Simulation-based roadmap for the integration of poly-silicon on oxide contacts into screen-printed crystalline silicon solar cells

Christian N. Kruse1,2, Sören Schäfer2, Felix Haase2, Verena Mertens2, Henning Schulte-Huxel2, Bianca Lim2, Byungsul Min2, Thorsten Dullweber2, Robby Peibst2,3 & Rolf Brendel1,2

We present a simulation-based study for identifying promising cell structures, which integrate poly-Si on oxide junctions into industrial crystalline silicon solar cells. The simulations use best-case measured input parameters to determine efficiency potentials. We also discuss the main challenges of industrially processing these structures. We find that structures based on p-type wafers in which the phosphorus diffusion is replaced by an n-type poly-Si on oxide junction (POLO) in combination with the conventional screen-printed and fired Al contacts show a high efficiency potential. The efficiency gains in comparison to the 23.7% efficiency simulated for the PERC reference case are 1.0% for the POLO BJ (back junction) structure and 1.8% for the POLO IBC (interdigitated back contact) structure. The POLO BJ and the POLO IBC cells can be processed with lean process flows, which are built on major steps of the PERC process such as the screen-printed Al contacts and the Al2O3/SiN passivation. Cell concepts with contacts using poly-Si for both polarities (POLO2-concepts) show an even higher efficiency gain potential of 1.3% for a POLO2 BJ cell and 2.2% for a POLO2 IBC cell in comparison to PERC. For these structures further research on poly-Si structuring and screen-printing on p-type poly-Si is necessary.

Passivating contacts enable selectivities with low recombination currents and contact resistances and therefore allow higher cell efficiencies than conventional diffused contacts, as can be found in passivated emitter and rear cells (PERC). One prominent passivating contact scheme, the heterojunction between amorphous (a-) H-rich Si and crystalline (c-) Si, provides the base for highly-efficient industrial double-side contacted cells with efficiencies up to 25.1%3 and is implemented in a lab-type interdigitated back contact (IBC) cell with the current Si world record efficiency of 26.7%4. Another prominent passivating contact scheme, which we denote as “Poly-silicon on oxide (POLO) junctions”, consists of a stack of an interfacial oxide and a doped (partially) crystalline Si layer. Its potential advantage as compared to a-Si:H/c-Si heterojunctions is that it is high-temperature stable and thus compatible with the manufacturing processes of today’s mainstream solar cell technology. Solar cell concepts featuring POLO or related junctions achieved high efficiencies up to 26.1% (26.0%) in lab-type IBC (double-side contacted)3,4 cells. In comparison, solar cell manufacturers are mainly producing mono-facial PERC and bifacial PERC+ solar cells5 with efficiencies around 22.5%-9. However, although numerous manufacturers have started (pilot) production of solar cells featuring POLO junctions or are producing similar cell concepts on rather small scale since some years with impressive results7-9, these approaches are in strong competition with PERC and PERC+ as the “conventional” current mainstream. Some reasons for this hard standing are an increased process complexity and an uncertainty on the efficiency potential of the respective new cell structures. In this paper, we analyze different options for a lean POLO implementation into new and existing solar cell structures and compare

1Institute for Solid State Physics, Leibniz Universität Hannover, Appelstrasse 2, 30167 Hannover, Germany. 2Institute for Solar Energy Research Hamelin (ISFH), Am Ohrberg 1, Emmerthal 31860, Germany. 3Institute Electronic Materials and Devices, Leibniz Universität Hannover, Schneiderberg 32, 30167 Hannover, Germany. *email: brendel@isfh.de

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These alternatives by means of a simulation-based study. Structures featuring POLO junctions have already been the subject of experimental and simulation-based studies\(^4\──^6\). However, due to the different experimental equipment, simulation approaches, input parameters and focus of the studies, the efficiency values extracted from these studies are not comparable. For some specific cell structures and specific realization routes, experimental building blocks have been evaluated, e.g. in Ref.\(^1\). Here, we focus on good comparability of the simulated efficiency between all analyzed cell structures and discuss the main processing challenges to give an impression of the expected process complexity. Besides the systematic screening of numerous cell structures, we also compare them in-depth based on their full Synergistic Efficiency Gain Analysis (SEGA). In addition, the simulation base is used to assess the sensitivity of the discussed solar cell structures to bulk lifetime variations. The goal of our study is to identify cell structures that show a high efficiency potential along with potentially lean process flows. From this discussion, a possible roadmap for industrializing p-type c-Si solar cells using POLO junctions is derived. (The experimental realization of the most promising cell structures will be the subject of future work.).

**Simulation method**

Two important aspects when evaluating new cell structures for their potential economic benefit in large-scale production are the process complexity and the efficiency gain compared to proven concepts. To estimate the potential efficiency gain, we perform numerical device simulations using the Quokka\(^13\) implementation of the conductive boundary model\(^14\). These simulations require the knowledge of the resistive and recombination properties for the bulk, the passivated surfaces and the contacts as well as a charge-carrier generation profile for each structure. For a good comparability between all analyzed cell structures we choose equal input parameters for equal or comparable properties of the respective structures. For example, the recombination at a 200 nm thick n-type poly-Si junction is always assumed to be \(J_0 = 3 \text{fA} / \text{cm}^2\) although in reality this \(J_0\) can vary for different cell structures due to the influence of other process steps on the passivation quality. In general, we choose best-case measured parameters achieved in lab-type production with industrial-type processes. We confine our analysis to screen-printed contacts, because most industrial solar cells are produced using screen-printing. Combining all these best-measured parameters into a hypothetical simulated cell gives an estimate for cell efficiencies to be expected in the next years for cells with POLO junctions and for PERC+ cells. All input parameters are detailed in Table 1.

For the optical performance of each structure we employ ray-tracing simulations using the SUNRAYS ray-tracer\(^15\) with optical parameters chosen according to Refs.\(^16\──^18\). This program does, however, not allow simulating the complex unit cell shown at the top of Fig. 1. Therefore, we resolve the complex optical structure into simple unit-cells shown in the second row of Fig. 1 and simulate them separately. Each of these simulations yields a charge-carrier generation rate. Underneath the metal no optical generation is assumed. The generation profiles are averaged according to their area fractions. It should be noted that we assume an external mirror with ideal specular reflectance on the rear side in the simulations to represent a measuring chuck or white module back-sheet. The effect of this mirror on the simulated cell efficiencies is comparable for all structures because they all have a textured front and a planar rear side. Note that while the area-averaging approach accounts for differences in front surface metallization and parasitic absorption in the poly-Si, it lacks the impact of multiple internal reflections at, e.g. metal contacts. However, we estimate the additional parasitic absorption at the contacts to be

### Table 1. Electronic input parameters for the device simulations. All parameters without a reference are ISFH internal measurements.

| Parameter | Recombination | Resistive |
|-----------|---------------|-----------|
| p-type (B) bulk | \(\tau_0 = 2000 \mu s\) \(\tau_{\text{PA}} = 20,000 \mu s\) | \(\rho_s = 0.9 \Omega \text{ cm}\) |
| n-type (P) bulk | \(\tau_0 = 22,000 \mu s\) | \(\rho_s = 3 \Omega \text{ cm}\) |
| Pass. on p-Si (plan.) | \(J_0 = 1 \text{A} / \text{cm}^2\) | n.a. |
| Pass. on p-Si (test.) | \(J_0 = 3 \text{A} / \text{cm}^2\) | n.a. |
| Pass. on n-Si | \(J_0 = 3 \text{A} / \text{cm}^2\) | n.a. |
| emitter P-diff. | \(I_0 = 22 \text{fA} / \text{cm}^2\) | \(R_s = 133 \Omega / \Omega\) |
| selective P-diff. | \(I_0 = 100 \text{fA} / \text{cm}^2\) | \(R_s = 95 \Omega / \Omega\) |
| emitter B-diff. | \(I_0 = 14 \text{A} / \text{cm}^2\) | \(R_s = 135 \Omega / \Omega\) |
| selective B-diff. | \(I_0 = 110 \text{fA} / \text{cm}^2\) | \(R_s = 95 \Omega / \Omega\) |
| n-type POLO | \(I_0 = 3 \text{fA} / \text{cm}^2\) | \(R_s = 50 \Omega / \Omega\) |
| p-type POLO | \(I_0 = 5 \text{fA} / \text{cm}^2\) | \(R_s = 200 \Omega / \Omega\) |
| Ag on P cont. | \(I_0 = 1400 \text{fA} / \text{cm}^2\) | \(\rho_s = 1.5 \mu \text{m} \Omega \text{ cm}^2\) |
| Al on P cont. | \(I_0 = 400 \text{fA} / \text{cm}^2\) | \(\rho_s = 1.3 \mu \text{m} \Omega \text{ cm}^2\) |
| Ag on B cont. | \(I_0 = 740 \text{fA} / \text{cm}^2\) | \(\rho_s = 2 \mu \text{m} \Omega \text{ cm}^2\) |
| Ag on n-POLO cont. | Same as n-POLO | \(\rho_s = 0.9 \mu \text{m} \Omega \text{ cm}^2\) |
| Ag on p-POLO cont. | Same as p-POLO | \(\rho_s = 5 \mu \text{m} \Omega \text{ cm}^2\) |
below 10% of the total parasitic absorption within the cell. In addition, all structures have similar light trapping schemes and, thus, no structure is optically favored.

**Investigated solar cell structures**

Most of the recently installed industrial solar cell production lines produce PERC and PERC+ solar cells, which have a high conversion efficiency and allow low production costs. However, the diffused contacts and front contact shading suggest an efficiency limitation at around 24%\(^{19}\). Figure 2a shows a synergistic efficiency gain analysis (SEGA)\(^{20}\) of an industrial-type PERC+ cell with best-measured input parameters given below. It should be noted that we only consider busbar-less cells in this work. The SEGA yields potential efficiency gains in solar cells by switching off the respective loss channel in the cell simulation and calculating the difference in conversion efficiencies. Synergistic effects between multiple loss channels are analyzed by switching off multiple loss channels simultaneously. The simulated efficiency is 23.7%, which agrees with the efficiency to be expected from PERC+ production lines within the next years given today’s efficiencies of around 22.5% and a 0.5% per year learning curve according to Ref.\(^{21}\). The efficiency gains due to a reduction of the front finger shading and a reduction of recombination in the emitter and at the front contact are the largest individual gains possible besides the synergistic gain between the extrinsic recombination channels. Therefore, the Ag contacts and the phosphorus diffusion are the bottlenecks on the road towards higher efficiencies. Here n-type POLO junctions offer better electrical performance. The so far most common approach for their implementation is the industrialization of the TOPCon\(^{8,12}\) cell structure. The corresponding SEGA is shown in Fig. 2b. For the input parameters assumed in this study, the resulting simulated efficiency is 24.4%, i.e., 0.7% higher than that for PERC+. This difference is mainly a consequence of the mitigation of the contact recombination at the electron-collecting contact, the lower \(J_0\) of the B-doped front emitter, and the synergy between both loss channels. We acknowledge that our simulated efficiency for industrial TOPCon has already been surpassed experimentally: To our knowledge, the highest ISO 17025/IEC 6094 calibrated efficiency reported so far is 24.8%\(^{9}\). Obviously, it is possible to optimize single components further than assumed in Table 1. However, as discussed below, it still has to be shown that all possible measures for the optimization of a record cell (e.g. extremely deep emitters) are compatible with mass production. The median efficiency in production is currently around 23%\(^{8}\). It is still not yet clear whether TOPCon is the best suited cell structure for an integration of POLO junctions into industrial production.

Therefore, we investigate six alternative structures featuring POLO junctions and compare their performance potential and processing complexity with that of PERC+ cells and with a TOPCon\(^{8,12}\) cell structure. It should be noted that these are cell structures we regard as interesting candidates for industrial integration after preliminary simulations and discussion, which are not shown here. There are, of course, many other possible cell structures and our list does not aim at being all-encompassing. In particular we focus our discussion on p-type doped Si solar cells, because it allows the re-utilization of established production steps. The six cell structures investigated are shown in Fig. 3 along with two reference structures. (a) the reference PERC+ cell, (b) the reference TOPCon cell based on n-type Si with a diffused boron emitter and a full-area n-type POLO junction on the rear, (c) the PERC+ POLO structure, which is a PERC+ cell with n-type POLO under the front contacts, (d) the POLO B
structure, which features Al-alloyed (Al-p+) contacts on the front and a full-area n-type POLO on the rear side,
(e) the POLO IBC structure, which is an IBC concept featuring Al-p+ and n-type POLO junctions,
(f) the POLO2 BJ structure, which features local p-type POLO junctions on the front side and a full-area n-type POLO emitter on the rear,
(g) the POLO2 IBC structure, which is an IBC concept with POLO junctions for both polarities, and
(h) the n-Si POLO2 BJ structure, which is the same structure as (f) but with reversed polarities.

The six cell structures (c) through (f) feature either only one n-type poly-Si contact and an Al-p+ contact (POLO cell) or two POLO junctions (POLO2 cells) in either front junction (FJ), back junction (BJ) or interdigitated back contact (IBC) configuration. We acknowledge the pioneering work of other groups (including companies) on many of these or related cell structures. For example, although not all details have been disclosed, the concept of TetraSun was reminiscent of the n-Si POLO2 BJ structure, and SunPower’s Maxeon technology seems to correspond to a n-Si POLO2 IBC structure.

It should be noted that for all structures either the structured layers underneath the contacts (selective emitter or POLO) are wider than the metal contact or the metal is wider than the LCO in case of Al contacts. This is due to alignment tolerances required for screen-printing, which we consider in our simulation study. The Ag contacts for which we assume a printing width of 30 µm are printed on 75 µm-wide stripes of either POLO layers or the laser-doped selective emitter. For the Al contacts we assume an LCO width of 13 µm and 50 µm-wide Al fingers. For all structures we individually optimize the front contact pitches. For the IBC cells we assume an emitter coverage of 80%. For the rear side, smaller contact pitches are in most cases preferable over larger pitches due to reduced resistance contributions. We, therefore, choose a rear contact pitch of 0.5 mm for all cells.

Electronic input parameters

The electronic input parameters chosen in this work are values measured mostly on reference samples manufactured with lab-type processes that can be transferred to industrial-scale production. Where possible we choose values measured at ISFH because this assures the best consistency between the different parameters. All input parameters are shown in Table 1. Values without a reference are unpublished ISFH internal measurements. At ISFH we measure the lifetime of non-metallized samples using a WCT-120 lifetime tester by Sinton instruments and determine the surface recombination using the in-built Kane–Swanson analysis tool. The recombination at metallized surfaces is determined by comparing samples with different metallization fractions and fitting the respective lifetime, measured by photoluminescence emission, using Quokka. Bulk and sheet resistances are determined using a four-point probe setup whereas contact resistivities are determined using the transfer-length-method (TLM). It should be noted that some values differ from the ones used in our recently published paper in Ref., because they were either just recently measured or a publication exists for a similar value, which makes it easier for the reader to check the experimental details of the input parameters.

The boron-doped Czochralski-grown material that we use in our analysis was measured at ISFH with \( n_b = 0.9 \) Ω cm and mid-gap Shockley–Read–Hall lifetime parameters \( \tau_{\text{RD}} = 2000 \mu s \) and \( \tau_{\text{QH}} = 20,000 \mu s \). The phosphorus-doped n-type material was also measured at ISFH with \( n_b = 3 \) Ω cm and an injection-independent SRH lifetime of \( \tau_b = 22,000 \mu s \). For the surfaces passivated with an Al2O3 /SiN stack we determine a \( J_0 \) of 1 fA/cm² on planar

![Figure 2](https://example.com/figure2.png)

Figure 2. SEGA for (a) the reference PERC+ cell and (b) the TOPCon cell as a further benchmark for the alternative cell structures discussed in this study. Bars show potential efficiency gains due to complete suppression of different loss channels. Red bars are linked to recombination, green bars to optics, and blue bars to resistance. The gray bars show synergistic gains.
and 3 $\text{fA/cm}^2$ for the passivation on textured $p$-type material. For $n$-type material we assume a passivation quality equal to that on $p$-type. For the phosphorus-diffused surfaces we determine $J_0 = 22 \text{fA/cm}^2$ and $R_s = 133 \Omega/\square$ for the inter-finger region$^{25}$. For the selective doping underneath the front contacts we determine $J_0 = 100 \text{fA/cm}^2$ and $R_s = 95 \Omega/\square$. For the planar $n$- and $p$-type POLO junctions we assume 200 nm thickness for the calculation of the optical performance. We assume this thickness to be required for a screen-printed POLO contact to retain its recombination properties. Lower thickness values are possible in combination with other metallization schemes that are not considered here. For planar $n$-type POLO layers we measure a $J_0$ of 3 $\text{fA/cm}^2$ and a sheet resistance of 50 $\Omega/\square$. For $p$-type POLO layers we measure a $J_0$ of 5 $\text{fA/cm}^2$ and a sheet resistance of 200 $\Omega/\square$. We determine the contact recombination of Ag contacts on a $p$-diffusion on $J_0 = 1400 \text{fA/cm}^2$ in low injection. For the Ag on $P$ contact we measure a contact resistivity of $\rho_c = 1.5 \text{m}\Omega \text{cm}^2$. For the Al-$p^+$ contacts we determine $J_0 = 400 \text{fA/cm}^2$ and $\rho_c = 1.3 \text{m}\Omega \text{cm}^2$. For the Ag contact on boron diffusion we again refer to the literature with $J_0 = 740 \text{fA/cm}^2$ and a $\rho_c = 2 \text{m}\Omega \text{cm}^2$. For contacts on $n$-type POLO formed with firing-through pastes we observe that the contacts do not contribute to the recombination. These contacts were measured with a resistivity of $\rho_c = 0.9 \text{m}\Omega \text{cm}^2$. For $p$-type POLO the firing-through pastes yield high $J_0$-values of around 250 $\text{fA/cm}^2$ as published by Mack et al.$^{36}$ However, non-firing-through pastes on $p$-type POLO lead to insignificant recombination similar to $n$-type POLO contacts with a resistivity of $\rho_c = 5 \text{m}\Omega \text{cm}^2$ as measured at ISFH$^{33}$.

Results and discussion

Table 2 shows the simulated I–V parameters for each of the cell structures shown in Fig. 3. The results show clear trends in terms of increasing $V_{OC}$ and $J_{SC}$: $V_{OC}$ increases with the reduction of recombination losses at conventional contacts and diffusions. Consequently, the POLO$^2$ concepts show the highest $V_{OC}$ followed by the POLO
In the following, we discuss the different cell structures in detail: a) Our reference PERC+ cell shows a front-side diffusion for introduction of n+-type doping, which is an expensive process without the necessity of any waste management. A further technological advantage of the PERC+ structure is that Al alloying during firing forms the contact for the second polarity (holes) inexpensively. The rear side metallization furthermore only contains a limited amount of silver (for soldering pads, not for fingers). Thus, the PERC+ structure poses a tough benchmark (with respect to cost per watt, but also to Levelized Costs of Electricity) for any other cell concept.

Table 2. Simulation results.

| Cell concept | η [%] | J SC [mA/cm²] | V OC [mV] | FF [%] |
|--------------|-------|---------------|-----------|-------|
| a) PERC      | 23.7  | 41.3          | 697       | 82.5  |
| b) TOPCon    | 24.4  | 41.2          | 712       | 83.1  |
| c) PERC+ POLO| 24.1  | 40.9          | 712       | 82.8  |
| d) POLO BJ   | 24.7  | 40.6          | 736       | 82.7  |
| e) POLO IBC  | 25.5  | 41.7          | 733       | 83.4  |
| f) POLO BJ   | 25.0  | 40.6          | 742       | 83.0  |
| g) n-Si POLO BJ | 25.9 | 41.7        | 742       | 83.5  |
| h) n-Si POLO BJ | 25.1 | 40.9        | 737       | 83.3  |
b) A widely evaluated concept is the TOPCon concept based on an n-type Si wafer with n-type POLO on the rear and a diffused boron emitter on the front. We include this concept as a second reference. In our simulation study, it yields a front-side efficiency potential of 24.4%, i.e., 0.7% higher than that of the PERC+ cell. The bifaciality of industrial TOPCon cells is reported to be in the range of 80-90%, i.e., comparable to that of PERC+.

In terms of technology, the process sequence for the fabrication of a TOPCon cell strongly depends on the availability of real single-side Si deposition techniques (without wrap-around) and on the availability of firing-only contacts. Without the latter, two high temperature processes—one for BBr3 diffusion and one for POLO junction formation—are required. The reason is that the thermal budgets of both processes are typically quite different (unless “thick” interfacial oxide is used), and that the boron silicate glass and the boron doping from the rear has to be removed prior to the poly-Si deposition. Replacing the BBr3 diffusion by single-side boron doping techniques like AP-CVD deposition of boron-doped glasses could eliminate the latter necessity. The boron doping profile can be driven-in several micrometers in order to mitigate contact recombination on the front - in expense of process time. Selectively doped boron emitters are less mature than their phosphorus-doped counterparts, and most work in this direction is so far based on etch-back approaches rather than on laser doping. If the poly-Si deposition yields an intolerable wrap-around, the poly-Si has to be removed selectively from the already present boron front-side emitter. To enable this step, an alkaline-resistant etch-stop layer on the front side (deposited prior to poly-Si deposition) could be required. This layer could also serve as a diffusion barrier when doping the poly-Si on the rear in-situ via POCl3 diffusion. A cost disadvantage of the TOPCon structure—as compared to the PERC+ cell—is that the Ag consumption on the rear is increased due to the necessity to print Ag fingers for contacting the poly-Si without alloying through it. Besides cost aspects, it is worth to mention that contacting of n+ type poly-Si with firing-through Ag pastes without degrading the passivation quality works fairly well. Since there are controversial statements on cost perspectives for n-type wafers from the industry, it is difficult from our perspective to arrive at a clear answer on this issue. Also the relevances of n-type material with respect to some metal contaminations is difficult to assess. Under “clean” process conditions, the bulk lifetime in p-type material (and, more important, the diffusion length) can be comparable to that of excellent n-type material (see our results above), but the conditions in industrial environment might be different. Last but not least, the absence of Light Induced Degradation is currently an advantage of n-type material, which might vanish in the near future due to the availability of low [O] B-doped or of Ga-doped p-type material, as well as due to the readiness of industrial feasible permanent BO deactivation schemes.

c) The PERC+ POLO cell replaces the selective emitter of the PERC+ cell with a local n-type POLO layer. This brings an efficiency gain of 0.3% due to a higher open-circuit voltage (+15 mV) and fill factor (+0.3%) because of the reduced recombination and contact resistance. This positive effect overcompensates the loss in the short-circuit current density (−0.4 mA/cm²) due to absorption and reflection by the planar POLO layer in the finger regions.

In terms of technology, the process sequence for the fabrication of a PERC+ POLO cell is identical to that of a PERC+ cell for passivation and all steps afterwards. However, the necessity to structure the poly-Si on the front can imply a significant increase in process complexity. The straightforward approach adds the steps interfacial oxide growth, full-area deposition of poly-Si, print of an etch mask, poly-Si removal in the inter-finger regions, removal of the etch mask + cleaning and, for organic barriers, implies the necessity of waste treatment. Laser processing, e.g. laser oxidation, might avoid the latter aspect and even the necessity to deposit or to print an etch mask. The most attractive option would be to deposit the poly-Si already structured, e.g., by using shadow masks, laser transfer techniques or by utilizing local print of Si inks or liquid Si. All of these approaches are non-standard processes, which need development before becoming an option for industrial production. The rather small simulated efficiency gain of 0.3% questions the overall advantage of this cell structure.

d) The POLO BJ cell yields an efficiency potential that is 1.0% above the reference PERC+ process. The SEGA of this structure is shown in Fig. 4. The larger shading losses of 0.7% (Jsc: −0.7 mA/cm²) due to the 50 µm Al fingers are overcompensated by the gain in the open-circuit voltage (+39 mV) due to avoiding the recombination at the conventional P-diffusion and Ag contacts (0.45% and 0.35%, respectively). The disadvantage of larger shading will be reduced in the module, because a significant fraction of the light reflected from the finger will be redirected to the cell. Besides the shading loss, the resistance due to the large optimal finger spacing, and the synergies between the remaining recombination channels are the main bottlenecks of this structure. Regarding bifaciality, similar values as for TOPCon (i.e., 80%) are expected and we measured values in this range, which will be published separately.

In terms of technology, this cell can possibly be manufactured with a lean process flow since no structuring of the poly-Si is required and the hole contact is formed by Al alloying during firing (as for the PERC+ cell). The latter aspect also implies comparable metallization costs for the POLO BJ cell as for the PERC+ benchmark. A small wrap-around of nominal single-sided poly-Si deposition techniques might be less detrimental than for e.g. the TOPCon structure since the poly-Si will not be in contact with highly doped regions of the opposite polarity. If in-situ doped firing-only contacts are available and the bulk lifetime is already sufficiently high even without any gettering step, a process sequence without any high temperature step in a quartz tube furnace might be possible. Achieving or maintaining a sufficiently high bulk lifetime, which is essential for all cell concepts and in particular for all back-junction concepts, is one of the technological challenges. To estimate the impact of different final lifetimes we run a variation of the bulk lifetime for all cell concepts based on p-type Si, which we show below. Printing narrow Al contacts (here 50 µm width) with good electrical properties and aligning them with respect to the laser contact openings in industrial production is another challenge, and requires further development. It is interesting to note that Al fingers as narrow as 50 µm were recently reported. A further challenge is the reliable and, under industrial environment (automated handling, no cleanroom atmosphere), robust passivation of a lightly doped textured surface. Finally yet importantly, the inter-connection of the cells for module integration has to be developed. Besides geometrical constraints for possible soldering pads on the
front due to shading, there are constraints on the SiNx thickness in order to still fulfill antireflection purposes. Thus, metal spiking (i.e., by the Ag soldering pads) through the passivation layer cannot be mitigated by simply increasing the SiNx thickness (as on the rear of a PERC+ cell). The comparably high efficiency potential of the POLO BJ structure comes with a potentially lean process flow and thus makes managing the above mentioned technological challenges a worthwhile research target.

e) Conceptually, the POLO IBC has the same advantages as the BJ version discussed above, yielding a high gain in open-circuit voltage of +36 mV compared to PERC+. The SEGA for this structure is shown in Fig. 5. When comparing the SEGAs of the POLO BJ and IBC structure, we see that the loss due to shading (0.71%) is completely avoided in the IBC design. In contrast the loss due to recombination at the Al-p+ contacts is increased

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**Figure 4.** SEGA for the POLO BJ cell discussed in this study. Red bars show potential efficiency gains due to recombination, green bars gain due to optical losses and blue bars gain due to resistive losses. The gray bars show synergistic gains.

**Figure 5.** SEGA for the POLO IBC cell discussed in this study. Red bars show potential efficiency gains due to recombination, green bars gain due to optical losses and blue bars gain due to resistive losses. The gray bars show synergistic gains.
because of the reduced contact spacing (+ 0.2%). The reduced contact spacing, however, also yields a reduction of the loss due to the lateral resistance between the contacts (−0.24%). All effects combined yield an efficiency gain of 0.8% compared to the POLO BJ structure, which is 1.8% higher than for the reference PERC+ cell. The bifaciality of this cell structure can be expected to be slightly reduced from that of the abovementioned structures since both metal grids are on the rear. We expect a bifaciality factor of 70%.

In terms of technology, this cell structure also has the advantage of an inexpensive and elegant formation of the hole collecting contact by Al alloying. Unfortunately, it also combines challenges from the PERC+ POLO cell (necessity for structuring of the poly-Si) with challenges from the POLO BJ cell (requirement for high bulk lifetimes, passivation of lightly doped textured surfaces, avoidance of metal spiking through passivation layers, adapted cell interconnection scheme). Again, the increase of the SiNx thickness on the rear is not a perfect measure for the mitigation of metal spiking (e.g. from the soldering pads) since the Ag paste still has to fire through the stack in the emitter (n+ POLO) regions. Assuming that these technological challenges can be solved, the high efficiency potential of this structure can also be combined with a lean process flow.

f) and g) The POLO IBC concept is limited to 25.5% mainly by the screen-printed Al-p+ contacts. The logical next step is thus to also replace the Al-p+ contacts by p-type POLO junctions, promising an efficiency gain of 0.38%. We study both, f) the POLO² BJ and g) the POLO² IBC cell as improvements over the POLO BJ and IBC cells, respectively. Both cells are identical to the POLO BJ and IBC concepts except for the base contacts, which are replaced by p-type POLO junctions. The POLO² BJ structure shows an efficiency gain of 1.3% in comparison to PERC+ and 0.3% in comparison with the POLO BJ concept. In comparison to PERC+ this efficiency gain originates mainly from the large voltage gain of +45 mV and in comparison to POLO BJ from the reduced shading and recombination at the Al-p+ contacts. The IBC version of the POLO² concept yields the highest efficiency in our comparison with 25.9%, which is another 0.4% higher than the POLO IBC concept by avoiding the recombination at the Al-p+ contacts and 2.2% higher than PERC+. The SEGA for the POLO² IBC cell is shown in Fig. 6. We see that the main losses are due to recombination within the bulk and at the textured front surface. A further efficiency increase requires, thus, extensive research on Si material with reduced SRH defects and better surface passivation. All efficiency gains yield a final efficiency of 27.7%. The remaining margin to the efficiency limit of 29.6% is due to parasitic absorption, front surface reflection and imperfect light trapping, which do not show up in our analyses. Also the cell thickness and doping do not have the values required for the 29.6% limit. Bifaciality is expected to be lowest for POLO² IBC (60%), since the rear side of this structure features the metallization for both polarities as well as parasitically absorbing poly-Si on almost the entire area of the planar surface. Nevertheless, this cell structure will have the highest energy yield for any monofacial application such as rooftop, vehicle integrated PV and others.

In terms of technology, these POLO² structures add further challenges to those already mentioned for POLO IBC: the necessity to dope the poly-Si locally or at least single-sidedly, the necessity to contact p+-type poly-Si with high temperature screen-printing without alloying through it and, for POLO² IBC, the necessity to electrically separate the n+-type and p+-type poly-Si regions. For the first aspect, we proposed in Ref. the overcompensation concept, which can be applied for both POLO² BJ and POLO² IBC. Alternative approaches would be the single-sided or even structured deposition of doping sources (e.g. boron or phosphorus containing glasses, doping inks, masked ion implantation) on initially intrinsic poly-Si, the single-sided or even
structured deposition of in-situ doped poly-Si (structuring e.g. via shadow masks or inkjet printing of liquid silicon), or the extensive use of (structured) diffusion barriers. Many of the latter approaches are still in a very early stage of their development and have not proven their viability yet. For the second additional challenge (contacting of p+-type poly-Si via screen-printing), it will be essential to what extent the paste manufacturers can improve their products. We have reported our current status ($J_0 = 500$ fA/cm$^2$, $\rho_c = 5$ m$\Omega$cm$^2$) in Ref.11, and these results are consistent with those of other groups36. Since it could be a fundamental aspect that a certain amount of Al in the paste is required to form a low contact resistance to p+-type (poly-) Si72 while the Al always yields a certain degradation of the passivation quality, alternative metallization concepts like plating or Physical Vapor Deposition (PVD) might become more attractive for POLO$^2$ structures than they used to be for PERC+ cells. For the third additional challenge (separation of n+-type and p+-type poly-Si for POLO$^2$ IBC), different approaches can be used: oxidation of the poly-Si between the fingers73, introduction of a trench between the fingers74–77, or introduction of an initially intrinsic poly-Si region4,66–68. To summarize, both POLO$^2$ concepts are likely to require more time for industrial integration than the POLO BJ and IBC concepts. (h) The cell structures (c) through (g) discussed above use p-type Si wafers as does today’s mainstream PERC+ technology. Switching to n-type material brings a slight efficiency advantage due to the higher lifetime: The n-Si POLO$^2$ BJ concept shows an efficiency potential of 25.1%, 0.1% higher than in the p-Si concept, that is otherwise identical. The benefit of integrating n-type Si into cell concepts is small due to the slightly higher recombination at the p-type POLO junction (5 fA/cm$^2$ compared to 3 fA/cm$^2$ for n-type POLO).

In terms of technology, the challenges for n-Si POLO$^2$ BJ are—except for the fact that another passivation layer might be required for the lightly doped inter-finger regions on the front—comparable to those for the p-Si POLO$^2$ BJ cell.

Sensitivity of the efficiency to bulk lifetime variations

The bulk lifetime is an important parameter for finding the best cell type with POLO junctions for industrial production. In general, the bulk lifetime depends on the cell process due to gettering effects and defect kinetics. This is especially true for p-type Si wafers. We, therefore, vary the bulk lifetime of the various cell types to analyze the sensitivity of the efficiency to bulk lifetime variations. The cell geometries are those optimized for high lifetimes. The optimized front finger pitch, however, depends mainly on the trade-off between resistance contributions and shading. Therefore, the variation of the optimum finger pitch with varying lifetime is small. Figure 7 shows the efficiencies of all cell types as a function of a fixed, i.e. injection-independent, SRH bulk lifetime. The lifetime has only a minor effect on the efficiencies of the PERC+ and PERC+ POLO concepts of less than 0.5% for lifetimes between 500 $\mu$s and 10,000 $\mu$s. The benefit of integrating POLO into BJ or IBC p-type Si cells, however, depends strongly on the bulk lifetime. For a low lifetime of 500 $\mu$s the benefit of the POLO BJ and IBC are only 0.22% and 1.09%, respectively; and for the POLO$^2$ BJ and IBC concepts 0.48% and 1.26%, respectively. Compared to the expected efficiency gain at $\tau_b = 10,000$ $\mu$s, this is ca. 1.2% lower for the POLO concepts and 1.4% for the POLO$^2$ concepts and probably hardly justifies the increased process complexity. Thus, the economic feasibility relies on good quality Si and efficient gettering. In this context, it is interesting, that POLO layers also serve as gettering layers in high temperature steps78.

Roadmap for future cell development

The results presented above can serve as a roadmap for a step by step improvement of crystalline Si solar cells using POLO junctions. Figure 8 shows this roadmap. The POLO BJ and IBC concept show a high efficiency gain of 1.0% and 1.8%, respectively, provided the high lifetimes measured on test structures can be transferred to the finished cell. Both concepts have potentially lean process flows with no and with one structuring process for the
poly-Si, respectively. Potential process flows of both cells contain many process steps similar to that of the PERC+ process, which certainly eases the industrial integration. Therefore, starting from the PERC+ process, working towards the two POLO cells seems promising in terms of a fast integration of POLO junctions into industrial cells. Also there are large synergies between experimental developments of the POLO IBC and the POLO BJ cell. We actually started experimental work in this direction. For POLO BJ, we achieved an independently confirmed efficiency of 22.3% within a (on our scale) very short development time of 1/2 year⁷⁹,⁸⁰. For POLO IBC, we achieved so far an open-circuit voltage of 711 mV and an efficiency of 22.6% on small area⁸¹,⁸². This $V_{OC}$ is still below the simulated value (due to a non-optimized Al-p+ contact), but already the highest experimental $V_{OC}$ reported so far for p-type solar cells with Al-p+ contacts. We think that these first experimental achievements are promising and show that our simulation-based roadmap is reasonable. A promising next development target are industrial-type POLO² cells based on p-type Si both as IBC and BJ structure. We already fabricated a POLO² IBC cell with 26.1% efficiency⁷⁷ using lab-type processes and evaporated contacts. The POLO² concepts are ultimately limited by their optical performance and material quality and of course the intrinsic transmission and thermalization losses. To address the latter loss channel, a further valuable research goal is the development of perovskite tandem cells with the Si POLO² concepts as bottom cells for 2- and 3-terminal devices. The analysis of these tandem cells is not part of this study but we estimate an efficiency of above 33% by adding an approximate 21% efficiency for the top and half of the efficiency simulated here for the bottom cell. The roadmap shown in Fig. 8 is one possibility for approaching highest efficiencies. It is, however, also possible to directly develop POLO² structures or implement tandem cells in earlier development stages like the POLO concepts or even at the current PERC+ state.

**Conclusion**

In the simulation study presented here we analyzed various cell structures for their suitability for industrial integration. We simulated eight cell types: a PERC+ reference, a TOPCon reference, three cell concepts with n-type POLO and Al base contacts, and three cell concepts with POLO junctions for both polarities, two of which are based on p- and one based on n-type Si. The input parameters are chosen to represent measured best-case parameters achieved with processes relevant to industrial production. We simulated an efficiency gain compared to the reference PERC+ cell of 1.0% and 1.8% for the POLO BJ and IBC concepts, respectively, which feature n-type POLO emitters and Al base contacts. We further simulated an efficiency gain of 1.3% and 2.2% for the POLO² BJ and IBC concepts, respectively. We also analyzed the concepts based on p-type Si for their sensitivity to the bulk lifetime showing a 1.2% efficiency gain difference for the POLO concepts and 1.4% for the POLO² concepts for SRH bulk lifetimes ranging from 500 to 10,000 µs. For the integration of POLO junctions into industrial production in the near future the POLO BJ and IBC concepts appear attractive. Both utilize the base contacts and passivation surfaces from PERC+ and can, therefore, build on process steps from existing production lines. The next development step is to replace the Al base contacts with p-type POLO junctions also in either

**Figure 8.** Roadmap for further cell development: Starting from our PERC+ cells we focus on the POLO IBC structure for industrial Si cells in the near future. We also exploit the large synergy between the POLO IBC and BJ concepts to also realize the BJ concept as it has the benefit of being compatible with conventional cell interconnection. As a further development step we work on the POLO² concepts, which require development in terms of POLO structuring and screen-printing on p-type poly-Si. The development step after the POLO² concepts is to combine the POLO² as bottom cells to perovskite tandem cells.
BJ or IBC configuration. Even though the industrial integration of these concepts may be a few years down the road, the building blocks for their realization are currently under development.\textsuperscript{12,28} Perovskite-Si tandem cells are still in a research state where their economic feasibility is not yet proven. Nevertheless, perovskite-Si tandem cells using the Si POLO\textsuperscript{2} as bottom cells are an option for further improvement after the power losses in Si cells approach the intrinsic limit.

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fraction of the ‘results and discussion’ part of the paper. R.B. provided useful information regarding all simulations and background information regarding the POLO BJ concept. T.D. was mainly responsible for the definition of the nomenclature and overall presentation of the proposed cell concepts. B.M. provided input parameters and useful information on possible potential processing challenges. B.L. was responsible for the work and writing the paper. S.S. performed all optical simulations. F.H. conceived the POLO IBC structure and provided input parameters for POLO junctions. V.M. performed the initial evaluation of the process complexity to be expected when manufacturing the hypothetical cell concepts. H.S.-H. conceived the predecessor of the POLO 2 BJ concept and provided useful information on possible potential processing challenges. B.L. was responsible for the nomenclature and overall presentation of the proposed cell concepts. B.M. provided input parameters and useful background information regarding the POLO BJ concept. T.D. was mainly responsible for the definition of the PERC+ reference case and the PERC+ POLO structure as well as the evaluation of suitable alignment tolerances for screen-printing. R.P. conceived the initial idea for this work and was mainly responsible for the discussion of the challenges to be expected with industrial processing of the hypothetical concepts. He also wrote a large portion of the ‘results and discussion’ part of the paper. R.B. provided useful information regarding all simulations and the overall concept of the work. He was further responsible for the creation of the roadmap (Fig. 8) based on discussions with the other co-authors. All authors reviewed the manuscript.

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Author contributions
C.N.K. performed all electrical simulations and was responsible for the overall concept of the work and writing the paper. S.S. performed all optical simulations. F.H. conceived the POLO IBC structure and provided input parameters for POLO junctions. V.M. performed the initial evaluation of the process complexity to be expected when manufacturing the hypothetical cell concepts. H.S.-H. conceived the predecessor of the POLO^2 BJ concept and provided useful information on possible potential processing challenges. B.L. was responsible for the nomenclature and overall presentation of the proposed cell concepts. B.M. provided input parameters and useful background information regarding the POLO BJ concept. T.D. was mainly responsible for the definition of the PERC+ reference case and the PERC+ POLO structure as well as the evaluation of suitable alignment tolerances for screen-printing. R.P. conceived the initial idea for this work and was mainly responsible for the discussion of the challenges to be expected with industrial processing of the hypothetical concepts. He also wrote a large portion of the ‘results and discussion’ part of the paper. R.B. provided useful information regarding all simulations and the overall concept of the work. He was further responsible for the creation of the roadmap (Fig. 8) based on discussions with the other co-authors. All authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Correspondence and requests for materials should be addressed to R.B.

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