INDUSTRIAL TOPCON SOLAR CELLS REALIZED BY A PECVD TUBE PROCESS

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ABSTRACT: These days low-pressure chemical vapor deposition (LPCVD) is commonly used by the photovoltaic industry to deposit Si layers for tunnel oxide passivated contact (TOPCon). This work summarizes the development of an alternative TOPCon deposition process using a tube plasma-enhanced chemical vapor deposition (PECVD) tool. In the first part, the main results of the German federal project “Upgrade Si-PV” are summarized. The goal of this project was the transfer of the TOPCon process to an industrial-proven tube PECVD system. In the second part, recent progress on the implementation of this process into a TOPCon cell is presented. It is demonstrated that the PECVD system is capable of producing screen-printing compatible TOPCon layers yielding recombination currents of \( \approx 3 \) fA/cm² and \( \approx 200 \) fA/cm² in the passivated and metallized region, respectively. The best TOPCon cell which was realized with the PECVD process yielded the following current-voltage parameters: \( V_{oc} = 700 \) mV, \( FF = 79.6\% \), \( J_{sc} = 41.2 \) mA/cm², and \( \eta = 22.95\% \).
Keywords: PECVD, passivation, TOPCon, solar cell, silicon

1 INTRODUCTION

Poly-Si-based passivating contacts (herein referred to as TOPCon [1]) have enabled laboratory solar cells to surpass the 26% efficiency threshold [2, 3]. To date, research institutes and manufacturers are working towards the implementation of TOPCon into a lean and industrial-viable process flow. Most work has been dedicated to realizing the so-called industrial TOPCon (i-TOPCon) solar cell featuring a boron-diffused emitter, a TOPCon rear contact, and screen-printed contacts on both sides. Although the efficiency has climbed from 20.7% [4] through the lower 23% range [5, 6] to a record efficiency of 24.58% [7], there are still a few hurdles to overcome to convince the majority of cell makers to migrate from PERC to i-TOPCon. Especially the rather complex process flow, which typically requires additional high-temperature processes (BBr3 diffusion for the emitter and POCl3 diffusion to dope the intrinsic poly-Si layer), wet-chemistry (single-side etching of poly-Si at the front) as well as the use of silver-containing pastes at front and back increase the all-in cell costs compared to a PERC cell [8]. One step into the direction of a lean process flow is the replacement of the LPCVD technology by a deposition technology exhibiting higher throughput and up-time, capability of in-situ doping, and preferably single-sided deposition.

This manuscript is divided into two parts. Firstly, the main results of the federal project “Upgrade Technologies for Silicon Photovoltaics (Upgrade Si-PV)” which received funding from the German Ministry of Economic Affairs and Energy research [9] is summarized. Secondly, recent progress on the i-TOPCon cell featuring PECVD TOPCon is presented.

2 SUMMARY OF PROJECT UPGRADE SI-PV

The project “Upgrade Si-PV” was dedicated to promote understanding of the regeneration process for PERC solar cells (please see [10]) as well as advancing the industrialization of the TOPCon deposition technology by using a tube PECVD system of the company centretherm.

2.1 Tunnel oxide

One work package was focusing on the investigation of alternative methods to form the tunnel oxide. To this end, the work on ozone-based (O3) oxides published in [11] was continued and, additionally, a thermal tunnel oxide (referred to as TO) process was developed.

![Figure 1: IVoc plotted over contact resistivity measured on n-type 5 Ωcm FZ-Si wafers passivated by n-TOPCon featuring different tunnel oxides. The open symbols refer to TOPCon with a slightly reduced dopant concentration. The annealing temperatures are placed adjacent.](Image)

Fig. 1 shows the implied open-circuit voltage \( IV_{oc} \) vs. contact resistivity \( \rho_1 \) measured by the transfer line method (TLM) of \( n \)-TOPCon featuring different tunnel oxides. Both, the O3 oxide and TO show a superior surface passivation quality and a higher thermal stability compared to the nitric acid (HNO3) oxide. The latter can be explained by the fact that both O3 oxide and TO block phosphorus atoms diffusing from the poly-Si into the c-Si more effectively and presumably are less susceptible to balling-up of the oxide layer. Such distinctive behavior of different tunnel oxides upon annealing has recently been associated with the oxide’s stoichiometry and thickness.
[12]. Since TOPCon featuring the TO suppressed recombination most effectively and enabled reasonably low contact resistivity values of $\approx 2 \text{m}\Omega\text{cm}^2$, this oxide became the standard process.

The interaction between tunnel oxide and dopant diffusion has been studied in more detail by using plasma grown and plasma nitrided tunnel oxides [13]. The doping profiles which are depicted in Fig. 2 were recorded by time-of-flight secondary ion mass spectroscopy (ToF-SIMS) and phosphorus was found to pile up at the poly-Si/SiO$_2$ interface while boron readily diffused through the oxide thereby forming a less shallow junction which adversely affects surface passivation. By using a nitrided oxide boron diffusion was suppressed successfully but a degradation of surface passivation quality was observed as well. Moreover, 1D and 3D numerical process simulation describing the diffusion profiles revealed that boron diffusion through pinholes contributes little to the aggregate diffusion profile in the c-Si which is accessible via electrochemical capacitance-voltage profiling.

![Figure 2: ToF-SIMS profiles of n-type and p-type TOPCon structures featuring a plasma oxide and a plasma nitrided oxide, respectively. The $V_{oc}$ values provided in the graph’s legend were determined after furnace anneal and hydrogenation.](image)

2.2 Development of PECVD tube process

The main objective of the given project was the transfer of the TOPCon deposition process from the laboratory PECVD tool Roth&Rau AK-400M to the industrial PECVD tool centrotherm cPLASMA 2000. The main development was performed using a horizontal boat configuration as displayed in Fig. 3. Following up on the first encouraging results that were published in [14], a process was developed that is characterized as follows:

- **In-situ** doping via addition of phosphine (PH$_3$)
- High deposition rate of $\approx 12 \text{ nm/min}$
- Low H-content in layers permits growth of films thicker than 200 nm
- Excellent surface passivation quality ($J_0 \leq 1 \text{ fA/cm}^2$ on planar and $J_0 \approx 3 \text{ fA/cm}^2$ on textured surfaces)
- Homogeneous deposition and passivation quality over the whole boat (see Fig. 3)

As reported in [15], the deposition process was successfully transferred to a vertical boat which resembles a boat configuration used in production.

![Figure 3: (Top) A photograph of a horizontal PECVD boat. (Bottom) A side view schematic of the horizontal boat displaying the $V_{oc}$ and $iFF$ values determined by QSSPC on the respective wafers.](image)

2.3 Hydrogenation of TOPCon

The saturation of dangling bonds at the Si/SiO$_2$ interface [16] and presumably the elimination of defects within the poly-Si layer [17] via hydrogen is a key process step enabling record-high surface passivation quality. At Fraunhofer ISE, a process called remote plasma hydrogen passivation (RPHP) [18] soon became the standard hydrogenation process for small wafer sizes. In the third work package alternative hydrogenation processes were investigated.

One important finding was that processes which solely supply molecular hydrogen are less effective than processes based on atomic hydrogen [19] (see also [20]). The latter can be released from atomic layer deposition (ALD) Al$_2$O$_3$:H, PECVD SiN$_2$:H, and even hydrogen-containing transparent conductive oxide (TCO) layers [21] during a low-temperature anneal at about 400°C or a fast-firing process. The process sequence PECVD SiN$_x$ and firing was studied in more detail within the framework of the project “PV-BAT400”. As presented at the 36th EU-PVSEC [15], SiN$_x$ layers with a higher refractive index (and thus more Si-H bonds) enable a superior passivation quality after firing compared to SiN$_x$ layers with a refractive index of less than 2.0. Using an optimized process sequence, $J_0$ values of about 3 fA/cm$^2$ can be achieved on rough surfaces. More details can be found in [22].

2.4 Free carrier absorption

In contrast to our 25.8%-efficient lab-type TOPCon cell, industrial TOPCon cells make use of thicker (150-200 nm) heavily-doped poly-Si layers which adversely affect the infrared response of silicon solar cells via free
carrier absorption (FCA). The influence of poly-Si doping level was studied experimentally on lab-type solar cells featuring a 140 nm thick poly-Si layer at the rear. Fig. 4 plots the $J_{oc}$ losses attributed to FCA over poly-Si dopant concentration. The $J_{oc}$ losses were calculated from the cells’ external quantum efficiency (EQE) data in the infrared region. For more information, the reader is kindly referred to [23]. The dashed and dotted lines refer to FCA-induced $J_{oc}$ losses obtained from numerical device simulation using the parameterization for FCA in c-Si which was determined by Baker-Finch et al. [24]. It can be seen that FCA losses increased significantly with dopant concentration and that numerical device simulation using the mean value or upper limit of the FCA parameterization can reproduce the experimentally obtained data points. From this graph, it is obvious that thick poly-Si layers with $N_{D,\text{poly-Si}} \geq 1.5 \times 10^{20} \text{cm}^{-3}$ lead to significant FCA losses and, therefore, should be avoided.

2.5 TOPCon Cells with an area of 10x10 cm$^2$

Within the project “Upgrade Si-PV” 10x10 cm$^2$ solar cells were processed using 250 μm thick n-type 1 Ωcm FZ-Si wafers and the results were published in [25]. The solar cells featured a homogeneous boron emitter with an emitter sheet resistance $R_{sh} = 140 \Omega$/sq and a TOPCon rear contact with full-area physical vapor deposited (PVD) silver metallization. A three busbar H-pattern layout was realized at the front employing either one of the two following process sequences:

i) Photolithography (PL) and PVD metallization with a metallized area fraction of $\approx 0.5\%$ (non-contacting busbars)

ii) laser contact opening (LCO) and plating of NiCuAg with a metallized area fraction ($A_{\text{met}}$) of $\approx 1.5\%$ (contacting busbars)

The current-voltage (I-V) parameters of the best cells are shown in Table I. The cell featuring PVD metallization achieved a high $V_{oc}$ of 713 mV which is very close to the maximum $V_{oc}$ measured internally on cells featuring a homogeneous emitter. Significantly higher $V_{oc}$ values can only be obtained by introducing a selective emitter scheme or TOPCon underneath the front metallization. On the other hand, the cells with plated metallization showed a lower $V_{oc}$ of 697 mV due to enlarged $A_{\text{met}}$ and thus increased contact recombination. The latter can be attributed to laser-induced damage of the emitter bulk in the near-surface region which was not cured by a high-temperature process such as firing. Nevertheless, these results indicate, that the $V_{oc}$ of a TOPCon cell will be limited by recombination at the front surface in most cases.

Table I: I-V parameters of best 10x10 cm$^2$ solar cells independently confirmed by Fraunhofer ISE CalLab PVCells.

<table>
<thead>
<tr>
<th>Front metallization</th>
<th>$V_{oc}$ [mV]</th>
<th>$J_{sc}$ [mA/cm$^2$]</th>
<th>FF [%]</th>
<th>pFF [%]</th>
<th>$\eta$ [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>PL + PVD</td>
<td>713</td>
<td>41.4</td>
<td>83.1</td>
<td>84.8</td>
<td>24.5</td>
</tr>
<tr>
<td>LCO + Plating</td>
<td>697</td>
<td>41.4</td>
<td>81.2</td>
<td>84.3</td>
<td>23.4</td>
</tr>
</tbody>
</table>

3 RECENT TOPCON SOLAR CELLS

3.1 Experimental Details

Industrial TOPCon solar cells were realized on M2-sized n-type 1 Ωcm Cz-Si wafers which were supplied by LONGI. The process sequence is depicted in Fig. 5. After saw damage removal, texturing, and cleaning, three different boron emitters were realized by atmospheric pressure BBR$_3$ diffusion. The boron emitters are characterized as follows:

- $1) R_{sh} \approx 110 \Omega$/sq, $J_{sc} \approx 27$ fA/cm$^2$
- $2) R_{sh} \approx 120 \Omega$/sq, $J_{sc} \approx 22$ fA/cm$^2$
- $3) R_{sh} \approx 150 \Omega$/sq, $J_{sc} \approx 15$ fA/cm$^2$

More information on the processes can be found in [26]. After wet-chemical edge isolation, a tunnel oxide layer was grown thermally and 150 nm in-situ phosphorus-doped a-Si was deposited by a centrotherm cPLASMA PECVD tool. Since the deposition process produces a minimal wrap around at the edges, a wet-chemical process was employed to remove the a-Si layer from the front surface. Thereafter, TOPCon was activated by a furnace anneal. The front surface was passivated by a stack of ALD Al$_2$O$_3$ and PECVD SiN$_x$:H. The rear received a SiN$_x$:H layer with a slightly higher refractive index – as outlined above. A busbarless grid design was applied by screen-printing a silver-aluminum (AgAl) paste on the front and a silver (Ag) paste on the rear side. The cells were fired in an industrial conveyor belt furnace at peak set temperatures ranging from 800°C to 840°C. The cells were measured using a Halm inline flash tester which is equipped with a Pasan PCB-Touch unit featuring a black, non-conductive chuck.

Figure 5: Process sequence of i-TOPCon cells featuring a tube PECVD process.
3.2 Interplay between TOPCon and metallization

The contact properties (i.e., $J_{\text{met}}$, $\rho_c$) of screen-printed fire-through contacts were studied on 100-200 nm thick TOPCon layers. Fig. 6 plots the $J_{\text{met}}$ values over nominal poly-Si thickness in dependence of firing temperature. The blue data points refer to data from literature [27, 28]. The results suggest that a poly-Si thickness of 100 nm or less is still critical for screen-printed and fired contacts as $J_{\text{met}}$ values above 1000 fA/cm² were determined from photoluminescence (PL) images. On the other hand, reasonably low $J_{\text{met}}$ values of about 200 fA/cm² were achieved on both 150 nm and 200 nm thick poly-Si contacts. In addition, low contact resistivity values of $\approx$ 1 mΩcm² were obtained for the metal/poly-Si interface.

For comparison with the solar cell’s front side it is interesting to note that investigations on test structures featuring a boron-diffused emitter (not shown here) yielded $J_{\text{met}}$ values in the range of 1000-2000 fA/cm² and showed that contact resistivity decreased from 10 mΩcm² down to 2-3 mΩcm² with increasing firing temperature.

![Figure 6: $J_{\text{met}}$ values plotted over nominal poly-Si thickness for three different set peak firing temperatures $T_{\text{set}}$. Please note that some data points are adjacent placed for clarity.](image)

3.3 Solar cells

Fig. 7 plots the light $I-V$ parameters of the TOPCon solar cells over the emitter sheet resistance. The differently colored boxes refer to different set peak firing temperatures. One can see that the firing temperature had a strong impact on both the cells’ $V_{oc}$ and $FF$. $V_{oc}$ decreased with increasing firing temperature by about 10 mV. Such loss can mainly be ascribed to an increased recombination rate at the front metal contacts. The application of an emitter with a higher $R_{sh}$ yielded a slight increase in $V_{oc}$ from 694 mV ($R_{sh} = 110$ Ω/sq) to 698 mV ($R_{sh} = 150$ Ω/sq). Compared to the $V_{oc}$, the $FF$ shows a quite different dependence on firing temperature. At the lowest firing temperature of 800°C, the lowest $FF$ values were measured within each group. In case of the emitter exhibiting $R_{sh} = 110$ Ω/sq, a mean $FF$ of 78.2% was obtained. The cells of this group which were fired at 820°C and 840°C, respectively, yielded substantially higher $FF$ values above 80.0%. The groups featuring more resistive emitters exhibited similar $FF$ values in the range of 80.0% to 80.7% for both firing temperatures 820°C and 840°C. The increase in $FF$ with firing temperature went along with a reduction in series resistance from $\approx$ 0.8 Ωcm² ($R_{sh} = 110$ Ω/sq / 800°C) to $\approx$ 0.45 Ωcm² ($R_{sh} = 110$ Ω/sq / 840°C). This effect can be explained by improved contact formation at the front. The cells’ $J_sc$ averaged to 40.2 mA/cm² and did not show any significant dependence on emitter process or firing temperature.

The most efficient cell was then coated with magnesium fluoride (MgF₂) to further reduce reflection losses and has been measured by Fraunhofer ISE CalLab PVCells using a reflective chuck and 30 wires. The independently confirmed light $I-V$ parameters are displayed in Table II and an efficiency of close to 23% was obtained.

![Figure 7: Light $I-V$ parameters of TOPCon solar cells as a function of emitter sheet resistance.](image)
Table II: The I-V parameters of the champion solar cell were independently confirmed by Fraunhofer ISE CalLab PVCells. The solar cell was measured on a reflective chuck using 30 wires.

<table>
<thead>
<tr>
<th>V&lt;sub&gt;OC&lt;/sub&gt; [mV]</th>
<th>J&lt;sub&gt;SC&lt;/sub&gt; [mA/cm²]</th>
<th>FF [%]</th>
<th>η [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Champion</td>
<td>700</td>
<td>41.2</td>
<td>79.6</td>
</tr>
</tbody>
</table>

4 SUMMARY AND OUTLOOK

The work within the project “Upgrade Si-PV” laid the foundations for advancing the industrialization of centrotherm’s cPLASMA tube PECVD system. The TOPCon PECVD process is characterized by the following key features:
- Excellent surface passivation quality ($J_0 \leq 1$ fA/cm² on planar and $J_0 \approx 3$ fA/cm² on rough surfaces)
- In-situ doping via addition of PH₃
- High deposition rate of $\approx 12$ nm/min
- Layers are compatible with screen-printed metallization.
These layers were then successfully integrated into TOPCon solar cells achieving an independently confirmed efficiency of 22.95% and a $V_{OC}$ of 700 mV.

Our results show that the limiting factor of our cell is the recombination at the interface between metal and homogeneous boron emitter. Hence, a more advanced emitter structure is the key to achieve higher efficiencies.

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6 REFERENCES


