A novel silicon heterojunction IBC process flow using partial etching of doped a-Si:H to switch from hole contact to electron contact in situ with efficiencies close to 23%

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Abstract
We present a novel process sequence to simplify the rear-side patterning of the silicon heterojunction interdigitated back contact (HJ IBC) cells. In this approach, interdigitated strips of a-Si:H (i/p+) hole contact and a-Si:H (i/n+) electron contact are achieved by partially etching a blanket a-Si:H (i/p+) stack through an SiOx hard mask to remove only the p+ a-Si:H layer and replace it with an n+ a-Si:H layer, thereby switching from a hole contact to an electron contact in situ, without having to remove the entire passivation. This eliminates the ex situ wet clean after dry etching and also prevents re-exposure of the crystalline silicon surface during rear-side processing. Using a well-controlled process, high-quality passivation is maintained throughout the rear-side process sequence leading to high open-circuit voltages (VOC). A slightly higher contact resistance at the electron contact leads to a slightly higher fill factor (FF) loss due to series resistance for cells from the partial etch route, but the FF loss due to J02-type recombination is lower, compared with reference cells. As a result, the best cell from the partial etch route has an efficiency of 22.9% and a VOC of 729 mV, nearly identical to the best reference cell, demonstrating that the developed partial etch process can be successfully implemented to achieve cell performance comparable with reference, but with a simpler, cheaper, and faster process sequence.

KEYWORDS
amorphous silicon, dry etch, H2 plasma, heterojunction, in situ processing, interdigitated back contact (IBC), NF3/Ar plasma, process simplification

Symbols and abbreviations: η, efficiency; ρc, contact resistivity; ρc,n, contact resistivity at the electron contact; ΔRs, increase in series resistance; ΔFFRs, fill factor loss due to series resistance; FF, fill factor; J02, recombination current density; Jmp, current density at maximum power point; Jsc, short-circuit current density; Rs, series resistance; Voc, open-circuit voltage; Ar, Argon gas; a-Si:H, hydrogenated amorphous silicon; a-Si:H (i/n+), stack of intrinsic and n-doped a-Si:H; a-Si:H (i/p+), stack of intrinsic and p-doped a-Si:H; AR, aspect ratio; Cu, Copper; Cz, Czochralski; DIW, deionised water; DLTS, deep-level transient spectroscopy; ARDE, aspect ratio-dependent etching; EVA, ethylene-vinyl acetate; F*, fluorine radicals; FZ, float zone; H2, hydrogen gas; HF, hydrofluoric acid; HJ, heterjunction; HM, hard mask; IBC, interdigitated back contact; ITO, indium tin oxide; IV, Current–voltage; KOH, potassium hydroxide; N2, nitrogen; NF3, nitrogen trifluoride; O3, ozone; PECVD, plasma-enhanced chemical vapour deposition; PL, photoluminescence; QSSPC, quasi steady-state photoconductance; SiNx, silicon nitride; SiOx, silicon oxide; SOM, sulphuric acid-ozone mixture

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The silicon heterojunction interdigitated back contact (HJ IBC) solar cell technology has enabled the achievement of solar cell power conversion efficiencies above 25%\(^1,2\), with the world record efficiency standing at 26.7% for single-junction crystalline silicon solar cells.\(^3,4\) Hydrogenated amorphous silicon (a-Si:H) used as passivating contacts in this technology is key to reaching high open-circuit voltages (V\(_\text{OC}\)) up to 750 mV\(^5,6\) by suppressing surface recombination current, while the IBC architecture ensures high short-circuit current densities (J\(_\text{SC}\)) above 42 mA/cm\(^2\)\(^7\) by eliminating optical shading and parasitic absorption losses at the front side.

Our baseline HJ IBC process flow is described in Figure 1. The main challenge for this cell technology is the reduction of the process complexity on the rear-side, which requires the realisation of both the interdigitated pattern of a-Si:H strips of opposite polarities as well as interdigitated metal fingers contacting these strips. Lab-scale approaches typically involve photolithography\(^2,7\) for patterning, 2 to 3 alignment steps, several cumbersome wet processes, and a large number of steps, which reduce the throughput and processing reliability and increases the cost of solar cell fabrication. Therefore, several groups investigating this cell architecture focus on not only on performance improvement but also simplification of the rear-side process sequence. For industrial viability, the target is to attain a lithography-free process sequence, consisting of a minimum number of steps, preferably without using many dangerous chemicals leading to lower processing costs, greater process reliability, and higher throughput.

While well-known techniques such as screen-printing and inkjet printing can be used for metal finger patterning,\(^8\) it is the patterning of the a-Si:H strips that is the most challenging. The simplest approach that has been investigated is the use of mechanical shadow masks to deposit, in an additive fashion, successively the two juxtaposed finger patterns of opposite polarity in just 2 steps.\(^8,11-13\) As a further simplification, a novel “tunnel-IBC” structure, requiring just a single alignment step, has been implemented with great success.\(^7\) In addition, the entire patterning sequence is completed in situ and without wet chemical treatments during the patterning. While this is very advantageous, the industrial compatibility of using reusable shadow masks is debatable. Moreover, depositions through masks lead to tapered profiles of deposited films and require proximal contact of wafers with the masks.\(^14\)

Another popular “litho-free” alternative to patterning, which is contact-less and also drastically reduces the number of wet chemical steps, is laser ablation-assisted patterning.\(^15-20\) In this approach, the process is often directly structured onto a sacrificial mask layer on top of the a-Si:H stack to be patterned. While there is a risk of laser-induced thermal damage to the crystalline silicon and the heterocontact in the laser-ablated areas,\(^19,20\) significant strides have been made recently in tackling this issue.\(^17\) In this approach, etching of the underlying a-Si:H stack is needed, and hence, it is a subtractive route, just like in photolithography. Often, wet etching is used, which is not well-controllable and can lead to processing issues.\(^18,21,22\) Dry etching can considerably improve process reliability.\(^21\)

One of the strategies to simplify our baseline process sequence (depicted in Figure 1, left) into a simpler one that is litho-free, almost all-dry, and inexpensive is shown in Figure 1 (middle). We have shown in our recent work the successful replacement of lithography by laser ablation\(^2\), and wet etching by dry etching.\(^21\) In this paper, the development of the partial dry etch route, which eliminates the ex situ wet clean step, will be discussed. Such a sequence can be completed fully in situ, requires no use or disposal of dangerous wet chemicals such as hydrofluoric acid (HF), has higher throughput and is also more reliable for the processing of glass-bonded thin silicon (depicted in Figure 1, bottom right).

A part of the HJ IBC rear-side process sequence is depicted schematically in Figure 2. In the baseline process flow, after hard mask patterning, the areas with exposed a-Si:H (i/p\(^+\)) are completely etched to reveal the crystalline Si surface using NF\(_3/\)Ar plasma-based dry etching in the same PECVD tool used to deposit a-Si:H.\(^21\) After dry etching, the surfaces are cleaned ex situ using an HF dip to remove residual contaminants on the surface before being inserted back into the PECVD tool for the deposition of a-Si:H (i/n\(^-\)), thereby producing interdigitated strips of a-Si:H (i/p\(^+\)) and a-Si:H (i/n\(^-\)), following a subsequent lift-off step.
When processing glass-bonded HJ IBC devices, achieving and maintaining excellent rear surface passivation in the presence of bonding agents (silicones or ethylene-vinyl acetate [EVA]) is a challenging integration problem. This is because bonding agents contain volatiles that can outgas during the passivation process resulting in poor passivation. To address this, an extended outgassing step and in the case of silicones, an Ar plasma treatment step are applied to ensure reproducible high-quality passivation, as shown in Figure 3B. These processes, however, are not permanent fixes for the outgassing issue. Thus, any re-exposure of the crystalline silicon surface during subsequent processes (e.g., after dry etch) can lead to problems with the repassivation of the re-exposed areas, because of the same organic contamination issue, as depicted in Figure 3A. Figure 3C shows the PL image after repassivation of dry-etched areas, having an outgassing pattern from the edges, which is indicative of organic contamination from the exposed silicone at the wafer periphery.

The partial etching route (Figure 2, bottom) can solve these issues, whereby dry etching is carefully controlled to remove only the doped layer (p⁺ a-Si:H), leaving behind the intrinsic a-Si:H layer from the first passivation and not re-exposing the crystalline silicon surface. In the
2 | EXPERIMENTAL METHODS

2.1 | Samples for etch rate evaluation

Mirror-polished, 200 mm diameter, p-type, (100) Czochralski (Cz) Si wafers were cleaned in a bath containing sulphuric acid-ozone mixture (SOM) at 50°C for 10 minutes to remove organic contamination from the wafer surface. Subsequently, a short hydrofluoric acid (HF) dip is done to remove the chemical oxide from the previous step to produce a hydrophobic surface. Various stacks of hydrogenated amorphous silicon (a-Si:H), both intrinsic (i) and doped (p+ or n+), are deposited on the surface of these wafers at 175°C using plasma enhanced chemical vapour deposition (PECVD). The thickness of the a-Si:H film is varied using the deposition time. Nominal thicknesses of the p+ a-Si:H and n+ a-Si:H layers used for etch rate evaluation are approximately 21 nm and 27 nm, respectively, which correspond to the actual thicknesses used during solar cell fabrication. Spectroscopic ellipsometry is used to investigate the dielectric properties of the a-Si:H films in the wavelength range of 250 to 800 nm. By fitting the resulting spectra using the Tauc-Lorentz model, the a-Si:H film thickness and optical band gap can be determined.

In this work, plasma dry etching of a-Si:H films is performed in the same PECVD tool using NF3 and Ar as precursor gases at the same temperature of 175°C. A power density of approximately 110 mW/cm² and a pressure of 1.2 mbar were used. The NF3 and Ar gas flows are varied in the range of 10 to 200 sccm and 10 to 500 sccm, respectively, in order to vary the etch rate. Different etch times between 10 to 120 seconds were used to etch n- and p-doped a-Si:H films for each gas flow ratio. The remnant a-Si:H film thickness is evaluated using spectroscopic ellipsometry, as above. Using a linear fit of the remnant a-Si:H thickness versus time, the etch rate can be determined under different gas flow conditions.

2.2 | Samples for surface passivation quality evaluation

N-type, 200 μm thick, 150 × 156 mm² semi-square, 3 to 5 Ω.cm, (100) Cz wafers were first etched in a hot potassium hydroxide (KOH) bath to remove the saw damage from the wafering process. Subsequently, the wafers were cleaned in a bath containing ozone and deionised water (DIW:O₂) for 5 minutes, followed by an HF dip, to produce hydrophobic surfaces. The non-investigated side of the wafers was passivated with a stack of a-Si:H (i/n+) using PECVD at 175°C that results in excellent surface passivation. The investigated surface of these wafers was passivated using a stack of either a-Si:H (i/n+) or a-Si (i/p+). Partial etching was investigated on both polarities of doping. Passivation quality was evaluated using photoluminescence (PL) and injection level-dependent minority carrier lifetime measurements by means of quasi-steady state photoconductance (QSSPC).

2.3 | Samples for contact resistivity measurements

Selected samples used for passivation tests above were further used to evaluate contact resistivity. The rear-side a-Si:H (i/n+) passivation was removed using standard NF3/Ar plasma-based dry etching in 3 minutes at 175°C. The wafers were then diced into 3 × 3 cm² pieces and cleaned using an HF dip. Subsequently, a 200 nm thick aluminium layer was deposited on the side without a-Si:H using e-beam evaporation, to produce a low-resistance ohmic rear-contact. On the side with a-Si:H, circular pads of a stack of indium tin oxide (ITO) and copper (Cu) were deposited using sputtering and e-beam evaporation, respectively, through a shadow mask. The diameter of these circular pads was varied between 200 and 2000 μm in order to vary the contact resistance as a function of the contact area. Current–voltage measurements were performed using a Keithley K4200 system to determine the total resistance through the structures. The contact resistivity values, ρc, were then determined from the slope of the fits of the difference in total resistance and spreading resistance versus inverse of the contact area, as explained in detail by Cho et al.

2.4 | Solar cell fabrication

N-type, 200 μm thick, 150 mm diameter, round, 3 Ω.cm, (100) float zone (FZ) Si wafers were used for device fabrication. Our baseline process flow is outlined in Figure 1. The partial etch sequence developed in this work is integrated into this baseline process flow. For comparison, reference wafers following the conventional process sequence were also co-processed in the same run. The details of the different solar cell fabrication processes used in this work have already been described in detail in our previous publications. Here, only the salient features and key differences between the two splits are highlighted.

Although it is possible to pattern either the a-Si:H (i/p+) hole contact or the a-Si:H (i/n+) electron contact first, we chose to deposit and pattern the hole contact first. A hard mask consisting mainly of PECVD silicon oxide (SiOₓ) is deposited on the a-Si:H (i/p+) stack. Photolithography, dry etching, and lift-off are used to produce interdigitated strips of a-Si:H of opposite polarities on the rear side. For the conventional route, the a-Si:H (i/p+) hole contact areas exposed by the hard mask is etched off completely using NF3/Ar plasma dry etching with an NF3/Ar flow ratio of 200:100 sccm. A slight over-etch of about 100 to 200 nm of the crystalline Si surface is performed. Subsequently, an ex situ wet clean using an HF dip is performed for 1 minute to remove the residual surface contamination from the dry etch process. The repassivation of the etched surfaces is then done by depositing a stack of a-Si:H (i/n+). In contrast, for the partial etch route, mainly the p+ a-Si:H layer, plus an additional 1 to 2 nm of intrinsic a-Si:H are removed, keeping the bulk of original intrinsic a-Si:H in
place, by using a dilute NF<sub>3</sub>:Ar gas flow ratio of 25:500 sccm, which ensures a controllable and low etch rate. Subsequently, the etched a-Si:H film surface is cleaned in situ, using a short H<sub>2</sub> plasma etch with a power density of approximately 90 mW/cm<sup>2</sup> for about 30 seconds. Finally, an n" a-Si:H layer is deposited, again in situ, to form the a-Si:H (i/n") electron contact. All other process steps are identical for the two copy (DLTS).21,31,32 Thus, NF<sub>3</sub>/Ar plasma study.21,30

Two wafers were processed for each split. A total of 16 cells (2.5 x 2.5 cm<sup>2</sup>) were produced from each wafer by blade dicing. The cells were annealed at 150°C for 30 minutes in N<sub>2</sub> ambient to cure the damage resulting from the ITO sputtering process. The illuminated current–voltage (IV) characteristics of the resulting cells were measured under AM1.5G spectrum at 25°C through an aperture area of 3.97 cm<sup>2</sup>.

3 DEVELOPMENT OF PARTIAL DRY ETCHING OF A-Si:H

For a reliable and successful partial etch process, the following criteria must be achieved:

1. The etched thickness must be well-controllable (ie, low etch rate of < 0.5 nm/s and good spatial uniformity).
2. High surface passivation quality must be attained (ie, no plasma damage or contamination).
3. Contact resistivity values must be comparable with that of reference samples (ie, the quality of the contact layers should not be adversely affected by the partial etch process)

3.1 Selection of partial etch conditions

The standard dry etch process for removing a-Si:H in the baseline flow (Figure 1) is based on NF<sub>3</sub>/Ar plasma with a flow ratio of NF<sub>3</sub>: Ar = 200:100 sccm.23 This results in an etch rate of approximately 1.5 nm/s, which is too high to achieve a well-controllable partial etch process. On the other hand, with H<sub>2</sub> plasma etching, a well-controllable etch rate of approximately 1 nm/min is attainable. However, the low etch rate and the consequently prolonged etch durations in the order of several minutes would result in severe plasma damage to the crystalline silicon bulk even at low power, as revealed by minority carrier lifetime measurements and deep-level transient spectroscopy (DLTS).21,31,32 Thus, NF<sub>3</sub>/Ar plasma-based dry etching is the preferred choice for the partial etch development.

During NF<sub>3</sub>/Ar plasma dry etch, F* radicals are formed in the plasma, which react with Si atoms on the wafer surface. Ar* ions assist in providing energy for these reactions via physical bombardment. In this way, Si atoms are removed as SiF<sub>4</sub> gas during the etch process.33 Therefore, the etch rate can be reduced by increasing the NF<sub>3</sub> gas dilution in the chamber, by reducing the NF<sub>3</sub> gas flow (less reactant atoms) and by increasing the Ar gas flow (more diluent atoms). An exhaustive study of the influence of all other process parameters (eg, pressure and temperature) were not performed. A minimum power density of approximately 110 mW/cm<sup>2</sup> needed to sustain the plasma in the chamber was used.

The etch rates under different gas flow ratios for the etching of both n-doped a-Si:H and p-doped a-Si:H were determined with the help of thickness measurements using spectroscopic ellipsometry and fitting using the Tauc-Lorentz model. In search of a low and well-controllable etch rate, the NF<sub>3</sub> and Ar gas flows were individually varied. The influence of the different gas flow ratios on the etch rate of n-doped a-Si:H is shown in Figure 4. As expected, the etch rate decreases strongly with decreasing NF<sub>3</sub> flow rate (Figure 4A), but only slightly with increasing Ar flow rate (Figure 4B). Moreover, the error in the etch rate determination also becomes lower for conditions with low etch rates of < 0.5 nm/s. A similar exercise was performed for the etching of p-doped a-Si:H. Finally, for the partial etching of n-doped a-Si:H, a flow ratio of NF<sub>3</sub>:Ar = 10:100 sccm giving an etch rate of approximately 0.14 nm/s was chosen. For the etching of p-doped a-Si:H, a flow ratio of NF<sub>3</sub>:Ar = 25:500 sccm, which resulted in a well-controllable etch rate of approximately 0.32 nm/s, was chosen.

Since the above etch rates were determined on blanket samples, the etch rate uniformity was evaluated using a patterned sample having relevant features of different widths, namely, the BSF bus bar (approximately 1.9 mm) and the BSF finger (approximately 360 μm), as depicted in Figure 5 (left). This sample had the same cross-section as depicted in the left schematic of Figure 2, whereby an a-Si:H (i/p") stack, with a total measured thickness of approximately 29.3 nm is exposed by the hard mask in the electron contact areas. The developed partial etch process, with a gas flow ratio of NF<sub>3</sub>: Ar = 25:500 sccm, was carried out on such patterned samples to completely etch off the exposed p" a-Si:H layer, with an estimated thickness of approximately 21 nm, plus 1.5 nm of the underlying intrinsic a-Si:H layer to ensure the complete removal of the p" a-Si:H layer. With an etch rate of 0.32 nm/s (determined using blanket samples), the expected a-Si:H thickness at the end of the partial etch
process of 70 seconds in duration on a blanket sample is approximately 6.9 nm, which is indicated by the dashed line in Figure 5 (right). The remnant a-Si:H thicknesses at different locations in the finger and bus bar areas after the partial etch process were again estimated using spectroscopic ellipsometry, and plotted in Figure 5 (right). The median thickness of the remnant a-Si:H in the wide bus bar area was determined to be 7.4 nm, while that in the narrow finger areas was 8.5 nm. This indicates a slightly lower etch rate in narrower features, down to 0.29 to 0.30 nm/s in some finger areas.

Spatial non-uniformity during plasma dry etching has been attributed to phenomena such as microloading and aspect ratio-dependent etching (ARDE). Microloading refers to the reduction of the etch rate observed in areas of high pattern density, for the same feature size, while ARDE refers to the reduction of etch rate with increase in the aspect ratio (AR), which is defined as the ratio of height over width of a feature. While the observations in our case are similar to ARDE, this phenomenon is typically observed for feature sizes below 1 µm and for AR > 1. In our case, the feature sizes are in the order of hundreds of microns and the AR (= hard mark thickness/feature width) is well below 0.004, even for the narrow finger regions. Thus, the mechanisms leading to ARDE cannot explain the observed differences.

A lower etch rate could be the result of either a reduced influx of reactant species to the etched surface, a reduced reaction rate at the surface, or a slower desorption rate of reaction products from the surface. It is known that NF₃-based plasma produces F⁻ radicals, which can etch SiOₓ to produce volatile SiF₄ and O₂ products. About 80% of the finger areas is protected by the SiOₓ hard mask. Therefore, we suggest that it could be the local presence of O₂ in these areas that could be hampering the etching reaction at the a-Si:H surfaces in the open finger areas. The suppression of silicon etching because of the presence of O₂ by forming an oxide-like layer on the reaction surface has been reported previously. Nevertheless, further investigation is needed to unravel the mechanisms leading to the observed etch rate differences. Importantly, it should be noted that the etch rate differences are not drastic and the spatial non-uniformity during the etching of approximately 22 nm of a-Si:H is only approximately 1 nm, ie, < 5% variation. Such differences across a device are not expected to have a strong impact on device performance. Obviously, the absolute thickness variation would increase with increase in etch time if thicker p⁺ a-Si:H layers would be used, but typical p⁺ a-Si:H layers used for HJ IBC devices are below 20 nm. This reduced etch rate in patterned areas was taken into account during device fabrication by increasing the etch duration accordingly.

3.2 Evaluation of surface passivation in the partial etch route

As mentioned before, an important criterium for the rear-side patterning process is that the rear-side passivation quality should not be adversely affected such that high-quality passivation is maintained throughout the patterning sequence. There are several risks associated with partially etching only the doped layer of a stack of intrinsic and doped a-Si:H. Firstly, the statistical process control of the etch process should be tight enough that variations in the thickness of the remnant a-Si:H at the end of the process do not degrade device performance. Therefore, a buffer of a few nm of a-Si:H to account for over-etching and slower etching is necessary. Secondly, there could potentially be plasma damage to the underlying crystalline Si substrate and the remnant a-Si:H left at the end of the process. Thirdly, there is the risk that the intrinsic a-Si:H layer could be contaminated with the dry etch species from the partial etch process that could adversely modify the structure, and therefore, the transport and passivation properties of the intrinsic a-Si:H layer.

These risks were evaluated using minority carrier lifetime measurements and PL imaging. A set of wafers with a stack of a-Si:H (i/p⁺) on one side and a-Si:H (i/n⁺) on the other side were prepared, as depicted in Figure 6. These wafers were etched on the side with a-Si:H (i/p⁺) for different times targeting remnant a-Si:H film thicknesses of 2.7 nm, 5.0 nm, and 7.6 nm. From our past experience in optimising the intrinsic a-Si:H layer for high-quality surface passivation, we know that the minimum thickness needed for achieving good passivation is
approximately 5 nm. The typical intrinsic a-Si:H layer thickness used in the baseline process for the rear-side a-Si:H stacks is approximately 8.5 nm. This gives us a thickness window of approximately 3.5 nm. A thickness variation of intrinsic a-Si:H in this range is not expected to have a significant impact on the fill factor (FF).39,40 The minority carrier lifetime at an injection level of 10^{15} cm^{-2} is given inside the PL images, for all cases, except for the ones with lifetime below 100 μs, where the highest lifetime reached in low-injection is mentioned [Colour figure can be viewed at wileyonlinelibrary.com]

![FIGURE 6 Schematics on the left depict the three stages of the process sequence: A, after passivation, B, after partial etch of the p-doped a-Si:H layer on the front, and C, after deposition of the n-doped a-Si:H layer. The PL images corresponding to these stages are given on the right, where the partial etch time is varied from 70 seconds to 85 seconds, resulting in the estimated remnant a-Si:H thickness ranging from 7.6 nm to 2.7 nm. The minority carrier lifetime at an injection level of 10^{15} cm^{-2} is given inside the PL images, for all cases, except for the ones with lifetime below 100 μs, where the highest lifetime reached in low-injection is mentioned [Colour figure can be viewed at wileyonlinelibrary.com]](Image 113x489 to 482x736)

As expected, when the thickness of the remnant a-Si:H after the partial etch process is only 2.7 nm, the surface passivation is completely lost and is not recoverable, following the n^- a-Si:H deposition. With a targeted thickness of 5 nm of remnant a-Si:H, the partial etch process is at the border of the process window, whereby, any over-etching during the partial etch process could quickly degrade the passivation quality. In this case, while the minority carrier lifetime drops from 13 ms to 2.5 ms after the partial etch process, the lifetime value indicates that good quality surface passivation still exists at the end of the process. Upon deposition of the n^- a-Si:H, the lifetime improves significantly to approximately 8.0 ms, which is certainly high enough for high efficiency devices. However, from the non-uniformity of PL images of this wafer after partial etch, it can be discerned that the surface passivation in an area on the top left of the wafer is poor.

This could be because the spatial variation of the partial etch process could have resulted in a slightly thinner a-Si:H in this area. Even upon redeposition of the n^- a-Si:H, the surface passivation in this area remains poor. Thus, targeting 5 nm remnant a-Si:H after partial etch is too close to the border of the process window.

On the other hand, the wafer with an estimated remnant a-Si:H layer thickness of 7.6 nm shows high-quality passivation throughout the partial etch process. The drop in lifetime from 13 ms to 4.3 ms is attributed to the removal of the p^- a-Si:H layer. The lifetime recovers to nearly the same high value after n^- a-Si:H deposition, reaching 14 ms. This lies close to the other end of the process window, whereby a slightly slower etching could leave some unetched p^- a-Si:H layer at the end of the partial etch process, before the deposition of the n^- a-Si:H layer, which would lead to nonfunctional devices.

Overall, the process window of the developed partial etch process for a 21 nm-thick p^- a-Si:H layer on blanket samples is between 65 seconds (just enough to remove the entire p^- a-Si layer) and 78 seconds (leaving behind just enough intrinsic a-Si:H layer to maintain good passivation), which corresponds to about 13 seconds (roughly ~20% of the total process time). We believe that this process window is large enough for a reliable partial etch process. For device processing, we target approximately the middle of the process window. As mentioned earlier, since we have a buffer of 3.5 nm as allowance for variation in the remnant intrinsic a-Si:H thickness, an etch rate variation between 0.30 and 0.34 nm/s on blanket samples can be tolerated when targeting the middle of this process window, which is well within what we have observed so far experimentally.

3.3 Unexpected blistering of doped a-Si:H deposited on partially-etched a-Si:H

A problem that arose during the partial etch development is the unexpected and spontaneous blistering and flaking of the doped a-Si:H deposited on top of the partially-etched a-Si:H. As shown in the optical microscopy image of Figure 7A, when a sample with a stack of a-Si:H (i/n^+) is partially etched to remove the n^- a-Si:H layer and subsequently has a p^- a-Si:H layer deposited on top, and is then immersed...
into a dilute HF bath, blisters and flakes are observed in the partially etched area but not in the area protected by the hard mask during the partial etch process. Note that the particles seen in the non-etched areas have been redeposited from the blistered areas and the a-Si:H stack in the nonetched areas actually remains intact. Spectroscopic ellipsometry analyses at the locations marked "A" and "B" in Figure 7A revealed that the thickness at location "A" corresponded to the remnant a-Si:H after partial etch, while that at location "B" corresponded to the total thickness of the remnant a-Si:H plus the p+ a-Si:H layer deposited on top. This spontaneous blistering happens also when the partial etch process is carried out on a stack of a-Si:H (i/p+) with subsequent deposition of an n+ a-Si:H layer. As indicated in Figure 7B, the fractions of blistering events after a partial etch process is approximately 80%, which is very high.

Although it is not fully clear why this blistering occurs, it has been shown in our previous work that NF3/Ar plasma etching process leads to a residual layer on the surface at the end of the process. This layer is dissolvable in an HF solution. It is hypothesized that this residual layer could be the root cause of the observed spontaneous blistering. To verify this, partially-etched samples were dipped in a HF bath for 1 minute to remove the dry etch residual layer before deposition of the second doped layer. This ex situ HF clean solves the issue, thereby no incidents of blistering were observed on such samples, as shown in Figure 7B. However, since one of the goals of the partial dry etch process is to perform the entire sequence in situ, without taking the wafers out of the PECVD tool, a dry clean method was introduced. A short H2 plasma treatment, which does not contain contaminating species, was used to etch off the residual layer at a moderate power density of approximately 90 mW/cm² for 30 seconds. As shown in Figure 7B, this in situ clean also solves the blistering issue, thus enabling a fully in situ partial etch process. To verify that the H2 plasma treatment inserted after the partial etch does not affect the lifetime, a passivated wafer was subjected to the full in situ partial etch process. As shown in Figure 7C, the short H2 plasma treatment does not degrade the lifetime of the wafer.

### 3.4 Evaluation of contact resistivity

Since the a-Si:H (i/doped) stack is actually part of the contact, any modification of the a-Si:H structure by the partial etch process may affect the charge carrier transport across the a-Si:H layers, even though the surface passivation might be excellent. Contact resistivity measurements, performed as described in Section 2.3 and by Cho et al. on the a-Si:H (i/n−) electron contact formed using the partial etch sequence as well as the conventional route are compared in Figure 8. The average contact resistivity of the electron contact, $\rho_{c,n}$ from the partial etch process is slightly higher at 118 mΩ.cm² compared with 85 mΩ.cm² for the reference electron contact. Since the electron contact comprises only approximately 17% of the device area, this slight increase in contact resistivity would lead to an increase in the series resistance ($\Delta R_s$) at device level of approximately 200 mΩ.cm². The additional fill factor loss because of this increased series resistance can be calculated according to the following.

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**FIGURE 7** A. Optical microscopy image showing the blistering of p-doped a-Si:H deposited on remnant a-Si:H after the partial etch of n-doped a-Si:H, when immersed in HF solution at the end of the process. B. Plot of the fraction of blistering incidents as a function of surface treatment after partial etching. C. Photoluminescence (PL) images before and after the fully in situ partial etch process sequence, including the short H2 plasma dry clean [Colour figure can be viewed at wileyonlinelibrary.com]
from the partial etch split are plotted together in Figure 9. The IV characteristics of the best cell from the reference split and that determined from light IV cell measurements are summarised in Table 1.

The developed in situ partial dry etch route was implemented in our HJ IBC process flow on 200 μm-thick, n-type (100) FZ Si wafers, as described in Figure 1. For comparison, wafers following the conventional process sequence, with a full dry etch and an ex situ wet clean, were co-processed as the reference split. The average and best cell parameters determined from light IV cell measurements are summarised in Table 1. The IV characteristics of the best cell from the reference split and that from the partial etch split are plotted together in Figure 9.

First and foremost, functional cells were produced for the first time using a completely in situ partial etching route. The PL image of a wafer, consisting of a total 16 cells, after the rear-side patterning of a-Si:H strips is shown in the inset of Figure 9. Uniform passivation has been achieved across the entire wafer. The lack of significant contrast in the PL intensity between the electron and hole contact areas within a single cell implies that similar quality passivation has been achieved in both areas, where the n-doped area had undergone the partial etch process. As shown in Table 1, the reference process resulted in excellent $V_{oc}$ values close to 730 mV, indicating high-quality surface passivation. Cells from the in situ partial etch split also showed high-quality passivation with $V_{oc}$ values comparable with reference. This corroborates well with the lifetime results described in Section 3.2 and the PL image in the inset of Figure 9.

In general, all cells suffer from moderate FF values around 75%. A FF loss analysis, performed according to,41 shows that the FF of our reference cells is typically limited by series resistance (approximately 5% absolute) and $J_{02}$-type recombination (approximately 4% absolute) losses. In addition to the FF values, the series resistance at device level obtained using the Bowden method42 and the fill factor loss due to series resistance calculated using Equation 1 are also given in Table 1. As expected from contact resistivity measurements in Section 3.4, the series resistance of cells from the partial etch split are higher, which should result in a lower FF for these cells. Indeed, when comparing the best cells from each split, the fill factor loss due to series resistance, $\Delta FF_{R_s}$, for the partial etch split is 7.0%, which is 1.2% higher than that for the reference split. Nevertheless, the actual difference in FF is only 0.3%. This is because the FF loss because of $J_{02}$-type recombination was observed to be lower for cells from the partial etch split compared with those from the reference split. The zone at the juncture of the electron and hole contact areas could be the origin of higher $J_{02}$-type recombination observed in the reference cells, where the fully-etched open areas need to be repassivated. One can imagine that this border region might be difficult to repassivate very well. In the partial etch split, however, the rear-side passivation is maintained throughout the rear-side processing sequence, since the intrinsic a-Si:H layer from the first passivation stack is never removed. This may consequently lead to lower FF losses due to $J_{02}$-type recombination. Similar observations and conclusions regarding FF losses can be made by comparing the average FF values of the two splits, as given in Table 1, except that the difference in FF is larger at approximately 1%. This seems to indicate that the partial etch process may have greater spatial variation at wafer-level, which is not captured by the small-area contact resistivity samples. Further investigation is needed to confirm the suggested origin of the observed differences.

$$\Delta FF_{R_s} = \frac{J_{mpp}^2 R_s}{V_{oc} J_{sc}}$$

(1)

where $\Delta FF_{R_s}$ is the fill factor loss due to series resistance, $J_{mpp}$ and $J_{sc}$ are the current densities at maximum power point and at short-circuit, respectively, while $V_{oc}$ is the open-circuit voltage. It is assumed that $J_{mpp}$ does not change significantly with small changes in $R_s$. Taking typical values of 41.5 mA/cm$^2$ for $J_{sc}$, 40 mA/cm$^2$ for $J_{mpp}$ and 725 mV for $V_{oc}$, the additional loss in FF due to the increase in series resistance, $\Delta R_s$, of approximately 200 mΩ/cm$^2$ would be approximately 1%, which is expected to result in a slight drop in efficiency of approximately 0.3% compared with our reference cells. We believe that improvements to the in situ hydrogen plasma treatment after the partial dry etch would help to reduce the contact resistivity, and will be the focus of future improvements to this process.

4 CELL INTEGRATION OF THE PARTIAL ETCH PROCESS

The developed in situ partial dry etch route was implemented in our HJ IBC process flow on 200 μm-thick, n-type (100) FZ Si wafers, as described in Figure 1. For comparison, wafers following the conventional process sequence, with a full dry etch and an ex situ wet clean, were co-processed as the reference split. The average and best cell parameters determined from light IV cell measurements are summarised in Table 1. The IV characteristics of the best cell from the reference split and that from the partial etch split are plotted together in Figure 9.

### Table 1: Cell results of two different splits of heterojunction interdigitated back contact (HJ IBC) cells

| Split                  | $J_{sc}$ [mA/cm$^2$] | $V_{oc}$ [mV] | FF [%] | $\eta$ [%] | $R_s$ [Ω.cm$^2$] | $\Delta FF_{R_s}$ [%] |
|-----------------------|----------------------|---------------|--------|------------|-----------------|-----------------------|
| Reference             |                      |               |        |            |                 |                       |
| Best                  | 41.6                 | 728           | 75.7   | 22.9       | 1.2             | 5.8                   |
| Avg. of 4             | 41.4                 | 726           | 75.1   | 22.6       | 1.1             | 5.2                   |
| Partial etch route    |                      |               |        |            |                 |                       |
| Best                  | 41.7                 | 729           | 75.4   | 22.9       | 1.4             | 7.0                   |
| Avg. of 4             | 41.6                 | 723           | 74.0   | 22.2       | 1.5             | 7.3                   |
Overall, the best efficiency achieved with the reference split is 22.9%. With the newly-developed in situ partial etch route, the same best efficiency of 22.9% is attained for the best cell, thus proving that a similar performance to reference can be achieved with a simpler process sequence. In future, cell performance could be further enhanced by optimising the partial etch process with improved in situ hydrogen plasma treatment and better spatial uniformity.

5 | CONCLUSIONS AND OUTLOOK

We have developed a novel in situ partial dry etch process sequence for HJ IBC solar cells, whereby the doped a-Si:H on the rear-side is switched by etching off the exposed p⁺-a-Si:H (or n⁺-a-Si:H) only and depositing an n⁺-a-Si:H (or p⁺-a-Si:H) on top of the remaining intrinsic a-Si:H from the first passivation, without re-exposing the crystalline silicon surface. This simplifies the rear-side patterning of HJ IBC cells by doing away with the need to perform the ex situ clean after dry etching, as in the baseline process flow. In addition, since this route prevents re-exposure of the crystalline silicon surface during rear-side patterning, this is advantageous for processing glass-bonded samples, where outgassing of organics from bonding agents can lead to poor passivation.

The partial etch process was based on NF₃/Ar plasma with high dilution of NF₃ in Ar of 10:100 sccm for n⁺-a-Si:H and 25:500 sccm for p⁺-a-Si:H, which leads to controllable etch rates of approximately 0.14 nm/s and 0.32 nm/s, respectively, on blanket samples. Lifetime measurements showed that high-quality surface passivation is achievable with a reasonably broad and controllable process window. An unexpected and spontaneous issue with the blistering of the deposited doped a-Si:H layer on top of a partially-etched a-Si:H surface was encountered. This was solved by introducing an additional in situ clean step based on H₂ plasma, which eliminated this blistering problem, without degrading the lifetime. Contact resistivity measurements showed values that are only slightly higher compared with reference, indicating that the charge carrier transport across the a-Si:H contacts is not drastically affected by the newly-developed process.

Finally, the partial etch route was implemented in the HJ IBC process flow, resulting in excellent $V_{oc}$ values close to 730 mV, which can be attributed to the high-quality passivation attainable with the partial etch process. While the FF loss due to series resistance appears to be higher for the partial etch split, FF loss due to $J_{02}$-type recombination is lower. This provides room for further improvement of the developed process.

Overall, a best efficiency of 22.9%, identical to the reference split, was achieved, proving that the partial etching route can attain similar cell performance as the reference route, while simplifying the rear-side process sequence. The process simplification implemented in this work eliminates the need to remove the wafers from the PECVD tool after dry etch and the need to use a cumbersome wet cleaning step in between dry etching and repassivation. Moreover, use of dangerous chemicals such as HF and the associated chemical waste disposal can be minimised. Such a process sequence is also beneficial for processing of glass-bonded samples (in module-level cell processing concepts), since crystalline silicon surface re-exposure is avoided, thus maintaining high-quality surface passivation throughout the rear-side process sequence. This approach not only increases the processing simplicity but also improves the throughput. The next step of this work will focus on further optimisation of the partial etch process to improve the contact resistivity and subsequently on the implementation of the partial etch sequence together with laser ablation patterning. The efficacy of this approach on fully and partially textured rear-side will also be evaluated.

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