Rapid and Ultrasensitive Detection of Mutations and Genes Relevant to Antimicrobial Resistance in Bacteria

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The worldwide emergence of multidrug-resistant (MDR) bacteria is associated with significant morbidity, mortality, and healthcare costs. Rapid and accurate diagnostic methods to detect antibiotic resistance are critical for antibiotic stewardship and infection control measurements. Here a cantilever nanosensor-based diagnostic assay is shown to detect single nucleotide polymorphisms (SNPs) and genes associated with antibiotic resistance in Gram negative (Pseudomonas aeruginosa) and positive (Enterococcus faecium) bacteria, representing frequent causes for MDR infections. Highly specific RNA capture probes for SNPs (ampR<sup>D135G</sup> or ampR<sup>C154R</sup>) or resistance genes (vanA, vanB, and vanD) allow to detect the binding of bacterial RNA within less than 5 min. Serial dilutions of bacterial RNA indicate an unprecedented sensitivity of 10 fg µL<sup>-1</sup> total RNA for SNPs and 1 fg µL<sup>-1</sup> total RNA for vanD detection equivalent to single bacterial cell sensitivity.

1. Introduction

Multidrug resistant (MDR) bacteria pose an important threat to human health and achievements of modern medicine. In a recent report[1] (WHO, UN ad hoc interagency coordinating group on antimicrobial resistance), the World Health Organization estimates that bacteria with antibiotic resistance account for at least 700 000 deaths globally per year. A figure that could increase to ten million cases of deaths globally by the year 2050.[2] There is a clear danger that in a post-antibiotic era, common infections, such as pneumonia, blood stream infections, and sexually transmitted diseases, literally will become untreatable and surpass cancer-related deaths.

Bacteria harbor a broad range of resistance mechanisms against antimicrobial substances, many of which are genetically encoded. Therefore, determination of genotypes can be transferred to phenotypic behavior during antibiotic treatment. Single nucleotide polymorphisms (SNPs) may alter antibiotic target binding sites or expression of, e.g., porins resulting in antibiotic drug resistance. Resistance can also be acquisition of mobile genetic elements such as plasmids and transposons, harboring genes encoding for modified target binding sites or enzymes able to cleave antibiotic substances.[3] Bacteria with these resistance mechanisms have emerged worldwide due to a lack of antibiotic stewardship in veterinary and human medicine.[4]

In a clinical context, rapid and reliable detection of MDR bacteria is critical for treatment adaptations, antibiotic stewardship, and infection control. Current available technologies for screening and identification of MDR bacteria have been recently reviewed.[5] A key problem of phenotypic resistance tests is the requirement to culture and subculture specimen in order to reach single colonies for subsequent processing on, e.g., broth dilutions. The process to determine antibiotic resistance breakpoints usually takes 48–72 h.[6] Based on antimicrobial resistance (AMR) profiles, treatments can be tailored to more effective drugs, which is associated with patient mortality.[7] However, slow growing bacterial species as well as fastidious or anaerobic bacteria pose challenges for rapid AMR testing (Figure 1). Molecular diagnostic testing such as PCR or isothermal amplification[8] may overcome these obstacles by rapid amplification of DNA and specific detection. Current molecular diagnostics to detect gene or SNP associated with antibiotic resistance are commonly used and allow rapid detection within a few hours. However, these technologies are often expensive and detection of resistance mechanisms directly from specimen may be challenging due to inhibitory substances and sensitivity issues.[9,10]

A recently introduced method based on an atomic force microscopy (AFM) cantilever may overcome the previously mentioned obstacles.[11–15] A variation of this technology measures the activity of bacteria in real time,[16,17] whereby the determination of low-frequency fluctuations (<1 kHz) allows to
distinguish living from dead bacteria. Antibiotic susceptibility can be inferred based on fluctuation changes while adding different concentrations of antibiotics. The method still requires single bacterial colonies from previous cultures. The method would require preselection of the bacteria and requires placement of bacteria onto a single cantilever only without specifically immobilizing them.

Here, we applied nanomechanical microcantilever arrays using a fundamentally different approach, providing internal references for differential readout to eliminate thermal drift and nonspecific binding (Figure 2). Such devices represent ultra-sensitive sensors for the detection of biochemical interactions in liquid environments. Extracted RNA from isolates or samples are directly measured without pre-concentration. Each nanosensor is coated with probes (gene specific oligonucleotide) on a sensitive layer for molecular recognition. The binding of the bacterial RNA sequence is mechanically transduced to the cantilever surface, resulting in bending of the cantilever. In this study, we first focus on the identification

Figure 1. A schematic representation highlights the differences between the standard workflow and the nanosensor workflow. The enrichment and cultivation step can be omitted and shortens the procedure drastically from days to few hours. Particularly cultivation of fastidious or anaerobic bacteria, such as M. tuberculosis can take several weeks. Furthermore, the sample preparation step, extraction of DNA/RNA can be included into the sample collection stage.

Figure 2. Microfabricated arrays (a) scanning electron microscopy picture of an array used in this study). Arrays of eight identical silicon cantilevers with 250 µm pitch, a length of 750 µm, a thickness of 500 nm were provided by the Micro- and Nanomechanics group at the IBM Zurich Research Laboratory and coated with a layer of 3 nm titanium and 20 nm gold. A cantilever array is functionalized using a Microdrop inkjet printer, incubated for 1 h at room temperature and then mounted in a microfluidic cell. The same time total RNA can be extracted from bacterial samples and is then diluted in 0.03× SSC. Different concentrations between 1 fg µL⁻¹ and 40 ng µL⁻¹ of total RNA were injected sequentially at a flow rate of 5 µL min⁻¹ for a total of 100 µL volume by pulling the liquid through the liquid cell with the help of a syringe pump and a multiway valve. All measurements were performed at 28 °C in a temperature controlled box. Time-multiplexed vertical-cavity surface-emitting lasers (VCSEL) with regulated power supply (operated at 1 Hz, wavelength 760 nm, Avalon Photonics, Zurich, Switzerland) were used in combination with adjustable optics to yield a pitch of 250 µm. A 2D linear position-sensitive detector, PSD (SiTek Electro Optics, Partille, Sweden) was used for beam deflection readout of each cantilever with an accuracy of 0.1 nm. The signals were preamplified, and the data were acquired using a National Instruments (Austin, TX) PCMCIA 16XE-50 (16 bit, 200 kS s⁻¹) data-acquisition card. The instrument is driven by LABVIEW software to control liquid exchange via a syringe pump (GENIE, Kent Scientific, Torrington, CT), and a multiway valve selector (Rhoedyn, Rohnert Park, CA) and data acquisition (b) binding of target molecules to the sensor surface). Data processing is done using Origin Software.
of SNPs in bacterial genes (ampR<sup>D135G</sup>, ampR<sup>G154R</sup>)<sup>[20]</sup> that are involved in modulating gene expression and secondly on whole gene detection of typical resistance genes (vanA, vanB, vanD genes).

2. Results and Discussion

2.1. Detection of SNPs

First, we focus on single nucleotide polymorphisms in the bacterial genome. In order to assess the assay’s sensitivity to detect SNPs in the bacterial genome, we have designed probes able to bind specific mutations. As a proof of concept, we have selected the ampR gene, an important transcriptional regulator<sup>[20]</sup> in Gram negative bacteria, which is able to alter gene expression. AmpC is a beta-lactamase enzyme able to cleave many commonly used beta-lactam antibiotics. Two previously described and common mutations, i.e., ampR<sup>D135G</sup> and ampR<sup>G154R</sup> were used and compared to wild type ampR<sup>wt</sup>. These mutations are associated with phenotypic resistance against a series of beta-lactam antibiotics. We used four different oligonucleotides as probes, 19 bases in length, where the location of the point mutation is in the center to increase hybridization specificity (Table 1).

Based on previous nanosensor experiments<sup>[24]</sup> we found that this position yields optimized results. The four oligonucleotides consisted of wild type, ampR<sup>D135G</sup>, ampR<sup>G154R</sup>, and a different nucleotide at the position of the mutation as a reference termed ampR<sup>wt</sup>. We were able to distinguish ampR<sup>wt</sup> (Figure 3a), ampR<sup>D135G</sup> (Figure 3b), and ampR<sup>G154R</sup> (Figure 3c) using corresponding matching sequences within a few minutes. Multiple mutations are probed using the same array to discriminate two different mutations and the wild type, literally toggling between the different sequences.

We performed experiments at 28 °C at various concentrations of total RNA from P. aeruginosa ampR<sup>D135G</sup> and ampR<sup>G154R</sup> (Figure 3d,e) in the range from 1 fg µL<sup>−1</sup> up to 10 ng µL<sup>−1</sup> to investigate the limit of detection (LOD) for SNPs using a logistic fit function (Equation 1). We obtained average signals in between 5 ±1 nm up to 1202 ±10 nm. Single point mutation specificity was demonstrated in these experiments using arrays with cantilevers of 750 µm length and 500 nm thickness. 1 pg of total RNA was injected, corresponding to a sensitivity of detection of merely ten bacterial cells.

### Table 1. Sequences of AmpR oligonucleotides used in this study. The different SNPs are highlighted by capitalized letters.

| Designation   | Sequence                  | Position range | Position | T<sub>m</sub><sup>47</sup> |
|---------------|---------------------------|----------------|----------|-----------------------------|
| ampR<sup>wt</sup> | gcggcgatgTgcgcgcgggt     | 394–413        | 404      | 27.3 °C                     |
| ampR<sup>D135G</sup> | gcggcgatgCgcgcgcgggt    | -               | -        | 28.0 °C                     |
| ampR<sup>G154R</sup> | gcggcgatgAcgcgcgggt     | -               | -        | 27.1 °C                     |
| ampR<sup>D135G</sup> | gcgggcggcgatgcgcgcggc    | 452–470        | 460      | 28.1 °C                     |
| ampR<sup>G154R</sup> | gcgggcggcgatgcgcgcggc    | -               | -        | 27.9 °C                     |
| ampR<sup>G154R</sup> | gcgggcggcgatgcgcgcggc    | -               | -        | 27.1 °C                     |

<sup>47</sup>T<sub>m</sub> stands for melting temperature.

Second, we approached specific detection of antibiotic resistance genes, e.g., coding for genes producing enzymes able to cleave antibiotics or resulting in reduced binding affinities of an antibiotic to its target.<sup>[23]</sup> Three different and common vancomycin resistance genes<sup>[24,25]</sup> called vanA, vanB, and vanD were investigated. The three genes are distinguished by multiple mutations within the genes. We chose 19 to 21 long oligonucleotides (Table 2) from the 3’ terminal end as probes with similar hybridization temperatures for the experiments. In these experiments, a polyAC sequence of 19 bases was used as a reference sequence. We detected vanA, vanB, and vanD using total RNA extracted from Enterococcus faecium.

We initially used RNA concentrations of 100 fg µL<sup>−1</sup> for reliable and rapid detection of resistance genes (Figure 4a–d) and injected in each case overall 10 pg total RNA, which is equivalent to approximately 100 bacterial cells.<sup>[36]</sup> All three E. faecium RNA samples show a similar response of 80 nm indicating similar expression levels of the different vancomycin genes.

A series of experiments with different concentrations in the range of 1 fg µL<sup>−1</sup> to 40 ng µL<sup>−1</sup> (Figure 5) was conducted. Table 3 summarizes the corresponding cross reactivities between the probes on the sensors and the injected targets. The fact that cross reactivity signals from noncorresponding probes are only a few percent in magnitude of gene recognition signals from corresponding probes underlines the high specificity of this method.

The upper limit in the differential deflection signal at 40 ng µL<sup>−1</sup> is a result of the number of probes on the cantilever surface and the percentage that can bind to a target. These values were estimated to be 1.5 × 10<sup>6</sup> oligonucleotides per nanosensor and a maximum of 10% have been shown to bind.<sup>[27]</sup> Respectively, 40 ng RNA will contain an estimated number of 1.6 × 10<sup>9</sup> targets which corresponds to an occupancy of 10%. Therefore, as 10% of all possibly accessible probes are occupied, we almost reached the theoretical limit for binding.

Determination of Gibbs free energy (Figure 6) using Equation 2 for the hybridization of the vanD gene resulted in a value of ΔG = −58.38 kJ mol<sup>−1</sup>. The Gibbs free energy calculations gave a value of ΔG = −61.5 kJ mol<sup>−1</sup> for ampR<sup>D135G</sup> and −62.65 kJ mol<sup>−1</sup> for ampR<sup>G154R</sup>. Values of ΔG for the point mutations are larger than for the whole gene detection. This finding justifies the observation of the higher equilibrium binding signal of around 1100 nm versus the 400 nm differential deflection signal for the vanD gene.

$$D(c) = D_{\text{min}} + \frac{D_{\text{max}} - D_{\text{min}}}{\frac{1}{c^{b}} + \frac{1}{c}}$$  \hspace{1cm} (1)

$$D(c)$$ is equal to the deflection at a particular concentration c, $$D_{\text{min}}$$ corresponds to the minimum deflection and $$D_{\text{max}}$$ to the maximum deflection. It corresponds to the Hill coefficient at $$c_0$$ equal to half $$D_{\text{max}}$$. s is a control factor.

2.2. Detection of Bacterial Genes

### Table 2. Corresponding cross reactivities between different SNPs and 40 ng µL<sup>−1</sup> total RNA.

| Designation | Gene | Cross Reactivity | Transition | Sensitivity |
|-------------|------|------------------|------------|-------------|
| ampR<sup>D135G</sup> | vanD | 80 nm | 90% | 90% |
| ampR<sup>G154R</sup> | vanD | 80 nm | 10% | 10% |
| ampR<sup>wt</sup> | vanD | 80 nm | 5% | 5% |

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$$D(c) = D_{\text{min}} + \frac{D_{\text{max}} - D_{\text{min}}}{\frac{1}{c^{b}} + \frac{1}{c}}$$  \hspace{1cm} (2)

$$D(c)$$ is equal to the deflection at a particular concentration c, $$D_{\text{min}}$$ corresponds to the minimum deflection and $$D_{\text{max}}$$ to the maximum deflection. It corresponds to the Hill coefficient at $$c_0$$ equal to half $$D_{\text{max}}$$. s is a control factor.
$D(c)$ is equal to the deflection at a particular concentration $c$, and $D_{eq}$ corresponds to the equilibrium deflection. The constant $k_a$ is the hybridization equilibrium constant.

A total of 100 fg total RNA at the LOD (1 fg $\mu$L$^{-1}$) was injected, which corresponds to a single bacterial cell. This finding demonstrates the ultrahigh sensitivity of our method, which will be beneficial in measurements from samples of body compartments with very low bacterial cell concentrations such as blood during sepsis or cerebrospinal fluid during meningitis.

### Table 2

Showing sequences of the four different oligonucleotides used for vancomycin resistance experiments. Capitalized letters indicate the differences in the vanB and in the vanD gene compared to the vanA sequence, whereby polyAC serves as reference.

| Designation | Sequence | Position range | $T_m$ |
|-------------|----------|----------------|-------|
| vanA        | tcaccccttaacgctaat | 1014 to 1032 | 27.5°C |
| vanB        | tcacctttaacgcgaat | 1011 to 1029 | 28.0°C |
| vanD        | ttacctccGtAaGttcGc | 1012 to 1032 | 27.9°C |
| polyAC      | acacacacacacacac | – | – |

*$T_m$ stands for melting temperature.

### Figure 3

Differential responses to injection of 1 pg $\mu$L$^{-1}$ total RNA. Differential measurements: a) Shown in red: difference of mutation D135G response and reference cantilever response. b) Shown in black: *P. aeruginosa* total RNA response from *ampR* wild type minus reference cantilever response. c) Shown in green: difference of mutation G154R response and reference cantilever response. Response magnitude is about $28 \pm 3$ nm at a concentration of 1 pg $\mu$L$^{-1}$. Current limit of detection (LOD) for total RNA of *P. aeruginosa* d) *ampR* (wild-type) and e) *ampR* (mutant) strains. The insets show the LOD of the measurement system of 10 fg $\mu$L$^{-1}$ as determined from the logistic fit. Cross sensitivity is less than 2 nm.
located inside the gene and not at its ends. The overhangs reduce the signal due to steric hindrance, but do not compromise binding to the cantilever.

Second, genes associated with vancomycin resistance were targeted. For detection of a gene, we intentionally designed the probe location in such a way that it is at the end of the gene, thus further improving sensitivity due to the fact that nanomechanical responses are larger the closer they occur relative to a surface. In particular, we demonstrated detection of vanD RNA at only 1 bacterial cell per milliliter of fluid with high specificity. A key advantage, in addition to the very high sensitivity and specificity, is the rapid response time of our system in the order of 5 min. The magnitude of the signals for the Langmuir plot and LOD analysis were obtained 20 min after the injection. In certain compartments of the patient only a few hundred bacterial cells per ml fluid may be present. A low detection limit generates the potential to diagnose bacteremia and meningitis.

Table 3. Cross-reactivity matrix. The responses of the sensors functionalized with a specific probe to different target samples is shown. The corresponding signals are in bold. A ten to 20-fold signal magnitude is observed compared to noncorresponding probes.

| Probe      | Injection of vanA | Injection of vanB | Injection of vanD |
|------------|-------------------|-------------------|-------------------|
| vanA probe | 75.3 ± 7.0 nm     | 5.6 ± 2.9 nm      | 2.3 ± 2.8 nm      |
| vanB probe | -2.9 ± 2.3 nm     | 75.4 ± 0.0 nm     | 4.4 ± 3.1 nm      |
| vanD probe | -3.6 ± 3.7 nm     | -4.6 ± 4.2 nm     | 55.0 ± 7.0 nm     |

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directly from patient specimen. Our method presents a novel way to detect antibiotic resistance mechanisms at unprecedented speed and sensitivity. The technique is label-free and allows nonamplified detection of resistance markers in total RNA from as little as 100 fg (corresponding to 1 bacterial cell) at a concentration of 1 fg $\mu$L$^{-1}$ for vancomycin resistance and 1 pg (corresponding to ten bacterial cells) at a concentration of 10 fg $\mu$L$^{-1}$ in SNP analysis of the ampR gene. These findings represent vast progress in rapid detection of antimicrobial resistance and open new avenues to investigate other infectious diseases or conditions where evolution of specific genes needs to be tracked at very low concentrations, e.g., sepsis, rare genetic variants. No significant influences were found so far in preliminary studies investigating bacterial resistance markers in blood based clinical samples to assess possible matrix background noise effects. This will show the feasibility of the method to investigate total bacterial RNA isolated from complex samples, such as blood.

4. Experimental Section

Clinical Isolates: The bacterial isolates (Pseudomonas aeruginosa and Enterococcus faecium) used were from the biobank of the division of Clinical Bacteriology and Mycology at the University Hospital Basel. Strains identification was confirmed using MALDI-TOF mass-spectrometry (Bruker). All strains were cultured on standard blood agar plates (5% sheep blood, bioMérieux) over night and a suspension with 0.5 McFarland was generated from which RNA was extracted using the RNeasy Mini Kit (Qiagen). Extracted RNA was quality controlled using Invitrogen Qubit 3.0 and Nanodrop 2000 (Thermo Fisher Scientific). Serial dilutions were prepared to determine the spectrum of sensitivity of the test system. All strains were sequenced using whole genome sequencing on a MiSeq (Illumina) platform at mean 30-fold coverage and 2 $\times$ 300 nucleotide paired ends to confirm the molecular resistance mechanism.[29] Sequencing information was used to determine specific probes for the SNP experiments. All genomes are available at NCBI GenBank (P. aeruginosa AE004091.2 and E. faecium AA9000003.1).

Nanosensor Array—Thiol Modified Oligonucleotide Probes: Thiol-modified Oligonucleotide probes were ordered from Microsynth AG (Balgach, Switzerland) at a concentration of 100 $\times$ $10^{-6}$ m without Dithiothreitol (DTT). Prior to functionalization oligonucleotides were diluted to a concentration of 40 $\times$ $10^{-6}$ m in HPLC grade water containing 50 $\times$ $10^{-3}$ m tri-ethyl-ammonium-acetate (TEAA) buffer and 1 $\times$ $10^{-3}$ m tris(2-carboxyethyl)phosphine (TCEP). TCEP was used to reduce disulphide bonds of thiolated oligonucleotide probes. Chemicals and buffers were purchased from Sigma-Aldrich.

Nanosensor Array—Nanosensor Array Preparation: Thinner and longer cantilevers yield higher sensitivity representing a major technical advantage compared to previous sensors. Before Au coating, the arrays were cleaned in Piranha solution (30% H$_2$O$_2$:96% H$_2$SO$_4$ = 1:1 v/v) for 10 min, rinsed three times with water followed by ethanol and dried in air. The upper sides of cantilevers were coated with a 3 nm-layer of Ti followed by a 20 nm thick Au layer. While extracting DNA/RNA, nanosensor arrays were functionalized with thiol-modified oligonucleotides at a concentration of 40 $\times$ $10^{-6}$ m using a modified ink-jet spotting system (MDP705L, Microdrop Technologies, Norderstedt, Germany).[30] Upon
dispensing and subsequent incubation for an hour at room temperature, the array was mounted in the measurement chamber containing 0.03 x SSC (saline sodium citrate buffer, prepared using 20 x SSC from Sigma Aldrich). Arrays can be functionalized individually (typically three specific probe sensors, two wild type thiol oligonucleotide probes and three specific reference probes). It is of vital importance that differential measurements are performed. External factors such as nonspecific interactions and thermal drift are eliminated by calculating differential responses of probe and reference sensors (in this study derivatized with a nonspecific oligonucleotide of the same length as the probe sequence or, in some experiments, with a wild-type sequence).

Total RNA was dissolved and serial dilutions were carried out in DEPC treated water (0.1% diethylpyrocarbonate). For the experiments total RNA was finally diluted in 0.03 x SSC. All measurements were performed on a home-built sensor instrument. The bending of cantilevers was detected by reflection of an external laser beam focused at the cantilever apex. The instrument enables monitoring the deflection of all eight sensors in parallel in a time-multiplexed manner. All measurements were performed at 28 °C in a temperature-controlled box with a steady buffer flow of 5 µL min⁻¹, which was found optimal for this type of experiment. Data acquisition hardware, temperature regulation, and a syringe pump for buffer and sample injection were controlled by dedicated LabView software.

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Conflict of Interest
The authors declare no conflict of interest.

Author Contributions
D.L. prepared the bacterial isolates and extracted the RNA. V.H. characterized the samples. D.W. analyzed the whole genomes. F.H., H.P.L., C.G., E.M., and A.E. conceived and designed the experiments. F.H. and H.P.L. prepared the nanosensor arrays. F.H. functionalized the sensors, performed the experiments and analyzed the data. F.H., H.P.L., C.G., E.M., and A.E. wrote the manuscript, critically advised and contributed in interpreting the results. A.E., E.M., and C.G. were in charge of the overall direction and planning of this study, and all the authors commented on the manuscript.

Keywords
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