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ORIGINAL ARTICLE

Biofunctionalized all-polymer photonic lab on a chip with integrated solid-state light emitter

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A photonic lab on a chip (PhLOC), comprising a solid-state light emitter (SSLE) aligned with a biofunctionalized optofluidic multiple internal reflection (MIR) system, is presented. The SSLE is obtained by filling a microfluidic structure with a phenyltrimethoxysilane (PhTMOS) aqueous *sol* solution containing a fluorophore organic dye. After curing, the resulting xerogel solid structure retains the emitting properties of the fluorophore, which is evenly distributed in the xerogel matrix. Photostability studies demonstrate that after a total dose (at λ =365 nm) greater than 24 J cm⁻², the xerogel emission decay is only 4.1%. To re-direct the emitted light, the SSLE includes two sets of air mirrors that surround the xerogel. Emission mapping of the SSLE demonstrates that alignment variations of 150 µm (between the SSLE and the external pumping light source) provide fluctuations in emitted light smaller than 5%. After this verification, the SSLE is monolithically implemented with a MIR, forming the PhLOC. Its performance is assessed by measuring quinolone yellow, obtaining a limit of detection (LOD) of (0.60±0.01) µM. Finally, the MIR is selectively biofunctionalized with horseradish peroxidase (HRP) for the detection of hydrogen peroxide (H₂O₂) target analyte, obtaining a LOD of (0.7±0.1) µM for H₂O₂, confirming, for the first time, that solid-state xerogel-based emitters can be massively implemented in biofunctionalized PhLOCs.

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INTRODUCTION

Since the introduction of the µTAS concept by Manz and co-workers in 1990,¹ there has been a steady development of this research line, from the definition of a wide range of architectures and systems² to the application of new materials^{3,4} and the implementation of versatile production technologies.⁵ The reduction of size resulted not only in a decrease in the analysis time and reagent consumption, but also in an enhancement of the sensitivity and possibility of parallelizing the analysis without dramatically increasing both the complexity and overall size of the system. Then, the µTAS concept was further expanded to the lab on a chip (LOC),⁶ where all (or most of) the required steps in an analytical protocol can be pursued. These systems have increasingly been applied in many different fields such as drug discovery and development,⁷ genomics,⁸ clinical diagnosis⁹ and cellomics.¹⁰ In most of these examples, an inexpensive, disposable system with superior performance in terms of sensitivity to a specific analyte is required. Reduction of fabrication costs is generally associated with the use of polymeric materials and simple fabrication techniques, among which poly(dimethylsiloxane) (PDMS) and soft lithography¹¹ are excellent examples, respectively. PDMS has widely been applied in the development of LOCs due to its fabrication simplicity. This elastomeric mater-

ial can be easily patterned by soft lithography. Furthermore, PDMS is mechanically robust, exhibits a low Young's modulus (300-800 kPa), exhibits a low refractive index (n=1.41) and is transparent in the UV-NIR range. However, PDMS is also associated with some important drawbacks when being applied for biochemical analysis that are related to its hydrophobic character, which makes the material very prone to serious nonspecific absorption processes of different organic molecules and biomolecules. Here, several surface modification processes have been reported, such as oxygen plasma¹² or UV/ozone treatments,¹³ which produce silanol groups (Si-OH) on the material surface and allow tuning of its hydrophilic/hydrophobic balance. These groups are dynamic, and the surface recovers its hydrophobicity with time. Therefore, subsequent chemical processes must be readily applied to introduce other functional groups to which different (bio)chemical compounds can be firmly attached.¹² In a different approach, hvdroxyl (-OH) groups are introduced on the PDMS surface by physisorption of polyvinyl alcohol (PVA) polymer¹⁴ and further silanization¹⁵ of the resulting surface. This process allows the introduction of functional chemical groups that enables the eventual covalent attachment of biomolecules, 16-18 with the final aim of implementing a microbioreactor (understood as a biofunctionalized microfluidic element) in the LOC

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that provides the desired selectivity for a specific analyte while minimizing interference due to non-specific adsorption of other species.

Equally important is defining the most appropriate transducer to transform this biorecognition event into a quantifiable signal. Here, different transduction mechanisms, such as electrochemical,¹⁹ magnetic²⁰ and optic/photonic,²¹ have been integrated into LOCs. Among these options, optical transducers have been demonstrated to be one of the most sensitive analytical and selective detection methods in cell biology.²² Transference of this know-how from the macro- to the microscale has already been demonstrated with the so-called photonic lab on a chip (PhLOC).²³ Micro-optical elements, such as microlenses and mirrors,²⁴ can be implemented in the vicinity of a microfluidic network to provide the entire system with the required optical readout. Here, an open issue is to provide a mechanically robust, optically efficient and technologically simple method that allows light incoupling. Alignment between the light source and optical system is by far the most critical aspect in the PhLOC because small variations (less than 5%) of their relative positions cause unacceptable light coupling variations. This key aspect likely hampers the massive implementation of the PhLOC concept and restricts its application to research labs, where the alignment can accurately be controlled with the help of micropositioners. One first step toward tackling this issue is to define self-alignment microchannels for the clamping and positioning of fiber optics.²⁵ Although effective, this approach requires trained users and special tools to define high-quality fiber optic facets, not to mention their inherent fragility. The optimal configuration would be to have a disposable light source directly implemented (and hence aligned) in the PhLOC. Here, two different strategies could be pursued, namely, either heterogeneous or monolithic integration. In the first case, the final system is obtained by the combination of several elements defined in different substrates. An example could be the integration of (organic) light-emitting diodes, (O)LEDs,²⁶ or laser dyes.²⁷ Despite their good performance, such devices are normally multilayered, requiring several clean-room fabrication steps. In addition, the alignment issue is not solved, resulting in limited fabrication yield and a dramatic increase of the production costs. Conversely, monolithic integration provides inherent aligned structures because of the accurate positioning of the different elements at the photomask level. Despite being a more elegant technology compared with heterogeneous integration, the challenges associated with this approach are much greater. Silicon substrates have proven to be unsuitable for defining optical elements working in the visible range: the high absorbance in the UV-Vis range limits their application to photonics working in the near infrared (above 1100 nm), whereas the indirect bandgap hampers their use for implementing light sources. Hybrid monolithic integration can be an elegant alternative, as demonstrated by Balslev and co-workers,²¹ who defined a complete PhLOC, whereby a light source was integrated using a liquid dye laser and waveguides and fluidic microchannels were defined with a photostructurable polymer. Nevertheless, liquid dyes are extremely toxic, and thus, their combination with life science applications is not trivial. This issue is especially evident when using PDMS. The integration of liquid dyes with PDMS proves to be difficult, as this material suffers from severe swelling against most organic solvents and is permeable to both gases and small molecules (such as dyes). Therefore, alternatives to liquidbased light sources must be sought.

Solid-state light emitters (SSLEs), which are mechanically and chemically stable, could be an alternative to liquid dyes. Here, different materials, such as hybrid organic-inorganic polymers, have proven to be good candidates.²⁸ Among the candidates, those fabricated using

sol-gel technology consist of solid backbone networks into which active chemical species can be easily incorporated²⁹ while preserving their (bio)functionality. The optical properties of such materials (refractive index and spectral response) can be tailored as a function of the nature of the different monomers and their ratio.³⁰ Moreover, the material processing can be tuned to be compatible with microfabrication techniques, and micro- and nanostructures can be defined with smooth and crack-free surfaces. This last point becomes essential for the fabrication of high-quality photonic elements and systems.³¹ Not surprisingly, sol-gel technology has already been used to define waveguides³² and microlenses;³³ however, to the best of our knowledge, this approach has not been used to implement sol–gel-based light emitters in LOC devices.

In this work, we present the hybrid monolithic integration of a xerogel-based SSLE and an optofluidic multiple internal reflection (MIR) to produce a PhLOC. Although having an advanced configuration, the entire PhLOC only requires a single mask and single substrate technology. To verify the proposed approach, Atto 390 fluorophore has been used to define the SSLE, whereas either non-functionalized or enzymatically functionalized MIR has been implemented. This is the first time, to the best of our knowledge, that feasible integration of this type of light source and its successful performance for bioanalytical applications has been presented using an all-polymer technology.

MATERIALS AND METHODS

Chemicals

A PDMS Sylgard 184 elastomer kit was purchased from Dow Corning (Midland, MI, USA) and used according to the datasheet. SU-8 2050 and PGMEA (propylene glycol methyl ether acetate) were acquired from Microresist (Berlin, Germany). The 99% PVA, 99% triethylamine, Tween 20, phenyltrimethoxysilane (PhTMOS), dimethyl sulfoxide, horseradish peroxidase (HRP type VI), 2,2'azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) (ABTS), and quinoline yellow dye were purchased from Sigma-Aldrich Co. (St Louis, MO, USA). The 90% 11-triethoxysilyl undecanal was purchased from ABCR GmbH & Co. KG (Karlsruhe, Germany). The Atto 390 fluorophore (λ_{abs} =390 nm; λ_{fl} =479 nm, in water) was purchased from Atto-Tec GmbH (Siegen, Germany).

Design-I: SSLE

Figure 1a shows the SSLE design, which consists of a pear-shaped microfluidic element (height: 230 µm, total volume: 2.12 µL), two diamond-shaped internal reservoirs and inlet/outlet ports. A set of two air mirrors was also defined to enhance the efficiency of the coupling to an arbitrary optofluidic system (also included in the PhLOC, as discussed below). The SSLE concept is as follows. Assuming that the microfluidic element is filled with a fluorophore (either in liquid or solid phase), when externally pumped with the appropriate wavelength, light will be emitted in 4π steradians (Sr). This behavior is suboptimal if the idea is to couple as much emitted light as possible in an optofluidic system close to the SSLE. Thus, the role of the two air mirrors is twofold. First, the mirrors allow multiple interactions between the excitation wavelength and the SSLE, thus enhancing the photonic conversion. Second, the mirrors redirect the emission towards the SSLE tip, allowing a more efficient light coupling to the optofluidic system. The working principle of the air mirrors is illustrated in Figure 1b. Considering the refractive indices of PDMS (n=1.41) and air (n=1.00), if the light reaches the flat air mirror #1 at an angle higher than the critical angle ($\theta_1 > \theta_c = 45.17^\circ$), the light undergoes total internal reflection and will be redirected back to the SSLE.



Figure 1 (a) Design of the SSLE using PDMS (gray) as the structural material, fluorophore-doped xerogel (blue), and air (white) for defining the micro-optics elements. (b) Scheme of the working principle of the two air mirrors, considering incidence angles larger (θ_1, θ'_1) and smaller (θ_2) than the critical angle (θ_c) . (c) Ray tracing of the light emitted from the SSLE and its focusing at the tip. (d) Design of the PhLOC, including the SSLE, the MIR, the interconnecting waveguide, a biconvex microlens and a self-alignment microchannel for inserting the fiber optics readout. The optical path is shown in red. PDMS, poly(dimethylsiloxane); PhLOC, photonic lab on a chip; SSLE, solid-state light emitter.

Smaller angles $(\theta_2 < \theta_c)$ are transmitted through the first mirror according to the Fresnel laws, reaching the teeth-shaped mirror #2. The faces of each tooth in mirror #2 work jointly to reflect angles transmitted through mirror #1. One of the faces is tilted to ensure total internal reflection at the light incidence point $(\theta'_1 > \theta_c)$, whereas the second face back-reflects the light into the SSLE. In addition, the relative angle of each tooth varies radially along the SSLE to focus the light towards its tip, as observed in the ray-tracing simulation of the teeth-shaped mirrors working as a whole depicted in Figure 1c.

Design-II: PhLOC

The disposable PDMS-based PhLOC is illustrated in Figure 1d and consists of the previously described SSLE directly aligned to a MIR

optofluidic system²³ interconnected with the help of a waveguide. Both the MIR and waveguide make use of air mirrors whose working principle is identical to that of mirror #1, the single difference being that the light propagation inside the MIR follows a zigzag path aided by focusing air mirrors (allowing the optical path to be enlarged and thus enhancing its sensitivity without a dramatic increase of the required sample volume).²⁴ Flat air mirrors are located at both sides of the bulk PDMS, forming an interconnecting waveguide between the SSLE and MIR. Finally, transmitted light is collected at the end of the MIR with the help of two PDMS microlenses (implemented using only air and PDMS as constituent materials) and fiber optics positioned in a self-alignment microchannel.

Fabrication

The PhLOC was fabricated using PDMS and soda-lime glass substrates. The process comprised three main steps, mainly the definition of the master using a photopatternable polymer, replica molding of the master using PDMS, and implementation of the SSLE using a fluorophore-doped organic-inorganic polymer by sol-gel technology.³⁴ The first two steps make use of standard techniques, and a detailed description can be found elsewhere.³⁵ Briefly, a 230-µm-thick layer of SU-8 2050 was spun over a substrate. After the soft bake, it was exposed to UV light using the appropriate mask. Then, a postexposure bake was performed to crosslink the polymeric structure. Once finished, the wafers were immersed in PGMEA and rinsed vigorously until fully developed. To enhance the mechanical robustness of the masters and to heal possible microcracks generated by internal stresses, a hard bake was performed for 2 h at 120 °C in an inert atmosphere. PDMS was prepared by mixing the base elastomer and curing agent in the standard ratio of 1:10 (ν/ν). Vigorous stirring and applied vacuum assured a bubble-free pre-polymer. After pouring the PDMS onto the master, a second vacuum step was applied to ensure that no air bubbles remained entrapped between the master and PDMS. The PDMS was cured at 80 °C for 20 min. and then peeled off using tweezers. At this point, all the fluidic ports were opened using a dispenser tip ($\varphi 0.5$ mm). Then, the PDMS and a flat pre-cleaned glass substrate were activated with oxygen plasma (PVA TePla AG, Munich, Germany) at 500 W and 0.8 mbar for 18 s. Afterwards, they were placed in contact using deionized water as a lubricant. By thermal processing at 80 °C for 30 min, both the PDMS and glass substrate were irreversibly sealed. Once the bonding quality was verified by eye inspection, the SSLE was ready to be filled with the fluorophore-doped pre-polymerization sol solution, which is an adaptation of the micromolding in capillaries soft-lithographic technique.¹¹

Among the myriad different fluorophores suitable for this application, Atto 390 was selected because of its high molecular absorption (24.000 cm⁻¹ M⁻¹) and quantum yield (0.90) as well as its large Stokes shift (λ_{abs} =390 nm, λ_{fl} =479 nm, in water). Atto 390 is slightly hydrophilic and is soluble in polar solvents such as DMF or dimethyl sulfoxide. Solubility was important to ensure compatibility with different chemical environments, which is crucial to obtain a homogeneous dispersion of the fluorophore molecules in the polymeric material. Nevertheless, the technology presented here is not restricted to Atto 390; other fluorophores or dyes with similar properties could also be used.

The fluorophore-doped hybrid organic-inorganic polymer was synthesized using sol-gel technology.³⁴ The process involved three main steps: mixing, gelation and drying. During the mixing step, the pre-polymerization sol solution was prepared by vigorously mixing 500 µL of PhTMOS monomer with 300 µL of a 500 µM Atto 390 solution prepared in H₂O/dimethyl sulfoxide 5:1 (ν/ν) pH 3 (adjusted with diluted HCl). During this step, the monomer was hydrolyzed and started to condense, forming siloxane bonds. An acidic medium was needed to accelerate and favor the hydrolysis of the PhTMOS monomer. At this point, the sol solution was formed. Because the sol is a lowviscosity liquid, filling of the SSLE fluidic structure was achieved by capillary forces, thus avoiding the use of external pumps. Once this microstructure was filled, it was protected from light and left undisturbed to allow the polymer to dry at room temperature for 72 h. During gelation, the viscosity increased sharply, and the solvents in the solution evaporated and diffused through the PDMS stamp. Consequently, the final xerogel had a significantly smaller volume compared with the sol solution. When not properly addressed, this effect led to a partial emptying of the SSLE. In our case, the diamondshaped internal reservoirs assured additional material supply during the gelification step, thus impeding the SSLE emptying. Once the solgel polymeric material dried, a xerogel matrix with entrapped Atto 390 fluorophore was obtained.

Fluorophore-doped polymer characterization

Fluorophore homogeneous dispersion in the SSLE is of key importance for obtaining a PhLOC with identical performance. In this context, confocal microscopy analysis was performed using a Leica TCS SP2 confocal microscope (Leica Geosystems AG, St Gallen, Switzerland) including an HC PL APO CS $10.0 \times$ UV objective with a 0.40 numerical aperture and a UV 405 nm diode laser as a light source. The scanned area was $600 \ \mu\text{m} \times 600 \ \mu\text{m}$ (width×length), and images were captured every 5 μ m in depth with a resolution of 8 bits. Up to 50 pictures were taken to cover all the SSLE thickness.

Photostability

The SSLE photostability was analyzed by irradiation of the structure with UV light at controlled irradiation doses while acquiring emission spectra. The recorded spectra were then compared with the reference spectrum, and the percentages of the initial emission intensity at the maximum were calculated. The set-up for this study included a mask aligner as a light source (a MA1006 contact mask aligner equipped with a 350-W mercury lamp, 2-µm resolution and a split field microscope with 5× and 10× objectives, exposure dose of 9 mJ cm⁻² at λ =365 nm (i-line); Suss MicroTec, Munich, Germany) and a 230µm-diameter multimode optical fiber (Thorlabs Inc., Dachau, Germany) for collecting the light, coupled to a microspectrometer (QE 65000-FL; Ocean Optics, Dunedin, FL, USA). Each spectrum was acquired with an average of 10 scans, and the integration time was fixed at 100 ms. Because the power density is constant in the mask aligner, different irradiation doses were achieved by irradiating the sample at 21 different time intervals between 10 and 2700 s (equivalently, to optical doses between 90 mJ cm⁻² and 24.3 J cm⁻²).

Mapping emission

The optimal excitation point and alignment tolerances at which the emitted light intensity has its maximum value at the SSLE tip are key in validating the emitter configuration proposed in this paper. To this effect, the SSLE was also fabricated in a separate substrate, consisting of only the pear-shaped structure, the two air mirrors, the interconnection waveguide, the biconvex microlens and the output self-alignment system (a schema of the structure is provided in Supplementary Fig. S1). Once the doped xerogel was obtained inside the SSLE, a blue laser (λ =405 nm, Laser module NANO 250-532-100; Linos Photonics, Eschbach, Germany) was coupled to a multimode 230 µm optical fiber, which was positioned on a motorized XY stage (VT-80, PiMicos, Eschbach, Germany, Thorlabs, Dachau/Munich, Germany) perpendicular to the plane of the SSLE at a fixed distance of z=1 mm. The readout comprised an identical optical fiber inserted into the output self-alignment microchannel, coupled to a microspectrometer (QE 65000-FL; Ocean Optics, Dunedin, FL, USA). This configuration allowed geometric localization of the spots in the SSLE and simultaneous acquisition of the spectral response. Each spectrum was acquired with an average of 10 scans, and the integration time was fixed at 250 ms. This process resulted in an efficiency map of the SSLE emission as a function of the excitation position with a resolution of 150 µm. The study was repeated with four SSLE structures with different Atto concentrations (10, 50, 100 and 500 μ M).

Biofunctionalization

The immobilization of HRP as a biorecognition element on the PDMS walls of the MIR structure was performed following the protocol described in Ibarlucea *et al.*¹⁶ Briefly, the inner walls of the MIR were modified by direct adsorption of PVA linear polymer. This process introduced hydroxyl groups onto the PDMS surface, which were further silanized using the 11-triethoxysilyl undecanal silane containing an aldehyde end-group.³⁶ The covalent interaction between this group and a primary amine moiety of the protein structure enabled the selective robust attachment of the HRP enzyme on the PDMS surface.

Experimental set-up for the PhLOC and protocol for absorbance measurements

Light emitted from the blue laser (λ =405 nm, Laser module NANO 250-532-100; Linos Photonics) was coupled with a multimode 230 µm optical fiber, which was positioned at the maximum emission region of the SSLE, in accordance with the previously described mapping study. The readout comprised an identical optical fiber, which was inserted into the output self-alignment microchannel of the MIR system and connected to the microspectrometer (QE 65000-FL; Ocean Optics, Dunedin, FL, USA). The acquisition time was fixed at 400 ms, and each recorded spectrum was the average of 10 scans.

The performance of the PhLOC was first tested by sequentially injecting quinoline yellow dye aqueous solutions in a concentration range from 1 to 25 μ M. The readout was recorded immediately after filling the PhLOC, and deionized water was used as a reference.

For final PhLOC validation, biofunctionalized MIRs were tested by sequentially injecting 30 μL of 0.1 M acetate buffer pH 5.5 solutions containing 0.5 mM ABTS and increasing concentrations of H_2O_2 in the range from 1.4 μM to 27.5 μM . The readout signal of the acetate buffer electrolyte solution was used as the reference. The measurements were made under quiescent fluid conditions (absence of flow) in all cases. The generation of the colored product was time dependent because a delay time was required for the catalytic reaction to occur and the resulting enzymatic product to diffuse to the bulk of the solution inside the MIR. A 10-min stabilization time was selected. Reproducibility studies were performed by calibrating three different biofunctionalized PhLOCs on three consecutive days.

The results of both analytical studies described above were analyzed following the protocol presented in Ref. 35. Once the corresponding absorbance *vs.* concentration calibration curves were obtained, the limit of detection (LOD) could be estimated following the 3σ IUPAC criteria.³⁷

RESULTS AND DISCUSSION

Fluorophore-doped polymeric SSLE

The results from the confocal microscopy lambda analysis are presented in Figure 2. Thirteen different emission spectra at different arbitrary points of the microstructure were recorded; the mean values for the maximum fluorescence emission intensity and wavelength were (219 ± 6) a.u. (3% coefficient of variation, number of spectra=13) and (451 ± 1) nm, respectively. These results indicate that the fluorophore was homogeneously dispersed in the polymeric material.

A 3D reconstruction of the SSLE emitted light intensity was performed, the projection of which is presented in Figure 2b. As observed, the fluorescence emission is constant across the structure, which means the doped xerogel completely filled the SSLE PDMS structure. This effect can again be associated with the homogeneous fluorophore distribution on a xerogel SSLE with a constant thickness.



Figure 2 (a) Emission spectra recorded at 13 different points of the SSLE. (b) Image of the projection of the confocal images recorded every 5 μ m across the entire SSLE structure. SSLE, solid-state light emitter.

Photostability studies of the SSLE

The photostability study revealed that at low exposure times/radiation doses, fluctuations in the intensity signal occurred. The signal was stabilized after 15 min of total exposition time (Supplementary Fig. S2). With a total dose of 24.3 J cm⁻², the decay in emission was only 4.1% of the initial value. These results corroborate the high photostability of Atto 390 and straightforwardly, of the SSLE presented here.

Mapping studies

The resulting map of maximum emitted intensities at 450 nm as a function of the coordinates of the external pumping light source spot is presented in the image capture shown in Figure 3a, in which the outline of the pear-shaped structure is indicated with a discontinuous line. As predicted by the numerical simulations, a symmetrical response was observed, with the maximum of coupled light obtained when exciting along the central longitudinal axis. The output intensity measured at 450 nm as a function of the Atto fluorophore concentration is shown in Figure 3b. The maximum intensity values increase with the fluorophore concentration. The linear trend observed confirms that autoabsorption phenomena did not occur at the concentrations used in this work.

The positioning tolerance of the external pumping light source over the SSLE is also of significant importance when considering integ-



Figure 3 (a) Experimental emission intensity map study performed with the SSLE prepared with the 500 μ M Atto 390 solution. The outline of the pear-shaped structure is indicated by the discontinuous line for better visualization. (b) Maximum emitted intensity recorded at the optimal excitation point (highest emission) vs. Atto concentration. SSLE, solid-state light emitter.

ration of the PhLOC into a portable reading platform or into a future mass production system. To this effect, we have defined the longitudinal and transversal confidence positioning interval (η_L and η_T , respectively), the distance over which the external pumping light source spot can be moved while maintaining the output intensity at higher than 95% of the maximum SSLE emission intensity. The results are presented in Table 1 for all the fluorophore concentrations used. It can be concluded that whereas η_L (longitudinal to the SSLE axis) presents high tolerance against variations of the pumping excitation spot (positioning changes between 600 and 1125 µm cause the emitted light intensity to vary <5%), the η_T tolerance is between half and one order of magnitude lower, which is consistent with both the mapping results presented above and the numerical results. Nevertheless, the η_T values (150 µm) are large enough to be attained using massproduction technologies.

Table 1 Longitudinal and transversal confidence positioning intervals for each concentration

Concentration (µM)	Longitudinal confidence positioning interval η_L (µm)	Transversal confidence positioning interval η_T (µm)
10	1125	150
50	600	150
100	600	150
500	1050	150

Analytical performance of the PhLOC

Once the SSLE working principle was validated, the whole PhLOC was fabricated using the above described technology. Figure 4 presents an image of the final PhLOC under homogeneous wavelength excitation of 352 nm. In this case, the MIR was filled with a dilution of 50 μ M fluorescein in PBS for better visualization. The extent of the homogeneity of the emission from the SSLE can again be observed. In addition, using the fabrication procedure described in this work, the SSLE is inherently self-aligned with both the interconnecting waveguide and the MIR, dramatically simplifying the implementation of the main components in any photonic system, namely, the light source, sensing region and detector (being in this case the SSLE, MIR and output fiber optics, respectively).

Absorbance measurements were performed using a non-biofunctionalized PhLOC. Quinoline yellow dye is a water-soluble acid dye that is extensively used as an additive in the food industry³⁸ as well as a



Figure 4 Fabricated PhLOC comprising the SSLE, air mirrors, MIR, biconvex lens and channel for fiber optics insertion. The MIR was filled with a PBS solution containing 50 μ M fluorescein only for visualization, and the entire system was placed under homogeneous 352-nm wavelength excitation. A 1-cent euro coin is also included for size reference. MIR, multiple internal reflection; PhLOC, photonic lab on a chip; SSLE, solid-state light emitter.

coloring agent in general.³⁹ This dye is also a potential biohazard, confirmed by genotoxicity tests using in vitro human lymphocytes.⁴⁰ Therefore, the measurement of its concentration is not only important in the food industry but also in the environmental water analysis because of its deleterious effects, particularly for infants.³⁹ Quinoline yellow dye has a maximum absorption wavelength at λ_{max} =440 nm, which falls inside the Atto 390 emission spectra. The calibration curve obtained (Supplementary Fig. S3) using this PhLOC was in the linear range between 1 and 14 µM with a LOD of (0.60 ± 0.01) µM. In comparison, Zhao *et al.*⁴¹ used a carbon nanotube-modified electrode to perform electrochemical characterization, obtaining a linear range between 2.74 μ M and 73.18 μ M and an LOD of 1.82 µM. Optical measurements were also provided by other authors, such as Sorouraddin and co-workers,⁴² who obtained a LOD of 3.5 µM. Only using a combination of chromatographic and spectrophotometric methods was a LOD of 0.4 µM obtained, but this required a much more complex and expensive approach. Thus, the PhLOC reported here represents a dramatic improvement in the analysis of this target analyte compared with the state of the art. In addition, the hybrid monolithic integration of the SSLE allows for a simplification of its use and a reduction of the fabrication costs.

Finally, the performance of the biofunctionalized PhLOC for the detection of H_2O_2 was assessed. The working principle was based on the catalytic action of the HRP enzyme in the presence of an ABTS colorless charge transfer mediator. Upon the reduction of H_2O_2 to water, this mediator was concomitantly oxidized to the green-colored ABTS⁺⁻ radical cation, which exhibited an intense absorption band from 390 nm to 445 nm.

Three identical systems were functionalized and measured on three consecutive days, and the corresponding calibration curves are presented in Figure 5. The three systems presented a monotonic increase in the absorbance at a wavelength of 435 nm in solutions containing increasing concentrations of H_2O_2 ranging from 1.4 μ M to 8.2 μ M, which is consistent with the behavior previously reported using a PhLOC with an external excitation light source.¹⁶ A linear fitting was performed in this range, and the estimated analytical parameters are presented in Table 2.

Good agreement is observed between the different measured PhLOCs, with a maximum variation of 10% in sensitivity and with absolute values similar to or better than those previously published for





Table 2 Analytical data extracted from the H₂O₂ calibration curves performed with various PhLOCs

	Sensitivity (A.U. μM^{-1})	LOD (µM)*	R^2
PhLOC #1	0.0121±0.0005	0.64±0.02	0.992
PhLOC #2	0.0125±0.0006	0.88±0.09	0.989
PhLOC #3	0.0112±0.0005	0.67±0.02	0.991
Mean	0.0119 ± 0.0007	0.7±0.1	

Abbreviations: LOD, limit of detection; PhLOC, photonic lab on a chip. * LOD calculated following the 3σ IUPAC criteria using the linear concentration range

from 1.4 μM to 8.2 μM for H_2O_2 detection. Absorbance recorded at $\lambda{=}435$ nm.

other optical biosensing approaches for H_2O_2 .^{16,43} These results confirm the feasibility of monolithically implementing a low-cost SSLE in a PhLOC without increasing the technological complexity while maintaining the performance of the other components of the system. In addition, even though this work was performed with Atto 390 fluorophore, any other molecule (or quantum dots) with photonic reemission could have been used instead, conferring to the proposed PhLOC an outstanding flexibility. Similarly, the use of HRP was selected to benchmark the PhLOC against a well-established enzymatic response. Nevertheless, other enzymes that are more clinically or biologically relevant could also be used in combination with this PhLOC approach.

CONCLUSIONS

We demonstrate the easy integration and successful performance of a PhLOC with a SSLE made of a fluorophore-doped hybrid xerogel material and a MIR as the main elements. The resulting photonic approach can be fabricated at low cost because the fabrication only requires one photolithographic step. Thus, the elements of such systems are inherently aligned. Coupling enhancement between the SSLE and MIR was achieved by defining air mirrors surrounding the SSLE to redirect emitted light toward the MIR. Confocal studies of the SSLE revealed homogeneous fluorophore dispersion. Using quinoline yellow as a colored target analyte, the feasibility of this approach was verified, obtaining a LOD of (0.60 ± 0.01) µM. Finally, the MIR was selectively biofunctionalized with HRP, and the PhLOC was benchmarked against hydrogen peroxide (H₂O₂) as a target analyte, providing a LOD of (0.7 ± 0.1) µM for H₂O₂. This detection strategy could be applied to other enzyme- or immunoassay-based analytical tools for the measurement of a wide range of target analytes and validates both the proposed PhLOC and its fabrication strategy.

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