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Insertion of a thin highly doped crystalline layer in silicon heterojunction solar cells: Simulation and perspectives towards a highly efficient cell concept

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An emerging cell concept based on silicon heterojunctions called hetero-homojunction is investigated by means of numerical simulations. Compared to the usual amorphous/crystalline silicon (a-Si:H/c-Si) heterojunction architecture, the hetero-homojunction cell contains an additional thin and highly doped (p^+ - or n^+ - c-Si layer at the front or back (i)a-Si:H/(n)c-Si interface, respectively. In this paper, we show the dependence of solar cell performance on the additional heavily doped c-Si layer parameters (thickness and doping) and a-Si:H/c-Si interface properties. Insertion of the (p^+)c-Si improves the cell power conversion efficiency by almost 1% absolute and lowers its sensitivity to a-Si:H/c-Si interface defects. Improved field effect passivation leading to higher open circuit voltage and fill factor is evidenced and the added layer is optimized with regard to hetero-homojunction cell efficiency. The (n^+)c-Si layer addition also decreases the recombination rate at the back hetero-interface but does not improve significantly the conversion efficiency. The latter result is finally discussed. © 2015 AIP Publishing LLC.

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I. INTRODUCTION

Amorphous/crystalline silicon heterojunction (HET) (a-Si:H/c-Si) solar cell is a very promising technology to achieve high conversion efficiency at low manufacturing cost.¹ The use of thin hydrogenated amorphous silicon (a-Si:H) as an emitter has allowed several research institutes and companies to reach open circuit voltage (V_{OC}) values above 720 mV (Refs. 2–5) while 24.7% power conversion efficiency was recently demonstrated on industrial size wafers² (25.6% for rear contact HET cell⁶).

In today's HET cells, emitter passivation was identified as one of the most important issues.^{7,8} Therefore, ways to further improve the quality of interfaces in the (p)a-Si:H/(i)a-Si:H/(n)c-Si stack should be investigated to enhance cells' performance. On the one hand, the density of interface recombination centers (D_{it}) can be reduced by surface cleaning or by improving the (i)a-Si:H passivation layer.⁸ On the other hand, a higher selectivity of the hetero-interface, i.e., the presence of only one type of charge carrier, decreases interface recombination⁹ leading to more efficient devices. The latter route, commonly called field effect passivation, could be realized through the addition of a thin and highly doped (p^+)c-Si layer underneath the (i)a-Si:H passivation layer.¹⁰ Because of the added (p^+)c-Si/(n)c-Si homojunction, the alternative cell architecture can be referred to as a front side hetero-homojunction (HHJ_{FS}). The HHJ_{FS} architecture is introduced in Figure 1.

So far, only a few studies have investigated the hetero-homojunction concept.^{10–12} Zhong *et al.* simulated the previously defined HHJ_{FS} cell but removed the (i)a-Si:H passivation layer.¹¹ The cell displays V_{OC} and fill factor (FF) improvement and a reduced interface defect (D_{it}) sensitivity (i.e., less performance reduction for increasing D_{it}). Field effect at the front interface

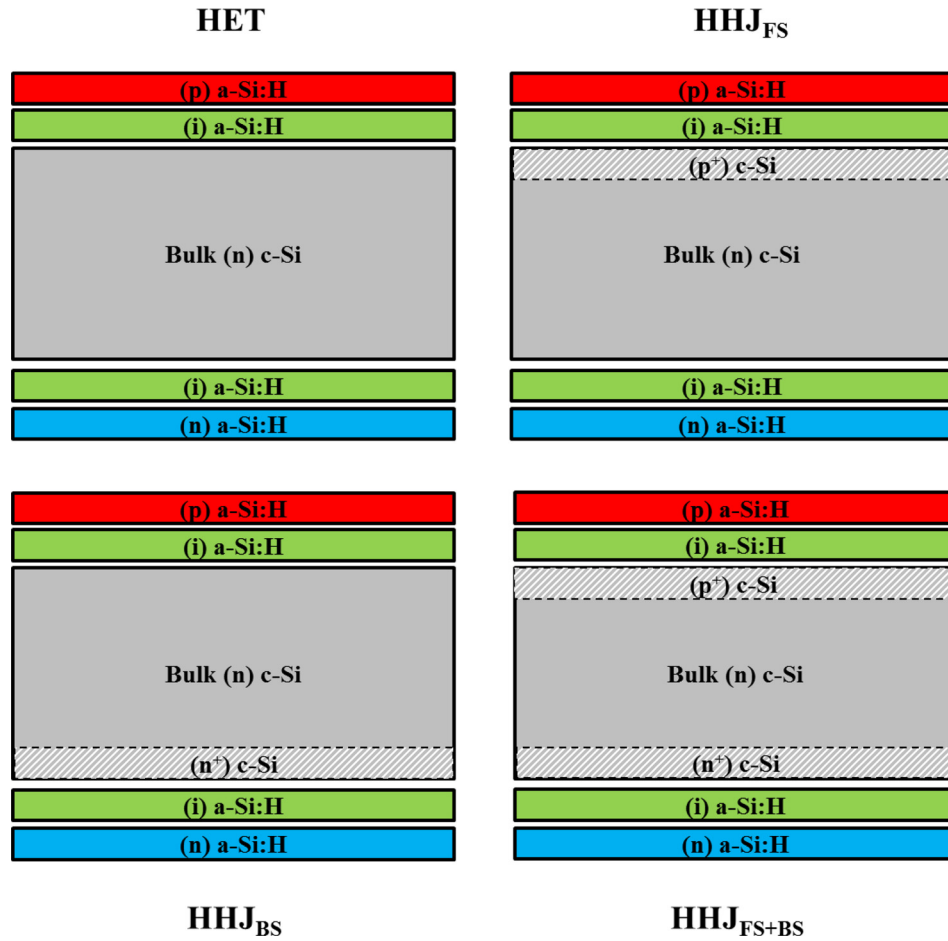


FIG. 1. Scheme of the HET cell and of the various HHJ cell architectures with the inserted (p^+) c-Si or/and (n^+) c-Si layers.

was evidenced as the passivation improvement mechanism. However, Zhong's calculations were performed with the same interface defect density as in the "traditional" (p) a-Si:H/ (i) a-Si:H/ (n) c-Si although the (i) a-Si:H passivation layer was removed, which is not realistic. In an experimental cell, the additional field effect brought by the (p^+) c-Si layer introduction may not be sufficient to compensate the loss of chemical passivation due to the (i) a-Si:H layer removal. In this case, all beneficial effects of the (p^+) c-Si layer would be lost.

Mikolášek *et al.* have simulated the HHJ_{FS} architecture but on a p-type substrate (with an (n^+) c-Si layer addition) and using a rear contact cell.¹⁰ They added the homojunction, while conserving the (i) a-Si:H passivation layer. With such cell architecture, they observed a V_{OC} improvement and a reduced interface defect sensitivity.

Hekmatshoar *et al.* manufactured a p-type HHJ_{FS} cell but without the (i) a-Si:H passivation layer.¹² However, the observed efficiency increase by 0.7% absolute (abs) (compared to the HET cell) cannot be directly attributed to front interface passivation improvement since the back side layer stack was also modified.

None of these studies focused on the role of the addition of the (p^+) c-Si layer alone, without removing the (i) a-Si:H passivation layer (required to achieve high chemical passivation, i.e., low D_{it}), on a n-type cell (which is nowadays used in industrial high-performance devices) and with a standard contact configuration.

This work aims at investigating the behavior of the hetero-homojunction cells as close as possible to experimentally achievable cells and comparing them to a state-of-the art HET cell. Thus, as described in Figure 1, we started from a standard n-type HET architecture to build the HHJ_{FS} cell by inserting the (p^+) c-Si layer. The (i) a-Si:H passivation layer was kept so that any

change can be attributed to the (p^+) c-Si layer insertion. Simulated cells are first described in Sec. II. They consist in a HET reference cell and in HHJ cells. In addition to the previously described HHJ_{FS} architecture, a hetero-homojunction with an added (n^+) c-Si layer at the back interface (HHJ_{BS}) and a heterojunction with both added (p^+) - and (n^+) -c-Si layer ($\text{HHJ}_{\text{FS}+\text{BS}}$) are also investigated. Cell parameters of the architectures are then compared in Sec. III. Finally, the cell performances are discussed in the light of recombination rates within the cells in Sec. IV. The HHJ_{FS} cell behavior is investigated and the (p^+) c-Si layer is optimized by varying the layer doping level and thickness. The inserted (n^+) c-Si layer benefits in HHJ_{BS} and $\text{HHJ}_{\text{FS}+\text{BS}}$ cells are assessed and compared to the (p^+) c-Si layer insertion.

II. SIMULATION MODELS

Numerical simulations are performed using the 1D simulation software “Automat FOR Simulation of HETerostructure” (Afors-HET) developed at the Helmholtz Zentrum Berlin.¹³ Simulated cell architectures with the position of the added layers are presented in Figure 1. The corresponding band diagrams at equilibrium are presented in Figure 2.

Layer parameters are described in Table I. The HET cell layers ((i)a-Si:H excepted) are taken from Varache’s work.¹⁷ The density of states (DOS) in the doped a-Si:H layers follows the defect pool model proposed by Powell and Deane.^{14–16} A simplified description of the DOS consisting in two deep Gaussian distributions was used in the (i)a-Si:H layers. No significant discrepancies could be found compared to the use of a DOS in (i)a-Si:H layers calculated according to the actual position of the Fermi energy (E_F) within the bandgap that is close to that in the adjacent doped a-Si:H layer due to the band bending at the hetero-interface which causes a DOS re-equilibration.¹⁵

At front and back hetero-interfaces, defects are introduced using a 1 nm-thick highly defective c-Si interfacial layer containing two Gaussian distributed density of states.¹⁷ The donor-like and the acceptor-like Gaussian distributions have their respective maximum at 0.56 eV and 0.76 eV above the valence band (E_V). Both distributions have a standard deviation of 0.2 eV and electron and holes capture cross sections (σ_e , σ_h) of $3 \times 10^{-15} \text{ cm}^2$. Their heights are adjusted to yield targeted equivalent interface defect densities (in cm^{-2}).

In the high quality crystalline substrate (bulk (n)c-Si), recombination centers are modelled with 10^9 cm^{-3} neutral defects located at 0.56 eV above E_V .¹⁷

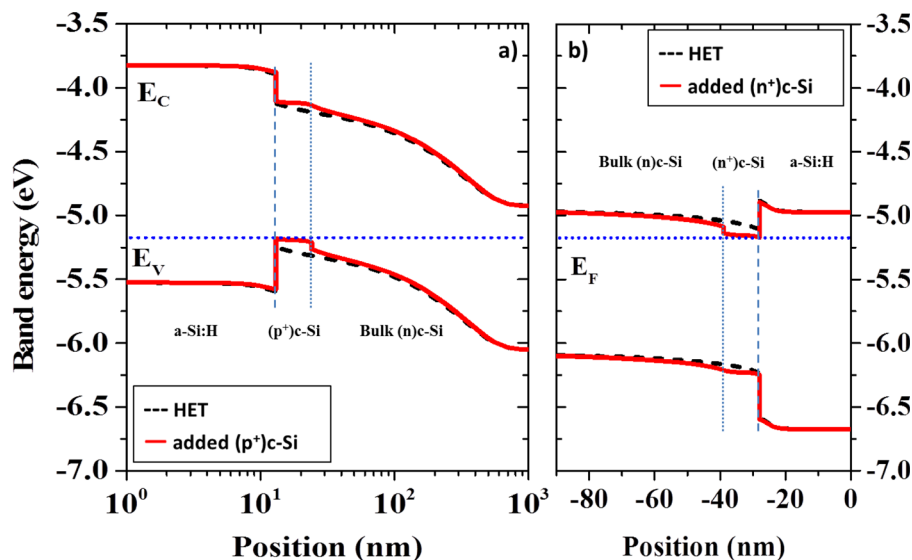


FIG. 2. Equilibrium band diagrams of the HET cell and the HHJ cells with the added layers at (a) the front interface and (b) at the back interface.

TABLE I. Layer parameter.

Layer parameters	Unit	(p)a-Si:H	(i)a-Si:H	(n)a-Si:H	Bulk (n)c-Si	Added (p ⁺)c-Si	Added (n ⁺)c-Si
Thickness	...	10 nm	3 nm	25 nm	150 μ m	0–1000 nm	10 nm
Optical and mobility gap	eV	1.7	1.7	1.7	1.124	1.074–1.124	1.124
Doping	cm ⁻³	6.1×10^{19}	0	1.7×10^{19}	2.0×10^{15}	1×10^{17} – 5×10^{18}	5×10^{18}
Relative dielectric constant	...	11.9	11.9	11.9	11.9	11.9	11.9
Electronic affinity	eV	3.8	3.8	3.8	4.05	4.05	4.05
Electron (hole) mobility	cm ² V ⁻¹ s ⁻¹	0.5 (0.5)	0.5 (0.5)	0.5 (0.5)	1321 (461)	225–1321 (96–461)	151.9 (164)
Total state density in CB (VB)	cm ⁻³	6.90×10^{20} (1.20×10^{21})	6.90×10^{20} (1.20×10^{21})	6.90×10^{20} (1.20×10^{21})	2.8×10^{19} (2.7×10^{19})	1.1×10^{19} (1.0×10^{19})	1.1×10^{19} (1.0×10^{19})
Neutral defect (at 0.56 eV) concentration	cm ⁻³	...	...	...	10^9	10^{14}	10^{14}
CB tail (VB tail) Urbach energy	eV	0.034 (0.060)	0.035 (0.050)	0.023 (0.060)	...	...	...
σ_e and σ_h for CB tail	cm ²	7×10^{-16}	7×10^{-16}	7×10^{-16}	...	...	...
σ_e and σ_h for VB tail	cm ²	7×10^{-16}	7×10^{-16}	7×10^{-16}	...	...	...
Maximum A- and D-like Gaussian state density	cm ⁻³ eV ⁻¹	1.20×10^{20}	1.00×10^{17}	2.25×10^{19}	...	...	...
Position of the A-(D-)like Gaussian (from valence band edge)	eV	1.450 (1.250)	1.025 (0.725)	0.850 (0.650)	...	...	...
Standard deviation of the A- and D-like Gaussian	eV	0.19	0.10	0.19	...	...	...
σ_e (σ_h) for A-like Gaussian states	cm ²	3×10^{-15} (3×10^{-14})	3×10^{-15} (3×10^{-14})	3×10^{-15} (3×10^{-14})	...	...	...
σ_e (σ_h) for D-like Gaussian states	cm ²	3×10^{-14} (3×10^{-15})	3×10^{-14} (3×10^{-15})	3×10^{-14} (3×10^{-15})	...	...	...

In the highly doped added layers, defects are also located at 0.56 eV above E_V but Shockley-Read-Hall (SRH) recombination dependence on doping is taken into account in the capture time constant: $\tau_{i0} = \tau_{i0_ini} * \left(\frac{10^{15}}{N_D}\right)^{0.4}$ ($i = p$ or n) and τ_{i0_ini} set at 10^{-7} s using the SRH parameters (thermal velocity, defect density, and capture cross section). The lifetime values obtained with this model are below the ones that can be found in literature from other models.^{18,19} As a consequence the potential beneficial effects of the $(p^+)c$ -Si layer insertion are not overestimated.

Regarding the carrier transport mechanism we considered drift-diffusion for homo-interfaces (c -Si/ c -Si and a -Si:H/ a -Si:H) and thermal emission with tunneling for hetero-interfaces (c -Si/ a -Si:H). Bandgap narrowing due to the high doping level of the added crystalline layers is taken into account. Interface defect densities (D_{it}) and external resistances were tuned so that the simulated HET cell displays performances comparable to that of state-of-the-art manufactured cells. D_{it} values are set at 5×10^9 and $1 \times 10^9 \text{ cm}^{-2}$ at the front and back hetero-interface, respectively. External resistances are set at 0.5 and $10^5 \Omega \text{ cm}^2$ for series and parallel resistances, respectively.

III. RESULTS

Figure 3 presents the HET and HHJ_{FS} cells' performance as a function of front interface defect density (D_{it}). Since the front contact limits the cells passivation, simplified architectures without back side (i) a -Si:H layer and interfacial defects are used for computing time issues. In the simplified architectures, D_{it} and series resistance are adjusted to $6 \times 10^9 \text{ cm}^{-2}$ and $0.3 \Omega \text{ cm}^2$, respectively, in order to keep the HET cell parameters.

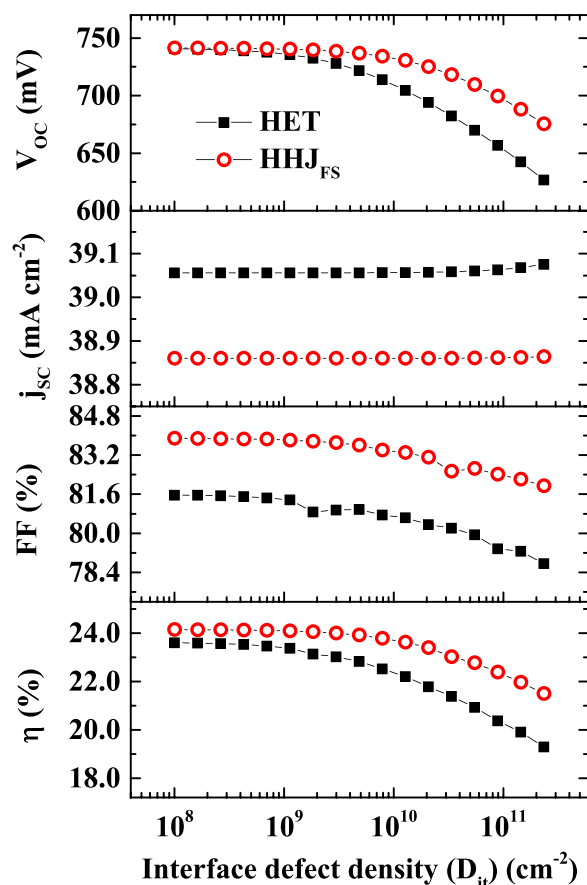


FIG. 3. Simplified HET (■) and HHJ_{FS} (red circle) cells output parameters versus D_{it} . In the HHJ_{FS} cell, the $(p^+)c$ -Si doping level and layer thickness are set at $5 \times 10^{18} \text{ cm}^{-3}$ and 10 nm, respectively.

In accordance with Zhong's results, V_{OC} and FF improvements are observed with the addition of the $(p^+)c\text{-Si}$ layer. Above $D_{it} = 10^9 \text{ cm}^{-2}$, V_{OC} decreases more slowly with increasing D_{it} for the HHJ_{FS} architecture, and FF is also slightly less sensitive to D_{it} than in the HET cell. The reduced D_{it} sensitivity seems to strengthen for high D_{it} . The short circuit current (j_{SC}) is only very slightly lower in the HHJ_{FS} cell. As a result, for any D_{it} , the HHJ_{FS} cell has higher power conversion efficiency than the HET cell. When both cells have equal V_{OC} (i.e., for low D_{it}), the HHJ_{FS} efficiency is still improved by 0.5% absolute (abs) thanks to the FF improvement.

The HHJ_{FS} concept is then extended by inserting a thin and highly doped $(n^+)c\text{-Si}$ layer at the $(i)a\text{-Si:H}/(n)c\text{-Si}$ back interface. Using the non-simplified architectures, the cell architectures (HHJ_{FS} , HHJ_{BS} , and HHJ_{FS+BS}) are compared in Table II. Both (n^+) - and $(p^+)c\text{-Si}$ layers have doping and thickness of $5 \times 10^{18} \text{ cm}^{-3}$ and 10 nm, respectively.

When adding a $(n^+)c\text{-Si}$ layer at the back interface of the HET and HHJ_{FS} cells (i.e., HHJ_{BS} and HHJ_{FS+BS} , respectively), V_{OC} and FF are only slightly improved and there is no j_{SC} decrease. As a result, the cells efficiency gain is very low as compared to the HET and HHJ_{FS} cells.

IV. DISCUSSION

A. Role of $(p^+)c\text{-Si}$ layer addition

From the band diagram presented in Figure 2, an improved field effect is observed with the highly doped layers addition. Therefore, V_{OC} and FF improvements brought by the $(p^+)c\text{-Si}$ layer addition at the front interface are expected to originate from an enhanced field effect passivation of this critical interface. Figure 4 introduces the recombination rate as a function of the position in the cell, near the front interface, for both simplified HET and HHJ_{FS} architectures in open-circuit conditions. In order to compare interface and bulk passivation, the recombination rate has been integrated over the layers thickness and expressed in $\text{cm}^{-2} \text{ s}^{-1}$ as shown in the inserted table in Figure 4.

Recombination at the HET cell front interface is much higher than in the $c\text{-Si}$ region and thus limits the passivation. At this critical interface, the recombination rate is substantially lower for the HHJ_{FS} cell. Therefore, V_{OC} and FF improvements in HHJ_{FS} are related to a lower recombination rate at the cell front interface. Since simulated HET and HHJ_{FS} cells have equal D_{it} , the improvement brought by the $(p^+)c\text{-Si}$ layer addition can be attributed to an enhanced field effect passivation. This is also confirmed by a lower electron concentration at the interface for the HHJ_{FS} cell ($2 \times 10^{13} \text{ cm}^{-3}$) compared to the HET cell ($8 \times 10^{13} \text{ cm}^{-3}$). The observed recombination rate increase in the HHJ_{FS} substrate is another indicator of the cell passivation improvement. Indeed, since there is less recombination at the interface, the free carrier concentration increases in the cell. On the one hand, a higher free carrier concentration in the substrate is related to a higher V_{OC} . On the other hand, recombination mechanisms (e.g., Auger or SRH)

TABLE II. Non-simplified HET and HHJ cell architectures parameter. Front and back interface have D_{it} equal to 5×10^9 and $1 \times 10^9 \text{ cm}^{-2}$, respectively.

	HET	HHJ_{FS}	HHJ_{BS}	HHJ_{FS+BS}
Added layers				
$(p^+)c\text{-Si}$ at front interface		X		X
$(n^+)c\text{-Si}$ at back interface			X	X
Cell parameters				
V_{OC} (mV)	713	730	716	737
j_{SC} (mA cm^{-2})	39.2	38.9	39.2	38.9
FF (%)	79.3	81.3	79.5	81.6
η (%)	22.2	23.1	22.3	23.4

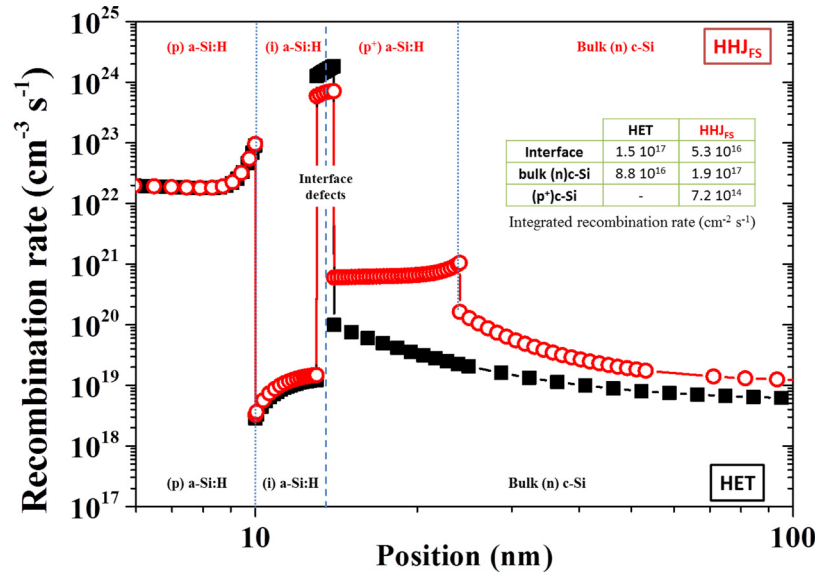


FIG. 4. Recombination rate in the simplified HET (■) and HHJ_{FS} (red circle) cells in V_{OC} conditions. The inserted table displays the integrated recombination rate at the interface and in the crystalline regions. In HHJ_{FS}, the (p⁺)c-Si layer doping density and thickness are set at $5 \times 10^{18} \text{ cm}^{-3}$ and 10 nm, respectively. For all structures D_{it} is taken equal to $6 \times 10^9 \text{ cm}^{-2}$.

are enhanced in the substrate. Thus, the observed recombination increase in the substrate is not a drawback but slightly moderates the (p⁺)c-Si layer benefits. However, due to its high doping level, the (p⁺)c-Si layer is more defective and introduces strong additional recombination.

The j_{SC} decrease with the (p⁺)c-Si layer addition is later discussed in Sec. IV C.

In an experimental HHJ_{FS} cell, the FF could be further enhanced by the (p⁺)c-Si layer which promotes carrier lateral transport within the inversion layer. Indeed, due to the high band bending in crystalline silicon (see Figure 2(a)), the c-Si substrate type is inverted near the hetero-interface and the p-n junction is located in the crystalline wafer.^{20,21} The resulting inversion layer is very conductive compared to amorphous silicon layers and thus might promote lateral transport.^{22,23} As a consequence, improving the doping in this region by the (p⁺)c-Si layer addition can enhance lateral transport in the inversion layer leading to even higher FF. This phenomenon cannot be directly observed in our simulations since Afors-HET is a 1D simulation model. However, from the simulated minority carrier density, the sheet resistance of the inversion layer was calculated. HHJ_{FS} inversion layer sheet resistance ($7.1 \times 10^3 \Omega/\square$) is twice lower than the HET inversion layer ($1.5 \times 10^4 \Omega/\square$) but one order of magnitude higher than the Indium Tin Oxide (ITO) layer on top of the cell ($\sim 1 \times 10^2 \Omega/\square$). Therefore, it can be concluded that the (p⁺)c-Si layer enhances the lateral transport in the inversion layer but should not contribute significantly to further FF improvement.

The reduced D_{it} dependence of the HHJ_{FS} architecture opens a new way to take advantage of the architecture: rather than enhancing the conversion efficiency, the HHJ_{FS} cell could be a way for cost savings. For instance reducing interface cleaning steps duration (or passivation layers thickness) might decrease the manufacturing costs while keeping similar cell efficiency.

From Figure 4, we have seen that the (p⁺)c-Si layer decreases recombination at the front interface but also that the layer is highly recombining. Though for a 10 nm-thick (p⁺)c-Si layer the added recombination within the layer ($7.2 \times 10^{14} \text{ cm}^{-2} \text{ s}^{-1}$) is much lower than at the HHJ_{FS} cell interface ($5.3 \times 10^{16} \text{ cm}^{-2} \text{ s}^{-1}$), it is nevertheless a counterpart to the added layer benefits. Additionally, with increasing layer thickness, recombination in the (p⁺)c-Si layer could also become predominant. Therefore, (p⁺)c-Si parameters such as doping level and layer thickness should be optimized to maximize the added layer benefits. This is precisely the objective of Sec. IV B.

B. (p⁺)c-Si layer optimization

In Figure 5, V_{OC} and recombination rates at the front interface and within the (p⁺)c-Si layer are plotted versus the layer doping level for simplified HET and HHJ_{FS} cells in open circuit voltage conditions.

(p⁺)c-Si doping higher than $5 \times 10^{18} \text{ cm}^{-3}$ could not be considered. Indeed, for higher doping the Fermi energy enters into the valence band (below E_V). In this situation, the Fermi–Dirac statistic that is not implemented in the software should be considered instead of Boltzmann's statistics.

From Figure 5, it is seen that improving the (p⁺)c-Si layer doping decreases recombination both at the interface and within the (p⁺)c-Si layer. Therefore, field effect passivation is beneficial to repel electrons from both the interface and the recombining (p⁺)c-Si layer. This explains the monotonous passivation increase with increasing the (p⁺)c-Si doping. No j_{SC} variation with doping was observed. Therefore, having a high (p⁺)c-Si doping is beneficial for the cell passivation and performance.

However, because of the high recombination rate within the (p⁺)c-Si layer, there is, for a given layer doping, an optimum layer thickness which maximizes the HHJ_{FS} cell efficiency. Simplified HHJ_{FS} cell output parameters are plotted versus the (p⁺)c-Si layer thickness in Figure 6. In Table III, recombination rates within the cell are presented for 10, 100, and 1000 nm (p⁺)c-Si layer thickness.

For a (p⁺)c-Si layer thickness of approximately 10 nm, the V_{OC} and FF of the HHJ_{FS} cell rise up to their maximum. At this point, recombination at the interface has become lower than in the substrate (bulk (n)c-Si) and recombination brought by the (p⁺)c-Si layer addition is very low. Then, due to the recombination brought by the added layer, these parameters slowly decrease for an increasing (p⁺)c-Si thickness. The decrease is slow because the passivation is now limited by recombination in the substrate.

j_{SC} decreases very slowly with increasing the layer thickness. The sudden FF drop around 150 nm could not be physically explained. Therefore, it is supposed to originate from calculation issues. It has nevertheless a small influence on the cell efficiency.

From these observations, optimum thicknesses calculated for the tested (p⁺)c-Si layer doping and the corresponding maximized HHJ_{FS} cell efficiency are presented in Figure 7.

When increasing the doping density, the maximum achievable efficiency increases. This is in accordance with the passivation improvement previously observed both at the interface and within the (p⁺)c-Si layer with increasing doping. However, the corresponding optimum thickness decreases for increasing doping. This results from stronger recombination within the (p⁺)c-Si layer at high doping level.

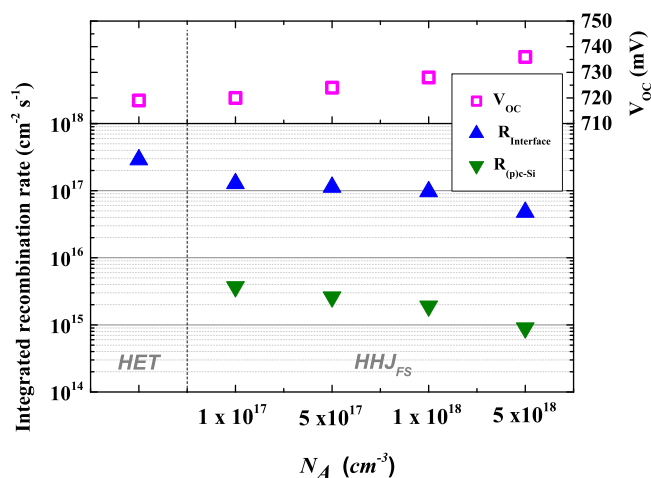


FIG. 5. V_{OC} and integrated recombination rates (R) of simplified HET and HHJ_{FS} cells at the interface and within the added (p⁺)c-Si layer versus layer doping level. In HHJ_{FS}, the (p⁺)c-Si layer is 10 nm thick.

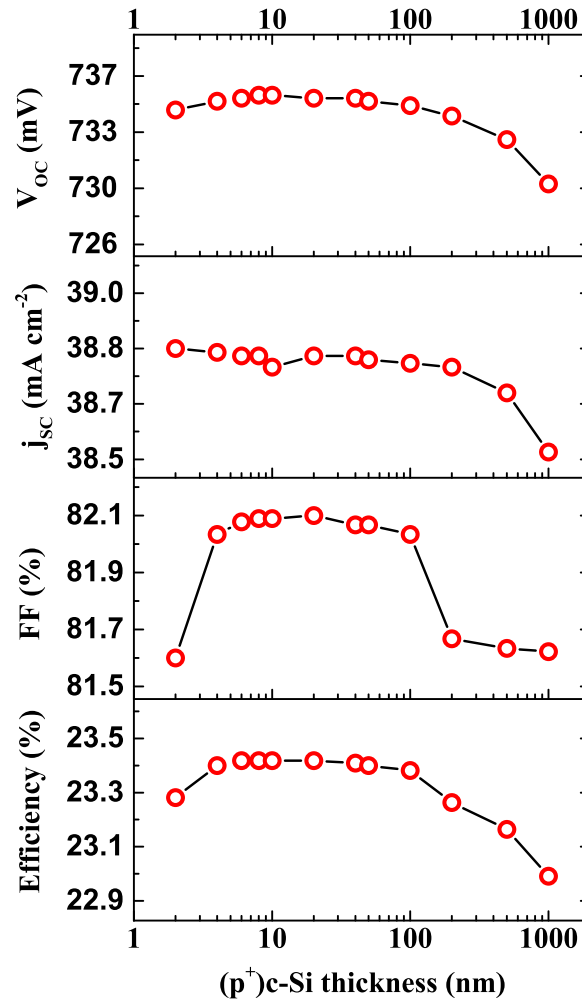


FIG. 6. Simplified HHJ_{FS} cell output parameters versus (p⁺)c-Si layer thickness. (p⁺)c-Si layer doping is set at $5 \times 10^{18} \text{ cm}^{-3}$.

From a practical point of view, such very thin optimized layers (<40 nm) would be difficult to experimentally achieve on large scale cells. However, from Figure 6, it is seen that the HHJ_{FS} cell performance decreases only slightly for (p⁺)c-Si layer thicknesses above its optimum. Therefore, under 100 nm thick, the (p⁺)c-Si layer will retain most of the architecture advantages. Such layer thickness is nowadays achievable at relatively low cost thanks to alternative ion implantation and fast annealing processes.^{24,25} However, it should be noticed that the (p⁺)c-Si layer created by ion implantation consists in a dopant depth profile rather than an homogeneous doping through the layer thickness, as simulated here. Thus, further work on dopant depth profile influence should be done.

TABLE III. V_{OC} and integrated recombination rate (R) at the interface, within the (p⁺)c-Si layer, and in the substrate for the simplified HHJ_{FS} cell with increasing (p⁺)c-Si layer thickness. (p⁺)c-Si layer doping is set at $5 \times 10^{18} \text{ cm}^{-3}$.

(p ⁺)c-Si layer thickness (nm)	V _{OC} (mV)	R _{Interface} (cm ⁻² s ⁻¹)	R _{(p+)-c-Si} (cm ⁻² s ⁻¹)	R _{substrate} (cm ⁻² s ⁻¹)
0	719	1.5×10^{17}	...	8.8×10^{16}
10	736	5.3×10^{16}	7.2×10^{14}	1.9×10^{17}
100	735	5.3×10^{16}	6.5×10^{15}	1.8×10^{17}
1000	730	4.4×10^{16}	5.3×10^{16}	1.4×10^{17}

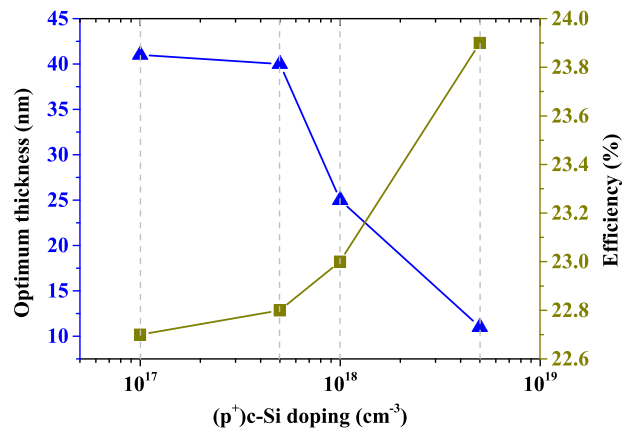


FIG. 7. (p⁺)c-Si layer optimum thickness and related maximized HHJ_{FS} cell efficiency as a function of the added layer doping.

The HHJ_{FS} cell studied in Sec. IV A corresponds to an optimized cell for a doping density of $5 \times 10^{18} \text{ cm}^{-3}$. As seen in the table of Figure 4, this cell is now limited by bulk recombinations since the bulk substrate has the highest integrated recombination rate. Therefore, substrate improvement is required to further enhance the HHJ_{FS} structure.

C. Addition of a (n⁺)c-Si layer at the back (n)c-Si/(i)a-Si:H interface

A main feature of the results presented in Table II is that the HHJ architectures only display significant passivation improvement (i.e., FF and V_{OC} improvements) if they have the added (p⁺)c-Si layer at the front interface (HH_{FS} and HH_{FS+BS}). The addition of the (n⁺)c-Si layer at the back interface does not improve significantly the cells performance. The recombination rates within the cell explain this behavior. Figure 8 presents integrated recombination rates at the front interface, back interface, and within the substrate (bulk (n)c-Si) and the corresponding V_{OC} for HET and HHJ cells. As previously seen, recombination within the added doped layers can be neglected since the layers are very thin (10 nm).

The recombination rate at the HET cell front interface is one order of magnitude higher than at the back interface. Thus, the HET cell performance is limited by the front interface. The HHJ_{FS} cell has its V_{OC} limited by substrate recombination. Therefore, though the (n⁺)c-Si layer addition in these cells decreases recombination at the back interface, the HHJ_{BS} and HHJ_{FS+BS} cells display almost no performance enhancement as compared to the HET and

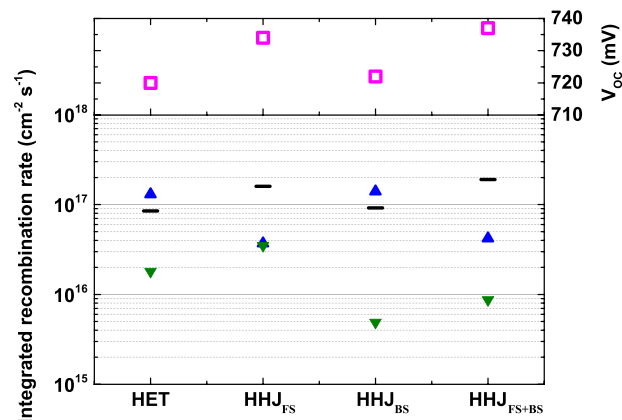


FIG. 8. V_{OC} and integrated recombination rates (R) at the front interface, back interface and within the substrate for HET and HHJ architectures. Thicknesses and doping densities are, for both added layers, set at 10 nm and $5 \times 10^{18} \text{ cm}^{-3}$, respectively.

HHJ_{FS} cells, respectively. The increase of recombination rate at the back interface with the addition of the (p⁺)c-Si is related to the same mechanism as that inducing the increase in substrate recombination (see Sec. IV A).

From these observations, it can be inferred that there is no great benefit of adding an (n⁺)c-Si layer at the back interface of neither the HET nor the HHJ_{FS} cell. Therefore, as previously discussed, further passivation improvement should be reached first by substrate improvement rather than back interface improvement.

In Table II, a j_{SC} decrease is observed with the front side layer addition only. This loss originates from light absorption in the added (p⁺)c-Si layer which is heavily doped and highly recombining. Therefore, carriers photo-generated in this region cannot fully contribute to the cell current density. Moreover, due to its high doping level, there is also a band gap narrowing²⁶ effect (see Figure 2) which promotes the light collection in the layer and thus participates to the current density reduction.

The added (n⁺)c-Si layer is located at the cell back side which is reached by less photons than the front side. Therefore, there are much less optical losses due to light absorption in the (n⁺)c-Si layer leading to no j_{SC} loss. This observation opens a new route for a better HHJ_{FS} use: an inverted HHJ_{FS} cell, with emitter at the cell back side,²⁷ would benefit from the passivation improvement without suffering from the j_{SC} losses. This idea was validated by our simulations. Because there is no more j_{SC} loss, the (p⁺)c-Si layer thickness in the inverted HHJ_{FS} cell should be optimized again.

V. CONCLUSION

An improved silicon heterojunction cell architecture where a (p⁺)c-Si layer is added at the front (i)a-Si:H/(n)c-Si interface has been studied by means of numerical simulations. The obtained HHJ_{FS} cell displays improved V_{OC} and FF. These improvements have been evidenced to originate from an enhanced interfacial field effect passivation. The added (p⁺)c-Si layer has been optimized. In order to maximize the hetero-homojunction power conversion efficiency, the (p⁺)c-Si layer doping should be as high as possible to promote field effect passivation at both the interface and in the (p⁺)c-Si layer. The corresponding optimum thickness, for the tested doping levels (up to $5 \times 10^{18} \text{ cm}^{-3}$), is always very thin (<40 nm). Therefore, we have evidenced the need of ultra-shallow junctions to benefit from such a layer addition. However, under 100 nm thick, the added layer is thin enough to retain the HHJ cell architecture advantages. Such thin layers should be experimentally achievable at relatively low cost. Finally, adding an (n⁺)c-Si layer at the back (n)c-Si/(i)a-Si:H interface was tested. This does not provide a great benefit because passivation of the HET and HHJ_{FS} cells are not limited by the back interface. Experiments are ongoing to experimentally achieve hetero-homojunction cells.

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