Postmetallization "Passivated Edge Technology" for Separated Silicon Solar Cells

Puzant Baliozian[®], Mohammad Al-Akash, Elmar Lohmüller[®], Armin Richter[®], Tobias Fellmeth, Anna Münzer[®], Nico Wöhrle, Pierre Saint-Cast[®], Hannah Stolzenburg, Alma Spribille, and Ralf Preu

Abstract—This article introduces a postmetallization "passivated edge technology" (PET) treatment for separated silicon solar cells consisting of aluminum oxide deposition with subsequent annealing. We present our work on bifacial shingle solar cells that are based on the passivated emitter and rear cell concept. To separate the shingle devices after metallization and firing, we use either a conventional laser scribing mechanical cleaving (LSMC) process or a thermal laser separation (TLS) process. Both separation processes show similar pseudo fill factor (pFF) drops of - 1.2% $_{\rm abs}$ from the host wafer to the separated state. The *pFF* of the TLSseparated cells increases by up to $+0.7\%_{abs}$ from the as-separated state after PET treatment due to edge passivation, while the pFF of LSMC-separated cells increases by up to $+0.3\%_{abs}$. On cell level, the combination of TLS and PET allows for a designated area output power density of $p_{out} = 23.5 \text{ mW/cm}^2$, taking into account an additional 10% rear side irradiance.

Index Terms—ALD, aluminum oxide, bifacial PERC, passivated edge technology, postmetallization, shingle solar cells, thermal laser separation (TLS).

I. INTRODUCTION

The REVIVAL of the shingling interconnection approach of solar cells [1] is an option to obtain higher photovoltaic module output power densities p_{out} . Shingling of solar cells is done by interconnecting the rear busbar of a cell to the front busbar of a neighboring one. The overlap and the interconnection of the busbars lead to: (i) the removal of cell spacing and therefore to an increase of active cell area within the module, (ii) the decrease of shading losses due to the absence of visible busbars and interconnectors, and (iii) the decrease of electrical resistance losses on the interconnection level. Shingling solar cells have first been used for niche applications such as satellite devices [2], electronic devices [3], and electrical vehicle prototypes [4]. The potential shown by the shingling approach leads to the recent increase in interest not only shown in publications [5]–[8]

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and patents [9]–[11], but also in already existing commercially available shingle modules [12], [13].

Similar to half-size cells [14], shingle cells are usually separated from metallized and fired host wafers. The separation process leads to a decrease in solar cell efficiency due to edge recombination induced by the separation [15]. Edge recombination becomes even more significant for small-sized cells with higher perimeter-to-area ratios, leading to losses mainly in fill factor *FF* (initially caused by losses in pseudo fill factor *pFF*) in addition to losses in open-circuit voltage $V_{\rm OC}$.

Edge recombination can be reduced by: (i) the depletion of charge carriers of a kind from the edge and/or (ii) the reduction of density of traps at the edge. As review of potential methods for edge passivation, we find that some previously investigated methods to reduce edge recombination of solar cells consist of introducing an emitter window commonly created by the use of diffusion barriers [16], [17]. The emitter window cuts off the emitter conduction towards the edge, thus reducing the support of minority charge carriers tremendously. Having the same aim, the removal of the emitter by creating passivated isolation trenches is proposed as another approach [18]. In this method, the trenches are formed by ablating the emitter, followed by wet-chemical etching, and finally, passivating the trenches by thermally grown silicon dioxide (SiO₂) or polysilicon. In Ref. [19], a strong doping at the separation path is suggested to create a repulsion of carriers from the edge by a surface field effect. It has also been reported that treating the edge wetchemically allows the growth of SiO₂ on the edge, which shows a passivation effect [20]. In Ref. [21], the surface passivation of two sawed slits tens of micrometers far from the cell active area is achieved by the deposition of aluminum oxide, capped by silicon nitride, and silicon oxide layers. The process is done after partially damage etching the slit surfaces.

However, fabricating such cells requires several additional premetallization process steps [22] or postmetallization chemical etching processes, which make the implementation on industrial fabrication scale challenging. Simulation results confirm the positive effect of passivated edges on shingle cells by reducing the effective surface recombination velocity at the edge, $S_{\rm eff,edge}$ [23], [24]. Nevertheless, no potentially industrial feasible process is found in literature.

This article introduces the concept of postmetallization "passivated edge technology" (PET). The PET is demonstrated on bifacial *p*-type silicon shingled passivated edge, emitter, and rear (*p*SPEER) solar cells, which have been initially reported without

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The authors are with the Fraunhofer Institute for Solar Energy Systems ISE, 79110 Freiburg, Germany (e-mail: puzant.baliozian@ise.fraunhofer.de; mohammad.al-akash@ise.fraunhofer.de; elmar.lohmueller@ise.fraunhofer.de; armin.richter@ise.fraunhofer.de; tobias.fellmeth@ise.fraunhofer.de; anna. muenzer@ise.fraunhofer.de; nico.woehrle@ise.fraunhofer.de; pierre.saint-cast@ise.fraunhofer.de; hannah.stolzenburg@ise.fraunhofer.de; alma.spribille @ise.fraunhofer.de; alma.spribille @ise.fraunhofer.de; alma.spribille

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PET in Ref. [25]. The bifacial *p*SPEER cell concept is based on the passivated emitter and rear cell (PERC) [26] architecture, where an additional laser-assisted separation process is applied to obtain the cell stripes. The separation processes investigated in this article are conventional laser scribing and mechanical cleaving (LSMC) and thermal laser separation (TLS) [27], [28]. The effect of the separation processes and the edge passivation are discussed together with the first solar cell results.

II. PASSIVATED EDGE TECHNOLOGY AND CELL CHARACTERIZATION

A. pSPEER^{PET} Solar Cell Concept

The separation process leads to an increase of edge recombination, especially when the emitter extends to the separated edges; see Fig. 1(a) for a schematic cross section of the pSPEER cell. The cells only have a natively grown SiO₂ layer at the edge. For that reason, an additional postmetallization/separation PET aims to decrease $S_{\rm eff,edge}$ by coating the edges with an intended dielectric passivation layer. The "pSPEERPET" solar cell is obtained after edge passivation by the PET; see Fig. 1(b). In this article, the PET consists of two steps: a postmetallization/separation deposition of an aluminum oxide (AlO_X) layer and a postdeposition annealing (PDA) for the activation of the deposited AlO_X passivation layer. Thermal atomic layer deposition (ALD) is used for the AlO_X deposition. This has several advantages: (i) ALD is known for a very conformal deposition, e.g., the wafer edge is well coated after the process; (ii) AlO_X can provide an excellent surface passivation because it combines enhanced chemical passivation with a strong field effect passivation induced by a high amount of fixed negative charges [29]; (iii) it is transparent for very thin films of thicknesses d ranging between 5 nm $\leq d \leq$ 15 nm [30]. OPTOS simulations done on module level confirm that such thin AlO_X layers deposited on the front and rear surfaces of the cells do not induce optical losses; (iv) the passivation effect of thermal ALD-deposited AlO_X is known to be activated at low annealing temperatures $T_{\rm ann}$ < 225 °C [31], [32]. The latter hinders thermally induced metal contact degradation which has been reported [33].

B. Solar Cell Fabrication

Industrial 6-inch gallium-doped Czochralski-grown silicon (Cz-Si:Ga) PERC precursors are used for the fabrication of the pSPEER and pSPEER^{PET} cells investigated in this article. Fig. 2 shows the corresponding fabrication process. The precursors' base resistivity ranges between 0.3 Ω cm $\leq \rho_B \leq 0.9 \Omega$ cm. The front phosphorous-doped emitter is passivated by a silicon nitride (SiN_X) layer; the rear silicon base is passivated by a layer stack consisting of AlO_X capped by SiN_X . The precursors are optimized for monofacial use, which explains the yellowish color of the rear side; see Fig. 3(a). The fabrication process performed in this article starts from the laser contact openings. The rear side silver busbars (external contacts) are screen printed first. Next, the rear-side aluminum contact grid and the front-side silver contact grid are printed. Contact firing is then performed in an industrial fast firing oven. The printed metallization layouts are designed to obtain six shingle devices from one host



Fig. 1. (a) Schematic cross section of a bifacial *p*SPEER solar cell. The structure is similar to the bifacial PERC concept with additional separated edges covered by native SiO₂. (b) After PET, the *p*SPEER^{PET} solar cell has an additional postmetallization dielectric passivation layer (i.e., AIO_X) all over the cell and, most importantly, at the separated edges (green layer). In both figures (a) and (b), the external contacts (rear-side silver busbars) are not shown.

6-inch industrial p-type Cz-Si PERC precursors
Laser local contact opening
Screen printing
Firing
Host wafers
Laser-assisted separation
pSPEER solar cells
Passivated edge technology (PET)
pSPEER ^{PET} solar cells

Fig. 2. Process flow of the *p*SPEER^{PET} solar cell's fabrication from industrialpassivated 6-inch *p*-type Cz-Si:Ga PERC precursors.

wafer [each exhibiting the dimensions $22 \text{ mm} \times 148 \text{ mm}$; see Fig. 3(b)]. Finally, six individual shingle cells are obtained after laser-assisted separation as shown in Fig. 3(c).

The performed separation processes in this article are (i) conventional LSMC and (ii) TLS, both implemented by using the *microDICE* tool from *3D-Micromac* [34]. The following items describe the processes:

- LSMC is done by a nanosecond infrared pulsed laser that scribes along the entire separation path and ablates up to one-third of the substrate's thickness. Consequently, the samples are manually and mechanically cleaved.
- TLS is performed by creating an initial scribe by means of the same pulsed laser used in the LSMC process to start a crack of length between 0.5 mm ≤ l ≤ 1 mm. For the



Fig. 3. Scanned images of the front side and rear side of (a) an industrial 6-inch PERC precursor, and (b) a metallized host wafer with the metallization layouts from which six cells can be separated. (c) pSPEER cells after laser-assisted separation with the dimensions 22 mm \times 148 mm. Microscope images show the metallization on the front and rear sides. The figure is adapted from Ref. [35].

cleave process, an infrared continuous wave laser in combination with a simultaneous water/air cooling jet is used to thermally induce a compressive stress and consequent tensile stress along the separation path. This leads to the cleavage of the substrate.

Planar view scanning electron microscope (SEM) images of exemplary cell edges for LSMC and TLS processed samples are shown in Fig. 4(a) and (b), respectively. The edge obtained by LSMC has a rough laser-scribed region which is about one-third of its thickness. The mechanically cleaved part of the edge is rather smooth. In contrast, the cell edge obtained by the TLS process shows an entirely smooth surface.

C. SunsV_{OC} and Current–Voltage Characterization

In this article, $SunsV_{OC}$ measurements [36], [37] are performed to characterize the effect of the separation and passivation processes. By this method, the effect of edge recombination on the solar cell performance is independent of possible changes





Fig. 4. SEM images of (a) cell edge separated by an LSMC process. The scribe-laser-induced roughness dominates around one-third of the thickness. (b) Cell edge separated by TLS with substantially smoother edges due to the complete thermal cleave as a result of the process.



Fig. 5. (a) Schematic of contacting positions during the Suns $V_{\rm OC}$ measurement of the host wafer before separation. Five positions are contacted per busbar (here busbar 3 is taken as an example). (b) Contacting of a shingle cell on the busbar at the same five positions. The separated cell is placed on the chuck at the position where it was positioned prior to its separation.

in series resistance r_S . From the measurements, pFF and $V_{\rm OC}$ values at an illumination intensity of 1000 W/m² are extracted. In order to exclude effects of local inhomogeneity (either from the host wafers or from the Suns $V_{\rm OC}$ measurement setup), each host wafer is measured at five different positions on each busbar; see Fig. 5(a). Separated cells are then placed at the same position on the Suns $V_{\rm OC}$ chuck as measured prior to separation within the host wafer; see Fig. 5(b). After each additional process step performed on the cell, it is contacted and measured in an



after annealing

Fig. 6 Experimental process and characterization steps to investigate the effect of PET on separated cells.

identical way. A reference cell not undergoing any process steps is measured continuously throughout the experiments to track the reproducibility of the Suns V_{OC} measurement.

The illuminated current–voltage (IV) measurement is done at standard test conditions on each side separately using an industrial cell tester calibrated by a reference *p*SPEER cell measured at Fraunhofer ISE CalLab PV Cells. During the IVtesting procedure, the cells are placed on a black, nonconducting background, which ensures negligible reflection well below 5% over the relevant spectrum. The contacting is performed at the front- and rear-side busbars by pin arrays.

III. EDGE PASSIVATION

A. Experimental Plan for PET on Cells

The aim of this experiment is to observe the effect of the separation processes from host wafer to separated state by using Suns V_{OC} measurements; see the experimental process flow in Fig. 6. In addition, the effect of the AlO_X deposition and subsequent annealing on cells is examined. LSMC is done 6 h before AlO_X deposition. The TLS-diced host wafers are separated also 6 h prior to the AlO_X deposition, while another group is TLS-diced 19 h prior to the AlO_X deposition. The different separation dates are to investigate the influence of the waiting time between separation and deposition. In air at room temperature, native SiO₂ growth occurs on exposed silicon [38]. Due to the logarithmic growth of SiO₂ as a function of exposure time to air, similar layer thicknesses are expected for both waiting times in this article.

The AlO_X deposition process is performed in a *FlexAL* reactor of *Oxford Instruments* [39] using trimethylaluminum and water vapor as precursors. Two deposited AlO_X layer thicknesses are investigated: $d_1 = 7$ nm and $d_2 = 14$ nm, as measured with spectroscopic ellipsometry on a planar silicon process control sample. Three to four separated cells have been processed in each deposition run (see Fig. 7). Height spacers are used to ensure a deposition on the rear side as well.

SunsV_{OC} measurements before and after deposition are completed by contacting at five measurement positions per busbar. Postdeposition hotplate annealing (PDA) is performed at temperature $T_{\rm ann} < 225$ °C. Control cells (without deposition) from each separation process are kept and are not coated with AlO_X



Fig. 7. Top view sketch of three shingle cells, an ellipsometer measurement sample, and the spacers during the PET deposition process in the *FlexAL*. The cells are placed with the front side facing up. The deposition on the front and rear sides is intended by spacers.

to observe the effect of the annealing process on the non-coated cells. Suns $V_{\rm OC}$ measurements are then done after PDA. To investigate the stability of the passivation effect, TLS-separated cells are measured again after 73, 99, and 167 h.

To understand the effect of the deposition and annealing processes and estimate the effective surface recombination velocity $S_{\rm eff}$, a symmetrical lifetime sample (*n*-type float-zone silicon, $\rho_{\rm B} \approx 1 \ \Omega$ cm, wafer thickness $W \approx 200 \ \mu$ m) is coated with $d_1 = 7 \ \text{nm AlO}_X$ layer.

Quasi-steady-state photoconductance (QSSPC) measurements [40] are performed to obtain the minority carrier lifetime τ_{eff} dependency on the excess carrier density Δn at the following states: (i) as-deposited and (ii) annealed. S_{eff} is then approximated by using [41]

$$1 / \tau_{\text{eff}} = (1 / \tau_{\text{bulk}}) + (2S_{\text{eff}} / W).$$
 (1)

Noting that τ_{bulk} is the intrinsic bulk lifetime and τ_{eff} is extracted at $\Delta n = 10^{15} \text{ cm}^{-3}$.

IV. RESULTS AND DISCUSSION

A. Effect on Charge Carrier Lifetime

The QSSPC measurement of the lifetime sample in Fig. 8 shows the injection-dependent lifetime obtained by means of QSSPC for the as-deposited and annealed states. After annealing, $\tau_{\rm eff} = 432 \ \mu s$ at $\Delta n = 10^{15} \ {\rm cm}^{-3}$ is measured.

Using (1), $S_{\text{eff}} = 22$ cm/s is obtained, showing the possibility of attaining low S_{eff} values by the current deposition/annealing method. Note that the planar S_{eff} is not necessarily equal to that of the edge due to different surface and geometrical conditions.

B. SunsV_{OC} Results

1) Effect of PET: From the SunsV_{OC} measurements, we extract $V_{\rm OC}$ values at an illumination intensity of 1000 W/m² and the *pFF*. The host wafers attain mean $V_{\rm OC} = 669$ mV. The changes in $V_{\rm OC}$ throughout the process steps are minor, with a maximum variation of $\Delta V_{\rm OC} = \pm 2$ mV. This small impact on $V_{\rm OC}$ is in agreement with simulation results for similar sample types and $V_{\rm OC}$ levels [23], [42].



Fig. 8. QSSPC measurement of the minority carrier lifetime $\tau_{\rm eff}$ dependent on excess carrier density Δn for the symmetrical *n*-type float-zone silicon lifetime samples. After annealing, the sample attains $\tau_{\rm eff} = 432 \,\mu {\rm m}$ extracted at $\Delta n = 10^{15} {\rm cm}^{-3} \approx 1000 {\rm W/m^2}$ illumination level).



Fig. 9. Suns $V_{\rm OC}$ measurement results of host wafers, after separation, asdeposited, and annealed. Control cells without deposition are also measured before and after annealing. The hours on the top of the plot indicate the approximate time gap between separation and deposition.

Hence, we focus on the *pFF* data which are shown in Fig. 9. The values are given for the four different sample states: (i) host wafer, (ii) separated, (iii) as-deposited, and (iv) annealed. They are divided into the different separation groups and AlO_X layer thicknesses. The results are discussed considering the state-to-state changes.

To start with, LSMC and TLS separation processes lead to a decrease in pFF ($\Delta pFF = -1.2\%_{abs}$) from the host wafer to the separated state due to similar edge recombination effects.

Noticeably, the introduction of the AlO_X layer with thickness d_1 or d_2 leads to an improved edge passivation and thus to a gain in *pFF*. The deposition on TLS-separated cells leads to the highest increase from separated to as-deposited state reaching $\Delta pFF = +0.4\%_{abs}$.

After annealing, the TLS-separated and coated cells show the highest increase, $\Delta pFF = +0.4\%_{abs}$ from as-deposited to annealed states. Such an increase after PDA is due to the activation of the passivation layers that coat the smooth edge surface. Annealing of control cells without deposition leads to an enhanced *pFF*. This is probably due to the grown SiO₂ on the edges, which is allowed by the absence of the AlO_X coating.



Fig. 10. pFF values attained by SunsV_{OC} measurements of TLS-separated cells before and after PET. The same cells are measured 73, 99, and 167 h after PET. Measurements of TLS-separated cells that are annealed without undergoing deposition are included. The reproducibility of the measurement is verified by a reference cell.

As an assessment of PET, the TLS-separated cells show higher *pFF* gains than the LSMC-separated ones when both PET process steps (deposition and PDA) are completed. The deposition on a smooth surface with reduced defects obtained by TLS and the activation of the deposited passivation layer lead to the highest *pFF* increase of $+0.7\%_{abs}$ and the highest measured *pFF* values. TLS-separated cells regain up to $50\%_{rel}$ of their initial *pFF* loss due to the separation process (observed for both deposition thicknesses and separation groups), and control cells that are not coated and are only annealed do not show such a regain. LSMC-separated cells regain around $16\%_{rel}$ after PET. This shows that PET leads to edge passivation for both separation processes; however, a larger effect is recorded for TLS-separated cells.

2) Edge Passivation Stability: Based on the *pFF* results, the passivation stability investigation (see Fig. 10) shows that the cells coated with $d_2 = 14$ nm remain almost constant with minimal ($\Delta pFF \approx 0.1\%_{abs}$) to no measured decrease. These results show a stable edge passivation over time for the cells coated with $d_2 = 14$ nm AlO_X, while cells with $d_1 = 7$ nm or without degrade with time. The explanation for the time-based passivation stability on the deposited layer thickness is under further investigation.

C. IV Measurement Results

Based on the results of the previous section, the favored processes for the fabrication of edge-passivated cells are used to fabricate cells for IV measurements. Host wafers are separated into shingle cells by means of TLS. The cells are then coated using an identical thermal ALD deposition process to the one used before. This time this leads to around 13-nm-thick postmetallization AIO_X layer. The cells are then annealed on a hotplate.

IV measurements of TLS-separated edge-passivated $pSPEER^{PET}$ solar cells are done in comparison to LSMC-separated pSPEER solar cells that do not undergo any further processes after separation. The front- and rear-side IV measurement data from the most efficient cells are summarized

TABLE I IV MEASUREMENT DATA FOR THE CELLS WITH HIGHEST OUTPUT POWER DENSITIES FOR TLS+PET-PROCESSED AND LSMC-SEPARATED CELLS IN ADDITION TO AVERAGE VALUES OF THE FRONT-SIDE MEASUREMENTS

Meas. side	η (%)	V _{OC} (mV)	$j_{\rm SC,des}$ (mA/ cm ²)	FF (%)	pFF (%)	$r_{\rm S}$ (Ω cm ²)	β (%)	p_{out} (mW /cm ²)	
TLS separated p SPEER ^{PET} solar cell									
Front Av.	21.8	668	40.4	81.0	82.6	0.38		-	
Front Peak	22.1	669	40.5	81.4	83.2	0.38	67	23.5	
Rear Peak	14.7	660	27.2	81.8	82.8	0.35			
LSMC separated <i>p</i> SPEER solar cell									
Front Av.	21.6	664	40.2	81.0	82.3	0.35		-	
Front Peak	21.7	667	40.3	80.9	82.4	0.35	66	23.1	
Rear Peak	14.4	657	27.0	81.0	82.0	0.35			

Characteristic data for the front- and rear-side measurements are shown. Front- and rear-side measurements are separately performed at $G = 1000 \text{ W/m}^2$. Since the busbar is intended to be covered in the module, a designated area $j_{\rm SC,des}$ excluding the busbar area is considered. $p_{\rm out}$ considers an additional 10% irradiance from the rear side.

in Table I. For the front-side measurements, average values (Av.) are also shown. The TLS-separated cell after PET treatment yields a designated front-side energy conversion efficiency $\eta_{\rm f} = 22.1\%$, featuring $V_{\rm OC} = 669$ mV, FF = 81.4%, and short-circuit current density $j_{SC,des} = 40.5 \text{ mA/cm}^2$. Due to the PET treatment, the cell features a high pFF = 83.2%. Even though PET is included as postfiring thermal process, a low $r_S = 0.38 \ \Omega \ \mathrm{cm}^2$ is achieved, hinting that no significant degradation at the metal contacts occurs. The LSMC-separated cell without PET attains $\eta_f = 21.7\%$. In comparison to the pSPEER cell, the pSPEER^{PET} cell features a higher front-side efficiency $\Delta \eta_f = +0.4\%_{\rm abs}$, which can be attributed to the higher short-circuit current density $\Delta j_{SC} = +0.2 \text{ mA/cm}^2$ and $\Delta FF = +0.5\%_{\rm abs}$. This difference in FF can be explained by the higher *pFF* of $+0.8\%_{abs}$, while featuring a slightly higher r_S .

Mainly, the improved pFF is a result of an enhanced edge quality obtained by the TLS process and the additional PET processes.

The rear-side measurement results also show similar trends, where the *p*SPEER^{PET} cell attains a designated rear-side energy conversion efficiency $\eta_r = 14.7\%$ that is $+0.3\%_{abs}$ higher than the LSMC-separated *p*SPEER cell that attains $\eta_r = 14.4\%$. The difference in $\Delta pFF = +0.8\%_{abs}$ is entirely projected on the $\Delta FF = +0.8\%_{abs}$ due to the identical rear side $r_S = 0.35 \ \Omega \ cm^2$ between the two compared cells. The TLS-separated and edgepassivated solar cell attains a bifaciality $\beta = \eta_r/\eta_f = 0.67$, whereas the LSMC-separated cell without PET attains $\beta = 0.66$. Considering an additional 10% rear-side irradiance $G_r = 100 \ W/m^2$, the edge-passivated cell attains an output power density $n_{rear} = 23.5 \ mW/cm^2$. This value is greater by

 $G_r = 100 \text{ W/m}^2$, the edge-passivated cell attains an output power density $p_{\text{out}} = 23.5 \text{ mW/cm}^2$. This value is greater by $\Delta p_{\text{out}} = +0.4 \text{ mW/cm}^2$ in comparison to the nonpassivated LSMC-separated cell.

The result of the TLS + PET processed cell shows its clear advantage in terms of efficiency. Therefore, we have demonstrated the functionality of the proposed PET postmetallization edge passivation approach. No changes in the premetallization processes in an industrial production line are required. The integration of PET into the fabrication after the separation of silicon solar cells (i.e. shingle cells, half-cut cells, or small-area cells) can, thus, lead to a significant boost in cell efficiency with a relatively reasonable integration within the already existing production infrastructure. This makes the proposed PET attractive for industrial application and incorporation in production lines.

V. CONCLUSION

This article introduces a postmetallization/separation PET for separated solar cells. The effect of PET on bifacial *p*-type silicon shingled passivated edge, emitter, and rear solar cells (*p*SPEER) is tested. After edge passivation, we call the PET-treated shingle cells "*p*SPEER^{PET}" solar cells.

The TLS process leaves the separated cells with visibly smoother edges in comparison to separation by LSMC. By just considering the separation processes without further PET, the separation processes lead to similar drops in pseudo fill factor pFF of $\Delta pFF \approx -1.2\%_{abs}$.

For the PET deposition process, low-temperature thermal ALD of AlO_X for two layer thicknesses $d_1 = 7$ nm and $d_2 = 14$ nm is performed. TLS-separated PET-treated pSPEER^{PET} cells have regained half the loss that is induced due to separation. The *pFF* increase from as-separated TLS-cut cells to edge-passivated cells is found to be up to +0.7%_{abs}, while LSMC-separated cells gain up to +0.3%_{abs} after PET. Thereby, the cells with $d_2 = 14$ -nm-thick AlO_X layer show constant *pFF* values within the experimental time frame of 167 h after passivation, thus indicating the stability of the passivation.

After TLS and PET, the best $p\text{SPEER}^{\text{PET}}$ solar cell attains a designated front-side energy conversion efficiency $\eta_f = 22.1\%$. Assuming an additional rear-side irradiance $G_r = 100 \text{ W/m}^2$, this cell features a total output power density $p_{\text{out}} = 23.5 \text{ mW/cm}^2$. TLS followed by PET leads to a higher output power density of $\Delta p_{\text{out}} = +0.4 \text{ mW/cm}^2$ compared to a PET-free LSMC-separated cell. The PET treatment introduced in this article shows a clear benefit and the thus processed and separated solar cells achieve higher p_{out} values.

The proposed PET is a promising postmetallization edge passivation treatment that might be realized in the industry without requiring adjustments in premetallization stages.

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Puzant Baliozian was born in Beirut, Lebanon, in 1993. He received the B.Sc. degree in physics from the American University of Beirut, in 2014, and the M.Sc. degree in renewable energy engineering and management from the University of Freiburg, Freiburg im Breisgau, Germany, for his thesis work completed at Fraunhofer IPM in the field of magnetocaloric energy conversion, in 2016. He is currently working toward the Ph.D. degree at Fraunhofer ISE, Freiburg im Breisgau.

His main research interests include development of *p*-type SPEER solar cells by focusing on the laser-assisted separation and edge passivation processes.



Mohammad Al-Akash was born in Amman, Jordan, in 1993. He received the B.Sc. degree in energy engineering from German Jordanian University, Madaba, Jordan. He received the M.Sc. degree in renewable energy engineering and management from the University of Freiburg, Freiburg im Breisgau, Germany, in 2019. His master thesis done at Fraunhofer ISE focused on the evaluation of different postmetallization edge passivation methods of *p*-type SPEER solar cells.



Nico Wöhrle was born in Breisach, Germany, in 1984. He received the graduation degree in physics from the University of Freiburg, Freiburg im Breisgau, Germany, for his work on optical simulation of silicon solar cells, in 2012. With a fellowship from the German Federal Environmental Foundation (DBU), he received the Ph.D. degree from the University of Freiburg, in 2016, for his work on simulation of PERC solar cells.

He is currently a Researcher with the Fraunhofer ISE, Freiburg im Breisgau. His research interests include simulation and inline characterization of solar cells as well as the conceptual development of *p*-type PERC solar cells.



Elmar Lohmüller was born in Tübingen, Germany, in 1983. He received the graduation in physics from the University of Tübingen, Tübingen, Germany, and Nelson Mandela Metropolitan University, Port Elizabeth, South Africa. He received the Ph.D. degree from the University of Freiburg, Freiburg im Breisgau, Germany, in 2015, for his work on the development of *n*-type MWT solar cells.

He is currently a Researcher, Project and Product Manager with the Fraunhofer ISE. His research interests include development of *p*-type PERC solar cells.



Pierre Saint-Cast was born in Lorient, France, in 1982. He received the M.Sc. and Engineering degrees from the Polytechnic Institute of Grenoble, Grenoble, France, in 2007, and the Ph.D. degree from the University of Konstanz, Germany, in 2012.

Since 2008, he has been with the Fraunhofer Institute for Solar Energy Systems (ISE), Freiburg, Germany. He is currently the Project Manager and Product Manager for PERC solar cells at the Fraunhofer ISE, Freiburg im Breisgau, Germany. His research interests include the development of passivation lay-

ers for solar cell applications, especially plasma-enhanced chemical vapor deposition of Al_2O_3 layers, and the analytical modeling of electrical transport in Si solar cells devices.



Armin Richter received the Ph.D. degree in physics from the University of Konstanz, Germany, in 2014, for the work on *n*-type silicon solar cells with frontside boron-doped emitters and the in-depth characterization of aluminum oxide-based silicon surface passivation.

His current research interests include atomic layer deposition of functional thin films (e.g., passivation layers, electron/hole transport layers, TCOs), in-depth characterization of dielectric surface passivation layers, as well as the development of high-

efficiency silicon solar cells along the whole process chain and 3-D device simulations.



Hannah Stolzenburg was born in 1993. She received the graduation degree in physics from the University of Göttingen, Germany. She received the M.Sc. degree in 2018 with the focus on material and solidstate physics while spending one semester studying physics at the University of Ghent. Currently, she is working toward the Ph.D. degree, on characterization and quantification of edge losses in silicon solar cells, with the Fraunhofer ISE, Freiburg im Breisgau, Germany.

Alma Spribille received the Diploma in energy and environmental management from the Europe-University of Flensburg, Flensburg, Germany.

She wrote her diploma thesis as an Industrial Engi-

Since December 2017, she has been heading

neer at Fraunhofer ISE about MWT solar cells in 2010

and joined the group MWT Solar Cells & Printing

the team process integration - structuring and



Tobias Fellmeth received the graduation degree in physics from the University of Konstanz, Konstanz, Germany, for his work on the development and characterization of MWT concentrator solar cells. He received the Ph.D. degree from the University of Tübingen, Tübingen, Germany, in 2014, for his work on silicon-based, low concentrator solar cells.

He is currently a Scientist and Project Manager with the Fraunhofer ISE, Freiburg im Breisgau. His research interests include development of bi-facial PERC and TOPCon solar cells.





Ralf Preu was born in Stuttgart, Germany, in 1967. He received the graduation degree in physics from the Universities of Freiburg, Germany and the degree in economics from the University of Hagen, Germany. He received the Ph.D. degree in electrical engineering.

He is currently the Director of the Division for PV Production Technology, Fraunhofer Institute for Solar Energy Systems (ISE), Freiburg, Germany. He joined Fraunhofer ISE in 1993 and has worked in different fields in PV, including system monitoring, silicon

solar cell and module technology, characterization and simulation. He teaches photovoltaics at the University of Freiburg, Freiburg im Breisgau, Germany. His research interests include development of advanced silicon solar cell technology and its transfer to industrial production.



Technology.



Anna Münzer was born in Stuttgart, Germany, in 1991. She received the B.Sc. degree in physics from the University of Tübingen, Tübingen, Germany, with a focus on semiconductor physics, in 2015. Afterwards, she studied renewable energy engineering and management at the University of Freiburg.

Since 2018, she has been an Engineer with the Fraunhofer ISE, Freiburg im Breisgau, where she is currently working on TLS process development in the research on shingled solar cells.