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## FDTD simulation of compositionally graded HgCdTe photodetectors

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### Abstract

The ability to simulate HgCdTe infrared detectors with realistic alloy composition profiles is essential for their optimization. We give practical guidelines for the realization of combined three-dimensional, realistic full-wave electromagnetic and electrical modeling for the description of detectors with compositionally-graded layers, suitable to most available numerical solvers. Following the known procedure to sample the compositionally-graded layers into a number  $N$  of sublayers, we assess the effects of different choices for  $N$ , both on calculation accuracy and computational cost. Quantum efficiency spectra calculated with the proposed approach are compared with those determined through two shortcuts: ray tracing, based on classical optics, and full-wave simulation where graded layers are replaced by constant, spatially-averaged optical properties. It is shown that the former is not generally a valid alternative, since it does not address interference effects due to back-reflections, while the latter can lead to incorrect estimates of the detector cutoff wavelength.

*Keywords:* HgCdTe, infrared detectors, LWIR, numerical simulation, compositional grading, focal plane arrays, FDTD

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## 1. Introduction

The outstanding performance of the II-VI alloy HgCdTe has inspired the development of sophisticated focal-plane array (FPA) infrared photodetectors with increasingly enhanced capabilities [1, 2]. Cryogenic cooling of HgCdTe IR detectors is typically  
 5 needed to decrease dark current and noise arising from several mechanisms associated with the narrow band gap of the alloy [3], and in the last fifteen years great efforts have been made to develop detectors with reduced cooling requirements. High Operating Temperature (HOT) detectors [4, 5, 6], obtained e.g. with  $nBn$  structures [7, 8, 9] or with  $P^+N^-n^-N^-N$  heterostructures [10], should ultimately lead to room-temperature  
 10 operation according to literature, avoiding or limiting the use of heavy and expensive cooling systems (here capital letters  $N(P)$  indicate donor (acceptor) doped layers with wider band gap than the donor low-doped  $n^-$  absorber layer).

The approach of carrier-depleting the HgCdTe absorber layer using appropriate composition and doping profiles [11, 12, 13, 10] has received attention as a method  
 15 of reducing the dark current, and the adoption of  $Hg_{1-x}Cd_xTe$  absorbers with fine-tuned compositional grading is becoming a standard option also in order to optimize the quantum efficiency (QE) [14]. Thus, the capability to numerically predict the performance of FPA infrared detectors with realistic state-of-the-art composition profiles is a present need and an essential step towards their development.

Most commercial three-dimensional (3D) device simulators allow to handle the  
 20 *electrical* modeling of photodetectors, even if compositionally graded. Nevertheless, in order to solve the electrical problem under illumination, the carrier photogeneration rate distribution  $G_{opt}$  ensuing from illumination must be evaluated. The propagation of electromagnetic waves in any medium is completely described by the solution of  
 25 Maxwell's equations, a task that in FPA detectors can be numerically addressed by full-wave techniques such as the Finite Difference Time Domain method (FDTD) [15, 16, 17]. When the electromagnetic solution is known,  $G_{opt}$  easily follows (see Sec. 3).

However, when compositionally graded photodetectors are considered, the full-wave calculation of  $G_{opt}$  may require the development of particular techniques, and  
 30 also in very recent works [18, 14] ray tracing [19] or even simpler methods based on

Beer's absorption law [20, Sec. 1.6.2] have been employed to this end, neglecting the wavelike nature of infrared radiation.

Scope of this work is to give practical guidelines in order to obtain an electromagnetic solution by FDTD for compositionally graded HgCdTe photodetectors, where the Cd mole fraction  $x$  gradually changes along the growth direction, investigating to what extent different handlings of the  $x$ -grading affect the accuracy of the solution. Our purpose is also to point out what can be lost when neglecting the detector compositional grading in the calculation of  $G_{\text{opt}}$ , for example by averaging  $x$  in the graded regions. A ray tracing implementation that correctly manages compositional grading and internal light back-reflections but is intrinsically unable to describe interference and diffraction [21] has been used as an additional reference.

The manuscript is organized as follows. In Section 2 a single pixel of a 3D detector is described as case study. Section 3 describes the method we propose to obtain the electromagnetic solution by FDTD. Simulated QE spectra are shown, compared and discussed in Section 4. Finally, Section 5 summarizes the main outcomes of this study.

## 2. Single pixel detector

We performed combined optical and electrical 3D simulations considering a long-wavelength infrared (LWIR) single pixel of a planar HgCdTe-based FPA photodetector, identified by a  $d \times d \times t$  cell, where  $d = 10 \mu\text{m}$  is the pixel size and  $t$  is the total epitaxial thickness.

Since the main scope of the present work is methodological, in order to test the proposed approach we chose an epitaxial structure inspired by the literature, very similar to [14]. A two-dimensional (2D) scheme of the adopted doping and composition profiles is shown in Fig. 1, where an inset showing the 3D single pixel layout is also reported. Above as CdTe substrate, a wide-bandgap  $n$ -Hg<sub>0.6</sub>Cd<sub>0.4</sub>Te layer was grown, doped with donor concentration  $N_D = 5 \times 10^{17} \text{ cm}^{-3}$ . It is followed by a  $5 \mu\text{m}$  thick, low donor-doped, narrow-bandgap HgCdTe absorber layer ( $N_D = 1 \times 10^{14} \text{ cm}^{-3}$ ), and by another wide-bandgap Hg<sub>0.6</sub>Cd<sub>0.4</sub>Te layer, with the same low donor concentration. The  $n-p$  photodiode junction was defined by simulating an ion implantation, yielding

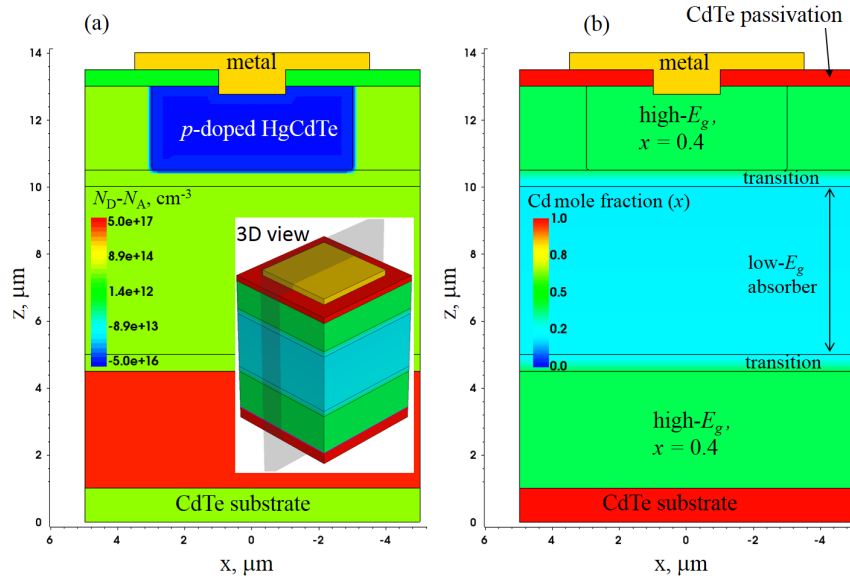


Figure 1: (a) 2D section of a single pixel with the doping profile  $N_D - N_A$ . In the inset, the 3D single pixel is shown. (b) 2D section of a single pixel, showing the  $z$ -dependent Cd mole fraction of the  $\text{Hg}_{1-x}\text{Cd}_x\text{Te}$  heterostructure, where  $z$  is the growth direction. In Fig. 2 more details are given.

60 an error-function-shaped acceptor density  $N_A$  ranging from  $5 \times 10^{16} \text{ cm}^{-3}$  just below the bias contact to virtually zero at the  $p$ - $n$  junction in  $\approx 2.5 \mu\text{m}$ .

In [14], the absorber layer was given a graded composition, varying the Cd mole fraction from  $x = 0.21$  to  $x = 0.19$ , from its lower to its upper interface, in order to maintain a constant peak wavelength at the temperature of operation. We adopted this choice not for this particular purpose, but just as a literature example of compositionally graded absorber. Still in [14], in order to increase realism two  $0.5 \mu\text{m}$  thick transition regions with linear compositional grading connect the two high- $E_g$  regions to the low- $E_g$  absorber layer (where  $E_g$  is the bandgap). In order to test the proposed model in more critical conditions, a variant was also considered, giving the absorber a steeper compositional grading, from  $x = 0.25$  to  $x = 0.19$ . For brevity, in the present work we identify the two variants respectively as the *standard* and the *steeper* one, and in Fig. 2, the Cd mole fraction and the dopant concentrations along a one-dimensional 70 (1D) vertical cutline at center pixel are shown, for the two considered variants. The bias

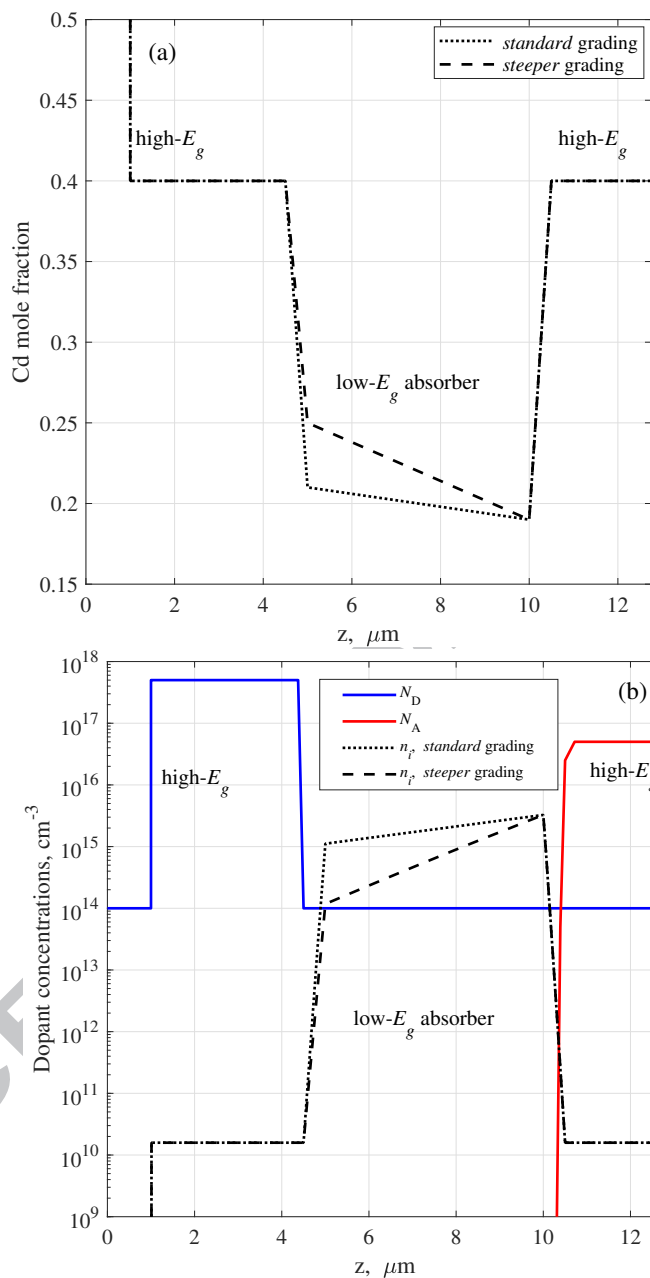


Figure 2: (a) Cd mole fraction for the *standard* and *steeper* variants of the absorber. (b) Dopant concentrations  $N_D$  and  $N_A$  along a 1D vertical cutline at center pixel. The intrinsic density  $n_i$  for  $T = 140\text{K}$  is also shown as black dotted and dashed lines for the two variants.

contact is connected to the  $p$ -doped region through a square metallic layer  $7\mu\text{m} \times 7\mu\text{m}$   
 75 wide, partly extending over a  $0.3\mu\text{m}$  thick CdTe passivation layer that covers the upper  
 face of the FPA's single pixel.

The geometry, doping and composition profiles were defined employing the Sen-  
 taurus 3D numerical simulator by Synopsys [22], also employed to perform the elec-  
 trical simulations as described in Sec. 4.

### 80 3. Electromagnetic simulations

In order to obtain the photogeneration rate  $G_{\text{opt}}$  for a given illumination, we adopted  
 the FDTD approach [17], solving Maxwell's equations for the electric and magnetic  
 fields  $\vec{E}$  and  $\vec{H}$ , discretized on a cubic grid known as the Yee grid [15]. For vali-  
 dation purpose, we employed and compared the results obtained by two commercial  
 85 codes, the Electromagnetic Wave (EMW) solver [22], and the RSoft FullWAVE [23],  
 both integrated in the same Synopsys simulation suite Sentaurus Workbench [22]. The  
 computational box includes air layers located above and below the pixel, and the opti-  
 cal boundary conditions (BC) along the upper and lower sides of the box are absorbing  
 (this is obtained with convolutional perfectly matching layers [16]), while periodic op-  
 90 tical BCs are applied along the lateral sides of the computational box, in order to exploit  
 symmetry and obtain a simulation equivalent to a FPA. In the FDTD model, the ma-  
 terial properties are represented by the electric permittivity and conductivity  $\varepsilon$  and  $\sigma$ ,  
 calculated from the material complex refractive index  $\hat{n} = n + i\kappa$  as

$$\begin{aligned}\varepsilon &= n^2 - \kappa^2 = n^2 - \left(\frac{\alpha\lambda}{4\pi}\right)^2 \\ \sigma &= \frac{n\alpha}{\mu_0 c},\end{aligned}\tag{1}$$

where  $\lambda$  is the wavelength,  $\mu_0$  is the vacuum magnetic permeability,  $c$  is the speed of  
 95 light in vacuum, and  $\alpha = 4\pi\kappa/\lambda$  is the absorption coefficient. The absorbed photon  
 density  $A_{\text{opt}}$  (number of absorbed photons per unit volume and time) can be evaluated  
 as the divergence of the time-averaged Poynting vector  $\langle \vec{S} \rangle$  [24, 25, 26, 27]

$$A_{\text{opt}} = -\frac{\vec{\nabla} \cdot \langle \vec{S} \rangle}{h\nu} = \frac{1}{2h\nu} \sigma |\vec{E}|^2,\tag{2}$$



where  $h\nu$  is the photon energy. The optical generation rate distribution  $G_{\text{opt}}$  into the pixel due to interband optical absorption is given by  $G_{\text{opt}} = \eta A_{\text{opt}}$ , where the quantum yield  $\eta$ , defined as the fraction of absorbed photons which are converted to photogenerated electron-hole pairs, was assumed to be unitary (see calculation examples for monochromatic [21, 28] and for broadband illumination [29, 30, 31], all following the FDTD method. Similar methods were also applied with success to other types of photodetectors, see e.g. [32, 33], and in simpler one-dimensional [34] and two-dimensional modeling descriptions [35].

In all simulations, we set a lattice temperature  $T = 140\text{ K}$ , the HgCdTe optical properties were described through the models and parameters reported in [28, Table I], and the contact metallizations were treated as Perfect Electric Conductors (PEC) [17].

### 3.1. Discretization of compositionally graded regions

The solution of the electromagnetic problem for detectors with arbitrary composition profile may not be straightforward, since not all simulators manage a compositional grading, neither as analytic function, nor as external file, requiring instead to define the detector geometry as a stack of layers each with uniform optical properties. However, since the considered composition profile only varies along  $z$  (the growth direction), the idea was to sample the profile, converting it to a staircase, discretizing consequently the detector itself into a large stack of layers, each with uniform values for  $n$  and  $\kappa$ .

In order to test the feasibility of this general idea, we implemented the method in both considered simulators, without taking advantage of particular solutions of the compositional grading problem, possibly implemented by the simulator manufacturer.

Exploiting the planarity of the structure, the simulation flow includes a Tcl [36] script that builds the device geometry, discretizing the low- $E_g$  absorber layer and the two low-to-high- $E_g$  transition regions into  $N$  sublayers each. This step is common to both simulators, since they are able to read the same geometry boundary files. Then, starting from the adopted expressions for the complex refractive index [28, Table I], another Tcl script converts the linear grading of the Cd mole fraction  $x$  to a staircase  $x_i, i = 0, \dots, N - 1$ , building  $N$  tables  $(\lambda_m, n_m, \kappa_m)_i$ , for values  $\lambda_m \in [1, 20] \mu\text{m}$ , separated by  $0.1 \mu\text{m}$ . Regarding the other regions with uniform composition (the sub-

strate, high- $E_g$ , and passivation regions), the usual approach was followed, building one  $(\lambda_m, n_m, \kappa_m)$  table for each of them. In this way, for a given operating wavelength  $\lambda \in \{\lambda_m\}$ , the  $N$  look-up tables  $(\lambda_m, n_m, \kappa_m)_i$  provide the desired discretized  $n$ - and  $\kappa$ -profiles. In the end, the described Tcl script produces a set of material libraries as ASCII files, one of each layer or sublayer, including them appropriately in the material library of both simulators, only respecting the required file formats they separately need.

### 3.2. Results

As an example and considering the *standard* variant, in Fig. 3(a-c) the discretized  $n$  and  $\kappa$  distributions are shown for  $N = 10$ , calculated for  $\lambda = 9\mu\text{m}$ . For appreciating more quantitatively their variation across the compositionally graded regions, in Fig. 3(d,e) the  $n$  and  $\kappa$  profiles are shown along a vertical 1D cutline. The optical grid was chosen with a spacing of 10 nm in the  $z$ -direction, and 25 nm in the  $x, y$ -directions.

Our starting point consisted in performing a FDTD simulation for an illuminating plane wave with  $\lambda = 9\mu\text{m}$ . The optical source was located in air, and the detector was illuminated from below, with wavevector orthogonal to the plane  $z=0$ , setting an optical flux  $P_{in} = 1 \text{ mW cm}^{-2}$ . The rate  $A_{\text{opt}}$  obtained for the case  $N = 10$  by the EMW solver is plotted in Fig. 4(a-c) on 2D cutplanes  $y = 0$  and  $y = 4$  on appropriate scales, to point up the reflections effect due to the metallization on the photogeneration distribution. Then, we repeated the simulation still with EMW, for  $N = 5, 10, 15, 30$  and, in order to validate the proposed computational method and discretization algorithm, the same simulation was also performed by FullWAVE for  $N = 10$ . The results are plotted in Fig. 4(d, e), along a vertical 1D cutline at  $x = y = 4\mu\text{m}$ . The four distributions obtained by EMW are apparently very similar, and the computational time does not depend much on the choice of  $N$  ( $\approx 30$  minutes on our servers), provided we employ the same spatial grid. The FullWAVE solver yields a solution differing a little more from them, but still following the same overall behavior.

155 **4. Electrical simulations**

For a given illumination, in the drift-diffusion framework  $G_{\text{opt}}$  enters as a source term in the continuity equations for the electron and hole current densities  $J_{n,p}$ , that can be solved as outlined e.g. in [21, 28, 30], in order to obtain the current under illumination  $I$  and in dark  $I_{\text{dark}}$ . The photocurrent follows as  $I_{\text{ph}} = I - I_{\text{dark}}$  and the

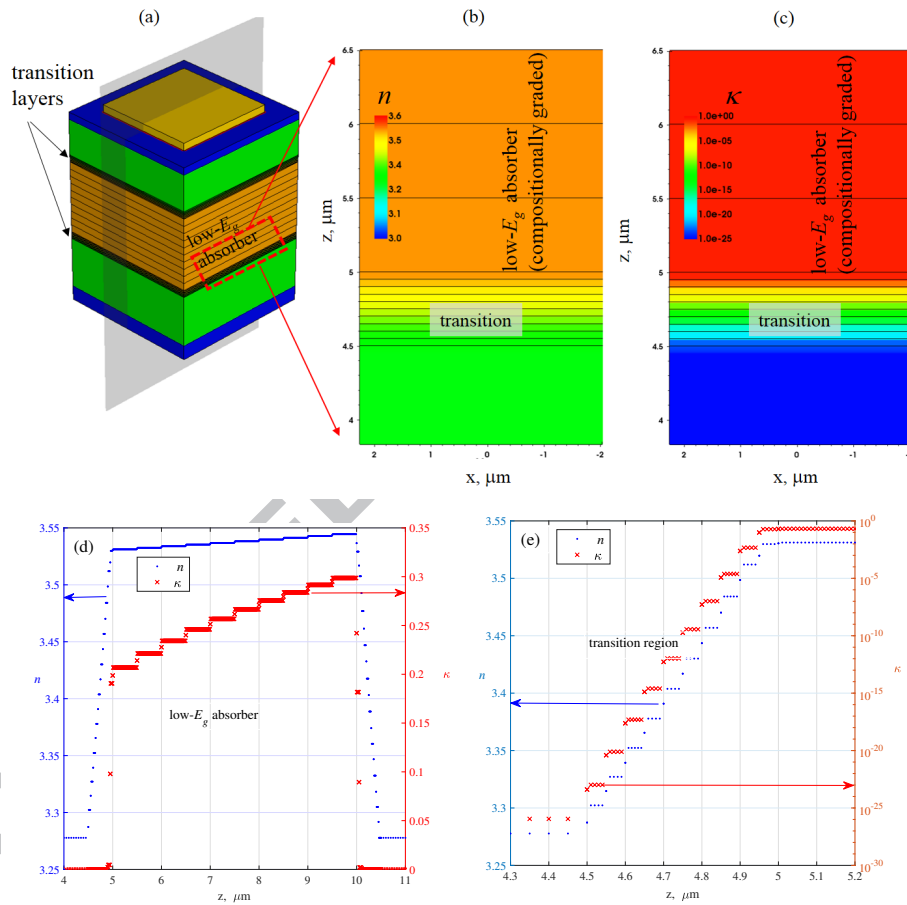


Figure 3: *Standard variant*: (a) 3D overall view of a single pixel, showing the discretization into  $N = 10$  sublayers of the low- $E_g$  absorber region and the two transition regions. In panels (b) and (c), the real and imaginary parts of the complex refractive index  $n$  and  $\kappa$ , calculated for  $T = 140\text{K}$  and  $\lambda = 9\mu\text{m}$ , are shown for a small portion of a vertical 2D cutplane (the dashed portion shown in panel (a)). In panels (d, e)  $n$  and  $\kappa$  are shown along a vertical 1D cutline at center pixel: panel (d) is centered across the low- $E_g$  absorber region, and panel (e) across one of the two transition regions.

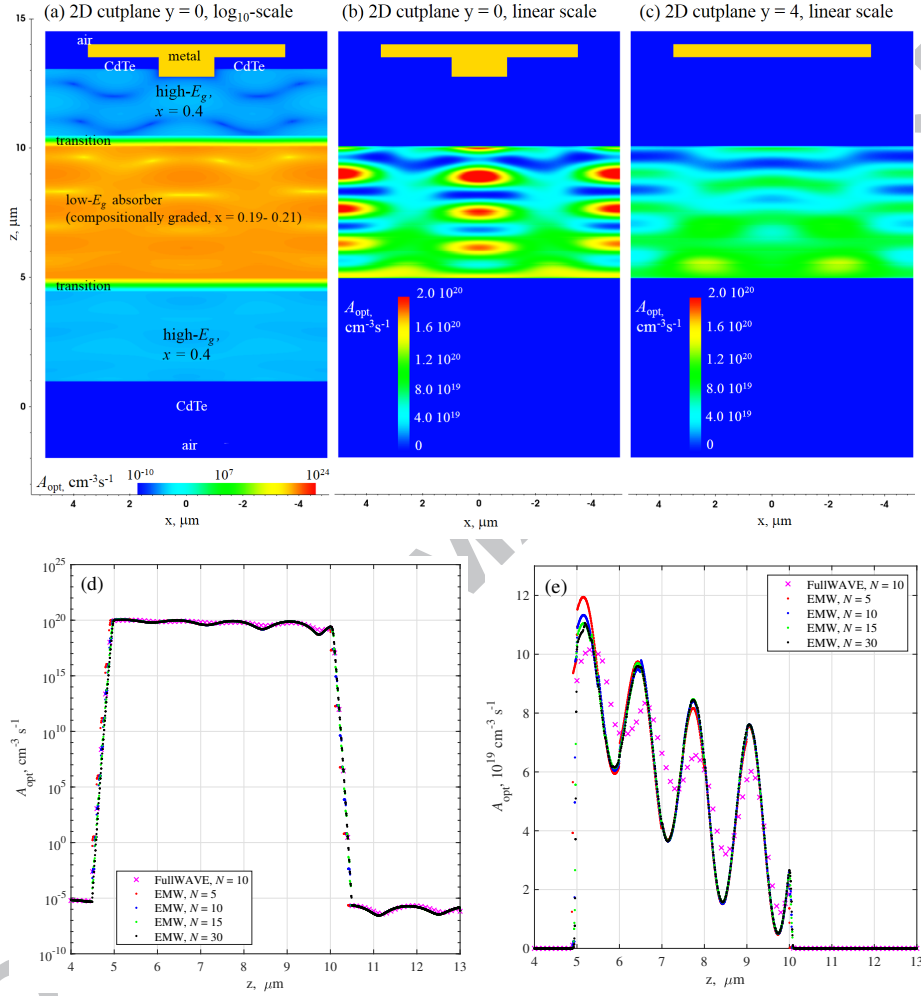


Figure 4: Standard variant:  $A_{\text{opt}}$  in the 2D cutplane  $y = 0$ , obtained for the case  $N = 10$  by the EMW solver for  $\lambda = 9 \mu\text{m}$ , in logarithmic (a) and linear (b) scale. In panel (c)  $A_{\text{opt}}$  is shown in the 2D cutplane  $y = 4$  in linear scale, to point up the reflections effect due to the metallization on the photogeneration distribution. In the panels (d) and (e),  $A_{\text{opt}}$  is shown on a logarithmic and linear scale along a vertical 1D cutline at  $x = y = 4 \mu\text{m}$ , for all the considered discretization cases  $N = 5, 10, 15, 30$  - EMW, compared with the case  $N = 10$  - FullWAVE, for the reciprocal validation of the method implemented in the two electromagnetic solvers.

160 quantum efficiency as  $QE = I_{ph}/(qN_{phot})$ , where  $q$  is the elementary charge and  $N_{phot}$  is the photon flux impinging the illuminated face. To this end, the detector in Fig. 1 was discretized into  $\approx 1.48 \times 10^5$  elements with a meshing tool which generates a denser grid in regions where gradients of current density, electric field, free charge density and material composition are present. The HgCdTe properties were described through the

165 models reported in [28], taking into account the composition, doping, and temperature dependence of the HgCdTe alloy. The Shockley-Read-Hall (SRH) recombination processes were modeled as in [37, 20, 38] considering a lifetime around  $2 \mu s$ , neglecting instead trap-assisted or band-to-band tunneling processes [39, 40]. Fermi-Dirac statistics and incomplete dopant ionization were taken into account, with activation energies

170 for HgCdTe alloys estimated according to [41, 42]. Electric contacts were treated as Ohmic with zero resistance, where charge neutrality and equilibrium were assumed. Ideal Neumann BCs were applied to the outer boundaries of the array, and the drift-diffusion equations were solved by the Finite Box (FB) method, setting  $T = 140K$  and driving the detector at  $0.5 V$  of reverse bias.

175 Compositional grading is not an issue for electrical simulations, since the nominal HgCdTe composition profile shown in Fig. 2(a) is evaluated at each point of the electrical grid, and the  $N$ -sublayers discretization is not required. On the other hand, since the grading imposed to develop particular techniques to solve the electromagnetic problem, it is interesting to compare the QE spectra obtained when  $G_{opt}$  is evaluated

180 a) following the described  $N$ -sublayers discretization method, and b) averaging  $n$  and  $\kappa$  in the graded regions, to see what we lose neglecting the grading in the electromagnetic modeling. In addition, c) the same calculation was done following the ray tracing method, and the obtained QE spectra are shown for the three cases in Fig. 5. The spectra for cases a) and b) are quite similar, due to the small variation of the

185 HgCdTe composition in the absorber. Conversely, the spectrum for ray tracing, case c), is significantly different close to the cutoff wavelength (usually defined as the  $\lambda$  value where the QE drops to one half of its maximum value). In fact, FDTD is able to describe interference effects, particularly relevant because of the large metallization of the bias contact, acting as a reflector to increase QE. On the contrary, ray tracing

190 does not address the wavelike nature of radiation: an extended discussion with several

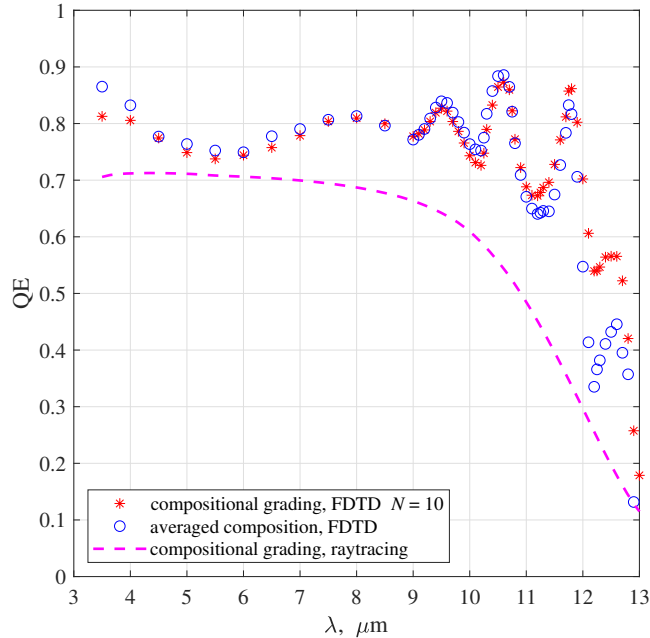


Figure 5: *Standard* variant calculated QE spectra, comparing: FDTD spectrum obtained with the described discretization scheme, with  $N = 10$  (stars), or averaging the  $n$  and  $\kappa$  in each graded region (circles). Dashed line, spectrum obtained with the ray tracing method.

comparisons between FDTD and ray tracing results can be found in [21].

Similar electromagnetic and electrical simulations were repeated for the *steeper* variant of the detector, in order to assess the effect of the compositional grading steepness in the absorber, and the obtained QE spectra are shown in Fig. 6, for the three  
 195 electromagnetic computational methods. In this case, it is evident that, neglecting the grading in the electromagnetic modeling (case *b*)), the overall cutoff wavelength would be wrongly estimated, although the gradient of the composition profile in the absorber is still moderate. Instead the two solutions are quite similar when the material is more  
 absorbing and the effect of the compositional grading loses importance, but this hap-  
 200 pens for  $\lambda < 8 \mu\text{m}$ , *i.e.* out of the LWIR window. The ray tracing spectrum lies in between the two FDTD solutions, but close to the cutoff wavelength the ray tracing method still underestimates the QE by a factor of two, precisely because it neglects any interference effects, particularly important when the material becomes less absorb-

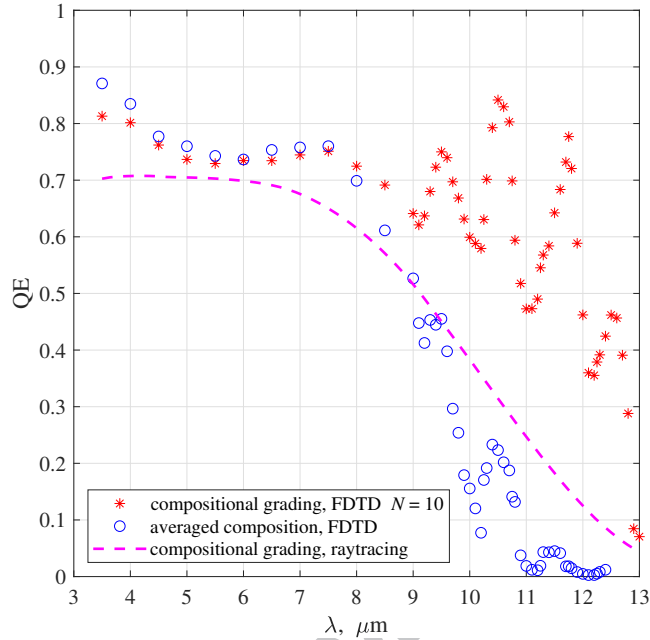


Figure 6: *Steeper* variant calculated QE spectra, comparing: FDTD spectrum obtained with the described discretization scheme, with  $N = 10$  (stars), or averaging the  $n$  and  $\kappa$  in each graded region (circles). Dashed line, spectrum obtained with the ray tracing method.

ing [21].

## 205 5. Conclusions

We investigated the optical and electrical response of compositionally graded HgCdTe planar photodetector by means of a combined FDTD-FB approach, giving practical guidelines for the implementation of a three-dimensional modeling method particularly suitable for electromagnetic solvers that do not manage compositional grading directly. In the present case, a linear grading was considered as an example, but in principle the method can be easily applied to a wide choice of profiles.

On this purpose, we assessed the effect of different choices for the number of sublayers  $N$  adopted to sample a compositionally graded region.  $N$  should be large enough to guarantee a small reflection coefficient  $R_i$  between adjacent sublayers in the absorber's graded region. It is easy to verify that, in the considered cases, its approximate

estimate as  $R_i \approx (n_{i+1} - n_i)^2 / (n_{i+1} + n_i)^2$  (Fresnel law), valid for normal incidence, assures  $R_i < 10^{-4}$  also for small values of  $N$ . We also observed that the value of  $N$  has a small impact on the computational cost of the simulation. Consequently, it is recommended to choose a value large enough to sample the absorption coefficient profile  $\alpha(z)$  properly, in order to get an accurate estimate of the detector's cutoff wavelength, to which end some tests are generally needed.

Another important point was to assess the possibility to consider simpler approximations, like averaging  $x$  in the region when the grading is present, or considering the ray tracing method to solve the electromagnetic problem. We pointed out that a  $\text{Hg}_{1-x}\text{Cd}_x\text{Te}$  composition profile cannot be approximated, in general, by simply averaging  $x$  in the graded region, since also moderate compositional gradients of  $x$  may lead to incorrect predictions of quantum efficiency spectra and cutoff wavelengths. In fact, if for the *standard* variant Fig. 5 shows that averaging  $x$  in the absorber is still an acceptable approximation, as soon as moderately steeper profiles are considered, the approximation loses precision, as shown in Fig. 6 for the *steeper* variant.

Regarding the ray tracing approach, it has not proved to be a valid alternative, not addressing the interference effects due to back-reflections. Similar conclusions were drawn in [25], which compared a FDTD calculation with a result obtained considering a simple exponential decay of the photon flux inside the device.

The greatest sources of back-reflections are the interfaces with metallizations, which are assumed to be totally reflecting. The effect is particularly relevant when large contact metallizations are present, leading to absorption enhancement close to cutoff wavelength owing to spatial resonances and cavity effects [21].

Future work will consist in simulating compositionally graded FPA detectors, single color or multi-color [1, 43, 44, 2], illuminated by extended, nonmonochromatic sources. Incoherent light, especially if produced by broadband (e.g. blackbody) source, results in less prominent resonance effects, as described in Ref. [30] for simpler photodetectors without compositional grading, and it would be interesting to extend that study to detectors with more general composition profiles. Finally, we stress that the presented modeling approach is suitable to be applied to any electromagnetic simulator not designed to directly manage compositionally graded regions.



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ACCEPTED MANUSCRIPT

- simulation of HgCdTe infrared detectors with realistic alloy composition profiles essential for their optimization
- realistic full-wave electromagnetic with compositionally-graded layers is difficult with commercial simulators
- we give practical hints for discretizing the grading into sublayers
- we test several different discretizations
- we assess when the grading may be neglected

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Declarations of interest: none

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