Angle insensitive enhancement of organic solar cells using metallic gratings

Aimi Abass,1,a Honghui Shen,2 Peter Bienstman,2 and Bjorn Maes3
1Department of Electronic and Information Systems (ELIS), Ghent University, Sint-Pietersnieuwstraat 41, B-9000 Gent, Belgium
2Photonics Research Group (INTEC), Ghent University-imec, Sint-Pietersnieuwstraat 41, B-9000 Gent, Belgium
3Micro- and Nanophotonic Materials Group, Institut de Physique, University of Mons, 20 Place du Parc, B-7000 Mons, Belgium

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We explore the optical enhancement of organic photovoltaic cells by incorporating a metallic grating as the back contact. We numerically demonstrate a strongly enhanced light absorption exploiting a complex interplay between multiple electromagnetic wave phenomena, among which surface plasmon polariton (SPP) resonances, waveguide mode resonances, Fabry–Perot modes, and scattering. We focus on a triangular grating structure and describe the particular opportunities to obtain a good angular performance. In addition we introduce a novel multiperiodic geometry that incorporates multiple types of SPP resonances. Our triangular structure shows an increased absorption of 15.6% with the AM1.5G spectrum in the 300–800 nm wavelength range. For the multiperiodic grating case a significant further increase to 20.7% is shown. © 2011 American Institute of Physics.

I. INTRODUCTION

Nowadays, organic solar cells are still restricted to thin films on the order of 200 nm for optimal electronic properties,1 which makes efficient photon management an important issue.2–4 Indeed the typical absorption spectrum of polymer films has room for improvement, usually in the infrared range. In addition, even thinner active layers, on the order of 100 nm, would be interesting as this leads to better electrical properties.

An interesting novel technique to boost the absorption of thin film solar cells, involves the use of metallic structures with surface plasmon polariton (SPP) resonances.5 These modes can achieve a strong confinement and scattering of the incoming light. In addition one can engineer the modes so that the absorption in the metal remains limited. Several metal geometries (gratings, particles6–9) and their locations in the cell (front, middle, or back) are under active investigation for different types of solar cells. Plasmonic grating structures5,10–15 form a particularly interesting class, as they involve the patterning of an available metallic electrode.

In this work, we focus on triangular grating structures for the metallic back contact (Fig. 1). These type of gratings are less studied than the square gratings, however we will show that the usage of these triangular gratings can give great absorption enhancement. Several lithographic methods (nanoimprint, optical or electron beam lithography) for metallic grating structures can be used to produce these types of gratings.16–18 The chosen metal back contact material is silver, as it allows for low metal absorption losses, and beneficial SPP resonance wavelengths. An example of the usage of silver square gratings to enhance silicon thin film solar cells is shown in recent publications.12,15 Here, we explore the characteristics further and we examine specifically the angular behavior of the excited modes for bulk heterojunction solar cells made of poly(3-hexylthiophene):[6,6]-phenyl-C61-butyric acid methyl ester (P3HT:PCBM) blends. The depth of the metallic grating is found to be crucial for a good wide-angle response for the case of normal triangular gratings. We show that an even greater absorption enhancement is achieved through more complex triangular gratings. We also investigate the use of two-dimensional (2D) pyramidal gratings. The design strategies we develop here are applicable to various cell materials.

Section II describes the numerical procedure and the initial strategy for boosting the absorption. Section III is the main part and discusses the use of various resonances in our devices.

II. METHOD

We simulate incoming light interaction with our solar cell using the finite element method, as implemented in the COMSOL MULTIPHYSICS software package. The bulk heterojunction P3HT:PCBM polymer is considered as a homogeneous material with a dispersive refractive index. Our refrac-
tive index data is obtained from ellipsometry measurements done at imec (Interuniversity Microelectronics Center, Leuven, Belgium). The refractive index data of the P3HT:PCBM blend is shown in Ref. 6. Due to the periodic nature of the problem, only one period needs to be simulated by using the proper boundary conditions. In order to study angular dependence, we employ Floquet–Bloch conditions on the cell boundaries.20 The top and bottom part of the computational domain is terminated with absorbing perfectly matched layer regions.

We are particularly interested in the absorption enhancement within the polymer itself. To quantify this we integrate the divergence of the Poynting vector $\vec{S}$ within the polymer region only and normalize it to the incoming power over one period $P_{inc}$.

$$A(\lambda) = \frac{\int (\nabla \cdot \vec{S}) dV_{\text{polymer}}}{P_{inc}}. \quad (1)$$

To quantify the absorption capability of our solar cell structure in the wavelength region of interest (300–800 nm), we define a normalized weighted integrated absorption for the AM1.5G spectrum:

$$A_{int} = \frac{\int \left[ P_{AM1.5G}(\lambda) \times A(\lambda) \right] d\lambda}{\int P_{AM1.5G}(\lambda) d\lambda}. \quad (2)$$

The choice of the wavelength region is connected to the P3HT:PCBM polymer blend absorption properties. For this material, we know that charge creation through photon absorption is still possible even up to the wavelength of 800 nm.21

Our approach in optimizing the triangular gratings starts from looking at planar solar cell structures. $A_{int}$ as a function of film thickness for normal incidence is shown in Fig. 2(a). The oscillation is due to interference or Fabry–Perot effects in the active layer. Several absorption spectra are shown in Fig. 2(b). The planar data provides us with approximative guidelines for the grating structure. First, the effective thickness decreases when introducing triangles, thus to obtain an approximate, optimum Fabry–Perot condition with the grating, we are guided to choose $T$ in the range 60–150 nm or in the range 200–250 nm. This will bring the effective thickness closer the first or second Fabry–Perot maximum, respectively. Second, the absorption spectra [Fig. 2(b)] indicate where we should position extra resonances. Indeed, between 620 and 800 nm the absorption is limited so a boost is needed.

III. ENHANCEMENT TECHNIQUES

We present several plasmonic grating enhancement techniques, spread over four categories. Section III A looks into the use of triangular grating SPP modes. Section III B employs polymer waveguide modes in thicker active layers. Section III C discusses a more complex grating device to access strongly localized SPP modes. Finally, Sec. III D shows results on pyramidal gratings to obtain a polarization independent absorption enhancement.

A. Using SPP modes

Here we show how one can strategically place plasmon resonances in order to achieve an angle independent enhancement. The resonant wavelengths of the modes depend strongly on the geometry of the grating. The grating periodicity induces folding of the band structure which determines strongly where the resonances are located.22,23 Full-field mode-solvers can be employed to find the bands, but in the case of shallow gratings we obtain a good estimate simply by folding the dispersion of the flat interface (silver–polymer). The folding procedure is performed in Fig. 3(a) with a periodicity $P=300$ nm.

This dispersion folding procedure should be compared with the rigorous finite element simulations in Figs. 3(b) and 3(c), where we show the absorption spectrum at different angles of incidence for a shallow grating structure with thickness $T=150$ nm, period $P=300$ nm, and height $H=60$ nm. The absorption spectra in this section are for TM polarization (meaning the magnetic field only has a component in the z direction of Fig. 1), unless mentioned otherwise. Figure 3(b) has fill-factor FF=1 (base length of triangle divided by the period). A clear correspondence is noticed between the crossings around 660 nm in Figs. 3(a) and 3(b).

The structure of Fig. 3(c), which has FF=0.5, shows a significantly different dispersion. There is strong splitting of the modes but the central wavelength is still around 660 nm. In addition, one of the modes (the “bright” mode) can be excited at normal incidence, while the other mode (the
The splitting is highly dependent on the grating geometry, more specifically on the fill-factor FF and triangle height H. The effects of the two geometrical parameters can be traced by examining the Fourier expansion of the grating profile.\(^{24}\) It can be shown that at FF=1 [Fig. 3(b)], the second harmonic of the profile Fourier expansion is 0 leading to less splitting of the first order crossing at \(k=0\). In the case shown in Fig. 3, the dark mode has a lower energy than the bright mode and thus appears at a longer wavelength. Depending on the structure in question, the vice versa situation is also possible.\(^{23}\)

As pointed out,\(^{24,25}\) increasing the grating height leads to a flatter SPP dispersion. Indeed, by increasing the grating height we get a more localized behavior of the resonances instead of a propagating behavior. This works well to our advantage in a solar cell design, since we would like to use a resonance that is not angle-dependent. We demonstrate this effect by increasing H in our structure (Fig. 5). Here, for clarity we show the absorption in the metal, instead of in the polymer. The splitting of the dark and bright mode also increases when the grating deepens, as this enhances the difference between the mode field profiles.

Thus, with the proper parameter choice we can utilize both dark and bright modes simultaneously for nonperpendicular incidence. In Figs. 6(a) and 6(b) two SPP peaks at 600 and 650 nm clearly boost the polymer absorption. The optimum incidence angle here is \(\theta=13.5^\circ\). These simulation results [Fig. 6(b)] show an absorption enhancement up to seven times due to the dark mode at 650 nm, from below 10% to nearly 70%. This clearly suggests that most of the SPP resonance absorption occurs in the polymer, instead of in the metal. It corresponds with the SPP mode field profile being well distributed in the polymer [inset in Fig. 6(b)]. Moreover, as the field is not concentrated near the metal interface, possible losses due to exciton quenching near metal surfaces should be reduced.\(^{26}\) Importantly, with the chosen geometric parameters, the absorption in the wavelength range <550 nm remains largely intact.
With larger periods, higher order plasmonic modes can be accessed to boost absorption. In that case the first Brillouin zone becomes narrower and higher order modes can exist in the wavelength region of interest. With larger periods it is also possible to have a diffraction edge in the neighborhood of our enhancement range. However, for our structures the SPP resonances seemed to offer better enhancement. For larger periods with strong interaction between SPP resonances and diffraction, such as in lattice resonances, further studies can be performed.

Figure 7 shows the absorption spectrum at 500 nm grating periodicity for normal incident light. The resonance peaks of the second order are also predicted quite well with simple band folding. Figure 7(b) shows the magnetic field profile at the two peaks indicated with A and B. As would be expected of higher order modes, the field profile of the second order mode is more confined to the metal interface. This makes it more prone to possible exciton quenching.

An optimum structure for wavelengths <650 nm and normal incidence which only utilizes the first order bright mode and Fabry–Perot considerations is given in Fig. 8. The grating structure has just a slight influence on the (already good) absorption below 500 nm. We note that there is also a significant enhancement for TE polarization, with which no plasmonic resonance can be excited. The integrated absorption efficiency $A_{int}$ for a planar unpatterned structure is 48%, while for a patterned structure we reach 63.6% for TM and 56% for TE polarization. We, therefore, see a significant 15.6% increase in $A_{int}$ for the TM case.

**B. Using polymer waveguide modes**

When the polymer thickness is above 200 nm, the grating structure can also excite polymer waveguide modes. This provides access to additional resonances to boost the absorption. The use of waveguide excitation has been demonstrated, however previous discussions mainly concerned silicon solar cells.

Figure 9(a) shows the absorption spectrum in such a thicker cell for $T=250$ nm, $P=400$ nm, $H=50$ nm, $FF=0.5$. The origin of the resonances can still be tracked by simple band folding of the waveguide dispersion curve, see Fig. 9(b) where the folded dispersion relation for a flat structure of the same thickness $T$ is shown. The line in Fig. 9(b) follows the SPP modes, the other modes have their field mostly in the polymer. The mode that starts around 750 nm, at point A in Fig. 9(a), is the bright first

**FIG. 6.** (Color online) Angularly optimized grating with $T=150$ nm, $P=300$ nm, $H=100$ nm, $FF=1$. (a) $A(\lambda)$ vs angle of incidence. The white, dashed line indicates $\theta=13.5^\circ$. (b) $A(\lambda)$ at $\theta=13.5^\circ$ (line with circles) compared to a planar structure at normal incidence (line). The insets show the magnetic field profile at the two peaks indicated with A and B.

**FIG. 7.** (Color online) Employing higher order plasmonic resonances. (a) $A(\lambda)$ of a structure with $P=500$ nm, $T=150$ nm, $H=120$ nm, $FF=0.5$ at normal incidence. (b) The $H_z$ magnetic field profile at the first order and at the second order resonance.

**FIG. 8.** (Color online) $A(\lambda)$ of an optimized structure for normal incidence with $T=150$ nm, $P=400$ nm, $FF=0.5$, and $H=120$ nm.
order plasmon mode. The mode starting at 650 nm and going to point C stems from a polymer waveguide mode. We observe an anticrossing interaction between the waveguide mode and the plasmonic mode, which can happen as they have the same TM polarization. This creates a gap in the dispersion and absorption diagram. It is therefore important to model these interactions, as they might cancel any enhancement in a certain wavelength range. The anticrossing effect is clear from the field profiles in Fig. 9:

For the structure in Fig. 10 we combine the dark and bright plasmonic modes together with a waveguide mode resonance, for $\theta=13.5^\circ$. The absorption spectrum in the wavelength range 600–720 nm is nicely filled by these three resonances, while the absorption below 550 nm remains intact. We obtain an integrated absorption $A_{\text{int}}=71.4\%$ for TM polarization, and 60.3\% for TE polarization. For a simple planar structure with the same $T=250$ nm, $A_{\text{int}}$ is only 56%.

The character of the modes is illustrated in Fig. 10(b). The polymer waveguide mode is peak A. The dark mode (peak B) only appears after a change in angle. Note that in this grating the antisymmetric dark mode appears at a smaller wavelength than the bright mode (peak C). We suspect this is due to the thicker polymer layer, leading to a less confined bright mode with smaller energy.

**C. Using strongly localized modes**

Up to this point we showed that the triangle grating structure gives access to a limited number of SPP and waveguide modes. More numerous and highly localized SPP resonances can be used by adding very small subwavelength features to the structure. Various multiperiodic grating structures have been studied before and are found to have multiple resonances with significant absorption capabilities. In our case we accomplish this by adding small triangular groove defects with a small period on top of our main triangular grating. The main grating periodicity is expected to induce band folding on a dispersion relation which now includes additional coupled localized SPP modes due to the defect grooves. We find that this procedure provides multiple resonances with strong absorption enhancements well into the desired enhancement range.

We show a proof of principle in Fig. 11. Figure 11(a) presents the absorption spectrum of a triangular grating solar cell structure with and without additional groove defects, the inset shows the geometry. We notice that the single plasmonic resonance (peak A) is replaced by four other reso-
It is desirable in solar cells to have a polarization independent efficiency. Therefore, we study (2D) pyramidal grating structures with rigorous three-dimensional (3D) simulations in this section, as opposed to the [one-dimensional (1D)] triangular gratings before where we only needed 2D simulations. Similar enhancement effects are obtained but various mode characteristics differ.

Figure 13 shows results for perpendicular incidence on two solar cell structures with pyramidal gratings of different FF (T = 150 nm, H = 120 nm, P = 320 nm). It is found that band folding still gives a good prediction in this case. The two peaks in Fig. 13 (for both gratings) stem from a first order crossing around 680 nm, see the inset in Fig. 13. Unlike in the 1D triangular grating case (where FF = 1 gives a minimum separation), here the smaller FF gives a smaller separation of the modes. In addition, both modes from the crossing are bright modes.

Figure 14 presents the absorption spectrum for two different periods (T = 150 nm, H = 120 nm, FF = 1). The first and second order modes appear around the wavelength determined by band folding as before. It can be seen that the dominant peaks in the graph occur at the larger grating periods of 550 nm. The peak around 600 nm wavelength in this structure is a second order peak. The peak stemming from
first order crossings (around 740 nm) is considerably weaker than the second order peak. The explanation of these differences is a topic of further study.

IV. CONCLUSION

The proposed grating designs combine several physical phenomena (such as SPP, Fabry–Perot and slab waveguide modes) to boost the absorption capability of a thin polymer film. We focus on enhancing the metallic backreflector with triangular structures in an organic solar cell, however, the strategies are applicable to other material systems as well.

We stress the importance of the control of bright and dark modes to obtain an angle-dependent optimized structure. The fill-factor and the grating height are crucial parameters to this end, as they control aspects such as band flatness, energy difference, and sequence of the bright-dark pair.

In addition, we demonstrate the potential of more complicated multiperiodic structures to obtain resonances that fill up a significant wavelength range. This is realized by multiple localized plasmonic resonances with slightly different frequencies, which nonetheless have most of their field in the polymer, and not in the metal.

Finally, 3D simulation results for pyramidal gratings demonstrate the extension of these ideas to polarization independent structures. Band folding, e.g., still gives a valid prediction of the resonance locations.

An enhancement of the absorption in the polymer up to seven times (from 10% to 70%) at a plasmonic resonance is predicted. For the triangular grating we find an increase of 15.6% for the integrated absorption efficiency (AM1.5G, 300–800 nm) compared to a planar structure. The multiperiodic structure gives a larger 20.7% increase. These results thus show that a major portion of the light absorption is engineered to occur in the active medium.

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