**Low-cost integrated biosensing platform based on SiN nanophotonics for biomarker detection in urine**

D. Martens\(^{1,2}\), P. Ramirez-Priego\(^3\), M. S. Murib\(^{1,2}\), A.A. Elamin\(^4\), A.B. Gonzalez-Guerrero\(^3\), M. Stehr\(^4\), F. Jonas\(^4\), B. Anton\(^5\), N. Hlawatsch\(^5\), P. Soetaert\(^6\), R. Vos\(^7\), A. Stassen\(^7\), S. Severi\(^7\), W. Van Roy\(^7\), R. Bockstaele\(^6\), H. Becker\(^5\), M. Singh\(^4\), L.M. Lechuga\(^3\), P. Bienstman\(^{1,2}\)

\(^1\) Photonics Research Group, Ghent University/imec, Technologiepark-Zwijnaarde 15, B-9052 Ghent, Belgium

\(^2\) Center for Nano- and Biophotonics (NB-Photonics), Ghent University, Ghent, Belgium.

\(^3\) Nanobiosensors and Bioanalytical Applications Group, Catalan Institute of Nanoscience and Nanotechnology (ICN2), CSIC, BIST and CIBER-BBN, Campus UAB, Bellaterra, 08193 Barcelona, Spain

\(^4\) LIONEX Diagnostics and Therapeutics GmbH, Braunschweig, Germany

\(^5\) microfluidic ChipShop GmbH, Stockholmer Str. 20, D-07747 Jena, Germany

\(^6\) Trinean NV, Dulle-Grietlaan 17/3, 9050 Gent, Belgium.

\(^7\) imec, Kapeldreef 75, 3001 Leuven, Belgium

**Abstract**

We present a low-cost integrated nanophotonic lab-on-a-chip platform suitable for point-of-care (POC) biomarker analysis. The sensor chip included in the platform contains multiplexed Mach-Zehnder interferometers with an on-chip optical spectral analyser consisting of an arrayed-waveguide grating. The sensor chip is fabricated in silicon nitride material, which renders it compatible with consumer-electronics-grade sources and detectors, leading to the possibility of low-cost instrumentation. The nanophotonic sensor chip exhibits a detection limit of \(6 \times 10^{-6}\) RIU (Refractive Index Units), which is in the same order of magnitude as the reported values for the evanescent wave sensors at the state-of-the-art. The sensor chip is biofunctionalised with specific bioreceptors and integrated in a polymer microfluidics cartridge. The POC instrumentation platform contains the optical excitation and read-out sub-systems and a dedicated on-board software for a real-time analysis of patients samples. To demonstrate the versatility of the platform, we present results both on the detection of an antigen related to tuberculosis directly in urine samples using a laboratory prototype and on the detection of a protein biomarker (CRP) related to inflammation using the integrated instrument.

**Introduction**

Integrated evanescent field photonic biosensors are promising candidates for real-time qualitative and quantitative evaluation of biomolecular interactions. In these sensors, a change in the effective refractive index caused by the binding of an analyte to its specific bioreceptor, is converted into a measurable quantity, most commonly a resonant wavelength shift, through an integrated photonic transducer. Various types of sensors have been developed: Young and Mach-Zehnder (MZI) interferometers, bimodal sensors, ring resonators, photonic crystal based sensors and various others [1–5]. In general, evanescent wave sensors provide sensitivity, selectivity and versatility, which renders them applicable in a broad range of disciplines, ranging from medicine over food quality control to environmental management. Furthermore, they offer great potential as lab-on-a-chip devices, performing point-of-care measurements at an affordable cost.
While practical applications have emerged, in many cases the readout of these sensors requires either an expensive tunable laser or an off-chip spectrometer, making them expensive and/or slow. Therefore, the instrumentation cost is typically high, rendering the system unfit for true point-of-care applications. We aim to overcome this important barrier by using a broadband light source in combination with an on-chip spectral filter as interrogation mechanism [6-8]. The filter divides the transducer spectrum into different wavelength channels with spatially separated output ports. By fitting a curve to this dataset, the central wavelength of the original sensor spectrum is determined with an accuracy that surpasses the resolution of the spectral filter itself by multiple orders of magnitude.

The manuscript is structured as follows. First, we present how such sensor chips can be designed and fabricated using mass-fabrication-compatible technology. Before moving on to a fully integrated platform with a dedicated microfluidics and readout components, we first characterised the performance of the sensor chip in a lab-based prototype with in-house microfluidics. We determine the bulk limit-of-detection and then move further to perform biosensing experiments to detect *Mycobacterium tuberculosis* lipoarabinomannan (LAM) in clinical samples. After that, we move from a lab-based prototype to a fully integrated platform that is closer to commercialisation. We discuss how these silicon nitride based sensor chips can be integrated in a standard polymer fluidics cartridge, and describe in more detail the custom readout instrument and the associated software. Finally, this integrated platform is used to detect the C-reactive protein biomarker, related to inflammatory processes.

**Photonic sensor design**

The most important property for an integrated photonic sensor to enable use as a point-of-care diagnostic device is compatibility with a cheap read-out system, requiring no expensive components such as tunable lasers or off-chip spectrometers. In our design, the approach is to interrogate the sensor through an off-chip broadband light source, an on-chip spectral filter and an off-chip camera, all of which are affordable within a point-of-care context. This is especially true for the very near-infrared wavelength range, which can be used due to the choice of the sensor material. The spectral filter employed is an Arrayed Waveguide Grating (AWG), which is a state-of-the-art design in the given wavelength range (850 nm) in terms of footprint as well as performance [9].

As a sensor for this interrogation scheme, the Mach-Zehnder Interferometer (MZI) was selected, with one of the arms isolated from and the other exposed to the analyte. The main merit of this MZI configuration is its large design freedom, which enables tuning the sensitivity and the spectral period independently. The latter is relevant, as the characteristics of integrated spectral filters are typically restricted by fabrication limitations [10], and their ratio has a large influence on the final performance of the sensor [6]. The light interfacing with the chip occurs through grating couplers. At the input side, the grating couplers are flood-illuminated by a broadband light source [11]. This method allows interrogating multiple sensors at the same time, and as such opens the door to multiplexing. At the output side, the transmitted power of a large number of grating couplers is measured in parallel with a camera. A picture of the complete design of the photonic sensing circuit is shown in Fig. 1.
The transduction mechanism is based on the evanescent field: a change in the environment of the exposed MZI arm, such as a binding event or a bulk refractive index shift, will result in a change in the effective refractive index of the exposed MZI arm. This, in turn, results in a phase shift in this arm, which causes a spectral shift in the whole spectrum of the MZI. This shift is tracked through continuous monitoring of the locations of the spectral peaks of the MZI. To determine the peaks of this spectrum, we illuminate the chip using a broadband light source. The spectrum of the sensor (Fig. 2a) is sent through the different channels of the spectral filter (Fig. 2b), yielding a discretised version of the sensor spectrum (Fig. 2c). The resolution of this spectrum is typically low, as it is limited by the number of available data points per spectral period, determined by the channel spacing of the on-chip spectral filter. To compensate for this drawback, the original spectrum is reconstructed out of the discretised data through fitting (Fig. 2d), and the peak of this reconstructed spectrum is tracked. This greatly increases the accuracy on the obtained peak wavelength values. This procedure is more elaborately described in [6].
The whole sensing chip contains 6 sensors, each with their MZI and AWG, as well as 30 output grating couplers per sensor, one for each of the AWG channels. The entire design fits in a chip area of less than 6 mm x 6 mm (Fig. 3).

Sensors fabrication

Photonic integrated circuits (PICs) sensors were fabricated based on the CMOS compatible SiN photonics developed at imec, using wafer-level processing on standard 725 μm thick 200 mm Si wafers [10]. First a 70 nm thick SiN anti-reflective coating was deposited using plasma-enhanced chemical vapor deposition (PECVD) using SiH₄, N₂ and NH₃ at 400°C (Fig. 4). The purpose of this layer is to decrease the back-reflections coming from the oxide-Si interface, which would reduce the light...
incoupling efficiency of the grating couplers. Subsequently, a 2.3 μm thick PECVD SiO$_2$ layer, which had been specifically engineered to compensate the stress level of the wafers, was deposited using SiH$_4$ and N$_2$O. The roughness of the layer was reduced by applying a short chemical-mechanical polishing (CMP) step. On top of this oxide layer, a 220 nm thick PECVD SiN layer was deposited using SiH$_4$, N$_2$ and NH$_3$ at 400°C. This waveguide material has been optimised to minimise the light propagation loss in the visible wavelength range [10]. Refractive indices at $\lambda = 850$ nm were $n_{\text{SiO}_2} = 1.48$ and $n_{\text{SiN}} = 1.88$ as measured by ellipsometry.

Grating couplers were defined using 193 nm deep ultraviolet (DUV) lithography followed by a shallow etch of 110 nm depth into the SiN to maximise light coupling into the waveguide. A fluorine-based inductively coupled plasma reactive ion etching process was used to minimise sidewall roughness and achieve the required sidewall slope. Next, waveguides were defined by a separate 193 nm DUV litho step and similar etch, but now the full thickness of the SiN layer was etched. Following this process, a 2 μm layer of PECVD SiO$_2$ was deposited and planarised by CMP. Finally sensing regions were defined by 365 nm i-line lithography followed by a selective removal of the top oxide cladding using a dry etching process followed by a wet etching step.

![Figure 4: Process flow for the SiN photonic chip fabrication.](image)

**Evaluation of the bulk sensitivity**

Sensors were initially characterised through bulk sensing in a lab-based setup, with equipment compatible with point-of-care sensing. The bulk sensing limit-of-detection (LOD) provides an excellent figure-of-merit as it can be compared between various sensor types regardless of surface biofunctionalisation. To evaluate this value, the sensor is exposed to different concentrations of a solution (i.e.: HCl) in alternation with water. An example of a resulting sensorgram, where 5 sensors are measured in parallel, is shown in Fig. 5.
To determine the LOD, the following procedure was used. The shifts of the spectra are tracked, and the sensitivity of each sensor is calculated as the slope of the obtained spectral shifts as a function of the corresponding changes in the bulk refractive index. The minimum detectable wavelength shift is calculated as 3 times the standard deviation on the signal after transients have passed. The ratio of the minimum detectable wavelength shift to the sensitivity yields the LOD. This experiment was done multiple times, with different chips from different wafers. The set of experiments yielded an average LOD of $6 \times 10^6$ RIU, with a standard deviation of $2 \times 10^6$ RIU. This indicates a high reproducibility (given that we used different chips) as well as results competitive with most integrated photonic sensors, despite the low-cost read-out [6].

![Sensorgram of parallel measurement of five MZI sensors, where the sensors are exposed to different concentrations of HCl in alternation with water.](image)

**LAM detection in urine (lab setup)**

Before building an integrated readout system with a dedicated microfluidic cartridge, we first tested the functionality of the sensor chip by performing LAM measurements in clinical urine samples. LAM is a biomarker indicative of tuberculosis (TB) infection, and we correlated the measurements with the disease status of the participants.

To prepare the chips, anti-LAM IgM mAb molecules were covalently immobilised on the SiN chips using a silane route [12, 13]. Prior to any functionalisation, the SiN chips were cleaned in acetone, isopropanol, and ultrapure water. Next, the chips were exposed to oxygen plasma for one minute using a reactive-ion etching (RIE) system to terminate the surface with hydroxyl groups. Afterwards, the silanisation of the chips was carried out by liquid-phase deposition of silanes in an organic solvent [13]. The chips were placed in a 600 mM solution of (3-amino-propyl) tri-ethoxysilane (APTES, 99%; Sigma–Aldrich, Steinheim, Germany) in toluene (≥99.9%, Sigma–Aldrich) for 15 hours in a nitrogen-filled glove-box. To wash off unbound APTES molecules, the chips were rinsed with toluene, followed by tetrahydrofuran (THF, ≥99.9%, Sigma–Aldrich). After drying the chips under nitrogen flow, they were cured for 30 minutes at 120°C, resulting in an especially strong adhesion of the APTES layer to the surface [13]. This way, the surface becomes modified with amines, allowing for the subsequent reaction with succinic anhydride (SA) in order to yield a carboxyl-terminated surface [12, 14]. Succinic anhydride (SA, ≥99.9%, Sigma–Aldrich) was used as a cross-linker molecule to ensure the binding of the amino group of APTES to the NH$_2$ groups of the anti-LAM by providing carboxylic groups at the surface. For this purpose, the chips were placed overnight in a solution of 800 mM of SA in acetonitrile (ACN) (anhydrous, 99.8%, Sigma–Aldrich), after which they were rinsed with pure acetonitrile and
dried under nitrogen gas. The amine-terminated surface reacts with one of the two carbonyl groups of the SA molecules, causing the anhydride ring to open and to form a covalent bond with this carbonyl group. The other carbonyl group is released to form a free carboxylic acid-terminated surface [15]. The anti-LAM molecules were covalently coupled to the carboxyl-terminated SiN chips using the zero-length crosslinker 1-ethyl-3-3-dimethylaminopropyl-carbodiimide (EDC) in 2-[N-morpholine]-ethanesulphonic acid (MES) buffer at room temperature. In order to lower the effect of non-specific binding, the chips were blocked with 0.5 mg/mL PLL(20)-g[3.5]-PEG(5) (SuSoS) and 10 mg/mL BSA-A7906 (Sigma-Aldrich).

Fig. 6 shows the experimental setup used to perform the evaluations of urine samples. In this set-up, an external syringe pump is employed to pump fluids. A dual mode injector valve is used to load the samples to be tested in line with the neutral PBS buffer. During the experiment, first the buffer is flown to establish the baseline, then the clinical sample, followed by the buffer again to wash away any nonspecific binding. The sensor chip is squeezed under a home-mode PDMS microfluidic channel. The light source is a 850 nm SLED. The signal is collected via a CMOS camera, which is connected to the computer to analyse the detected signal.

A series of healthy and non-healthy urine samples were selected for experimental test. Apart from testing the lab prototype, the use of actual urine samples also allows us to evaluate the effect of nonspecific binding. The samples were pumped in the microflow cell at a rate of 10 µL/min for 20 minutes. Results from four sensors on the chip were analysed.
Fig. 7 shows the real-time interaction of healthy and non-healthy urine samples with the anti-LAM molecules immobilised on the SiN sensors. First, 1x PBS (pH = 7) was administered to the sensors. After stabilisation of the signal, 200 μl of a healthy or non-healthy urine sample was injected onto the sensors.

In the case of healthy samples, initially a high spectral shift was encountered due to buffer exchange. This high spectral shift was only temporary and started to return back after a while since no LAM/anti-LAM binding was occurring. When 1x PBS was injected afterwards, the spectral shift moved back to a very low value (220 pm) as compared to the initial shifted value. The spectral shift did not completely return back to its starting value and this can be attributed to nonspecific binding of residuals present in the urine samples to the sensors surface.

In the case of non-healthy samples, a high increase in the spectral shift was encountered due to buffer effect. However, afterwards, when the 1x PBS was injected, a high permanent spectral shift compared to its initial value was encountered. This is due to the fact that LAM was present in the urine and it efficiently bound to the anti-LAM immobilised on the sensor surface. In Fig. 7b, the non-healthy sample caused a permanent spectral shift of about 6 nm.

<table>
<thead>
<tr>
<th>Sample Type</th>
<th>Spectral Shift</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy Sample # 1</td>
<td>0.75 nm</td>
</tr>
<tr>
<td>Healthy Sample # 2</td>
<td>0.85 nm</td>
</tr>
<tr>
<td>Healthy Sample # 3</td>
<td>0.22 nm</td>
</tr>
<tr>
<td>Healthy Sample # 4</td>
<td>0.50 nm</td>
</tr>
<tr>
<td>Non-Healthy Sample # 1</td>
<td>6.00 nm</td>
</tr>
<tr>
<td>Non-Healthy Sample # 2</td>
<td>5.50 nm</td>
</tr>
<tr>
<td>Non-Healthy Sample # 3</td>
<td>7.00 nm</td>
</tr>
<tr>
<td>Non-Healthy Sample # 4</td>
<td>4.00 nm</td>
</tr>
</tbody>
</table>

Table 1: List of the spectral shift due to series of healthy and non-healthy samples.

A series of four healthy urine samples and four non-healthy urine samples were measured. The results are summarised in Table 1. A maximum of 0.85 nm in permanent spectral shift was obtained from healthy samples. The non-healthy samples caused a minimum permanent spectral shift of 4 nm and a maximum permanent spectral shift of 7 nm, which depends on the concentration of the LAM present.
in the urine samples. The permanent spectral shift obtained due to non-healthy samples is very high compared to the permanent spectral shift obtained due to the healthy samples. Moreover, as can be noticed from Fig. 7a and 7b, the shape of the sensorgrams in both cases is also very different: while the sample is still being injected, the spectral shift already starts to decrease for the healthy sample, which is not the case for the non-healthy sample.

Finally, the sensorgrams also show that an assay time of 10 to 15 minutes is feasible.

Microfluidics cartridge

Microfluidic cartridge bodies were injection-molded in COC (cyclo-olefin-polymer, tradename Topas). This allows for a cost-effective large-scale production, so that the cartridges can be made cheap enough to be disposable.

For the integrated cartridge design, an injection molding platform of a microscopy slide format with footprint of 75.5mm x 25.5mm was chosen. This format provides an easy to handle size on which all relevant fluidic and detection functionalities were accommodated, allowing for a high integration density with various functional elements for such a disposable cartridge.

A photograph of a microfluidic cartridge with the integrated photonic chip is depicted in Fig. 8.

For the inlet of the urine sample and required reagents, the integrated cartridge includes female Luer interfaces being compatible with standard syringes, as commonly used.

An integrated filter ensures the retention of coarse components from the urine sample and an integrated air bubble trap retains undesirable air bubbles. At the end of the fluidic path, a reservoir for easy liquid waste handling is placed, followed by an air vent to allow air to escape from the cartridge when the waste reservoir fills up.

With respect to the integration of the sensor chip, there is an appropriately designed recess on the top, while the rest of the cartridge is designed to leave parts of the chip exposed for easy optical access. Indeed, light is coupled into and out of the chip through grating couplers, and these need to stay exposed.
Fig. 9 shows an intersection through the sensor area of the microfluidic cartridge. Note that the photonic chip is positioned upside down and that the optical detection is performed from the bottom side.

**Integrated readout instrument**

The realised readout instrument is shown in Fig. 10. The POC platform includes the optical readout, the pumping unit interfacing with the cartridge, and a touch screen with the user interface. The size of the instrument is determined by the choice of using vertically oriented syringes to load the samples and the buffers.
When designing the concept of this instrument, importance was given to flexibility, the possibility to test alternatives to specific parts, ease of assembly, ease of shipment and system installation at test sites. Likewise it should be noted that – although the readout architecture is selected to be low-cost - the prototype instrument design is also not yet optimised for cost. For example, the integration of the full Windows-PC including a high-res touch screen certainly adds up to the cost of the system, but also offers flexibility to update software in a late stage of the development (testing) cycle.

The optical readout unit (Fig. 11) contains the light source as well as the light sensor to interrogate the optical biosensor chip. The unit is mounted upside down in the bottom part of the instrument. The light source (SLED) and the detector (CMOS sensor) are both commercial off-the-shelf components, coupled to a custom lens system.
Three syringe pumps are integrated in the readout instrument (Fig. 12). Each pump has a stepper motor directly driving a piston up and down. The piston pushes the plunger of the syringe downwards, transferring the contents of the syringe (buffer, sample, …) to cartridge. As the distance between the syringes is small, the piston axis is not in line with the stepper motor axis, but parallel. A dry bearing system is used to guide the piston up and down. The piston also includes a pressure sensor, to detect mechanical contact between the piston head and the syringe plunger. To increase confidence in the measurements, an extra flow monitoring system was also installed, consisting of a separate camera system to image the fluidic channel to perform bubble detection.
Custom software was written to reliably identify the position of the output grating couplers in the images captured by the CMOS sensor. The intensities of these spots are then used in a fitting procedure, which allows us to track the position of the peak wavelength as a function of time.

Fig. 13 shows a simple proof-of-concept sensing experiment on the instrument using the actual microfluidic cartridge, which measures the physical adsorption of BSA on an unfunctionalised sensor chip.

**Biosensing with the integrated readout instrument**

Finally, this setup was used to perform biosensor experiments using functionalised chips. As a model system to test the prototype, the immnosensing of the C-reactive protein (CRP) biomarker was
chosen. CRP is a widely used biomarker in patients that plays an essential role as an early indicator of inflammation and infection, and is also used as an indicator of prognosis for cardiovascular diseases [16]. CRP concentration in serum in healthy conditions is lower than 10 μg/mL and it can increase up to 200 μg/mL in case of inflammation or infections. CRP can be detected through the interaction with its specific antibody.

Monoclonal antibody anti-human C-reactive protein C7 produced in mouse and human CRP was purchased from HyTest (Finland) and BBI Solutions (UK), respectively. All chemical reagents were purchased from Sigma-Aldrich (Germany).

A concentration of 50 μg/mL of anti-CRP in phosphate buffer saline (PBS; 10 mM phosphate, 137 mM NaCl, 2.7 mM KCl, pH 7.4) was immobilised onto the surface of the chip through physical adsorption, followed by a 10 mg/mL of bovine serum albumin solution (BSA) to block the free sensor surface areas. Milli-Q water (Millipore, US) was used as running buffer for the immobilisation and blocking steps. For the CRP protein detection, the running buffer was changed to PBS to ensure the optimal conditions for the immunosensing interaction, as well as to maximise the stability of the bioreceptor layer.

In a physical immobilisation strategy, it is not possible to remove the bound CRP protein to the antibody due to its weak attachment to the sensor surface and thus an additive immunosensing assay was employed to establish the analytical calibration curve of the CRP detection. To study the biointeraction between CRP and its specific antibody, successive dilutions of increasing CRP concentrations (50, 100, 250, 500 and 750 ng/mL) were flowed over the sensor area. A direct relationship between the concentration and the sensor response can be observed (Fig. 14).

Furthermore, the specificity of the immunoassay was analysed by injecting a control of 1000 ng/mL of BSA instead of CRP, resulting in a negligible response of the sensor, as shown also in Fig. 10.

As shown in Fig. 15, a detection curve was obtained by plotting the accumulated response signal as a function of the accumulated CRP concentration. The limit of detection (LOD) was determined from the calibration curve as three times the standard deviation of the signal. The resulting LOD was 19.478 ng/mL ($R^2 = 0.9701$) which theoretically corresponds to the lowest detectable CRP concentration. These results indicate that the integrated instrument and the designed photonic sensor could be used as an effective biosensor for CRP detection as the concentration of this biomarker when related to infection or disease is much higher than the one obtained.
Conclusion

We presented an integrated sensor platform that allows for low-cost detection of various biomarkers. The sensor chip with integrated spectral filter eliminates the need for an expensive tunable laser, such that a cheaper broad-band light source can be used. A technology to integrate this chip into a microfluidic cartridge was presented, together with the development of a dedicated readout instrument. We presented proof-of-concept results for detection of LAM in actual clinical urine samples as well as for the detection of CRP. These results show the feasibility of using this system as a promising platform technology to detect different kinds of biomarkers using a cheap readout instrument.

Acknowledgments: This work was funded by the European Commission under the FP7 project Pocket (grant 610389). The nanoB2A is a consolidated research group (Grup de Recerca) of the Generalitat de Catalunya and has support from the Departament d’Universitats, Recerca i Societat de la Informació de la Generalitat de Catalunya (2014 SGR 624). The ICN2 is funded by the CERCA programme/Generalitat de Catalunya. The ICN2 is supported by the Severo Ochoa programme of the Spanish Ministry of Economy, Industry and Competitiveness (MINECO, Grant no. SEV-2013-0295).

References


