Systematic Study of the Surface Plasmon Resonance Signals Generated by Cells for Sensors with Different Characteristic Lengths

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Abstract
The objectives of this study were to establish an in-depth understanding of the signals induced by mammalian cells in surface plasmon resonance (SPR) sensing. To this end, two plasmonic structures with different propagation and penetration distances were used: conventional surface plasmon resonance and long-range surface plasmon resonance. Long-range SPR showed a lesser sensitivity to the absolute number of round cells but a greater resolution due to its very narrow spectral dip. The effect of cell spreading was also investigated and the resonance angle of long-range SPR was mostly insensitive unlike in the conventional SPR counterpart. Experimental data was compared with suitable models used in the SPR literature. Although these simple averaging models could be used to describe some of the experimental data, important deviations were observed which could be related to the fact that they do not take into consideration critical parameters such as plasmon scattering losses, which is particularly crucial in the case of long-range SPR structures. The comparison between conventional and long-range SPR for cellular schemes revealed important fundamental differences in their responses to the presence of cells, opening new horizons for SPR-based cell assays. From this study, long-range SPR is expected to be more sensitive towards both the detection of intracellular events resulting from biological stimulation and the detection of microorganisms captured from complex biological samples.

Introduction
Mainstream applications of surface plasmon resonance (SPR) biosensors typically involve the detection of (bio)molecules adsorbed, specifically or not, on the plasmonic sensing surfaces. SPR signals generated by the presence of biomolecule adlayers are, in first approximation, linearly related to the thicknesses, as long as the latter is uniform and much smaller than the evanescent probing fields. However, SPR sensing has also developed into a powerful technology for the sensing of large biological entities such as cells and bacteria. For instance, SPR sensing has been successfully used to detect the binding of pathogens and cells or monitor their responses to external triggers such as drug or signaling biomolecules (cell-based assay) [1,2,3,4,5,6]. SPR technology provides indeed a powerful means of studying the cellular response to stimulants. The signal generated in such experiments originates in complex biological events that locally impact on the refractive index distribution. Subsequent experiments involving upstream and downstream inhibitors of the stimulation or complementarity techniques are therefore required to elucidate the biological meaning of the SPR response [7,8].

An important consideration is that the presence of microorganisms on solid surfaces leads to layers inherently larger than the characteristic dimensions of plasmonic evanescent waves and consequently, resulting signals are not trivial. For instance, cells and bacteria extend well-beyond the probing fields in the z direction. The propagation length of the plasmons (propagation in the x direction, intersection of the plane of incidence and the sensor surface) is also expected to be a crucial parameter in regards to how the plasmonic waves actually sense the complex medium composed of cells and cover solution, as it was approached in the literature [9].

With respect to morphology-induced SPR signal, the data presently available in the literature is contradictory. While some studies have considered that cell-covered and cell-free regions of the sensors contribute to the overall signal independently [10], other studies make use of an effective refractive index for the cover medium composed of the solution and the cells [11]. The latter approach is the most commonly used one in waveguide biosensing [12,13,14]. Experimentally, splitting of SPR dips in presence of cells has been observed, and these sub-dips were associated to the co-existence of cell-free and cell-covered areas [9,10]. These inconsistencies in both experimental and analytical reports are not surprising as to date, the effect of cells on the signal of SPR biosensors has not been systematically studied. A recent study also demonstrated that different parts of the SPR angular spectra reflect on different intracellular mechanisms (such as paracellular
and transcellular) [15]. However, the aim of the present study is to systematically elucidate the structure-activity relationship of SPR sensors in presence of microorganisms and in the absence of external stimuli. We focussed more specifically on the relationship between the surface cellular density or morphology and the SPR response.

To this end, two different SPR structures were used in this work. The first one, conventional surface plasmon resonance (cSPR), is characterized by short propagation and penetration dimensions. The second one, long-range surface plasmon resonance (LRSPR), is characterized by long propagation and penetration dimensions. Since increases in cellular coverage can originate from either increases in the number of cells on the surface or from cellular spreading of a fixed number of cells, two systematic studies were designed to address these two different situations. The first involved round cells attached on the surfaces at different cell surface densities, which can be readily translated into cell coverage. In the following sections this scheme is referred to as calibration. The second situation considered the spreading of cells on the sensor surface and is referred to as cellular spreading scheme. Although, it has been reported that the spreading of cells was not a prominent feature in SPR signal [16], previous studies have used optical biosensing to evaluate spreading and determine cellular phase [11], suggesting the relevance of such biological events in SPR cellular schemes. In order to elucidate the effect of cellular spreading on plasmonic signals, cells were seeded at low density to minimize cell-cell interactions. Such interactions could, otherwise, mislead the signal interpretation. Low cell density is also expected to minimize the appearance of TM0 waveguide mode which would significantly increase the complexity of the system under study [11].

The second main objective of this study was to rigorously compare cell-induced signals for cSPR and LRSPR. This is of interest since LRSPR structures possess larger penetration depths, therefore the sensing electromagnetic (EM) fields can reach deeper into the cellular medium. Penetration depths for cSPR structures are of the order of 100–200 nm, whereas those of LRSPR are typically 500–1000 nm [17]. On the other hand, cSPR has better angular sensitivity than LRSPR with respect to bulk refractive index changes [18]. However, it has been recently reported that, in the case of bacterial detection, LRSPR is more sensitive than cSPR [19,20]. To achieve a better understanding of the structure-activity relationship, a theoretical and experimental comparison of these two types of sensors is therefore provided in this study. Bridging this important knowledge gap will ultimately foster the application of SPR in the studies of microorganisms.

**Methods and Experimental**

**Preparation of devices for calibration experiments**

The 3T3 fibroblast cell line was used as a model in this work and was purchased from ATCC. 3T3 are widely utilized in laboratory research, and were established from disaggregated tissue of an albino Swiss mouse embryo. They were selected for their ability to spread rapidly, as required in the second experimental scheme. The cells were cultured in complete media Dulbecco’s modified eagle medium (DMEM) supplemented with glutamine and 10% (v/v) foetal bovine serum (FBS) at 37 °C in a 100% humidified incubator with 5% CO₂.

The sterilized SPR sensor chips were coated with poly-L-lysine (PLL) to enhance the cell-surface interactions. PLL was diluted to 0.5% (w/v) in phosphate buffer saline (PBS) (Sigma-Aldrich, St. Louis, USA). The PLL coating thickness was found to be approximately 2 nm on clean gold surfaces by SPR analysis. After mounting the PDMS chambers on the sensors, PLL solution was dispensed in the channel and left to adsorb for 30 minutes. The wells were then rinsed 3 times with PBS and then 3 times with the DMEM cell growth medium, and further incubated for an additional 60 minutes. Then the growth medium was replaced by PBS (3 rinsing steps). 3T3 cells were trypsinized and seeded into the custom-built wells at the desired concentrations calculated based on the targeted surface coverage. The cells were left to sediment on the sensors for 45 minutes at room temperature. The samples were then gently washed and fixed with a 4% formalin solution (Sigma-Aldrich) for 30 minutes. Both wells of the sensor devices were then gently washed 3 times with PBS. It can be noted that, from this procedure, the differences in adsorbed proteins and compounds on the control and cell surfaces were minimized in order not to bias the data interpretation.

**Cellular spreading experiments**

The samples were prepared as described above until the cell seeding step. 3T3 cells suspended in phenol-free growth medium were left to sediment at room temperature for 45 minutes on the SPR sensor surfaces at a density of 150 cells/mm². This corresponds to a surface coverage of 3%. The sample was then clamped into the SPR stage. The temperature was maintained at 37°C and the thermal stability was controlled by sequential measurements on the cell-free area of the sensor. After thermal equilibrium, the position was adjusted to place the cell-covered region in the beam’s illumination. Spectra were acquired every 30 minutes to determine the effect of cell spreading on the SPR signal. The laser beam was blocked in between to minimize light exposure to the cells. The surface coverage was typically approximately 15% at the completion of the cell spreading. The experimental design and time-frame of the study (<3 hours) was chosen to take advantage of the absence of significant cellular proliferation immediately after reseeding. This ensured that only negligible variation in the cell number occurred during the experiment, which was verified experimentally. In parallel to the SPR study, automated time-lapse microscopy imaging of the spreading of the 3T3 cells on PLL treated glass was carried out and used as reference.

**PDMS chambers for cell culture**

Custom-made poly(dimethylsiloxane) (PDMS) chambers were used to culture cells on the SPR sensors. PDMS was obtained from Dow Corning (Midland, USA). The fabrication process relied on standard master- replica fabrication. Two wells per chamber were utilized: one, in which the cells were seeded, the other one was used as a control for the SPR measurements. The surface area of each well was 154 mm². The height was approximately 4.5 mm. The PDMS chambers were sequentially cleaned with chloroform, acetone and water in a sonic bath to remove non-cross-linked, low-molecular weight, siloxane compounds which are toxic to cells [21]. The chambers were autoclaved prior to use.
Microscope and cell images

In order to calculate the density of cells on different samples, microscope images were taken and analysed either manually or with ImageJ software. The dimensions of the images were large enough to include many cells which enabled for reliable statistics. Typically at least 3 images were used and averaged. The cell sizes were calculated by measuring the number of pixels and using the pixel/micron conversion determined beforehand with a calibration grid. The density, initially expressed in number of cells/mm² was then converted into coverage (%). The coverage was calculated as the projected surface area of an average cell multiplied by the number of cells in the image and divided by the surface area of the whole image. Lastly, from round ST3 micrographs, the average diameter of a ST3 cell was calculated to be 8.05 ± 1.5 μm (± represent the extrema). For the cellular spreading scheme, the cell projected surface areas were calculated by contouring the cells on each image of the time-lapse video using ImageJ. The contours of each cell in each image were manually drawn since the complex shapes that cells can make couldn’t be dealt with by the software. Once the contours were drawn, the function measurement of ImageJ was used to determine the area within the contours. From these parameters the coverage could be calculated as described above. It was found, that when cells were spread on the surface, the average size of a spread cell was 722 μm², corresponding to an effective diameter of 30.3 μm. Assuming a constant cell volume, this yielded averaged layers to be 2.3 μm±0.5 μm in thickness (in the z direction).

SPR measurements and setup

SPR measurements were conducted using an optical setup based on the attenuated total reflection (ATR) method with Kretschmann configuration that was developed at the Max Planck Institute for Polymer Research in Mainz (Germany). As shown in Figure 1a, a light beam at wavelength of λ = 632.8 nm from HeNe laser (CVI Melles Griot, USA) passed through two polarizers (Bernhard Halle Nachfl. GmbH, Germany) and a chopper (PerkinElmer Instruments, USA) before it was coupled to an LASFN9 glass prism. The intensity of the laser beam reflected from the prism base was detected by a photodiode detector and a lock-in amplifier (Signal Recovery, USA). The SPR sensor chip was optically matched to the prism base by using a refractive index matching oil (Cargille, USA) and the custom made PDMS flow-cell chamber was pressed against its surface. The angle of incidence, θ, of the laser beam hitting the SPR sensor surface was controlled by a motorized rotation stage (Hans Huber AG, Germany) and the whole system was controlled by the software WasPlas developed at the Max Planck Institute of Polymer Research (Germany).

The spectra were measured at controlled temperature of the sensing medium in a reproducible way. After SPR spectra were acquired, the parameters half width half maximum (HWHM), intensity at resonance (Rₘₐₓ) and angle at resonance (θₘₐₓ) were extracted from the curves using the program Winspall (Max Planck Institute, Mainz, Germany) or an algorithm written in Maple from Maplesoft (Waterloo, Canada). The HWHM were calculated on the left line of the SPR dip, that is the angles used in HWHM are θₘₐₓ and the angle for which the intensity is half the intensity span and which is lower than θₘₐₓ. The values were taken for the dip of lower intensity in case two dips were present in the spectra. The HWHM was utilized for calculating the figure of merit as described in the first paragraph of the section Experimental results. To remove the bias introduced by minute fabrication differences between each sensor, the variation of these four parameters was used in this study. In the calibration scheme, the parameter values were subtracted by those for the PBS reference spectrum in the control well. In the cellular spreading scheme, the reference was the initial values obtained for non-spread cells (i.e. for coverage c≈3%).

In the following, the prism will be referred to as medium 0 which is in contact to a LRSPR or cSPR structure, itself in contact to a cover solution. The cover solution is either 30% C PBS in the calibration experiments or 37°C DMEM in the cellular spreading ones (see Figure 1b and 1c).

The refractive indices of the different layers were: n₉PM = 1.34662, n₉Cytop = 1.33675, n₉PBS(30°C) = 1.3384, n₉DMEM(37°C) = 1.3351.

Experimental Results

Comparison of cSPR and LRSPR in the calibration scheme

The experimental spectra obtained for both cSPR and LRSPR are presented Figure 2. A red shift was observed (resonance shifting towards higher angles) for both cSPR and LRSPR with increasing cell coverage. The intensity at resonance, Rₘₐₓ, displayed a more complex evolution for both cSPR and LRSPR, with an initial increase at low densities before a decrease at higher cellular densities. A comparison of the spectrograms indicates a sharper distinction of these features for LRSPR spectra in comparison to cSPR ones.

Figure 3a and 3b summarizes the evolution of the spectrum parameters, θₘₐₓ and Rₘₐₓ, as a function of the cell coverage, cₖ, the
other relevant parameters, half width at half maximum, HWHM, and the intensity at fixed angle \( R_{\text{cell}} \), are plotted in Figure S1a and S1b). The angular sensitivity of cSPR to cellular coverage was found to be significantly higher than that of LRSPR. This finding is of interest since a recent study reported greater sensitivity of LRSPR for the detection of bacteria adsorbing on the sensors [19]. Our observation is, however, in agreement with the fact that, for homogeneous bulk changes, cSPR sensors present a sensitivity of \( ~100%/RIU \) [18] whereas LRSPR sensors generally have a sensitivity of \( ~25%/RIU \) [22]. However, since the HWHM is much smaller in the case of LRSPR, the localisation of \( \theta_{\text{res}} \) can actually be more precise. When dealing with SPR sensitivity matters, it is therefore useful to consider the figure of merit (FOM), as it can be used as a tool to analyse the improvement in resolution of SPR-based sensors [22]. In this study the FOM is defined as \( \text{FOM} = \text{sens}_{\text{cell}} \times \text{HWHM} \), where \( \text{sens}_{\text{cell}} \) is the sensitivity of the sensors to the cell coverage on the surface. Although the full width at half maximum (FWHM) is often used, due to the asymmetry of SPR dips, using the HWHM is more precise and in turn can help minimizing potential discrepancies between FOM and the resolution calculated by refractometry taking into account the noise of blank signal [18].

The sensitivity to the cell coverage is defined in this study as the ratio: \( \text{sens}_{\text{cell}} = \Delta \theta_{\text{res}}/\Delta c \), where \( \Delta \theta_{\text{res}} \) and \( \Delta c \) correspond to angular shift and the increase in cellular coverage, respectively. Hence, FOM can be found directly by measuring the slope of \( \Delta \theta_{\text{res}}/\text{HWHM} \) function of \( c \) (curves plotted in the inset of Figure 3a). Assuming linear relationships, it was found that \( \text{FOM}_{\text{cSPR}} = 1.30%/^{-1} \) and \( \text{FOM}_{\text{LRSPR}} = 7.39%/^{-1} \), which corresponds to a 5.7-fold improvement of LRSPR over cSPR sensors towards the detection resolution of microorganisms such as cells and bacteria, although one has also to take into consideration the experimental noise associated with SPR modalities to accurately predict their limits of detection.

Comparison of cSPR and LRSPR in the cellular spreading scheme

Spectra for cSPR (a) and LRSPR (b) sensors obtained for various level of cell spreading are shown in Figure 4. Typical spreading time-lapse microscopic images can be found in Figure S2 as well as the associated trend of coverage versus spreading time, Figure S3. The coverage values determined by microscopy were used for the spectra of Figure 4.

Figure 3c and 3d summarizes the key spectrum parameters (Figure S1c and S1d show similar graphs for HWHM and \( R_{\text{fixed}} \)). Interestingly, the angle of resonance, \( \theta_{\text{res}} \), is sensitive to the spreading of cells only in case of cSPR. In the evolution of the \( R_{\text{min}} \) parameter, a similar trend is observed for cSPR and LRSPR at initial spreading but the cSPR \( R_{\text{min}} \) appears to saturate earlier than the \( R_{\text{min}} \) of LRSPR.

Modelling and Comparison with Experimental Data

Description of the models

To further understand and, in turn, predict the behaviours of the cSPR and LRSPR spectra in response to the presence of cells, theoretical curves from averaging models were compared to the experimental data.

Determining the refractive index (RI) of a cellular layer is a challenging endeavour, even though it has been done in some instances [23,24]. Such value is inherently highly dependent on the method used from an optical standpoint (evanescent wave technique, refractometry, interferometry, etc.) as well as on the biological point of view (spreading of the cell, presence of cell-cell interactions, growth phase, passage number, etc.). The RI values used in the relevant literature to represent cellular layers are typically in the range of 1.35–1.37 RIU [24,25,26]. Therefore in the present study the values 1.35, 1.36 and 1.37 were used to model the presence of cells on the sensor surface in order to assess their impact on the SPR signal.

Two different methods of simulating SPR spectra were considered. First, the contributions of cell-covered and cell-free areas were assumed to be independent and their intensities were averaged as a function of the cell coverage, \( c \). If \( R \) denotes the intensity of the normalized reflected p-polarized light, under the aforementioned assumption, one can write, following Ziblat et al. [10]:

\[
R_{\text{avg}}(\theta) = (1-c) R_{\text{cell}}(\theta) + c R_{\text{sol}}(\theta),
\]

where the subscript “avg” and “sol” stand for average and solution, respectively. This equation is only valid if the cell-covered and cell-free areas are independent with respect to the propagating surface plasmon waves. This means that the lateral dimension of
Figure 3. Comparison of cSPR and LRSPR spectra parameters for the two experimental schemes. Dependence of $\theta_{res}$ (a) and $R_{res}$ (b) with respect to the cell coverage (round cells). The inset in (a) represents $\Delta \theta_{res}/$HWHM, the slope of which is the figure of merit of the sensor. It was found that $FOM = 1.30\%^{-1}$ and $FOM = 7.39\%^{-1}$ for cSPR and LRSPR, respectively, yielding a 5.7-fold enhancement in the case of LRSPR. Dependence of $\Delta R_{res}$ (c), and $R_{res}$ (d) with respect to the cell spreading. The error bars represent the standard errors.

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the cells should be smaller than the characteristic propagation of the plasmons, i.e. the propagation length. According to Figure S4 and the discussion on propagation lengths in Appendix S1, it was found that this model should only be applied to cSPR structures.

In the second approach, an effective RI of the cover medium was assumed. It is comprised between that of the cover solution (e.g. PBS or DMEM) without cells, $n_{sol}$, and that of a cover medium composed of a contiguous monolayer of cells, $n_{cell}$. The condition to use the effective RI model is that the sensing waves feature long propagations compared to the cell lateral dimensions. Under this assumption, and based on the methodology first applied to waveguides, one can write [11,14]:

$$n_{eff} = n_{cell} + \frac{2}{L_p} \int_0^h (n_{cell} - n_{sol}) A(z) e^{-2z} dz,$$

where $A(z)$ is the cross section of the cells at the distance $z$ from the surface and $C_s$ the number of cells per surface unit. $L_p$ represents the penetration depth of the EM sensing fields into the medium (see paragraph S2 of Appendix S1). By using equation (2), one inherently assumes that the sensing EM fields penetrating into the cell-solution medium are evanescent, which is only true beyond the critical angle of the cellular medium, $\theta_{cell}$. In such a complex biological system, the critical angle is comprised between $\theta_{cell} = \sin^{-1}(n_{sol}/n_{prism})$, the critical angle of the solution, and $\theta_{cell} = \sin^{-1}(n_{cell}/n_{prism})$, the critical angle of the cell, with $\theta_{cell} < \theta_{cell}$ and a significant part of the SPR dip may not necessarily be beyond $\theta_{cell}$ (especially in the case of narrow SPR dips such as those of LRSPR structures). Additionally, $L_p$ varies substantially with the angle of incidence, $\theta$. Hence, the expression of $n_{eff}$ should strictly be a function of $\theta$ and should, moreover, be expressed by a different formula for angles smaller and greater than the critical angles.

The values for which $\theta < \theta_{cell}$ are complex to model with an effective-RI and are not relevant since the parameters that characterize the SPR dips use values for angles beyond $\theta_{cell}$. Therefore the model will focus on values for angles greater than $\theta_{cell}$. The part of the spectrum for which $\theta_{cell} < \theta < \theta_{cell}$ can be determined by weighting $n_{cell}$ and $n_{sol}$ by the amount of EM intensity as a function of $z$:

$$A = \int_0^h dz + \int_h^\infty e^{-\frac{2(z-h)}{L_p}} dz, \quad n_{eff}(\theta) = (1-c) n_{sol} + \frac{c}{A} \left[ n_{cell} \int_0^h dz + n_{sol} \int_h^\infty e^{-\frac{2(z-h)}{L_p}} dz \right],$$

where $h$ is the average height of the cells, which are assumed to remain cylindrical (thus, $\forall z > h_c : A(z) C_s = c$). Eq. (3) can rewrite:

$$A = h + L_{cell}/2 \quad \text{and} \quad n_{eff}(\theta) = (1-c) n_{sol} + \frac{c}{A} [n_{cell} h + n_{sol} L_{cell}/2].$$

When $\theta \geq \theta_{cell}$, one may use an equation featuring the weighting of $n_{cell}$ and $n_{sol}$ by the squared evanescent fields similar to Eq. (2). However, more rigorously, the penetration depths of each squared evanescent field – within the cell and beyond the cell – will be...
taken into account using $L_{\text{psol}}$, the penetration depth in the solution, and $L_{\text{pcell}}$, the penetration in the cell layer:

$$A = \int_0^h e^{-\frac{2 \pi z}{L_{\text{pcell}}}} dz + e^{-\frac{2 \pi h}{L_{\text{psol}}}} \int_h^\infty e^{-\frac{2 \pi (z-h)}{L_{\text{psol}}}} dz$$

and

$$n_{\text{eff}}(\theta) = (1-c)n_{\text{sol}} + \frac{c}{A} \left[ \int_0^h n_{\text{cell}} e^{-\frac{2 \pi z}{L_{\text{pcell}}}} dz + n_{\text{sol}} e^{-\frac{2 \pi h}{L_{\text{psol}}}} \int_h^\infty e^{-\frac{2 \pi (z-h)}{L_{\text{psol}}}} dz \right].$$

Equation (5) can rewrite:

$$A = \frac{L_{\text{pcell}}}{2} \left( 1 - e^{-\frac{2 \pi h}{L_{\text{psol}}}} \right) + \frac{L_{\text{psol}}}{2} \left( e^{-\frac{2 \pi h}{L_{\text{pcell}}}} \right)$$

and

$$n_{\text{eff}}(\theta) = (1-c)n_{\text{sol}} + \frac{c}{A} \left[ n_{\text{cell}} \frac{L_{\text{pcell}}}{2} \left( 1 - e^{-\frac{2 \pi h}{L_{\text{psol}}}} \right) + n_{\text{sol}} \frac{L_{\text{psol}}}{2} \left( e^{-\frac{2 \pi h}{L_{\text{pcell}}}} \right) \right].$$

According to the discussion on propagation lengths in Appendix S1, it was found that this model should only be applied to LRSPR structures.
Comparison of the experimental and averaged-intensity modelled cSPR spectra

1. **Calibration scheme.** Using Eq. (1) for three different values of \( n_{\text{cell}} \) (\( n_{\text{cell}} = 1.35, n_{\text{cell}} = 1.36 \) and \( n_{\text{cell}} = 1.37 \)), the simulation results show that by increasing the number of cells the characteristics of the spectra change significantly (see Figure S5). For all \( n_{\text{cell}} \), the increase of cell coverage leads to the angle of resonance (or apparent resonance), \( \theta_{\text{res}} \), to shift to higher angles. In addition, in the case of \( n_{\text{cell}} = 1.37 \), the modelled cSPR spectra happened to split up into two sub-dips, one related to the uncovered areas ("PBS dip") and one related to the cell-covered areas ("cell dip"). For \( n_{\text{cell}} = 1.36 \), the cSPR dips are getting misshaped as the coverage reaches \( \sim 35\% \). The dip never splits up but shows a prominent feature (higher resonance coupling), on the PBS-dip side before \( 35\% \) and on the cell-dip side after \( 35\% \) coverage. For \( n_{\text{cell}} = 1.35 \), although no "dip-splitting" is predicted, a broadening of the dip is modelled as well as a rise of the intensity at resonance (\( R_{\text{min}} \)), which both increase for low coverage values but then decrease for high coverage values. The resulting parameter analysis is presented in Figure 5a and 5b (Figure S6a and S6b show similar graphs for \( HWMW \) and \( R_{\text{fixed}} \)). The data for the averaged-intensity model for \( n = 1.37 \) significantly deviated from the experimental data and were therefore not included in the figure for clarity. It can be noted that, similarly to homogenous adlayer formations on cSPR sensors, \( \theta_{\text{res}} \) undergoes a red shift with increasing coverage. However, this shift doesn’t follow a linear trend and can rather be fitted by a sigmoidal curve (in case of no dip splitting).

Comparing the experimental and theoretical parameters of the cSPR spectra in Figure 5a and 5b, reveals a good correlation with respect to the \( R_{\text{min}} \) parameters. Furthermore, the trend of \( \Delta \theta_{\text{res}} \) could be sigmoidal as shown by quality of the fitting (\( R^2 = 0.976 \)). Finally, from the experimental data of the \( \Delta \theta_{\text{res}} \) it can be concluded that the refractive index of the cells lies between \( n_{\text{cell}} = 1.35 \) and \( n_{\text{cell}} = 1.36 \).

2. **Cellular spreading scheme.** Since the averaged-intensity model does not take into account the shape of cells, the spectra predicted for cellular spreading depends only on the cellular coverage and therefore follow the same trends as in Figure 5a and 5b but are limited to the coverage range from \( 3\% \) to \( 15\% \). The parameter analysis of cSPR spectra is presented in the Figure 5c and 5d (Figure S6c and S6d show similar graphs for \( HWMW \) and \( R_{\text{fixed}} \)). Quasi-linear behaviours are observed for most of the cases and any potential dip splitting has not occurred by the time the spread cells cover \( 15\% \) of the surfaces. The comparison between the experimental data and the model data is not fully satisfactory.

Comparison of effective-RI model with experimental LRSPR data

1. **Calibration scheme with LRSPR.** LRSPR spectrum simulations under the effective-RI model are plotted in Figure S7. The behaviour of the SPR spectra is radically different from that that was simulated for the averaged-intensity model as the dip splits up in two for RI as low as \( n_{\text{cell}} = 1.35 \) (graphs can be found in Figure S8). In these simulations, LRSPR dips shift almost linearly. Furthermore no splitting and very little misshaping of the dips are predicted for all of the three \( n_{\text{cell}} \) values. In addition, \( R_{\text{min}} \) does not vary significantly.

The comparison of the experimental data with the effective-RI model is shown in Figure 6a and 6b (Figure S9a and S9b show...
similar graphs for HWMW and R fixed). The simulations showed a good agreement with the experimental angular shifts as the data could be fitted with a linear trend (R² = 0.916). A poor correlation was, however, obtained for Rmin.

2. Cellular spreading scheme with LRSPR. The comparison of the experimental data of LRSPR monitoring cellular spreading and the modelled data is presented Figure 6c and 6d (Figure S9c and S9d show similar graphs for HWMW and Rfixed). It is worth noting that, according to the model, the Rmin parameter varies only minimally in response to the spreading of cells, whereas hres does. Yet, the experimental data reveal a drastically different behaviour to what is predicted by the model.

Discussion

Calibration scheme

In the calibration scheme, which deals with increasing number of non-spread cells on the surface, one can observe an angular red shift, a convex behaviour for both Rmin for both cSPR and LRSPR. However, according to this study the origin of these behaviours is expected to be different. The dip parameters of cSPR can be satisfactorily described with the averaged-intensity model for which one observes a sigmoidal variation of hres, a convex variation of Rmin. This confirms the underlying hypothesis of Eq. (1) and implies that plasmonic waves propagating through the boundaries of cell-PBS interfaces contributes only little to the overall SPR signal which is in agreement with the fact that the propagation length is of similar dimension than the cell size. The observed discrepancies can be attributed to the fact that the propagation length of cSPR is not much smaller than the lateral dimension of round cells which put some stress on the assumption that cell-covered regions and cell-free regions contribute independently to the total reflectivity.

Furthermore, in previously reported publications, a SPR dip splitting was observed, which was not the case in our experimental data. Such dip splitting is predicted in the averaged-intensity model in the case of large RI contrasts between cells and cover solution. This situation can of course only happen for Lprop of the same order of magnitude or smaller than the lateral size of the cells. The presented effective-RI model does not support any SPR dip splitting.

Contrary to the cSPR, the variation of the spectrum parameters in LRSPR is expected to originate from an effective refractive index since the long-ranging plasmonic waves propagate much farther than the lateral size of the cells and therefore encounter many cell-free and cell-covered areas. In addition, this effective RI should be complex to take into account scattering losses due to the cell-PBS interfaces [15], however in this study the focus was brought on a real part of the effective RI similarly to what was found in the literature. The scattering losses are expected to account for the convex behaviour of Rmin [11] but further in-depth studies are required to model these complex phenomena.

Nevertheless, assuming that scattering losses do not influence the position of θres, the fitting of Δθres yielded ncell - ncell = 0.022 in the case of LRSPR and ncell - ncell = 0.017 for cSPR. The apparent higher RI measured by LRSPR can be explained by the fact that the EM fields of the LRSPR sensors probe deeper into the cell and can sense more of the high-RI intra cellular structures and membrane. Additionally, a lesser percentage of the evanescent tail of the LRSPR overlaps the nanometric gaps between the cell membrane and the substrate, typically considered to be in the 50 to 100 nm range [27]. The tendency of cSPR HWHM and Rmin
to have their maximum close to the 50% coverage is in agreement with the prediction of the averaged-intensity model as well as with previous reports including findings made for optical waveguides [29,29]. Despite the theoretical challenges in understanding their complex behaviour, LRSPR sensors can be better tools to measure cellular concentration of round cell deposited on the surface as the narrowness of the dip translates into greater FOM over cSPR and the angular position is not affected by the spreading of cells.

The discrepancy between the present study’s findings with mammalian cells, conveyed by Figure 3a, and those published in the literature with bacteria most likely do not originate from the difference in the bio-microobjects detected but rather their distance from the surface [19,20]. Indeed LRSPR sensors have higher sensitivities towards the detection of material further away from the surface whereas cSPR perform best for material in contact to the surface. In this study, only minimalist surface chemistry was used to ensure cell adhesion to the gold surfaces. On the other hand, in the previously reported bacteria detection studies, complex surface chemistry could have contributed to displace the analyte further away from the surface with also a possibility of their being pushed away from the surface by their pili or flagella [30].

**Cellular spreading scheme**

In this scheme, rather drastically different behaviours were observed for the two types of SPR structures. It is still reasonable to state that the averaged-intensity model can describe the spreading of cells in the case of cSPR. However noticeable discrepancies have appeared towards the end of the spreading phase. $\theta_{res}$ still shifts even though there is no more increase in cell size as observed in optical microscopy. On the one hand, this could be attributed to a greater sensitivity of SPR-based measurements, able to probe with a greater sensitivity cellular spreading than what can be observed in microscopy as suggested by studies relying on SPR imaging [27]. But this could also be attributed to the fact that cells are releasing microexudates at this stage without spreading further [12]. This situation could then be modelled by the deposition of a nanometric homogeneous adlayer, which would translate into angular shifts with small broadening of the dip (i.e. increase of $HHWM$ – see Figure S6c) and no changes in $R_{min}$ which corresponds to what is indeed monitored from 8% to ~13% of cell coverage.

An interesting finding of this study is that $\theta_{res}$ is not sensitive to cellular spreading in the case of LRSPR. This could be explained by the fact LRSPR structures sense a proportion of cellular membrane (highest-RI component in the cell) that reduces significantly during the spreading of cells (due to the fact that LRSPR structures sense a large volume of the cells, including vertical edges of the cell). Hence, there might be more and more cellular material on the surface during the spreading, the LRSPR effectively considers this material with a lower and lower RI, and the two effects cancel out, leading to only minor fluctuations of $\theta_{res}$. Additionally, during cell spreading, the diffraction of the plasmons due to the cell-DMEM interfaces increases drastically because the cells do not spread in spherical caps (in fact the circularity drops down to ~50% with ST3 cells) which participates in the modifications of $R_{min}$ while having little impact on $\theta_{res}$.

No inference of $n_{cell}$ could be made for LRSPR. However, for cSPR, fitting the linear parts in Figure 5c yielded $n_{cell}$=1.340 (by quadratic extrapolation of the slopes). This means that $n_{cell} - n_{water}$=0.013 for cells spreading on cSPR structures whereas $n_{cell} - n_{water}$=0.017 for rounded cells. The apparent lower RI in the case of spreading cells most likely stems from the fact that the proportion of cell membrane over the cell volume is lower when cells are spreading.

**Conclusions**

This study has been carried out in order to improve the understanding of the effects of cells on plasmonic signal in SPR biosensing and ultimately towards achieving a detailed structure-activity relationship. The first aim was to assess the validity of simple models that have been used previously in the literature for related optical sensing schemes. For SPR configuration when $L_{prop}$ is of the same dimension as that of the cell lateral size (which is the case of cSPR sensors), the averaged-intensity model was found to provide a reasonably good prediction of SPR signals, taking in consideration some deviations in $R_{min}$ parameter. On the other hand, when $L_{prop}$ is much larger than the cell dimensions (LRSPR case) the model needed is based on the fact that an effective RI is sensed by the plasmonic waves. The effective-RI model could describe the angular shifting experimentally measured in the calibration scheme although it did not describe accurately the spreading of cells of the sensors. The difference between the two schemes was attributed to the importance of the cell membrane configuration and its apparent weight in the EM sensing fields, thereby bringing the focus on the issue of penetration depth in addition to the propagation length in the studied systems. However, even when the angular shifting was correctly described (calibration scheme), this model did not provide an accurate description of the parameters $R_{min}$ of the spectra, which can be related to scattering loss in the system. Indeed, the main source of discrepancy between the data and the models is thought to originate from the scattering losses of the plasmonic waves encountering scatterers [11]. The implementation of scattering contribution in existing model is very challenging, especially since the vertical dimension of the scattering edge can vary from a few hundreds of nm for filopodia, to the whole cell’s height. As a starting point for implementing plasmonic scattering into a model, one could consider making use of Rayleigh series’ to describe the EM fields [31] or studying the propagation losses induced by refractive-index steps [32].

The second aim of the study was to compare cSPR and LRSPR in regards to their ability to “sense” the presence and morphological changes of cells on SPR sensors. It was observed that the coupling parameter $R_{min}$ could be related to cellular coverage increase (associated to either cellular spreading or cell coverage increase), and is more sensitive in the case of LRSPR. Therefore, it can be concluded here that LRSPR can be used efficiently to monitor cellular spreading through the $R_{min}$ parameter. Importantly, $\theta_{res}$ is not perturbed by the spreading of cells and therefore LRSPR sensors are expected to be more reliably used for the detection of intra-cellular signaling event without any unwanted influence of the spreading and other macroscopic morphological changes. It should also be possible in a single experiment to determine both the cellular surface concentration with no bias from cellular spreading, as well as to measure simultaneously cellular spreading by monitoring the parameter $R_{min}$. This feature opens new horizons for cell-based assays. Finally, LRSPR offers a significantly better resolution with respect to the cell number (coverage of rounded cells) since the FOM is superior to that of cSPR by 5.7 fold. The latter together with the fact that the spreading of cells does not bias the measurement, justifies the use of LRSPR in biosensing approaches aimed at the detection of microorganisms from complex mixtures.
Supporting Information

Figure S1 Comparison of HWHM and \( R_{\text{fixed}} \) for cSPR and LRSPPR in the two experimental schemes. The parameter \( R_{\text{fixed}} \) represents the readable signal of typical biosensing experiments, it is obtained by monitoring the intensity of the signal for a constant angle chosen to be close to, but smaller than, \( \theta_{\text{res}} \), in the linear region of the SPR dip. Dependence of HWHM \( (a) \) and \( R_{\text{fixed}} \) \( (b) \) with respect to the cell coverage (round cells). Dependence of HWHM \( (c) \) and \( R_{\text{fixed}} \) \( (d) \) with respect to the cell spreading. The error bars represent the standard errors.

(TIF)

Figure S2 Sequential images of the spreading of 3T3 cells. \( t = 0 \) min corresponds to seeding in the wells. From the analysis of the surface area covered by the cells, the trend of increase in cell coverage was inferred and plotted in Figure S3. Scale bar is 100 \( \mu \)m.

(TIF)

Figure S3 Average cell coverage as a function of spreading time. The complex behaviour of cellular spreading is linked to the fact that cells are constantly spreading, retracting and migrating on the surface. Nevertheless, up to about 3 hours after seeding, cells undergo an attachment and spreading phase. From 3 hours to approximately 7.5 hours the spreading slows down, possibly because of reaching its maximum. After 8 hours the coverage increases again and this can be attributed to proliferation of cells on the surface which is beyond the scope of this study. Error bars represent the standard errors.

(TIF)

Figure S4 Plot of the propagation lengths of two LRSPPR structures. (▲) LRSPPR composed of glass substrate, 850 nm of Cytop polymer, varying thickness of gold and PBS as the cover medium and (△) 800 nm of Cytop polymer, varying thickness of gold and PBS as the cover medium.

(TIF)

Figure S5 Simulated cSPR reflectivity intensities from the averaged-intensity model as function of cell coverage increase. Different cell refractive indices were implemented: (a) \( n_{\text{cell}} = 1.35 \), (b) \( n_{\text{cell}} = 1.36 \) and (c) \( n_{\text{cell}} = 1.37 \). The short-dashed curve represents the situation of the cover medium composed only by PBS and the long-dashed curve shows the spectrum for a hypothetical monolayer of contiguous cells. The behaviours are close to those that one can find for bulk RI changes. The main contributor of the increase \( R_{\text{min}} \) is the RI mismatch between the cover medium and the polymer interlayer (Cytop) in the LRSPPR structure. This translates by the coupling efficiency of the light into the plasmons departing from its optimum with increasing values of the cell coverage.

(TIF)

Figure S6 Comparison of experimental cSPR HWHM and \( R_{\text{fixed}} \) to those predicted by the averaged-intensity cSPR model. The parameter \( R_{\text{fixed}} \) represents the readable signal of typical biosensing experiments, it is obtained by monitoring the intensity of the signal for a constant angle chosen to be close to, but smaller than, \( \theta_{\text{res}} \), in the linear region of the SPR dip. Dependence of HWHM \( (a) \) and \( R_{\text{fixed}} \) \( (b) \) with respect to the cell coverage. Dependence of HWHM \( (c) \) and \( R_{\text{fixed}} \) \( (d) \) with respect to the spreading of cells. The simulations are plotted in dashed curves \( (n_{\text{cell}} = 1.35: \text{diamond}, n_{\text{cell}} = 1.36: \text{squares}, n_{\text{cell}} = 1.37: \text{triangles}) \). Fitting the linear parts in \( (a) \) yielded \( n_{\text{cell}} = 1.3483 \) (by quadratic extrapolation of the slopes). Error bars represent the standard errors.

(TIF)

Figure S7 Simulated LRSPPR reflectivity intensities from the effective-RI model (as per Eq. 1) as a function of cell coverage increase. Different cell refractive indices were implemented: (a) \( n_{\text{cell}} = 1.35 \), (b) \( n_{\text{cell}} = 1.36 \) and (c) \( n_{\text{cell}} = 1.37 \). The short-dashed curve represents the situation of the cover medium composed only by PBS and the long-dashed curve show the spectrum for a hypothetical monolayer of contiguous cells. The behaviours are close to those that one can find for bulk RI changes. The main contributor of the increase \( R_{\text{min}} \) is the RI mismatch between the cover medium and the polymer interlayer (Cytop) in the LRSPPR structure. This translates by the coupling efficiency of the light into the plasmons departing from its optimum with increasing values of the cell coverage.

(TIF)

Figure S8 Simulated LRSPPR reflectivity intensities from the averaged-RI model (as per Eq. 1) as a function of cell coverage increase. Different cell refractive indices were implemented: (a) \( n_{\text{cell}} = 1.35 \) and (b) \( n_{\text{cell}} = 1.37 \). The short-dashed curve represents the situation of the cover medium composed only by PBS and the long-dashed curve show the spectrum for a hypothetical monolayer of contiguous cells. A dip splitting is observed for \( n_{\text{cell}} \) as low as 1.35, which precludes any rational use of this model and confirms the hypothesis that LRSPPR models should involve an effective refractive index.

(TIF)

Figure S9 Comparison of experimental LRSPPR HWHM and \( R_{\text{fixed}} \) to those predicted by the effective-RI LRSPPR model. The parameter \( R_{\text{fixed}} \) represents the readable signal of typical biosensing experiments, it is obtained by monitoring the intensity of the signal for a constant angle chosen to be close to, but smaller than, \( \theta_{\text{res}} \), in the linear region of the SPR dip. Dependence of HWHM \( (a) \) and \( R_{\text{fixed}} \) \( (b) \) with respect to the cell coverage. Dependence of HWHM \( (c) \) and \( R_{\text{fixed}} \) \( (d) \) with respect to the spreading of cells. The simulations are plotted in dashed curves \( (n_{\text{cell}} = 1.35: \text{diamond}, n_{\text{cell}} = 1.36: \text{squares}, n_{\text{cell}} = 1.37: \text{triangles}) \). The linear fitting in \( (a) \) yielded \( n_{\text{cell}} = 1.360 \). Error bars represent the standard errors.

(TIF)

Appendix S1 Relationship between the plasmonic waves and the cells.

(DOC)

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Author Contributions

Conceived and designed the experiments: RM BT. Performed the experiments: RM. Analyzed the data: RM BT. Contributed reagents/materials/analysis tools: BT. Wrote the paper: RM BT.
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References

1. Taylor AD, Yu Q, Chen S, Homola J, Jiang S (2005) Comparison of E. coli O157:H7 preparation methods used for detection with surface plasmon resonance sensor. Sensors and Actuators B: Chemical 107: 202–208.

2. Mijard R, Grieser HJ, Thierry B (2014) Optical biosensing for label-free cellular studies. TRAC Trends in Analytical Chemistry.

3. Milgram S, Bombora R, Livache T, Roupin M (2012) Antibody microarrays for label-free cell-based applications. Methods 56: 326–333.

4. Yanase Y, Hiragun T, Yanase T, Kawaguchi T, Ishii K, et al. (2012) Evaluation of peripheral blood basophil activation by means of surface plasmon resonance imaging. Biosensors and Bioelectronics 32: 62–68.

5. Stojanovic I, Lirtsman V, Golosovsky M, Davidov D, Aroeti B (2010) Real-time monitoring of multi-parametric cell profiling. Biosensors and Bioelectronics 25: 63–70.

6. Shevchenko Y, Camei-Ural G, Cuttica DF, Dokmeci MR, Albert J, et al. (2014) Surface plasmon resonance fiber sensor for real-time and label-free monitoring of cellular behavior. Biosensors and Bioelectronics 56: 359–367.

7. Michaelis S, Wegener J, Robelek R (2013) Label-free monitoring of cell-based assays: Combining impedance analysis with SPR for multiparametric cell profiling. Biosensors and Bioelectronics 49: 63–70.

8. Chabot V, Miron Y, Charrette PG, Grandbois M (2013) Identification of the molecular mechanisms in cellular processes that elicit a surface plasmon resonance (SPR) response using simultaneous surface plasmon-enhanced fluorescence (SPEF) microscopy. Biosensors and Bioelectronics 50: 125–131.

9. Golosovsky M, Lirtsman V, Yaselnysky V, Davidev D, Aroeti B (2009) Multifrequency surface-plasmon resonance: A novel biophysical tool for studying living cells. Journal of Applied Physics 105: 102006–102011.

10. Ziblat R, Lirtsman V, Davidev D, Aroeti B (2006) Infrared Surface Plasmon Resonance: A Novel Tool for Real Time Sensing of Variations in Living Cells. Biophysical Journal 90: 2592–2599.

11. Yaselnysky V, Lirtsman V, Golosovsky M, Davidev D, Aroeti B (2010) Real-Time Monitoring of Epithelial Cell-Cell and Cell-Substrate Interactions by Infrared Surface Plasmon Spectroscopy. Biophysical Journal 99: 4028–4036.

12. Ramdun JJ, Horvath R (2009) Optical biosensors for cell adhesion. Journal of Receptors and Signal Transduction 29: 211–223.

13. Ramdun JJ, Li SY, Heinzle E, Prenosil JE (1993) Kinetics of adhesion and spreading of animal cells. Biotechnology and Bioengineering 45: 939–945.

14. Ramdun JJ, Li SY, Heinzle E, Prenosil JE (1994) Optical method for measurement of number and shape of attached cells in real time. Cytometry 19: 97–102.

15. Vittal T, Granqvist N, Hallala S, Ravinta M, Viippiranta M (2013) Elucidating the Signal Response of Multi-Parametric Surface Plasmon Resonance Living Cell Sensing: A Comparison between Optical Modeling and Drug MDCK Cell Interaction Measurements. PLoS ONE 8: e72192.

16. Yanase Y, Suzuki H, Tatsuki T, Hiragun T, Kameyoshi Y, et al. (2007) The SPR signal in living cells reflects changes other than the area of adhesion and the formation of cell constructions. Biosensors and Bioelectronics 22: 1081–1086.

17. Mijard R, Dostalek J, Huang C-J, Grieser H, Thierry B (2013) Tuneable and robust long range surface plasmon resonance for biosensing applications. Optical Materials 35: 2507–2513.

18. Wang Y, Huang C-J, Jonas U, Wei T, Dostalek J, et al. (2010) Biosensor based on hydrogel optical waveguide spectroscopy. Biosensors and Bioelectronics 25: 1663–1668.

19. Huang C-J, Mi L, Jiang S (2012) Interactions of alginate-producing and -deficient Pseudomonas aeruginosa with zwitterionic polymers. Biomaterials 33: 3626–3631.

20. Vila M, Etheridge S, Rauch JA, Homola J (2009) Long-range surface plasmons for sensitive detection of bacterial analytes. Sensors and Actuators B: Chemical 139: 59–63.

21. Lee JN, Park C, Whitesides GM (2003) Solvent Compatibility of Poly(dimethyl-siloxane)-Based Microfluidic Devices. Analytical Chemistry 75: 6544–6554.

22. Dostalek J, Kaary A, Knoll W (2007) Long Range Surface Plasmons for Observation of Biomolecular Binding Events at Metallic Surfaces. Plasmonics 2: 97–106.

23. Liang XJ, Liu AQ, Lin CS, Ayi TC, Yap PH (2007) Determining refractive index of single living cell using an integrated microchip. Sensors and Actuators A: Physical 133: 349–354.

24. Bae R, Joseph S (1954) Refractometry of Living Cells: Part I. Basic Principles. Quarterly Journal of Microscopical Science 55-95: 399–423.

25. Beuthan J, et al. (1996) The spatial variation of the refractive index in biological cells. Physics in Medicine and Biology 41: 369.

26. Boss D, Kuhn J, Depursinge C, Magistrati J, Marquet P (2012) Quantitative measurement of absolute cell volume and intracellular integral refractive index (RI) with dual-wavelength digital holographic microscopy (DHM). 84270B–84270B.

27. Giebel KF, Bechinger C, Herminghaus S, Riedel M, Leiderer P, et al. (1999) Imaging of Cell/Substrate Contacts of Living Cells with Surface Plasmon Resonance Microscopy. Physical Journal 76: 509–516.

28. Voos J, Graf R, Knauf GL, Brinink M, Mayer J, et al. (2000) Feasibility study of an online toxicological sensor based on the optical waveguide technique. Biosensors and Bioelectronics 15: 423–429.

29. Cotter K, Horvath R (2008) Imageless microscopy of surface patterns using optical waveguides. Applied Physics B: Lasers and Optics 91: 319–327.

30. Jenkins JTA, Buckling A, McGehee M, interfer-constant RH (2003) Surface plasmon resonance shows that type IV pilis are important in surface attachment by Pseudomonas aeruginosa. Journal of The Royal Society Interface 2: 253–259.

31. Toma M, Knoll W, Dostalek J (2012) Bragg-Scattered Surface Plasmon Microscopy: Theoretical Study. Plasmonics 7: 293–299.

32. Berger CEH, Kooyman RPH, Greve J (1999) Surface plasmon propagation near an index step. Optics Communications 167: 183–189.