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Effects of photon reabsorption phenomena in confocal micro-photoluminescence measurements in crystalline silicon

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Confocal micro-photoluminescence (PL) spectroscopy has become a powerful characterization technique for studying novel photovoltaic (PV) materials and structures at the micrometer level. In this work, we present a comprehensive study about the effects and implications of photon reabsorption phenomena on confocal micro-PL measurements in crystalline silicon (c-Si), the workhorse material of the PV industry. First, supported by theoretical calculations, we show that the level of reabsorption is intrinsically linked to the selected experimental parameters, i.e., focusing lens, pinhole aperture, and excitation wavelength, as they define the spatial extension of the confocal detection volume, and therefore, the effective photon traveling distance before collection. Second, we also show that certain sample properties such as the reflectance and/or the surface recombination velocity can also have a relevant impact on reabsorption. Due to the direct relationship between the reabsorption level and the spectral line shape of the resulting PL emission signal, reabsorption phenomena play a paramount role in certain types of micro-PL measurements. This is demonstrated by means of two practical and current examples studied using confocal PL, namely, the estimation of doping densities in c-Si and the study of back-surface and/or back-contacted Si devices such as interdigitated back contact solar cells, where reabsorption processes should be taken into account for the proper interpretation and quantification of the obtained PL data. *Published by AIP Publishing.*

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I. INTRODUCTION

Photoluminescence (PL) spectroscopy has proven to be a powerful tool for characterizing silicon wafers and solar cells.¹ Its most important applications include: the analysis of defects,² the study of passivation layers,³ the determination of open circuit voltages,⁴ and the calculation of charge carrier diffusion lengths⁵ and charge carrier lifetimes.^{2,6} In fact, quasi-steady state photoluminescence (QSSPL) has been up to now one of the most employed techniques to determine the injection dependent charge carrier lifetime,⁷ a paramount parameter to assess the quality of c-Si wafers and photovoltaic (PV) materials. In 2006, Trupke reported a detailed and interesting study about the influence of photon reabsorption on QSSPL measurements on crystalline silicon (c-Si).⁸

Confocal micro-PL spectroscopy has become during the last few years an important experimental tool for developing novel and comprehensive studies of advanced solar cell concepts⁹ with micron resolution. In particular, micro-PL measurements have been extensively used to perform high-resolution spatially resolved studies of different properties and material features, including the analysis of micro-structural defects,¹⁰ the estimation of doping densities,^{11,12}

and the detection of precipitates and impurities.¹³ In confocal micro-PL measurements, a laser light is focused onto the surface of the sample to be studied by microscope objectives, and the backscattered PL signal is collected via a confocal aperture. Thus, according to the size of the selected pinhole aperture, only the photons that pass near the focal volume are collected.¹⁴ By definition, the reabsorption probability is dependent on the optical path length of spontaneously emitted photons before their collection, i.e., photons emitted further from the collection point have higher probability to be reabsorbed.⁸ Thus, since the detection volume in confocal systems depends on the experimental parameters and setup, the reabsorption is partially linked with the specific experimental conditions. As we will see, this fact has important implications in the study of c-Si.

In this work, we present a comprehensive study about the effect of photon reabsorption in micro-PL spectroscopy measurements in c-Si. The aim of the study is: (a) to analyze how the reabsorption phenomenon is affected by the different configurations and selected experimental parameters and (b) to study and quantify how reabsorption processes affect the emission line-shape of the resulting micro-PL spectra. The obtained experimental results are supported and strengthened by theoretical calculations, which estimate the flux of emitted photons escaping from a c-Si sample. Finally, in order to prove the importance of this study, we present

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two practical examples performed with micro-PL, namely, (a) the estimation of high doping densities in c-Si at room temperature and (b) the study of rear-surfaces of PV devices, where reabsorption phenomena must be taken into account for the proper interpretation and quantification of the obtained PL data.

II. EXPERIMENTAL DETAILS

Room-temperature confocal micro-PL microscopy experiments were carried out with a Witec Alpha300 system. A diagram of the Witec setup can be found elsewhere.¹⁵ A diode-pumped laser with a wavelength of 532 nm was used as the excitation source, resulting in a light penetration depth of about 1 μm . Two different Plan Achromat microscope objectives (Olympus 50 $\times$, Numerical Aperture (NA) = 0.8 and Olympus 20 $\times$, NA = 0.45) optimized for near-infrared (NIR) detection were used to focus the laser light beam onto the sample surface. The incoming power density on the sample surface using the 20 $\times$ objective was estimated to be $6 \times 10^4 \text{ W/cm}^2$ approximately. The backscattered PL signal was collected by the same lenses, and three different optical fibers (50, 200, and 1500 μm in diameter), acting as pinhole apertures, were employed to relay the collected PL signal to the monochromator and its associated detector. It is worth mentioning that when using the 1500 μm -diameter pinhole, confocal conditions are not fulfilled. The PL signal was recorded through a 300-mm spectrometer composed of two gratings (150 and 300 lines/mm) and a 1024×1 pixel linear InGaAs photodiode array optimized for spectroscopy applications in the range 1–1.7 μm .

Most of the measurements in this work have been performed on calibrated commercial p-type (boron doped) Czochralski (Cz) c-Si wafers. However, other more complex sample architectures have been also employed for specific measurements. For those cases, the details of the different samples are described at the time the measurements are presented along the document.

III. THEORY OF PL

Photoluminescence emission can be quantified in both direct and indirect semiconductors using the generalized Planck's law, which accounts for non-thermal radiation by assigning a chemical potential to the emitted photons.¹⁶ This chemical potential is related to the splitting of quasi-Fermi levels of electrons and holes ($\Delta\mu$). Thus, in semiconductors such as Si, the rate of spontaneous emission dr_{sp} per photon energy interval $d(\hbar\omega)$ and volume element that contributes to the PL photon flux is described by

$$dr_{sp}(z, \hbar\omega) = \frac{\alpha(\hbar\omega)(\hbar\omega)^2}{4\pi^3 c^2 \hbar^3} \times \frac{1}{\exp\left[\frac{\hbar\omega - \Delta\mu(z)}{kT}\right] - 1} d(\hbar\omega), \quad (1)$$

where $\hbar\omega$ is the energy of the emitted photons, $\alpha(\hbar\omega)$ is the absorption coefficient of the material at the emission energy, c is the velocity of light, k is the Boltzmann constant, and T

is the temperature. Note that the separation of the quasi-Fermi levels $\Delta\mu$ is spatially dependent within the sample.

For calculating the emitted photon flux dj_{em} , reabsorption effects on the way to the exiting surface must be considered, then an exponential absorption term should be added to Eq. (1) resulting in

$$dj_{em}(z, \hbar\omega) = dr_{sp} \exp[-\alpha(\hbar\omega)z] T_r, \quad (2)$$

where z is the distance between the volume element where dr_{sp} takes place and the surface from which the PL is detected, and T_r is the transmittance of such a surface.

Equation (2) describes the spontaneous emission rate that contributes to the photon flux in a positive z -direction. However, when the photons reach the exiting surface, part of them are reflected back in the negative z -direction. Once this reflected photon flux reaches the opposite surface the same effect occurs, part of the photons are transmitted and part of them are reflected back to the material, where they can be partially reabsorbed. This mechanism happens continuously and is known as multi-reflection.

A schematic representation of this process is illustrated in Fig. 1. Having this sketch in mind, the total multi-reflected photon flux density contributing to the detected PL can be determined by the sum of the individual flux densities according to

$$\begin{aligned} dj_{em,MR}(z, \hbar\omega) &= dr_{sp}(z, \hbar\omega) e^{-\alpha z} T_F (1 + e^{-2\alpha t} R_B R_F \\ &\quad + e^{-4\alpha t} R_B^2 R_F^2 + \dots) \\ &\quad + dr_{sp}(z, \hbar\omega) e^{-\alpha(2t-z)} T_F R_B \\ &\quad \times (1 + e^{-2\alpha t} R_B R_F + e^{-4\alpha t} R_B^2 R_F^2 + \dots) \\ &= (dr_{sp}(z, \hbar\omega) e^{-\alpha z} T_F + dr_{sp}(\hbar\omega) \\ &\quad \times e^{-\alpha(2t-z)} T_F R_B) \\ &\quad \times (1 + e^{-2\alpha t} R_B R_F + e^{-4\alpha t} R_B^2 R_F^2 + \dots), \end{aligned} \quad (3)$$

where t is the wafer thickness, T_F is the transmittance at the front surface, and R_F and R_B are the reflection coefficients at front and back side of the silicon wafer, respectively. Then, applying the conversion for a geometric progression sum: $a + ar + ar^2 + ar^3 = \frac{a}{1-r}$, we obtain

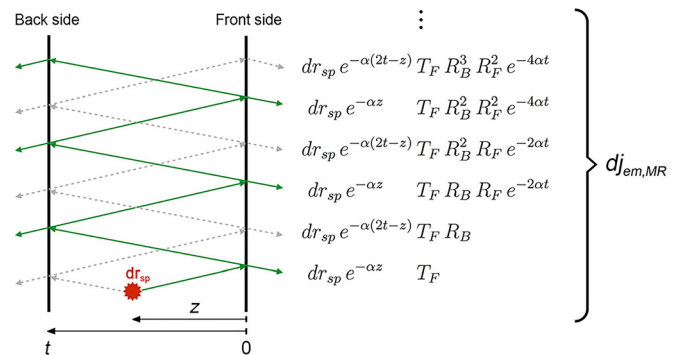


FIG. 1. Sketch illustrating the resulting photon flux of a spontaneous emission dr_{sp} situated at a distance z from the front surface in a wafer of thickness t . The total photon flux dj_{em} takes into account reabsorption and multi-reflection mechanisms.

$$dj_{em,MR}(z, \hbar\omega) = dr_{sp}(\hbar\omega) T_F \frac{e^{-\alpha z} + e^{-\alpha(2l-z)} R_B}{1 - e^{-2\alpha l} R_B R_F}. \quad (4)$$

Equation (4) describes the emission rate from a specific volume element located at a distance z from the front surface. The total emitted PL spectrum from all emission centers is obtained by integrating Eq. (4) over the thickness of the wafer.

IV. RESULTS AND DISCUSSION

A. Photon reabsorption in micro-PL measurements

Figure 2(a) shows the normalized PL emission spectra resulting from Eq. (2) for different z distances from the front surface (see Fig. 1). As it can be seen, the variation in reabsorption for the different spectra is evidenced by a decrease in PL intensity in the low wavelength range, i.e., 1000–1100 nm. As suggested by Trupke,⁸ this is correlated with the increase in the Si absorption coefficient at lower wavelengths (see the black curve of Fig. 2(a), which represents the Si absorption coefficient¹⁷ along the PL emission wavelength range). Thus, emitted photons with higher (lower) energies (wavelengths) have a higher probability to be reabsorbed. Moreover, another important observation in Fig. 2(a) is that the level of reabsorption increases with increasing z distances. Photons emitted at longer z distances with respect to the front surface (see Fig. 1) have associated

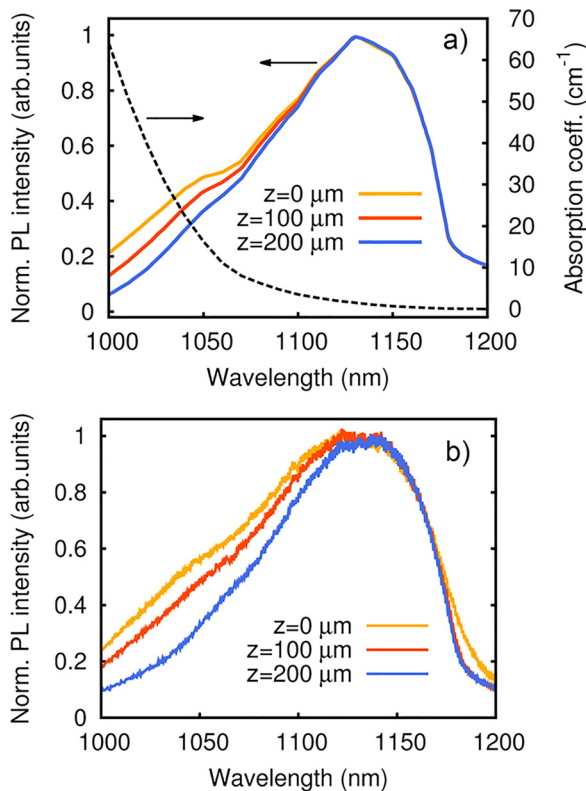


FIG. 2. (a) Normalized PL spectra calculated using Eq. (2) for different values of z distance (see Fig. 1). The absorption coefficient (black dashed line) for c-Si along the represented wavelengths is also shown. (b) Normalized confocal micro-PL spectra recorded from a commercial p-type c-Si wafer at different depth distances from the front surface.

a higher reabsorption probability, as they travel longer path lengths before collection.

Figure 2(b) shows the normalized confocal micro-PL spectra recorded in the backscatter mode from a p-type c-Si wafer by changing collection point distance from the front surface, i.e., $z = 0$ (on focus), 100, and 200 μm . The wide PL emission with a peak maximum at 1130 nm is linked to the phonon-assisted indirect radiative transitions in c-Si. As it can be seen, the reabsorption level increases with increasing the depth distance, and the PL spectra of Fig. 2(b) are qualitatively the same as the calculated spectra of Fig. 2(a). This proves the consistency of the theoretical model presented in Section III.

It is worth noting that the experimental results presented in Fig. 2(b) were obtained by changing the distance between the objective lens and the sample surface. This experimental procedure allows us, as previously mentioned, to collect the backscattered PL signal at different depths. In addition, it also impacts the illumination power density impinging on the silicon surface. The effect of the power density can be evidenced in the PL spectrum at lower wavelengths (1180–1200 nm), where the PL intensity slightly decreases when the objective lens, initially focalized at the sample surface ($z = 0 \mu\text{m}$), is moved out of focus ($z = 100$ and $200 \mu\text{m}$). However, it is important to stress that the change in power density taking place when the objective focal point is moved from $z = 0$ to 100 or 200 μm does not have any influence on the reabsorption variations observed in the range 1000–1100 nm, which are exclusively linked to the optical collection at different z locations. Indeed, this last assumption was confirmed by confocal micro-PL measurements (data not illustrated), where the collection depth was kept at $z = 0 \mu\text{m}$ and the laser power was reduced two orders of magnitude. No change in the collected PL spectra was evidenced in the short wavelength region (1000–1100 nm) after normalization of the spectra; however, as previously observed, only a decrease in the PL intensity was detected around 1180–1200 nm.

Photon reabsorption phenomena have been highlighted through the capabilities of the confocal microscope to collect the PL signal at different depths. Notably, the NA of the objective and the pinhole size describing the level of confocality play an important role for evidencing reabsorption spectral changes. Some of these points are going to be developed and illustrated below.

In micro-PL spectroscopy, the incident light is focused on the sample surface by microscopic lenses, resulting in typical beam spot sizes of few microns in diameter. The incident light is absorbed in the sub-surface region and notably for silicon the absorption depth varies between 100 nm and 10 μm for characteristic excitation wavelengths of 400 and 785 nm, respectively. As pointed out by Gundel *et al.*,¹⁵ due to the high carrier densities achieved in micro-PL measurements, the diffusion of photo-generated carriers is limited by Auger recombination, and therefore, the charge carrier profile is typically confined close to the illuminated surface.

The depth extension of the detection volume in confocal microscopy, the so-called confocal optical slice thickness (Δz), is given by¹⁸

$$\Delta z = \sqrt{\left(\frac{0.88 \lambda_{em}}{(n - \sqrt{n^2 - NA^2})}\right)^2 + \left(\frac{n \cdot \sqrt{2} \cdot PH}{NA}\right)^2}, \quad (5)$$

where λ_{em} is the emission wavelength, n is the refraction index of the measurement medium, NA is the numerical aperture of the objective, and PH is the pinhole size. The first squared term of Eq. (5), often called the wave-optical term, is constant for a given objective and emission wavelength. In contrast, the second squared term is influenced by the geometric-optical effect of the pinhole aperture and becomes prominent for larger pinhole sizes. Thus, for a given material with a specific λ_{em} and a given illumination configuration with a specific λ_{exc} and NA , the depth extension of the collection volume will be directly related to the selected PH . Having in mind that the reabsorption probability increases with increasing the path length of spontaneously emitted photons, changes in the pinhole size should affect the reabsorption probability.

Figure 3(a) shows normalized PL spectra recorded from a p-type c-Si wafer for three different pinhole sizes, i.e., 50, 200, and 1500 μm . Additionally, a PL spectrum recorded in the transmission mode is also shown. Fig. 3(b) illustrates an approximation of the excitation and detection volumes for different pinhole diameters. The excitation volume (dark red) remains constant, whereas the collection volume (gray) increases progressively for higher pinhole diameters. As it can be seen in Fig. 3(a), the spectrum obtained with the smaller pinhole, i.e., 50 μm , is the one that shows a higher PL intensity in the 1000–1100 nm wavelength range, evidencing a lower degree of reabsorption. In contrast, the

spectrum obtained using the 200 μm pinhole shows a lower PL signal in the mentioned wavelength range; hence, a higher degree of reabsorption takes place. This feature is further enhanced with the 1500 μm pinhole, where the relative reabsorption rate is further increased. As it can be easily deduced from the diagram, the increase in reabsorption is linked with the progressive extension of the collection volume for higher pinhole diameters, where the average travelled distance of the collected photons increases accordingly. The extreme case of this trend corresponds to the PL spectrum obtained in the transmission mode. Two optically aligned objectives were employed in this configuration, where the objective collecting the PL signal was located below the sample. Here, only the photons that cross the entire wafer thickness are detected; hence, the reabsorption probability reaches a maximum level. This last hypothesis is confirmed by measuring in transmission mode c-Si wafers with different thicknesses (data not shown). The result is that the higher the wafer thickness, the higher the reabsorption level.

We have seen so far that the reabsorption probability, and in turn, the line-shape of the obtained PL emission spectra is dependent on the selected experimental configuration. This dependence between the experimental setup and the spectral line-shape of the recorded PL data can have, in certain measurements, important implications. One of these examples is related to the study of hot carrier solar cells, where the so-called carrier temperature and the thermalization factor can be estimated by monitoring the broadening of the high energy tail of the PL band.¹⁹ Another example is the determination of doping densities in c-Si via micro-PL spectroscopy. This latter example is studied in more detail in Section IV B.

B. Doping density quantification via micro-PL spectroscopy

Over the last few years, there have been an increasing number of works focused on the high-resolution assessment of doping densities. This is linked to the interest in determining the doping features of different micro-structures such as selective emitters,²⁰ or laser fired contacts (LFCs),²¹ which are commonly used in advanced c-Si solar cell architectures.

Low temperature PL spectroscopy represents a suitable technique for studying impurity-related transitions and doping features,^{22,23} as for temperatures below a threshold thermal ionization of impurity-bounded carriers are suppressed and novel PL emission lines involving trapped charges set in. However, experiments at low temperature require special equipment such as a cryostat, which is commonly characterized by a small sample volume and often implies specific sample preparation steps and significant amounts of time for the cooling-down process. All these aspects make low-temperature PL studies not suitable for certain applications. Some works have already reported methods for quantifying doping densities in Si via room-temperature PL studies.^{11,24} In these studies, the recorded PL spectra are fitted in order to obtain the reduced band gap energy E_g , which then is correlated with the corresponding doping level. Despite the fact

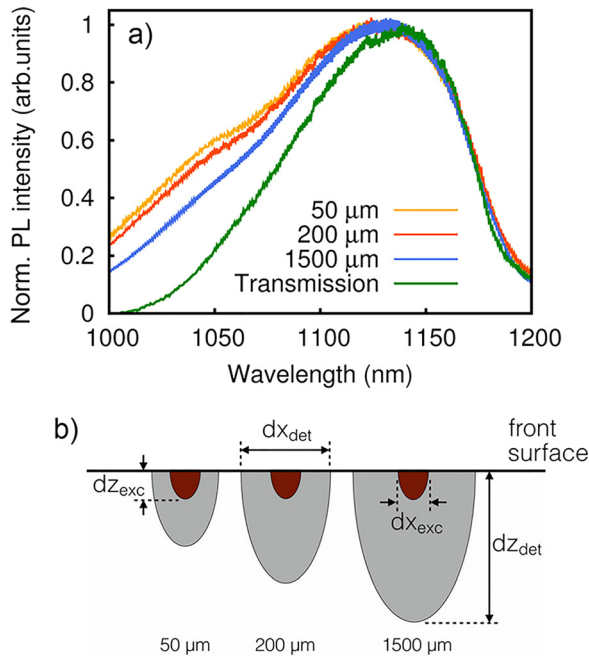


FIG. 3. (a) Normalized PL spectra recorded from a p-type c-Si wafer using three different pinhole diameters, i.e., 50, 200 and 1500 μm . In addition, a fourth PL spectrum obtained in transmission mode, i.e., with the detector located at the opposite surface compared to the excitation surface, is also represented. (b) Diagram representing the change in the detection volume (dx_{det} and dz_{det}) corresponding to three pinhole diameters. The excitation volume (dx_{exc} and dz_{exc}) is kept constant. The diagram is not drawn to scale.

that this methodology can be applied to room-temperature measurements, it is more precise for cryogenic temperature studies where the contribution of phonon energies to indirect radiative transitions is much lower and then the resulting narrower PL lines can be more precisely fitted. As presented elsewhere,¹² we propose an alternative and simple PL method for studying variations in doping densities at room temperature.

Figure 4 shows the normalized PL spectra recorded at room temperature from calibrated p-type c-Si wafers with different doping densities, i.e., 7×10^{17} , 5×10^{18} , 4×10^{19} , and $1 \times 10^{20} \text{ cm}^{-3}$. For doping densities below $7 \times 10^{17} \text{ cm}^{-3}$ (data not shown for the sake of clarity), the PL spectra do not suffer any change at all and they show exactly the same features as the $7 \times 10^{17} \text{ cm}^{-3}$ spectrum. Conversely, for doping densities around $5 \times 10^{18} \text{ cm}^{-3}$ and above, the PL emission band starts to widen progressively towards longer (lower) wavelengths (energies) as a result of a band gap renormalization known as band gap narrowing. An instructive observation is the small but progressive decrease of the PL intensity between 1000 and 1050 nm with increasing doping densities. This fact is linked to an increase in the Si reabsorption coefficient with increasing doping densities,²⁵ proving again the importance of photon reabsorption in confocal micro-PL experiments.

Due to the widening of the PL band for high doping levels, the doping density can be quantified by monitoring the center of mass (CM), i.e., intensity-weighted spectral position, of the PL emission spectrum. The CM is defined as

$$CM = \frac{\sum_{\lambda_1}^{\lambda_2} \lambda \cdot I_\lambda}{\sum_{\lambda_1}^{\lambda_2} I_\lambda}, \quad (6)$$

where λ_1 and λ_2 are the first and last points of the wavelength range for which the CM is calculated, λ is the specific wavelength, and I_λ is the corresponding PL intensity.

Figure 5 shows the dependence of the PL emission band CM as a function of the doping density for two objectives with different NA. The main plot is represented on a semi-log scale, whereas the inset shows the same data on a linear scale.

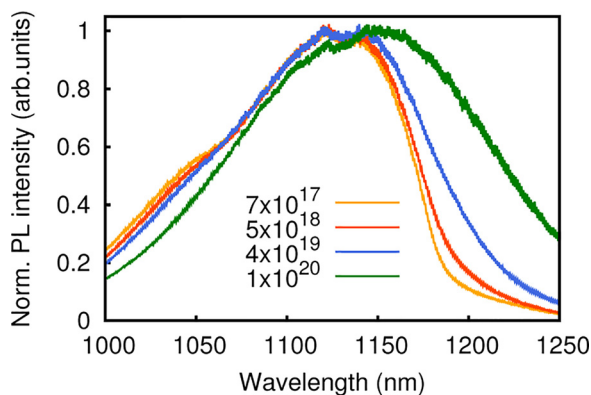


FIG. 4. Normalized PL spectra recorded from p-type c-Si wafers with different doping levels, i.e., 7×10^{17} , 5×10^{18} , 4×10^{19} , and $1 \times 10^{20} \text{ cm}^{-3}$.

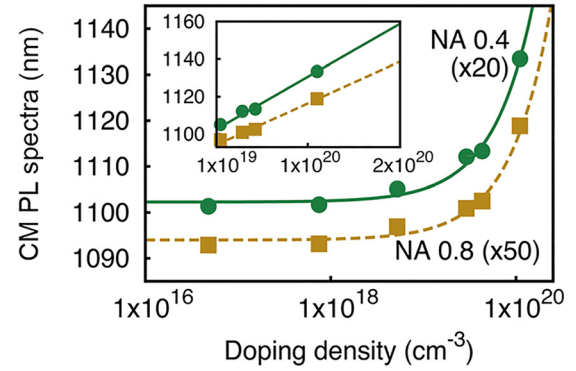


FIG. 5. Correlation between doping density and CM of PL spectra recorded from commercial p-type c-Si wafers for two different objectives, i.e., NA 0.4 and NA 0.8. The main plot is represented on a semi-log scale. The inset shows the same two curves on a linear scale.

scale. Following the trend observed in Fig. 4, we can point out a relatively constant CM for doping densities below 10^{18} cm^{-3} , and conversely an increase of the CM for high doping levels ($>10^{18} \text{ cm}^{-3}$) as a consequence of the expected bandgap narrowing effect. An important feature that should be noted concerns the fact that the two curves are shifted with respect to one another. In a first approach, the origin of this shift can be explained by the detection volume linked with the pinhole and the objective lens used for these measurements. Indeed, the objective lens with lower NA has associated a larger detection volume, and therefore, the higher degree of reabsorption moves the CM of the 0.4-NA curve towards longer wavelengths with respect to the 0.8-NA curve. Taking into account the previous explanation and that the absorption coefficient increases with the doping density,²⁵ it appears that the slope of the curves (see inset) is not identical. Notably, the slope obtained with the 0.4-NA objective lens, which has associated a larger detection volume, is greater than the one obtained with the 0.8-NA objective. The following results suggest that the quantification of doping densities through confocal PL is only valid for a specific experimental configuration, namely, a particular pinhole size, objective lens, and excitation wavelength.

In order to illustrate this point, we have used the two calibration curves depicted in Fig. 5 for quantifying the doping density of a laser doped region (LPR). Gundel *et al.*¹¹ have already employed confocal PL measurements to quantify doping densities in cross-sections of laser-processed point contacts. The PL analysis of the LPR presented here was performed using the two same objectives and experimental configuration employed to obtain the calibration curves presented above. LPRs were fabricated by a 1064 nm-wavelength laser in an $\text{Al}_2\text{O}_3/\text{SiO}_x\text{:B}$ -passivated n-type c-Si wafer. The power and duration of the laser pulse were 1.2 W and 400 ns, respectively. An image of the studied LPR is shown in Fig. 6(a), and the evolution of the CM along the white dashed line is depicted in Fig. 6(b). The two vertical dotted lines delimit the borders of the LPR region. The CM curves are relatively constant in the regions outside the LPR but increase remarkably in the regions inside the processed region, suggesting an increase in the doping density.

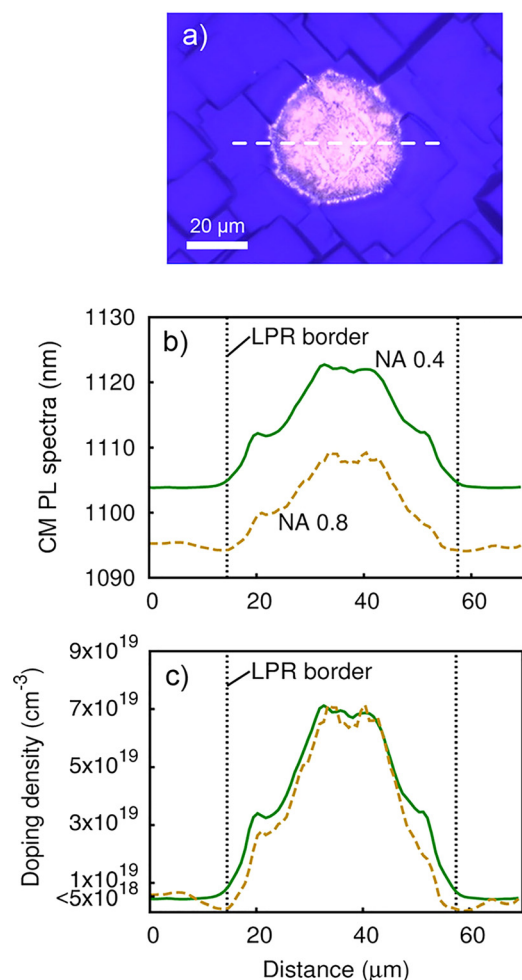


FIG. 6. (a) Picture of a single laser-processed region (LPR). The horizontal dashed line defines the section from which (b) the center of mass and (c) the doping density profiles were obtained. PL data were collected using two different objectives, i.e., NA 0.4 and NA 0.8.

As expected, we observe a shift between the two curves induced by the difference in reabsorption associated with the two objectives. Fig. 6(c) shows the doping density profiles across the LPR obtained by means of the calibration curves represented in Fig. 5. Both profiles show very similar results, notably a doping density value of $7 \times 10^{19} \text{ cm}^{-3}$ at the very center of the LPR, whereas in the regions outside the LPR the doping density remains below $5 \times 10^{18} \text{ cm}^{-3}$. Indeed, the calculated doping concentration outside the LPR region is expected to be the doping concentration of the bulk (n-type, 10^{15} cm^{-3}). However, the method for doping level quantification presented here is performed at room temperature and only allows to evidence high doping densities ($>5 \times 10^{18} \text{ cm}^{-3}$), as illustrated through the calibration curves in Fig. 5.

C. Reabsorption phenomena for the study of back surfaces

Surface reflectance can also impact the reabsorption phenomena and consequently the PL spectrum line-shape. For studying the influence of reflectance in reabsorption, we consider Eq. (4), which includes the effect of multi-reflection in the calculated PL emission. Fig. 7 depicts the PL spectra

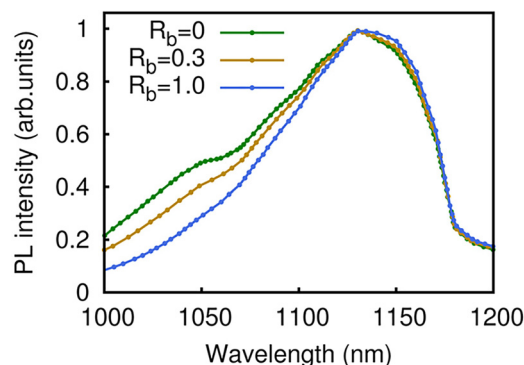


FIG. 7. PL spectra calculated by integrating Eq. (4) for three different values of back surface reflectance (i.e., 0, 0.3, and 1.0).

obtained by integrating Eq. (4) for three different values of the back surface reflectance, i.e., 0, 0.3, and 1. As it can be seen, higher values of reflectance lead to an increase of reabsorption, as revealed by the change in PL intensity between 1000 and 1100 nm. From the experimental point of view, a SiO_2 -passivated p-type c-Si wafer with a thin Au layer deposited onto the back surface was studied by micro-PL. The Au layer (150 nm-thick) was deposited by sputtering with the help of a mask. Fig. 8(a) shows a picture of the sample back surface with the circular Au film covering part of the sample. Note that for carrying out the measurements, the surface with the Au deposit was placed at the back side, while the backscattering PL signal was collected scanning the front side of the sample.

The integrated PL intensity map (1000–1200 nm spectral range) of the selected area under study is represented in Fig. 8(b). As it can be seen, the Au region has associated a higher PL intensity, indicating a higher collection of spontaneously emitted photons, which we correlate with a higher reflectance of the Au region in the near-infrared (NIR) compared to the Si region. The inset in Fig. 8(c) confirms this assumption through spectral reflectance measurements performed in the same sample under the same experimental configuration, i.e., Au layer placed at the back surface and illuminating the front side. The effect in reabsorption can be observed in the main plot of Fig. 8(c), where the normalized and averaged PL spectra corresponding to both regions, i.e., inside and outside of the Au region, are represented.

By comparing the PL intensity in the 1000–1100 nm spectral range, we can identify a different degree of reabsorption, in this case induced by changes in the back surface reflectance. Since the variation in reflectance between the Au and Si areas (see Fig. 8(c) inset) is smaller than the variation between R_b coefficients used to obtain Fig. 7 data (specially in the spectral range where the Si absorption coefficient is higher, i.e., 1000–1050 nm), the changes in reabsorption observed in Fig. 8(c) are slightly smaller than the ones obtained in Fig. 7.

Not only variations in reflectance at the rear surface influence the reabsorption rate in micro-PL measurements, other sample properties such as the surface recombination velocity (SRV) can also have implications in the reabsorption level. The effect of SRV on reabsorption is illustrated in Fig. 9,

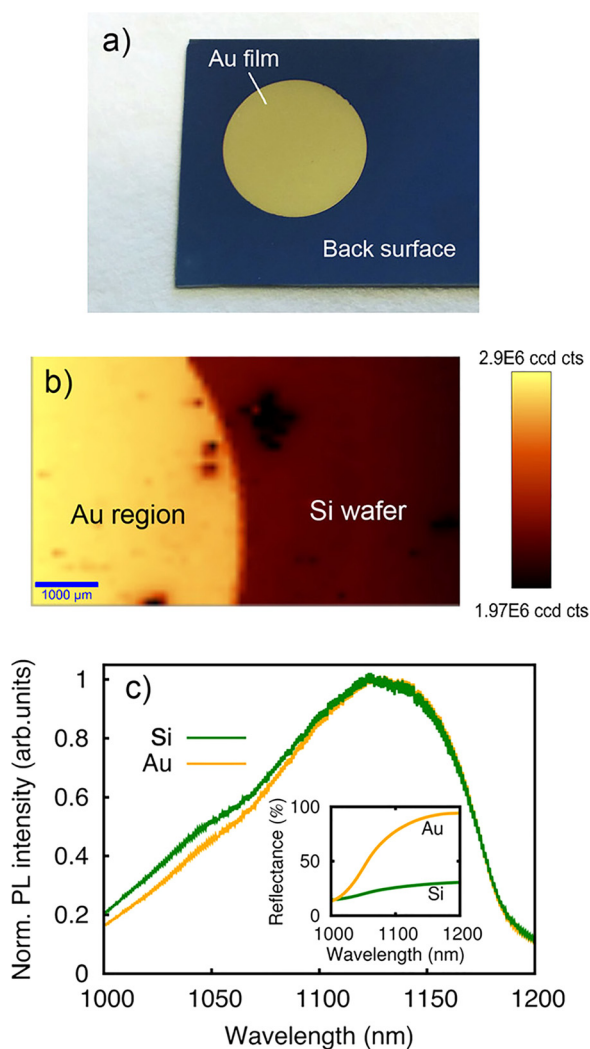


FIG. 8. (a) Picture of the back surface of the measured sample. (b) PL intensity map recorded from the front surface. (c) Normalized averaged PL spectra corresponding to the region inside the Au deposited layer and outside the Au region (c-Si wafer). The inset shows the reflectance of both regions for the spectral range under study.

where a micro-PL spectroscopy study of an Interdigitated Back Contacted (IBC) silicon solar cell is presented. The IBC solar cell was based on a 200 μm -thick 1–5 $\Omega\text{ cm}$ n-type float-zone (FZ) c-Si wafer. The texturized top side was passivated by a $\text{SiN}_x\text{:H/i-a-Si:H}$ stack, which was deposited by plasma enhanced chemical vapor deposition (PECVD) at 200 $^\circ\text{C}$. Both the back-surface field (BSF) and the emitter were processed at the rear side of the cell. An a-Si:H(i)/a-Si:H(n) stack was used to form the BSF, and an a-Si:H(i)/a-Si:H(p) stack was employed for the emitter. Both elements were deposited by the same PECVD system in an interdigitated pattern thanks to the use of a masking process. Between the deposition of the BSF and the emitter, a thin SiO_2 layer was deposited in order to avoid shunting effects. Finally, the IBC solar cell was completed by the transparent conductive oxide (TCO) and metal electrodes deposition. A scheme of the studied IBC solar cell is represented in Fig. 9(a).

Here, the measurements were done using a pinhole size of 1500 μm in order to be more sensitive to the back surface properties, and again, the excitation was done from the top

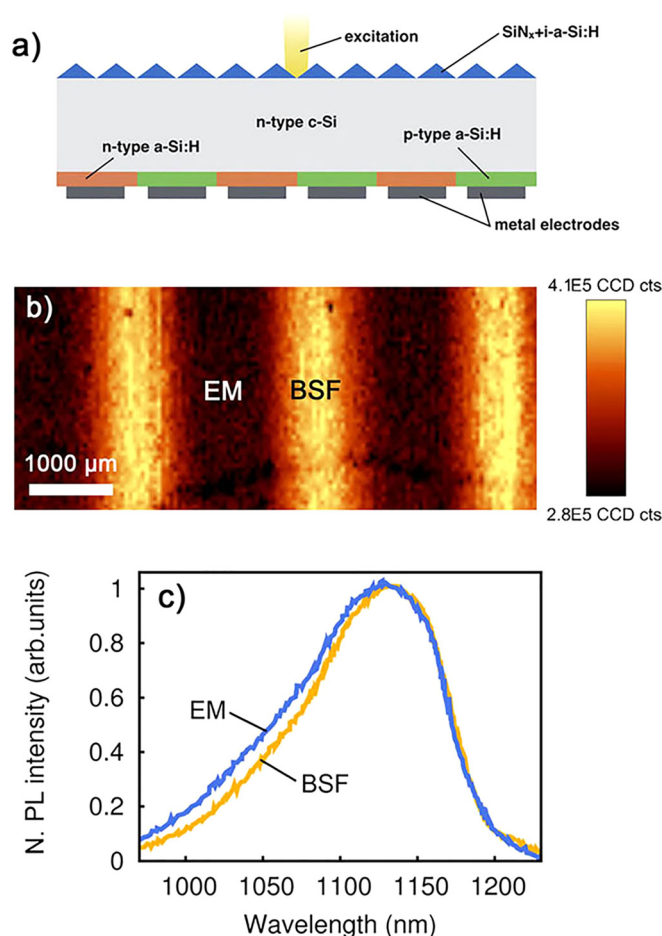


FIG. 9. (a) Scheme of the measured IBC solar cell. (b) Reconstructed PL intensity map of the selected surface area under study. Both emitter and BSF regions are indicated. (c) Averaged and normalized PL spectra corresponding to the emitter and BSF regions observed in (b).

and the PL signal was collected in backscattered mode from the front surface. The reconstructed PL intensity map of the selected IBC solar cell area is represented in Fig. 9(b), where two regions, i.e., emitter and BSF, are clearly identified. Since the front surface passivation is presumably uniform, and the bulk recombination is negligible in front of surface recombination (as it is the case in our high-quality FZ c-Si wafer), the change in PL intensity between the two regions is in principle linked to differences in SRV at the back surface. As it can be seen, BSF regions show a higher PL intensity, then we can conclude that BSF areas offer a better passivation and have associated a lower SRV than emitter regions. Fig. 9(c) shows the averaged and normalized PL spectra correlated with each one of the two regions, i.e., emitter and BSF. Again, PL spectra show variations in the 1000–1100 nm wavelength range, indicating that a higher reabsorption rate takes place while scanning the BSF regions. The lower SRV associated with the BSF regions leads to a decrease of non-radiative recombination and a subsequent increase of the radiative recombination, enhancing the collection of photons coming from the back-surface while BSF regions are scanned. These photons emitted at the back surface have to cross the entire wafer thickness for being collected, and therefore, the reabsorption probability tends to increase. A second mechanism, the so-called

free-carrier absorption, could also be linked with the observed increase of reabsorption. Since BSF regions have associated a lower SRV, the concentration of charge carriers while scanning BSF areas is higher, and then the probability for the emitted photons to be reabsorbed by free carriers should be more favorable. However, for photons with wavelengths within the spectral range of our interest, i.e., 1000–1100 nm, the intrinsic band-to-band absorption is greatly predominant,²⁶ and therefore, the impact of free-carrier absorption can be considered negligible.

V. CONCLUSIONS

A detailed and comprehensive study about the implications of photon reabsorption processes (also known as photon recycling) in confocal micro-PL spectroscopy measurements in c-Si is presented. On the one hand, we have shown that all those experimental parameters that affect the confocality level of a given PL apparatus, i.e., numerical aperture of focusing lenses and pinhole size, have an important influence on the reabsorption rate, leading to changes in the short wavelength range (i.e., 1000–1100 nm) of the PL emission spectra at room temperature. This was demonstrated in the study of highly doped LPRs in c-Si via micro-PL spectroscopy, where the doping level was quantified by monitoring the intensity-weighted spectral position of the recorded PL spectra. Only by taking into account reabsorption phenomena and using the specific calibration curve according to the used experimental setup, a same doping density in the order of $7 \times 10^{19} \text{ cm}^{-3}$ could be obtained from PL data recorded using different setups. On the other hand, we have also shown that certain sample features such as the reflectance or the back surface recombination velocity (SRV) have also implications in reabsorption phenomena, and therefore, influence the obtained PL spectra. This has been demonstrated in the study of an IBC silicon solar cell structure, where the variation in SRV between the emitter and BSF has been observed to induce a change on the reabsorption rate between both regions. This work proves that reabsorption is an intrinsic mechanism in confocal micro-PL spectroscopy measurements, and therefore, it should always be taken into account for the correct interpretation of confocal micro-PL data.

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