

Highlights

Modeling the Impact of Iron Defect Variability on Silicon Solar Cell Performance Across Different Scenarios

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Table 1

Parameters varied during the simulation

Parameter	Value
$d_p, (\mu)$	180 – 380
$N_B, (\text{cm}^{-3})$	$10^{15} - 10^{17}$
$N_{\text{Fe}}, (\text{cm}^{-3})$	$10^{10} - 10^{14}$
$T, (\text{K})$	290 – 340
Illumination	AM1.5G, 1000 W/m ² ; 940 nm 5 W/m ² ; 940 nm 10 W/m ²

the concentration of iron in SSCs based on changes in the ideality factor. Likewise, in [12], the concentration of iron in SSCs was determined using the open-circuit voltage. However, developing such approaches requires evaluating how a particular parameter changes when iron-containing pairs dissolve and determining whether it can estimate the iron concentration N_{Fe} . For example, the most evident conditions for utilizing a specific parameter include its change due to the transformation $\text{Fe}_i\text{B}_{Si} \rightarrow \text{Fe}_i + \text{B}_{Si}$, even by 10 %, along with a consistent dependence of these changes on N_{Fe} .

This study intends to determine the variations of J_{sc} , V_{oc} , η , and FF resulting from the decay of iron-containing pairs in boron-doped silicon solar cells. Previous similar calculations have been conducted [?], but the results presented in these works typically pertain to particular temperatures, illumination levels (often AM1.5), and solar cells with distinct structures. In this study, we performed calculations over a sufficiently wide temperature range (290-340 K) and for solar cells with varying base thickness (180-380 μ) and doping levels (boron concentration in the base ranging from 10^{15} cm^{-3} to 10^{17} cm^{-3}). The results obtained allow us to assess the feasibility and potential of using the main photovoltaic parameters of specific solar cells to estimate the N_{Fe} value across a range of temperatures, including conditions similar to those encountered in typical SSC applications. Furthermore, investigations have explored changes in photoelectric performance under solar illumination (AM1.5G) and low-intensity monochromatic light (wavelength of 940 nm, intensities of 5 W/m² and 10 W/m²).

In the first case, while it is customary to adhere to standard conditions, it's essential to consider that illumination at 1000 W/m² can lead to the disintegration of Fe_iB_s complexes. Therefore, measurements for cases requiring the presence of undissociated pairs in silicon must meet specific constraints. On the other hand, intentionally chosen monochromatic illumination penetrates the emitter with negligible losses and does not reach the rear side. In this context, the photoelectric parameters demonstrate a remarkable sensitivity to recombination processes occurring within the solar cell base, making them responsive to variations in iron concentration.

Finally, in our calculations, we attempted to use the latest literature data concerning the exact values of silicon parameters, including light absorption values [?] and coefficients characterizing intrinsic recombination [?].

2. Experimental Methodology

The study involved modeling IV curves of SSCs with an $n^+ - p - p^+$ structure, as illustrated in Figure 1. A notable feature observed in both Al-BSF cells (full area), which are gradually losing relevance, and PERC cells (locally), which are the most widely used in mass production, is the presence of a back surface field (BSF). We considered structures with a base uniformly doped with boron. The doping concentration N_B , and the base thickness d_p were varied during the modeling process, as detailed in Table 1. The concentration profiles of the dopants, their maximum values ($N_{p,\text{max}}^+$ and $N_{n,\text{min}}^+$), and layer thicknesses (refer to Figure 1) were selected based on the work of Fell et al. [?]

The simulation was conducted using the SCAPS3.3.11 code [?]. SCAPS-1D software, developed by the University of Gent, is founded on theoretical computations that involve solving Poisson's equation, continuity equations for holes and electrons, and drift-diffusion at each position within the solar cell, considering the boundary conditions. Despite its one-dimensional modeling approach, SCAPS is extensively used for modeling various types of solar cells [?] and for investigating the effects of defects on their performance [?].

As can be seen from Table 1, calculations spanned a broad range of temperatures and base doping levels. Therefore, to improve the accuracy of the calculations when inputting the initial parameters into SCAPS, temperature and concentration dependencies (where applicable) of the following silicon parameters were taken into account:

- bandgap according to Passler [17];

- doping induced bandgap narrowing according to Yan & Cuevas [18];
- effective density of states at conduction and valence band and intrinsic carrier concentration according to Couderc et al. [19];
- thermal carrier velocities according to Green [20];
- free carrier effective masses according to O'Mara et al. [21];
- carrier mobilities according to Klaassen's theory [22];

The values of surface recombination coefficients were considered equal to the thermal velocities of carriers [?]. The calculations addressed recombination processes within the structural volume, incorporating both intrinsic recombination and Shockley-Read-Hall recombination at iron-related defects. In the first case, processes of band-to-band radiation recombination were considered (where the calculation of the corresponding coefficient included the fraction of radiatively emitted photons reabsorbed via band-to-band processes according to Niewelt et al. [?]) and Auger recombination (where the coefficients considered the effect of Coulomb enhancement [?] and temperature dependence [?]).

When accounting for the influence of iron impurities, we considered that Fe atoms were uniformly distributed within the base and p^+ layer, with a total concentration of N_{Fe} (see Table 1). We considered two cases:

Case 1. The concentration of interstitial iron defects $[Fe_i] = N_{Fe}$ at each position throughout the solar cell, with no pairs present $[Fe_iB_s] = 0$. This case corresponds to the state of the structure immediately after intense illumination, for example.

Case 2. Iron atoms predominantly form pairs with acceptors, $[Fe_iB_s] \gg [Fe_i]$, but the exact concentration ratio depends on the position of the Fermi level and temperature [?] and varies from point to point within the solar cell. Further details about how we calculated the concentration profiles of Fe_iB_s and Fe_i are provided in [1]. This case corresponds to prolonged storage of the structure in darkness or under conditions of low-intensity ($< 0.01 \text{ J/cm}^{-3}$ [?]) illumination.

During the calculations, we assumed that Fe_i forms a single donor level, while the Fe_iB_s pair has a trigonal configuration and acts as an amphoteric defect. We obtained defect parameters (including level positions within the bandgap and electron and hole capture cross-sections) from relevant studies [?].

As previously mentioned, we modeled how the solar cell behaves under different illumination conditions, including solar light (AM1.5G) and monochromatic light (wavelength 940 nm, intensities of $W_{ill} = 5 \text{ W/m}^2$ or 10 W/m^2) – see Table 1. The calculations incorporated the light absorption values in silicon based on Green's study [Photovoltaics 30 164] [?].

The I-V characteristics were simulated for both Case 1 and Case 2 (see Fig. 2), and from each curve, the short-circuit current density, open-circuit voltage, efficiency, and fill factor were determined. Assessing the influence of iron defect variability relied on the relative changes in each photovoltaic conversion parameter:

$$\epsilon A = \frac{A^{FeB} - A^{Fe}}{A^{FeB}} \times 100\% \quad (1)$$

where A represents one of the parameters (J_{sc} , V_{oc} , FF i η), superscript "FeB" corresponds to the parameter value for coexistence of Fe_i and Fe_iB_s (Case 2), superscript "Fe" is related to the decay of all pairs (Case 1).

Impact of change of iron defects was investigated as a function of temperature from 290 K to 340 K, base depth from 180 μ to 380 μ , base doping level from 10^{15} cm^{-3} to 10^{17} cm^{-3} , and total impurity iron atom concentration from 10^{10} cm^{-3} to 10^{14} cm^{-3} . For each illumination scenario, calculations were carried out with 11 different temperature values (T) and 5 base depth values (d_p), evenly distributed within the specified ranges. The concentration values were distributed equally on a logarithmic scale with 4 (N_B case) and 6 (N_{Fe} case) steps per decade. As a result, for the AM1.5 illumination scenario, for instance, 24750 IV characteristics were simulated. An exception occurred with monochromatic illumination at $W_{ill} = 10 \text{ W/m}^2$, where we used only two base depth values.

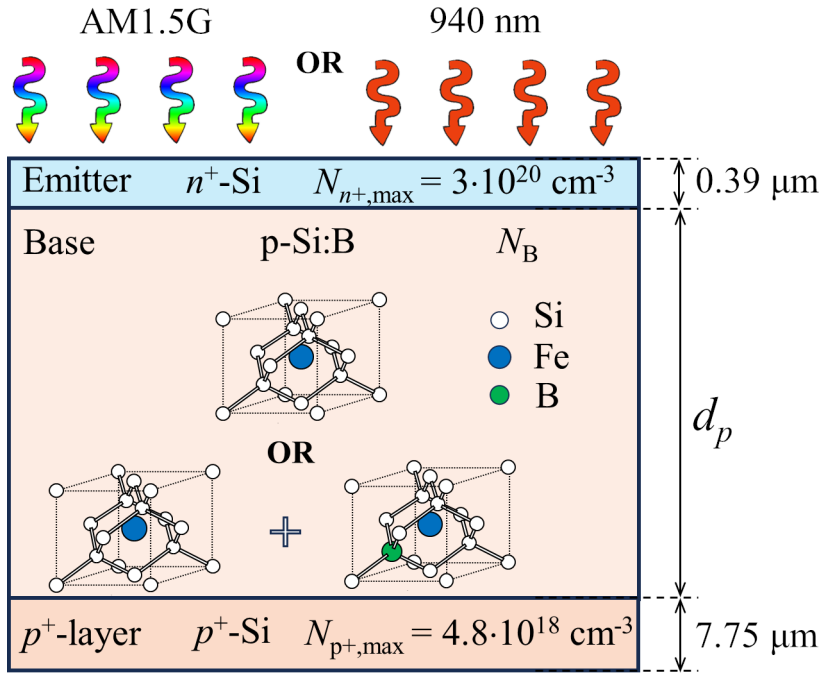


Figure 1: Schematic diagram of analyzed solar cells.

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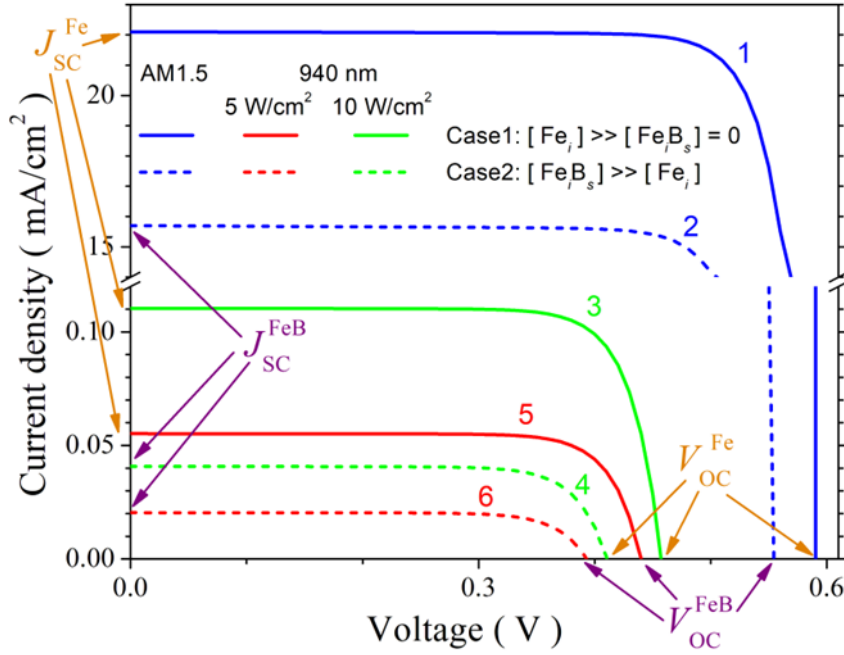


Figure 2: Typical IV characteristics, calculated for structure with $d_b = 180 \mu$, $N_B = 10^{17} \text{ cm}^{-3}$, $N_{Fe} = 10^{14} \text{ cm}^{-3}$ at $T = 290 \text{ K}$. Illumination: AM1.5 (curves 1, 2), 940 nm 10 W/cm² (3, 4) and 940 nm 5 W/cm² (5, 6). Solid and dotted lines correspond to Case 1 and Case 2, respectively.

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