Gain-Based Mechanism for pH Sensing Based on Random Lasing

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Here, we investigate the mechanism of a random-lasing-based sensor which shows pH sensitivity exceeding by 2 orders of magnitude that of a conventional fluorescence sensor. We explain the sensing mechanism as related to gain modifications and lasing-threshold nonlinearities. A dispersive diffusive lasing theory matches the experimental results well, and it allows us to predict the optimal sensing conditions and a maximal sensitivity as large as 200 times that of an identical fluorescence-based sensor. The lack of complex alignment and the high sensitivity make this mechanism promising for future biosensing applications.

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I. INTRODUCTION

Fluorescence-based sensing exploiting spontaneous emission is among the most widespread mechanism for biochemical detection [1,2]. Latest developments have focused on improving the biochemistry of the fluorescent binder [3] and on expanding the monitored functionalities [4], as well as on engineering nanoscale light fields via surface plasmons [5], microcavities [6], photonic crystals [7], or optical resonators [8] to enhance light-matter interaction.

Lasing instead, which is based on stimulated emission, has been largely overlooked as a sensing transducer, mainly because of the complexity of a conventional lasing architecture. Lasing has the potential to outperform fluorescence due to the signal amplification inherent to the lasing process, the increased signal-to-noise ratio, the narrow emission line, and nonlinear dynamics, as has been shown for laser-based interleukin sensing [9], explosives detection [10], and the remote identification of hazardous chemicals [11]. Just recently, biocompatible lasing architectures made with vitamins [12] and proteins [13,14] have been fabricated, indicating a path for laser-based biosensing inside living tissues [15].

While conventional lasing requires periodic geometries or carefully aligned cavities, random lasing (RL) occurs in disordered systems with optical gain [16] ranging from semiconductor powers [17,18] to biomaterials such as human tissue [19]. The lack of an optical cavity gives this structure resilience against deformation and makes it appealing for implantation in biological media. Despite the inherit randomness of RL, emission control has been achieved both spectrally [20,21] and directionally [22], and its rich modal properties have just started to be explored [23,24].

Sensing with RL has been limited so far to the detection of changes of the scattering strength of the matrix by a refractive index [19,25] or temperature [26] variation. Instead, the potential of targeted sensing via biochemical interaction at the gain level, affecting the amplification process, is largely unexplored.

We recently demonstrated a biocompatible random laser [14] reacting to pH changes which provides a preliminary sensing proof. In this paper, we explain the mechanism of

FIG. 1. Random-lasing sensing scheme. Light multiple scattering in the gain medium embedded in a photonic glass leads to amplification and lasing. This result is experimentally visible in the emission spectrum which shows a narrow-band emission [the red line in (a)]. For alkaline pH, the lasing emission is switched off, resulting in the broadband fluorescence emission and the lower intensity [the orange line in (b)].
sensing by random lasing based on gain variation upon interaction with the biochemical environment: the lasing action at neutral pH [Fig. 1(a)] is suppressed at alkaline pH [Fig. 1(b)]. The experiments are in very good agreement with the calculations of a dispersive diffusive lasing model without free parameters. We predict the optimal sensing conditions and we show that the random-lasing sensitivity can be up to 200 times that of fluorescence.

II. RESULTS

The random-lasing system is fabricated by self-assembly of an inverse silk photonic glass [27] with embedded laser dye (Rhodamine 6G), as detailed in Ref. [14]. The pores have a diameter of approximately 1.3 μm, which optimizes the optical scattering (the measured transport mean free path in water is $\ell_t \approx 14$ μm). The sample (thickness $L \approx 100$ μm) is excited at a fixed power $P = 840 \mu J/\text{mm}^2$ with 6-ns pulses of a Nd:YAG $Q$ switched (wavelength, 532 nm; spot diameter, approximately 2 mm) well above the lasing threshold ($T \approx 80 \mu J/\text{mm}^2$). The pH of a water solution surrounding the laser is controlled by varying the concentration of NaOH. The porous structure of the inverse photonic glass and the permeability of silk [28–30] allow for the molecules to be efficiently affected by the solution. As shown in Fig. 2(a), a progressive decrease in peak intensity (the blue circles) is observed for an increasing pH: beyond the value $pH = 13$, the lasing action is switched off. The peak intensity shows an overall approximately 100-fold intensity decrease, and the full width at half maximum (FWHM) (the red squares) instead increases smoothly from 14 nm at $pH = 7$, corresponding to the above-threshold linewidth [shown in Fig. 1(a)], to $pH \approx 13$, where it sharply reaches 54 nm, which is the FWHM of the fluorescence spectrum [shown in Fig. 1(b)]. This result indicates a shift of the threshold to a larger pump intensity. The error bars are calculated as the standard deviation of the average of ten repeated measurements, each by pumping with a single laser pulse.

This sensing dynamics can be predicted by a dispersive diffusive lasing model, built on light diffusion coupled to classical molecular rate equations, that includes spectral mode competition [31]. This model has no free parameters—it describes the realistic sample characteristics given the scattering and gain properties of the medium either measured or taken from the literature [14,32]—while the gain cross section ($\sigma_g$) is calculated by assuming the same scaling of the absorption ($\sigma_a$) [32]. The resulting theoretical predictions are shown in Fig. 2(b) and are in very good agreement with the experimental data. As expected [31], the predicted lasing linewidth is underestimated, as, in the model, the narrowing is limited only by the gain saturation.

Qualitatively, we can understand the lasing switching off as being due to a reduction of the optical amplification which increases the gain length, $\ell_g$ (the distance required for amplification of a factor $e$): when the critical lasing size

$$L_{cr} \propto \sqrt{\ell_t \ell_g}$$

required for lasing becomes larger than the sample size, the laser switches off. Here, the transport mean free path $\ell_t$ is unchanged by pH variations, whereas the gain length $\ell_g$ is instead pH sensitive. More quantitatively, in the approximation of a stationary and uniform system, the lasing threshold $T$ can be expressed as

$$T \propto \left[ (N \tau_e v \sigma_e - 1) \tau_i \Phi \sigma_a \right]^{-1},$$

where $N$ is the density of molecules, $v$ is the speed of light in the medium, $\tau_e$ is the Thouless time (the typical time it takes for a photon to escape the disordered medium), which accounts for the losses at the surface, and the relevant properties of the molecules providing optical gain are modeled with the stimulated-emission cross section $\sigma_e$, the absorption cross section $\sigma_a$ at the pump wavelength, the radiative lifetime of the excited state $\tau_i$, and the quantum efficiency $\Phi$. The latter two are related via the nonradiative decay rate $\Gamma_{nr} = 1/\tau_{nr}$, as $\Phi = \Gamma_i/(\Gamma_i + \Gamma_{nr})$.

A change in any of the molecular parameters in Eq. (1) would modify the lasing threshold and be detectable by the lasing sensor. The dye fluorescence parameters measured as a function of pH, which are the input of the lasing model, are shown in Fig. 3.
Dyes of the rhodamine family are employed for sensing owing to their photophysical properties, including large quantum yield, absorption, and photostability [33]. The molecular fluorescence is reduced or enhanced when the dye properties are unaffected in the pH range 7–10. Starting at pH ≈ 10, we observe a pronounced decrease of absorption and lifetime, and, from pH ≈ 11, a similar decrease of the quantum efficiency. The quantum efficiency and the lifetime are obtained from fluorescence studies of the porous and doped silk matrix with picosecond pulsed excitation (\(\lambda = 532\) nm, 40 MHz): \(\Phi\) is obtained as the variation of the fluorescence intensity when recording the light escaping from the sample with an integrating sphere, and \(\tau\) by fluorescence lifetime spectroscopy by time-correlated single-photon counting. The absorption is measured with a spectrophotometer and is consistent with the lifetime and quantum efficiency calculated by the Strickler-Berg relation [37]. No significant emission spectral shift is observed. It is evident when comparing Figs. 3 and 2 that the changes in the molecular properties are amplified by the lasing system, and this amplification results in a large intensity variation with a sharp transition of the lasing emission, which offers an opportunity for efficient sensing.

### III. DISCUSSION

The sensitivity capabilities and limits of RL sensing can be predicted by calculating the effect of the dye parameters on the lasing threshold. Although these parameters are typically coupled in real dyes, we consider them independently to isolate their role. We have chosen as the figure of merit the peak intensity (\(I\)) relative sensitivity, defined as

\[
S_\alpha = \left| \frac{dI}{d\alpha/\alpha} \right|
\]

where \(\alpha\) is the parameter examined. The linear response typical of the fluorescence regime would give \(S_\alpha = 1\). In Fig. 4, we compute \(S_\alpha\) for the various molecular parameters (\(\alpha = \sigma_\alpha, \tau, \Phi, \sigma_e\)) at different pump intensities. The color map highlights regions with a linear response (white) and a highly nonlinear response above \(S_\alpha = 1\) (red). The blue areas correspond to regions with a small or negligible effect on the measured intensity. The black dashed lines are the calculated lasing threshold, marking the boundary between the fluorescence and lasing regimes. For all parameters, there are regions with increased sensitivity when compared to fluorescence (\(S_\alpha > 1\)).

Figure 4 can be understood by considering the role of the different molecular parameters in the lasing process.
$\sigma_a$ describes the pump absorption, and therefore the excitation probability of the fluorophores. This is a typical property exploited in fluorescence sensing, as it induces a variation of the measured emitted light intensity. In the regime where RL has sizes exceeding the penetration depth of the pump, a change in pump absorption can be compensated for by an increase of the active volume inside the system, such that the total available gain is the same; i.e., for large absorption values [the right side of Fig. 4(a)], the RL is insensitive to changes in $\sigma_a$. Instead, for lower absorption values [the left side of Fig. 4(a)], when the pump absorption length is comparable with the system size, the absorbed intensity and the emission intensity are linearly related to $\sigma_a$ for both fluorescence and lasing (the white areas). Interestingly, around the lasing threshold (the black line), a twofold increase in the sensitivity $S_{\sigma_a} = 2.2$ (the red-light region) is predicted.

Similar behavior can be observed for $r$, which instead describes the lifetime of the population of the excited state and therefore is related to the ease of inducing population inversion. As shown in Fig. 4(b), the recorded intensity is largely insensitive to a change of $r$, for both fluorescence and lasing. Instead, a lifetime decrease induces a mild shift of the lasing threshold towards higher pump intensities, resulting in a roughly linear sensitivity, with $S_r = 0.9$ around the lasing threshold.

The quantum efficiency $\Phi$ is another quantity often exploited in fluorescence-sensing techniques, as it relates directly to the emitted intensity. The wide linear (white) region below the lasing threshold in Fig. 4(c) is the linear sensitivity of the fluorescence regime. The lasing-emission intensity well above threshold is marginally affected by the quantum efficiency because nonradiative decay processes are slower than the stimulated-emission process ($\Gamma_{\text{nonradiative}} \ll \Gamma_{\text{stimulated emission}}$), and therefore they become irrelevant once stimulated emission dominates. Instead, around the lasing threshold, the sensitivity peaks up to $S_{\Phi} = 201$, as shown by the red region. This result can be understood as the emission intensity increases rapidly as stimulated emission (unaffected by $\Phi$) takes over spontaneous emission (affected by $\Phi$).

The most direct way of tuning the lasing threshold is by controlling the gain value, i.e., altering $\sigma_e$ as shown in Fig. 4(d). This is a parameter unique to lasing which has no effect on fluorescence. As expected, the calculated fluorescence sensitivity is independent of $\sigma_e$ and, once again, the largest response is found around the lasing threshold. In this case, a decrease of $\sigma_e$ to roughly 10% of the original value results in the suppression of the lasing emission, regardless of the pump intensity. In these conditions, high sensitivity ($S_{\sigma_e} = 186$) is reached.

We can now discuss the experimental sensing profile reported in Fig. 2. The top bars in Fig. 4 identify the measured variation of the parameters. As expected, a $p$H variation affects all of them, but, most notably, the gain and quantum efficiency. The resulting experimental sensitivity extracted from the experimental data is $S_{pH} = 200 \pm 50$ at $p$H = 13, which is larger than the theoretical expectation of $S_{pH} \approx 60$. Finally, in the range $p$H = 12–13, we estimate a limit of detection (LOD) of LOD$_{pH} = 0.03$, defined as 3 times the signal-to-noise ratio.

Stimulated emission can, therefore, boost the sensitivity of a fluorescence sensor as well as provide an additional sensing parameter, i.e., $\sigma_e$. These advantages come at the expense of additional complexity. Lasing requires a nanophotonic architecture to promote stimulated emission, a disordered medium for RL, and a dye capable of providing net optical gain. The critical length limits the miniaturization of the lasing device to approximately ($10 \mu$m)$^3$ which implies that the RL sensor is not suitable for sensitivity at the single-molecule level. When compared to fluorescence schemes, RL requires a higher excitation intensity, in the microwatt range ($>1 \mu$ pulse energy) instead of the nanowatt range of conventional single-molecule spectroscopy. While this requirement could be a problem for in vivo sensing, preliminary results in living cells [38,39] show that these power ranges are below the damage threshold of the biological media.

IV. CONCLUSIONS

In this paper, we introduce a RL sensing scheme based on the lasing intensity and FWHM modification of the gain dye parameters, which we attribute to a threshold shift. We present a detailed description of the sensing mechanism and a theoretical model which matches very well the experiments on $p$H sensing by silk-based random lasing. We identify the most efficient sensing scheme, with a 2-order-of-magnitude enhancement with respect to fluorescence. Given the universality of multiple scattering, its robustness against stress and deformation, and the large availability of fluorescent and lasing dyes, we foresee possible applications for biological and chemical sensing in living tissues.

All data created during this research are provided in full in the results section. They are openly available from Ref. [40].

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