with MSNL-F cantilevers ($f = 110=120$ kHz, $k = 0.6$ N/m) with
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36 average tip radius of $2=12$ nm for topographical imaging in intermittent contact (AAC) mode.
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38 The AFM topography images were leveled, line-corrected and measured (height profiles)
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40 using Gwyddion, a free and open-source SPM (scanning probe microscopy) data visualization
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42 and analysis program.
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49 **Results**

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52 **Surface characterization and calculation surface coverage** Optical analysis of a SIP
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54 surface imprinted with *E. coli* clearly reveals a heterogeneous distribution of imprints with an
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56 average diameter of 0.5 to 2 μ m, corresponding to the dimensions of the template bacteria
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58 (figure 1a). The calculated surface coverage of $7.18 \times 10^6 \pm 8.54 \times 10^5$ imprints cm²
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60 corresponds to a total surface coverage of 14.13 ± 1.8 %. The microscopic analysis of the *S.*

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3 *aureus* SIP (figure 1b) shows a heterogeneous distribution of spherical imprints with a
4 diameter of $\pm 500 \text{ nm} - 800 \text{ nm}$. The imprint surface coverage of $5.82 \times 10^6 \pm 9.84 \times 10^5$
5 imprints cm^2 corresponds to a total surface coverage of $12.36 \pm 2.3 \%$.
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26 **Figure 1.** Microscopic analysis of an *E. coli* (a) and *S. aureus* SIP (b). Images were made at a
27 magnification of 1000 and Image J was used to calculate imprint surface coverage.
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30 To analyze the morphology of the imprints more thoroughly, an *E. coli* SIP was used for
31 topographical analysis with an atomic force microscope (Figure 2). The overview of a 20×20
32 μm^2 area clearly shows a heterogeneous distribution of rod-shaped imprints with lengths
33 ranging from 0.5 to $2 \mu\text{m}$ and widths of 0.2 - $0.5 \mu\text{m}$ which corresponds well to the shape and
34 size of the template bacteria (figure 2a). One of these imprints was analyzed more thoroughly
35 confirming that the horizontal dimensions of the template are faithfully transferred into the
36 SIP layer (figure 2b). However, the 3D image and depth profile (figure 2c and d) indicate that
37 the vertical dimensions are not transferred to the layer, as the imprint is quite shallow ranging
38 up to about 30 - 40 nm in depth. These data are in accordance with a previously performed
39 analysis on a MCF-7 SIP which demonstrated that the shallow nature of the imprints is
40 actually the key to their selectivity.⁴⁴
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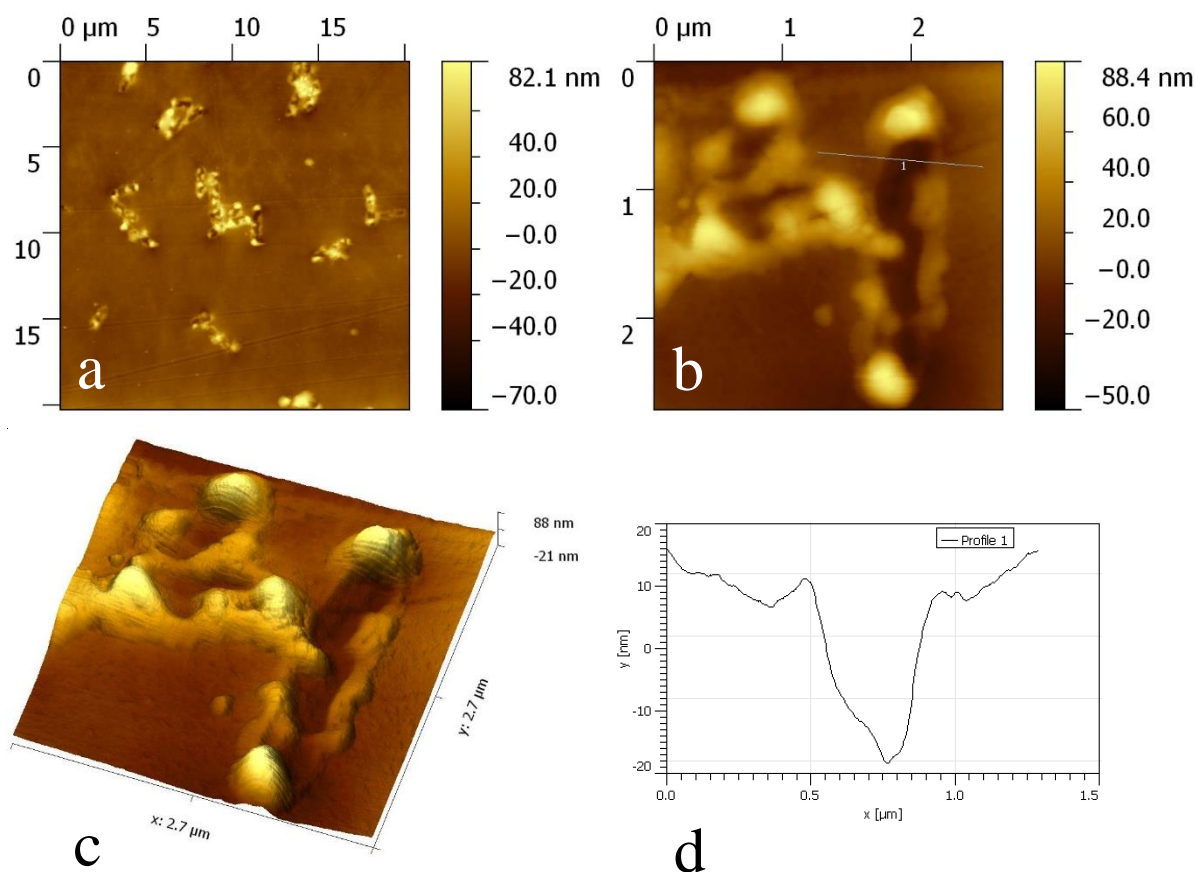


Figure 2. Topographical analysis of an *E. coli* SIP using an atomic force microscope. Rod-size imprints can be observed (a) with dimensions corresponding well to the size and shape of the template bacteria (b). The 3D-imprint (c) and depth (d) profile reveal that the imprints are quite shallow with depths up to 40 nm.

Bacterial detection and live/dead discrimination The setup's potential to discriminate between living and dead bacteria, SIP's were imprinted with living *E. coli* cells in PBS (concentration 1×10^7 CFU mL⁻¹) as described in Experimental Section 2.2. The SIP-coated aluminum was mechanically pressed with its non-coated, polished backside onto a copper block, ensuring an optimal thermal contact between chip and heat sink. The flow cell was filled with PBS and the R_{th} signal was allowed to stabilize for 60 minutes.

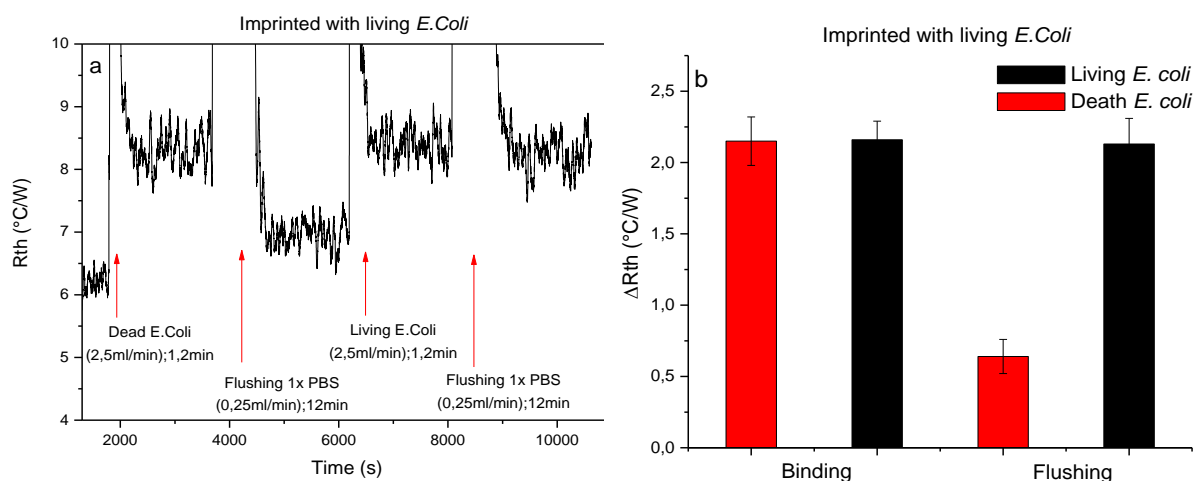


Figure 3. Time-dependent R_{th} data of a SIP imprinted with living *E. coli* cells after consecutive exposure to PBS solutions containing dead and live *E. coli* respectively (1×10^7 CFU mL⁻¹) (a). The results demonstrate that both exposure events result in an increase in thermal resistance at the solid-liquid interface. The increase associated with an addition of dead bacteria can be partially reversed by flushing with PBS, whereas the increase caused by adding living *E. coli* cells is irreversible. A boxplot is shown summarizing the data (b). Error bars indicate the standard deviation of the noise on the signal.

Dead bacteria are introduced into the flow cell at a flow rate of 2.5 mL min⁻¹ (72 seconds, 3mL). The flow is stopped and the signal is left to stabilize for 60 minutes, allowing the bacteria to sediment towards the SIP surface. Any unbound bacteria are removed by flushing the flow cell with PBS at a rate of 0.25 mL min⁻¹ (12 minutes, 3mL). After a 60-minute stabilization interval, the experiment is repeated with living *E. coli* cells. The results of this experiment are shown in figure 3 and demonstrate that the signal increases upon addition of a solution of dead bacteria in PBS by 2.15 ± 0.17 °C/W. Upon flushing the chamber with PBS the signal drops back with to a value 0.64 ± 0.12 °C/W above the baseline. After infusing the live bacteria into the measuring chamber the signal increases again to a value 2.16 ± 0.13

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3 °C/W. Flushing with buffer solution does not cause a measurable decrease in R_{th} as the signal
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5 remains at 2.13 ± 0.18 °C/W above the baseline.
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10 **Selectivity test: *E. coli* (Gram-negative) vs. *S. aureus* (Gram-positive)** To investigate the
11 selectivity of the proposed platform, SIP's were imprinted with *S. aureus* and *E. coli*
12 analogous to the experiment described in Section 3.2. The time-dependent R_{th} data acquired
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The data summarized in figure 4a indicate that exposing an *E. coli* SIP to a suspension of *S. aureus* cells in PBS (concentration 1×10^7 CFU mL⁻¹) increases the thermal resistance at the solid-liquid interface with 1.91 ± 0.22 °C/W. Rinsing the flow cell with PBS will return the signal back to baseline ($\Delta R_{th} = 0.06 \pm 0.15$ °C/W). Repeating the cycle with an *E. coli* solution with the same concentration will lead to an irreversible increase in R_{th} of 2.13 ± 0.14 °C/W (ΔR_{th} upon flushing = 2.08 ± 0.23 °C/W). A similar trend can be observed when performing the same experiment with a *S. aureus* SIP (figure 4b). Exposure to a solution of *E. coli* cells increases the R_{th} signal with 1.63 ± 0.15 °C/W but upon rinsing the flow cell with PBS the thermal resistance stabilizes at a value -0.14 ± 0.25 °C/W above the baseline. Exposing the SIP to a solution of target cells on the other hand, will lead to an increase in thermal resistance of 1.79 ± 0.14 °C/W. Flushing the cell with PBS will not significantly change the signal (1.77 ± 0.16 °C/W).

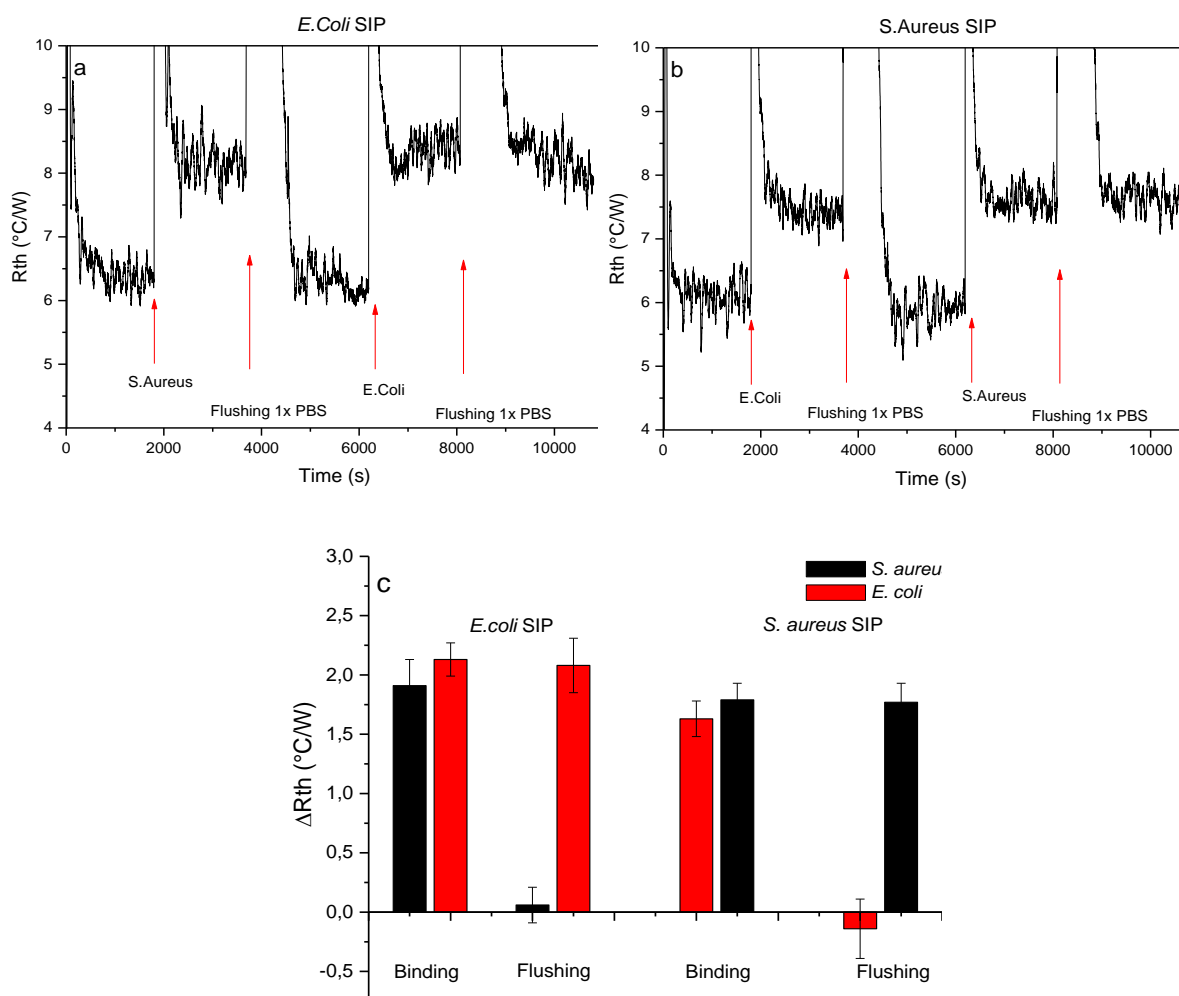


Figure 4. Time-dependent R_{th} measurements of SIPs imprinted with either *E. coli* (a) or *S. aureus* (b) during consecutive bacterial exposure events to analogue non-target bacteria and finally target bacteria. In both cases, addition of non-target bacterial species leads to an increase in thermal resistance, but the signal returns back to baseline upon flushing the flow cell with buffer solution. Binding of target bacteria to the SIP on the other hand, leads to an irreversible rise in R_{th} . The results of this experiment are summarized in a box plot (c).

Sensitivity test: dose-response curve In order to determine the limit-of-detection (LoD) of the sensor, the time-dependent R_{th} response of an *E. coli* SIP, exposed to an increasing concentration of target cells, was analyzed. To this extent a stock solution of *E. coli* cells in PBS with a concentration of 1×10^7 CFU mL^{-1} was diluted 1000, 500, 200, 100, 50, 20, 10 and

5 times and the SIP was consecutively exposed to an increasing concentration of target cells. In between each exposure step, the flow cell is rinsed with ethanol and, upon stabilization of the signal, PBS to ensure full removal of the bacteria from the SIP layer both steps were performed at a rate of 0.25 mL min^{-1} (12 minutes, 3mL). The results of this experiment are described in figure 5.

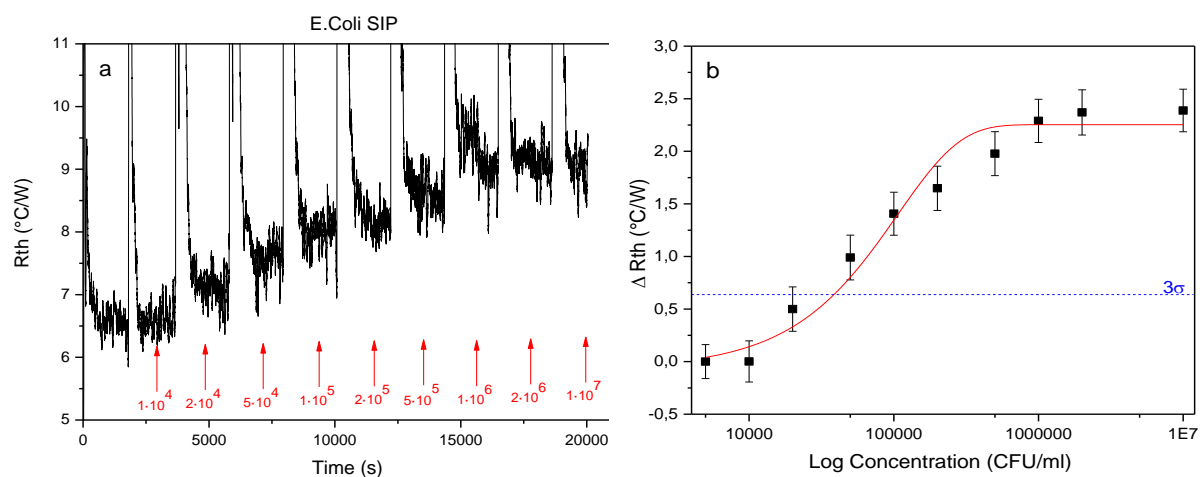


Figure 5. Dose-response experiment performed on a SIP imprinted with *E. coli*. A stock solution ($1 \times 10^7 \text{ CFU mL}^{-1}$) was diluted 1000, 500, 200, 100, 50, 20, 10 and 5 times and the SIP was exposed to an increasing concentration of target cells (a). Upon each exposure step, the layer was rinsed with ethanol and PBS to ensure full removal of the cells. The thermal resistance increases noticeably and the increase seems to be concentration-dependent. These results combined with the results from the previous experiment were used to establish a dose-response curve: response in R_{th} as a function of the added target-bacteria concentration (logarithmic). An exponential fit is drawn through the obtained data with an R^2 -value of 0.97. The dashed line corresponds to the limit-of-detection, defined as three times the highest error on the data (b).

The time-dependent thermal resistance data shown in figure 5a, indicate that exposing the SIP to a concentration of $1 \times 10^4 \text{ CFU mL}^{-1}$ does not result in a measurable increase in R_{th} . Upon

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3 addition of a concentration of 2×10^4 CFU mL⁻¹ the signal starts increasing in a
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5 concentration-dependent manner and saturates at a concentration of 2×10^6 CFU mL⁻¹. These
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7 data were used to obtain a dose-response curve (figure 5b). The dose-response curve nicely
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9 follows ($R^2 = 0.97$) an empirical, exponential fit function according to the formula:

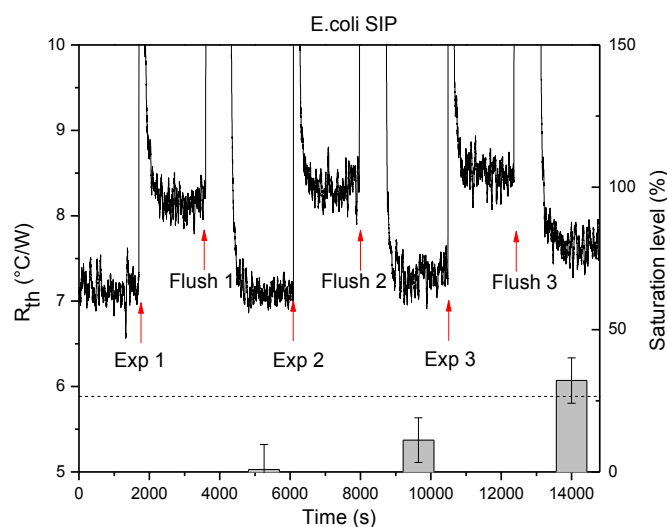
$$\Delta R_{th}(c) = A - B \cdot \exp\left\{-\frac{c}{C}\right\}$$

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16 The limit-of-detection was calculated using this dose-response curve and is defined as the
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18 intercept between the fitted curve and the dashed line in figure 5b corresponding to three
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20 times the biggest error on the data set.⁴⁴
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25 **Proof-of-application: detection of *E. coli* in a semi-complex matrix** The results obtained in
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27 the previous sections indicate that the sensor is indeed able to selectively and specifically
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29 detect bacterial cells in buffer. To examine if our sensor is able to detect its target in a more
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31 complex matrix a mixed cell solution was made containing both *E. coli* and *S. aureus* cells in
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33 a 1:99 ratio (total concentration of bacteria: 1×10^7 CFU mL⁻¹). This mixture was used in a
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35 progressive enrichment experiment, exposing a SIP imprinted with *E. coli* three consecutive
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37 times to the mixture, while flushing the layer with buffer between each exposure event. The
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39 results are shown in figure 6 and indicate that the signal does not significantly increase in
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41 comparison to the baseline after the first exposure event. The net change in thermal resistance
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43 appears to be positively correlated to the number of exposure steps as R_{th} increases after the
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45 second and third exposure step.
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51 To demonstrate this effect more clearly, the saturation level at each step was determined as
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53 the ratio of ΔR_{th} after exposure to the mixture and after flushing with buffer respectively. The
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55 LoD is illustrated as a dashed line and is defined as three times the standard deviation on the
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57 signal corresponding to 26.424 %. After the first two cycles the signal only reaches 0.8 ± 8.08
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3 % and 11.8 ± 7.82 %, well below the detection limit. After a third exposure round the signal
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5 reaches the limit of detection at a saturation level of 32.1 ± 8.00 %.



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26 **Figure 6.** Progressive enrichment experiment conducted on an aluminum chip covered with
27 an *E. coli* SIP. The SIP is exposed to a 1:99 mixture of *E. coli* and *S. aureus* cells. Each
28 exposure event consists of injection of the mixture, stabilization of the signal, rinsing the flow
29 cell with buffer and another stabilization period. This cycle is repeated three consecutive
30 times. The data show that the signal gradually increases after each exposure event. This was
31 summarized in a box plot of the saturation level, defined as the ratio of ΔR_{th} after flushing and
32 exposure respectively. The dashed line corresponds to the LoD, defined as three times the
33 standard error on the signal.
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48 Discussion

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50 The experiments on dead and living *E. coli* cells clearly demonstrate that the difference in
51 surface chemistry is sufficiently big to discriminate between both, despite the morphological
52 similarities. The thermal resistance profile in figure 3 shows a comparable response upon
53 initial exposure to dead and living bacteria, although the increase in R_{th} is somewhat lower for
54 dead cells. This can be explained by the fact that the morphology of the dead cells is
55 compatible to the dimensions of the microcavities on the SIP surface. Both living and dead
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3 cells block the heat flow through the microcavities of the SIP, thereby increasing the heat-
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5 transfer resistance at the solid-liquid interface. However, as the AFM analysis in figure 2
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7 demonstrates that the imprints are quite shallow in comparison to the diameter of the
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9 template, which corresponds to previous findings,⁴⁴ the morphological match must be
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11 complemented by a complementarity in the distribution of functional groups between imprint
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13 and template to ensure tight adhesion of bacteria to the surface. Previous research has
14
15 demonstrated that both protein expression and the presence of carbohydrate groups on the
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17 outer membrane of the cells are decisive elements in the recognition of cells by SIPs.⁴⁵ In
18
19 addition, Hayden *et al.* demonstrated that the recognition is driven by the formation of
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21 hydrogen bonds that were created between imprint and template during crosslinking of the
22
23 polymer in the imprinting procedure.³⁵ This explains why the signal returns back to baseline
24
25 after flushing with PBS in the exposure experiment performed with the dead *E. coli* cells. The
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27 bacteria were killed in 70% ethanol which denatures the membrane proteins of the cells and
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29 dissolves the phospholipids inside the membrane. Therefore, the dead bacteria are not tightly
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31 bound to the SIP and can be washed out easily. The small degree of cross-selectivity that is
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33 observed can be attributed to the fact that some carbohydrate patterns might still be present on
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35 the dead cells which, by chance, might result in a bond that is strong enough to withstand the
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37 shear force associated with flushing.
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46 The cross selectivity experiment described in figure 4 reveals a similar trend. In both cases,
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48 exposure of the SIP to a solution containing an analogue bacterial species, leads to an increase
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50 in thermal resistance. However, these cells can be washed away easily by rinsing the flow cell
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52 with PBS, while target cells remain bound to the layer, even after flushing. This can be
53
54 explained the fact that gram-negative *E. coli* and gram-positive *S. aureus* have a distinctly
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56 different outer membrane in terms of protein expression and the presence of carbohydrate
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58 patterns.^{48, 49} The bond between the analogue cells and the SIP can therefore be easily broken
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3 by flushing the flow cell with buffer. The target bacteria on the other hand, remain firmly
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5 bound to the layer and the thermal resistance remains at an elevated level even after flushing.
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8 The data in figure 5 reveal that the sensor does not only qualitatively respond to an elevated
9
10 concentration of target bacterial species in a sample but the response can also be quantified.
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12 At relatively low concentrations, the sensor's response stays within noise levels and can
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14 therefore not be regarded as significant. But starting from a concentration of 2×10^4 CFU mL⁻¹
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17 we see the signal significantly increasing to a value well above the baseline, indicating that a
18
19 sufficient amount of cells interacts with and binds to the microcavities on the SIP, blocking
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21 the heat flow through the layer and thereby increasing the heat-transfer resistance. This effect
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23 becomes more pronounced with an increasing concentration but the sensor seems to saturate
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25 at concentrations above 2×10^6 CFU mL⁻¹. Using the exponential fit to the data and defining
26
27 the detection limit as the concentration at which the signal-to-noise ratio is bigger than 3, we
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29 calculated the LoD at 3.5×10^4 CFU mL⁻¹. Although there are label-free platforms that are
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31 more sensitive,¹ HTM has the benefit of being very low-cost, user-friendly and can be easily
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33 scaled down in terms of point-of-care sensing. Furthermore, all data shown are raw, unfiltered
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35 data. The amount of noise on the signal that can be observed, originates from stringently
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37 controlling the temperature underneath the sample (T_1). However, the noise can be decreased
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39 by filtering, electronic noise reduction and careful re-designing of both the flow cell and the
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41 measuring device, which would significantly improve the sensitivity of the set up. The
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43 experiments demonstrated in this article describe a very first series of experiments that assess
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45 the platform's potential for bacterial identification.
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47 The final set of experiments provides a very first proof-of-application illustrating how the
48
49 sensor would respond to a more complex matrix as opposed to a cell solution containing only
50

51 one type of bacterial species. The results in figure 6 also demonstrate another possible method
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53 for improving the LoD by gradually exposing the SIP to target cells during multiple
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3 consecutive exposure cycles. After exposure to the bacterial mixture, the R_{th} signal initially
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5 increases to saturation which indicates that both target and analogue cells bind to the layer.
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7 After flushing the signal falls back to a value that does not significantly differ from the
8
9 baseline value. This can be explained by the fact the *E. coli* cells are overwhelmed by a 99-
10
11 fold excess of *S. aureus* cells which also bind to the microcavities upon addition of the
12
13 mixture to the flow cell. *E. coli* cells cannot bind to microcavities that are already occupied by
14
15 analogue bacteria. Due to steric hindrance the analogue bacteria also prevent the target
16
17 bacteria from interacting with the SIP. These findings indicate that the sensitivity of the
18
19 platform could decrease when analyzing bacteria in a complex matrix. This can be overcome
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21 by increasing the number of exposure cycles. With this enrichment strategy, the signal will
22
23 gradually increase with each cycle and eventually reach the LoD which enables detection of
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25 lower concentrations of bacteria in increasingly complex mixtures. In order to detect trace
26
27 amounts of bacteria in biological or environmental samples the device will need to be
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29 redesigned and probably combined with pretreatment of the samples under analysis.
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38 **Conclusion**

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40 The results obtained during this study indicate that the proposed sensor platform is able to
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42 detect bacteria in buffer with a high degree of selectivity. Although the device performance
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44 has only been assessed in buffer the presented manuscript offers a first proof-of-principle,
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46 illustrating the potential of the combination of SIP's and HTM for bacterial detection. Further
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48 research should be devoted at improving the sensitivity of the device. This can be achieved by
49
50 optimizing the imprinting procedure to obtain a higher and more homogenous surface
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52 coverage on the SIP surface which will lead to a larger effect size and improved LoD. In
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54 addition, the noise on the thermal resistance signal can be improved by optimizing both the
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56 measurement technique and flow cell. Finally, the results obtained within, the gradual
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3 enrichment experiment indicate that the sensitivity of the device can be improved by
4 maximizing the exposure between sample and receptor layer by *e.g.* developing a flow cell for
5 continuous exposure.
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23 24 **Author Contributions**

25
26 B.v.G. and K.E. contributed equally to this work. K.E. prepared all cell-imprinted polymer
27 layers and was responsible for culturing cells in cooperation with O.A. and E.S.R. O.A., S.E.
28 and A.K. were in charge of optical analysis of the surface. The heat-transfer device was
29 designed and built by B.v.G, H.D. and T.J.C. All heat-transfer measurements were performed
30 by B.v.G. in cooperation with K.E and O.A. Biological assistance and guidance on possible
31 medical/biotechnological applications was provided by S.E., A.K., E.S.R. and P.W., while
32 H.D., M.P. and T.J.C. provided input on SIP synthesis and template removal. B.v.G. and K.E.
33 interpreted the heat-transfer data in close cooperation with T.J.C. and P.W. O.D. and C.B.
34 performed the AFM analysis on the samples and assisted in interpreting the results. The
35 manuscript was jointly written by K.E. and B.v.G. All authors have given approval to the final
36 version of the manuscript.
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Table of Content Figure

