Research Article

In Situ LID and Regeneration of PERC Solar Cells from Different Positions of a B-Doped Cz-Si Ingot

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In order to investigate the light-induced-degradation (LID) and regeneration of industrial PERC solar cells made from different positions of silicon wafers in a silicon ingot, five groups of silicon wafers were cut from a commercial solar-grade boron-doped Czochralski silicon (Cz-Si) ingot from top to bottom with a certain distance and made into PERC solar cells by using the standard industrial process after measuring lifetimes of minority carriers and concentrations of boron, oxygen, carbon, and transition metal impurities. Then, the changes of their I-V characteristic parameters (efficiency \( \eta \), open-circuit voltage \( V_{oc} \), short-circuit current \( I_{sc} \), and fill factor FF) with time were in situ measured by using a solar cell I-V tester during the 1st LID (45°C, 1 sun, 12 h), regeneration (100°C, 1 sun, 24 h), and 2nd LID (45°C, 1 sun, 12 h). The results show that the LID and regeneration of the PERC solar cells are caused by the transition of B-O defects playing a dominant role together with the dissociation of Fe-B pairs playing a secondary role. The decay of \( \eta \) during the 1st LID is caused by the degradation of \( V_{oc} \), \( I_{sc} \), and FF, while the increase of \( \eta \) during the regeneration is mainly contributed by \( V_{oc} \) and FF, and the decay of \( \eta \) during the 2nd LID is mainly induced by the degradation of \( I_{sc} \). After regeneration, the decay rate of \( \eta \) reduces from 4.43%–5.56% (relative) during the 1st LID to 0.33%–1.75% (relative) during the 2nd LID.

1. Introduction

Although boron-doped (B-doped) Czochralski silicon (Cz-Si) solar cells possess the advantages of low cost, high efficiency, and mature production technology, they also have a problem of light-induced degradation (LID) which restricts their development. It is generally accepted that boron-oxygen (B-O) defects formed under illumination are mainly responsible for the LID in minority carrier lifetime of B-doped Cz-Si wafers thus the electrical performance of solar cells based on the Si wafers [1–3]. Since specific composition of B-O defects is still unknown [1, 2] and B-O defects related LID (BO-LID) can bring about huge losses in power generation to customers [4], the investigations on the LID of boron-doped Cz-Si solar cells and its elimination are not only of great scientific research value but also of great practical significance [4]. In order to suppress BO-LID, many research works have been done. In 2006, Herguth et al. proposed a method called regeneration to alleviate the recombination activity of B-O defects by illumination at elevated temperatures [5]. After regeneration treatment, the electrical performance of B-doped Cz-Si solar cells can nearly be recovered, and especially, they possess an anti-LID ability under solar cell working conditions. Thereafter, effectiveness of the regeneration reaction of B-O defects has been confirmed in B-doped Cz-Si wafers [1, 2] and even in p-type compensated Cz-Si solar
cells [6], and the regeneration treatment has been regarded as an effective way to suppress or eliminate the LID of B-doped Cz-Si solar cells [4].

However, most experiments on BO-LID and regeneration effect were conducted on lifetime samples. In contrast, there are few reports on the LID and regeneration of B-doped Cz-Si solar cells especially the new type PERC solar cells. Just as Basnyat et al. from the National Renewable Energy Laboratory (NREL) noted in their paper in 2015, "although a great deal of information is available on B-O effects in silicon, LID in solar cell has not been understood fully. LID effect on crystalline silicon solar cells is studied by a small number of research groups, using comparatively small sets of samples. Even the results reported in literature show large diversity" [7]. Different from the structures of the lifetime samples, Cz-Si solar cells have multiple functional layers and interfaces, therefore, to what extent can the findings obtained on the lifetime samples be applied to Cz-Si solar cells remains unknown [8]; thus, it is necessary to directly study the LID and regeneration of B-doped Cz-Si solar cells especially the PERC solar cells. Herguth et al. and Fertig et al. [9, 10] first reported the LID and regeneration of the PERC solar cells in 2015. However, their research results were based on the prototype PERC solar cells prepared in laboratory, which cannot represent those of real industrialized PERC solar cells [11]. In 2018, Modanese et al. studied the effect of Cu-LID on the PERC solar cells and found that copper contamination and silicon wafer quality are the key factors affecting the degree of Cu-LID [12]. Herguth et al. investigated a LID phenomenon of unknown source in PERC solar cells in 2018, which is characterized by a slight loss in $J_{sc}$, a strong loss in $V_{oc}$ and a great loss in FF and efficiency. It was found that degradation of front contact resistance results in significant degradation of series resistance, and back-surface-related degradation can give a consistent explanation on the experimental results [13]. In 2019, Fertig et al. reported that regenerated PERC solar cells suffer a severe LID of 19.1% (relative value) after a prolonged dark annealing (150 °C, 552 h); in contrast, the measured LID was only 5.6% (relative value) for the PERC solar cells fabricated without regeneration treatment [14]. In 2019, Helmich et al. reported that the regeneration rate constant of the B-O defects in PERC solar cells increases proportionally with the excess carrier concentration [15].

Although several research works have been done on the LID and regeneration of the PERC solar cells, much more research works are still needed to thoroughly understand the LID and regeneration of B-doped Cz-Si solar cells especially the PERC solar cells. Different from previous studies, this paper focuses on in situ investigation on the LID and regeneration of industrialized PERC solar cells fabricated by using the silicon wafers from different positions of a commercial B-doped Cz-Si ingot. The research results are helpful to clarify the effects of the silicon wafer positions and resultant impurity concentration difference on the LID and regeneration of the industrialized PERC solar cells.

2. Experiment

Five groups of 156.75 mm × 156.75 mm pseudosquare silicon wafers were cut from a commercial solar-grade B-doped Cz-Si ingot with a certain distance and were numbered group 1 to group 5 in order from the top to bottom. For each group of silicon wafers, the concentrations of transition metal impurities (iron, copper, and nickel) were measured by inductively coupled plasma mass spectrometer (ICP-MS), and the concentrations of interstitial oxygen and carbon were measured on 2 mm thick neighboring silicon wafers by a Fourier transform infrared (FTIR) spectrometer at room temperature according to ASTM F 121:1983 using a calibration factor of $2.45 \times 10^{15}$, while substitutional boron content was indirectly determined by measuring the resistivities of the wafers by using a four-probe tester after killing thermal donors using 650 °C annealing in Ar for 2 h. After removing about 8 μm damaged layer on each side by using 20% KOH solution at 80 °C and thermally growing about 5 nm SiO$_2$ film on each side at 760 °C, the minority carrier lifetimes of the wafers were measured by a SemiLab WT-2000 lifetime tester. Then, five groups of silicon wafers were made into PERC solar cells by using the standard industrialized process, which includes removing the damaged layer and texturing by KOH solution, POCl$_3$ diffusion at 850 °C to form an emitter about 85 Ω/□, removing phosphorosilicate glass (PSG) and back junction by HF/HNO$_3$ solution, depositing about 20 nm AlO$_x$/140 nm SiN$_x$ passivation layer on the rear side by PECVD at 400 °C, depositing about 80 nm SiN$_x$ antireflection film on the front side by PECVD at 450 °C, laser grooving on the back passivation layer to form electrical contact windows, screen printing the front and back electrodes, and final sintering with temperature up to 800 °C. It should be mentioned that no regeneration treatment was performed on the as-prepared PERC solar cells, and the efficiencies of the PERC solar cells (cell area = 244.67 cm$^2$) were in the range of 20.3% to 20.8% at the STC condition (AM1.5 spectra, 1000 W/m$^2$, 25 °C). It should be emphasized that the measured efficiencies are lower than the real values because the solar simulator was calibrated to output 1 sun light intensity but the shading loss caused by the five-row-probe cartridge above the tested solar cell sample was not taken into account.

Finally, the as-prepared PERC solar cells were treated by the first 12-hour LID (45 °C, 1 sun), 24-hour regeneration (100 °C, 1 sun), and the second 12-hour LID (45 °C, 1 sun) in sequence on a IVT VS-6821M solar cell I-V tester. At the four time nodes (before the 1st LID, after the 1st LID, after regeneration, and after 2nd LID), we performed full-area LBIC scan on the PERC solar cells using 2 μm step by a SemiLab WT-2000 lifetime tester at room temperature. However, the LBIC mapping results are not shown in this paper as some of LBIC maps contain a large proportion of blank areas formed by void data points which may result from limitations of the instrument itself. Although the results obtained from LBIC scan including two-dimensional LBIC (light beam-induced current) mapping diagrams and DL (diffusion length) distribution diagram are unreliable and invalid and cannot be used for explaining our experimental results, LBIC scan step offered the conditions for formation of Fe-B pairs as the process was carried out in a dark box for about 2.5 h. Since the $I$-$V$ characteristic
parameters (efficiency \(\eta\), short-circuit current \(I_{sc}\), open-circuit voltage \(V_{oc}\), and fill factor FF) of the PERC solar cells were in situ measured by the VS-6821M solar cell I-V tester at regular time intervals (10 s in the first hour, 1 min from the 1st hour to the 3rd hour, and 5 min after the 3rd hour) during the 1st LID, regeneration, and 2nd LID processes, the in situ LID and regeneration curves of the I-V characteristic parameters of the PERC solar cells were obtained. According to the claim of the manufacturer (IVT Corporation of Singapore) of the VS-6821M I-V tester, the uncertainties of measurement results of \(I_{sc}\), \(V_{oc}\), FF, and \(\eta\) in the confidence level of 95.4% are 4.99%, 0.51%, 0.48%, and 5.00%, respectively.

3. Basic Knowledge about Solar Cell Device Physics

To facilitate discussion, we give a brief review on fundamentals of solar cell device physics. According to single diode model, the I-V characteristic equation of a solar cell can be expressed as [16]

\[ I = I_{ph} - I_D - I_{sh} = I_{ph} - I_0 \left( e^{\frac{V + IR_s}{R_{sh}}} - 1 \right) - \frac{V + IR_s}{R_{sh}}, \tag{1} \]

where \(I\) and \(V\) are the current and voltage on the load, respectively; \(I_{ph}\) is the photogenerated current; \(I_D\) is the leakage (or dark) current through the diode; \(I_{sh}\) is the shunt current through the shunt resistance \(R_{sh}\); \(I_0\) is the reverse saturation current of the diode in the dark; \(q\) is the elementary charge; \(R_s\) is the series resistance; \(n\) is the ideality factor of the diode; \(k\) is the Boltzmann constant; and \(T\) is the device temperature. For a practical solar cell with finite dimensions, \(I_0\) can be written as [17]

\[ I_0 = qA \frac{n_i^2}{N_D} \left( \frac{D_p}{n_iN_D} \cdot F_N + \frac{D_n}{n_iN_A} \cdot F_p \right) \]

\[ = qA N_c N_v \left( \frac{D_p}{N_D} \cdot F_N + \frac{1}{N_A} \cdot F_p \right) e^{-\left(\frac{E_gkT}{q}\right)} \tag{2} \]

where

\[ F_N = \frac{S_p \cdot \cosh \left( \frac{W_N}{L_p} \right) + (D_p/L_p) \cdot \sinh \left( \frac{W_N}{L_p} \right)}{(D_p/L_p) \cdot \cosh \left( \frac{W_N}{L_p} \right) + S_p \cdot \sinh \left( \frac{W_N}{L_p} \right)} \]

\[ F_p = \frac{S_n \cdot \cosh \left( \frac{W_p}{L_n} \right) + (D_n/L_n) \cdot \sinh \left( \frac{W_p}{L_n} \right)}{(D_n/L_n) \cdot \cosh \left( \frac{W_p}{L_n} \right) + S_n \cdot \sinh \left( \frac{W_p}{L_n} \right)} \tag{3} \]

where \(N_c\) and \(N_v\) are the effective density of states in the conduction band and valence band, respectively; \(\tau_p\) and \(\tau_n\) are the lifetimes of holes and electrons as minority carriers, respectively; \(S_p\) and \(S_n\) are the surface recombination velocities of the \(n\) region and \(p\) region, respectively; and \(W_N\) and \(W_p\) are the thickness of the \(n\) region and \(p\) region, respectively. Obviously, for our case, the reverse saturation current \(I_0\) is mainly determined by the properties of silicon, the cross-sectional area of the \(p-n\) junction, thickness, surface recombination velocities, crystalline quality \((\tau_p, \tau_n, I_{ph}, I_{sh})\), and doping concentration \((N_D, N_A)\) of the \(n\) region and \(p\) region as well as temperature.

When a solar cell is at the short-circuit state, \(V = 0\) and \(I = I_{sc}\), the dark current \(I_D\) through the diode is very low and can be neglected; from equation (1), we have

\[ I_{sc} = \frac{I_{ph}}{1 + (R_s/R_{sh})}. \tag{4} \]

For the general case, \(R_{sh} > R_s\); thus, \(I_{sc} \approx I_{ph}\) holds.

When a solar cell is at the open-circuit state, \(I = 0\) and \(V = V_{oc}\), assuming that the \(R_s\) is large enough so that the \(I_{sh}\) can be neglected, from equation (1), we have

\[ V_{oc} = \frac{nRV_s}{q} \quad \text{ln} \left( \frac{I_{ph}}{I_{0}} + 1 \right). \tag{5} \]

Fill factor FF is defined as the ratio of maximum power \(P_m\) output by a solar cell to the product of \(V_{oc}\) and \(I_{sc}\), i.e.,

\[ FF = \frac{P_m}{V_{oc} I_{sc}}. \tag{6} \]

For an ideal solar cell in which the effect of \(R_s\) and \(R_{sh}\) can be neglected, its fill factor FF can be expressed as [17, 18]

\[ FF_0 = \frac{V_{oc} - \text{ln} \left( \frac{V_{oc} + 0.72}{V_{oc} + 1} \right)}{V_{oc} + 1}, \tag{7} \]

where \(V_{oc}\) is the dimensionless voltage \(V_{oc} = qV_{oc}/nkT\). Equation (7) shows that the ideal fill factor \(FF_0\) increases with the \(V_{oc}\). After considering the effect of \(R_s\) and \(R_{sh}\), the FF of a \(p-n\) junction solar cell can be given by [17, 18]

\[ FF = FF_0 \left[ 1 - \left( \frac{V_{oc} + 0.7}{V_{oc} + 1} \right)^{r_s \cdot r_{sh}} \right]. \tag{8} \]

Here,

\[ FF_c = \frac{FF_0 \left[ 1 - (1.1R_s) + \left( \frac{r_s^2}{5.4} \right) \right]}{1}, \tag{9} \]

where \(FF_c\) is the fill factor of a solar cell in which only the effect of \(R_s\) is considered and \(r_s\) and \(r_{sh}\) are the normalized resistances, which are equal to \(R_s\) and \(R_{sh}\) divided by \(V_{oc}/I_{sc}\), respectively. From equations (7)–(9), we can see that FF is affected by multiple factors such as \(R_s\), \(R_{sh}\), \(V_{oc}\), \(n\), \(I_{0}\), and \(I_{sc}\). 


in a complex way. To improve FF, one can decrease $R_s$ and $I_0$ and increase $R_{sh}$, $V_{oc}$, and $I_{sc}$ meanwhile.

Efficiency $\eta$ is defined as the ratio of maximum power $P_m$ output by a solar cell to the incident light power $P_{in}$ on the device, i.e.,

$$\eta = \frac{P_m}{P_{in}} = \frac{V_{oc} \cdot I_{sc} \cdot FF}{P_{in}}. \quad (10)$$

Equation (10) shows that the $\eta$ is determined by the product of $V_{oc}, I_{sc}$, and FF for the given incident light intensity and area of a solar cell.

4. Results and Discussion

4.1. Impurity Concentration and Lifetime of Silicon Wafers.

Table 1 shows the measured lifetimes and resistivities as well as boron (B), oxygen (O), carbon (C), iron (Fe), copper (Cu), and nickel (Ni) contents of the five groups of silicon wafers. As shown in Table 1, from the top to bottom of the Cz-Si ingot, O content gradually decreases while B, C, Fe, Cu, and Ni contents gradually increase. With the increase of B content, the lifetime and resistivity gradually decrease. It should be mentioned that O content is approximately one order of magnitude higher than C content, two orders of magnitude higher than B content, three orders of magnitude higher than Fe content, and four orders of magnitude higher than Cu and Ni contents.

For the impurities such as B, C, Fe, Cu, and Ni, their distributions along the solidifying direction of the Cz-Si ingot are basically determined by their segregation coefficients in silicon; herein, the segregation coefficient is defined as the ratio of the solubility of the impurity in the solid phase at the liquid-solid-interface to the solubility in the liquid phase [19]. Since B, C, Fe, Cu, and Ni impurities all possess segregation coefficients less than 1 in silicon [19, 20], their contents gradually increase towards the tail of the Cz-Si ingot. For O impurity, its distribution along the solidifying direction of the Cz-Si ingot is determined not only by its segregation coefficient in silicon but also by the contact area between the quartz crucible and the silicon melt, and the latter may play a more important role because O impurities in the Cz-Si ingot mainly come from the dissolving quartz crucible wall under corrosion of the silicon melt at high temperature. As the growth proceeds, the contact area between the quartz crucible and the silicon melt is gradually reduced, which results in decreasing O incorporation in the silicon melt and thus decreasing O content towards the tail of the ingot [21]. Furthermore, this distribution is enhanced by the segregation coefficient of O which is larger than 1 [19]. Obviously, our test results on concentrations of the impurities can be given a good explanation from the point of view of impurity segregation and major origin of O. As compared with impurity distribution reported in Cz-Si, our O distribution along Cz-Si ingot is consistent with ref. [21] and our transition metal impurity distributions are in good agreement with ref. [22]. Since different groups of the Si wafers show little difference in C content and C content has little effect on lifetime of Cz-Si wafers [23], we will neglect the influence of C content on the LID and regeneration of the PERC solar cells in the following.

4.2. PERC Solar Cells.

Figure 1 shows the in situ changes of the I-V characteristic parameters of five groups of PERC solar cells with time during the 1st LID (45°C, 1 sun, 12 h). As shown in Figure 1, $\eta$, $I_{ph}$, $V_{oc}$, and FF of most samples exhibit the typical characteristics of BO-LID, which is characterized by a double exponential decay corresponding to a “fast decay” and a “slow decay,” respectively [2, 24]. Furthermore, P3 has the highest $\eta$ and $V_{oc}$, whereas the bottom samples (P4 and P5) have larger LID in $V_{oc}$.

Under the condition of the 1st LID, the changes of $R_s$, $R_{sh}$, and surface passivation can be neglected, therefore, the decays of I-V characteristic parameters of the PERC solar cells are mainly caused by the LID of bulk minority carrier lifetime $\tau$. For the as-prepared PERC solar cells, $R_s$ is about 0.2 $\Omega \cdot \text{cm}^2$ while $R_{sh}$ is above 2000 $\Omega \cdot \text{cm}^2$, which means $R_{sh} >> R_s$, thus, $I_{sc} = I_{ph}$ holds according to equation (4). It can be seen from equations (2)-(5) that the LID of the $\tau$ can result in the decrease of the collection efficiency thus decay of $I_{ph}$ or $I_{oc}$ on the one hand [25] and the increase of dark saturation current $I_0$ thus the decay of $V_{oc}$ on the other hand. Furthermore, the decay of $I_{ph}$ can also induce the degradation of $V_{oc}$ according to equation (5). It is easily known from equations (7)-(9) that the decay of FF is mainly determined by that of $V_{oc}$ when the changes of $R_s$ and $R_{sh}$ are neglected. From equation (10), the decay of $\eta$ is the product of the decays of $V_{oc}$, $I_{sc}$, and FF. In a word, the degradations of $V_{oc}$, $I_{sc}$, FF, and $\eta$ during the 1st LID result from the LID of $\tau$ and degradation of $\tau$ under LID conditions has been reported by ref. [1, 3]. The larger LID in $V_{oc}$ of bottom samples (P4 and P5) may result from their higher B and Fe contents. Ref. [26] has shown that the higher B content means the higher B-O defect concentration when the O content is in a range of 5.0 × 10^{17} \text{cm}^{-3} to 8.0 \times 10^{17} \text{cm}^{-3}. In addition, the Coulomb attraction between Fe$^+$ and B reduces the efficiency of phosphorus diffusion gettering (PDG) [27], resulting in higher Fe-B pair concentration in the samples with higher B and Fe content, and the dissociation of Fe-B pairs would result in LID of most I-V characteristic parameters of a solar cell [28]. P3 has the highest $\eta$ and $V_{oc}$, which may be related to its moderate B and O content and lower transition metal impurity content.

Figure 2 shows the in situ changes of the I-V characteristic parameters of five groups of PERC solar cells with time during the regeneration (100°C, 1 sun, 24 h). As shown in Figure 2, the $\eta$ of P1-P5, $V_{oc}$ of P3-P5, and FF of P2-P4 first decay then rise and finally reach saturation, with the extent of the rise much larger than the extent of the decay. Furthermore, the decay and rise extent in $\eta$, $I_{sc}$, and $V_{oc}$ as well as the time required to reach saturation basically increases with B content. Different from above regeneration curves, the $V_{oc}$ of P1-P2 and FF of P1 first rise linearly and then tend to reach saturation. In addition, the $I_{sc}$ of P1-P3 first decreases, then increases, and finally decreases slowly, while the $I_{sc}$ of P4-P5 first decreases, then increases, and finally tends to reach saturation.

We note that FF shows similar regeneration curves to $V_{oc}$, which implies that under our regeneration condition,
the change of FF is basically determined by that of $V_{oc}$, and the changes of $R_s$ and $R_{sh}$ can be neglected. In addition, $\eta$ also shows a similar regeneration curves to $V_{oc}$, indicating that the change of $\eta$ is also dominated by that of $V_{oc}$. In contrast, $I_{sc}$ shows different regeneration curves from those of $\eta$, $V_{oc}$, and FF, with decreased $I_{sc}$ values after the regeneration except P5. The reason why $I_{sc}$ shows very different regeneration curves from $V_{oc}$ is as follows: (1) the lifetimes limited by B-O defects or Fe-B pairs (or interstitial Fe ion $Fe^{i+}$) are both injection dependent; however, $V_{oc}$ and $I_{sc}$ are measured
at very different excess carrier concentrations ($\Delta n$), which results in the very different behavior of $V_{oc}$ and $I_{sc}$ [29]. Taking the dissociation of Fe-B pairs as an example, the recombination activities of a Fe-B pair and Fe$_i^+$ change with injection level of minority carriers. When the injection level is lower than excess carrier concentration corresponding to the crossover point ($\Delta n_{COP}$), the recombination activity of a Fe-B pair is lower than that of Fe$_i^+$, which means that dissociation of a Fe-B pair will result in LID of minority carrier lifetime $\tau$; when the injection level is higher than $\Delta n_{COP}$, the recombination activity of a Fe-B pair is higher than that of a Fe$_i^+$, which means that dissociation of a Fe-B pair will give rise to photo-induced enhancement of $\tau$ [29]. The $I_{sc}$ measured at injection levels always below the $\Delta n_{COP}$ would degrade when FeB pairs dissociate, while the $V_{oc}$ may decrease, or remain constant, or even increase, depending on the position of excess carrier concentration under open-circuit condition relative to $\Delta n_{COP}$ [29]. (2) $V_{oc}$ and $I_{sc}$ have different dependencies on $\tau$. Specifically speaking, $I_{sc}$ is proportional to $\tau^{1/2}$, while $V_{oc}$ is proportional to $ln \tau$. That is to say, with the decrease of $\tau$, $I_{sc}$ decreases more remarkably than $V_{oc}$. In other words, $I_{sc}$ is more sensitive to the change of $\tau$ than $V_{oc}$. (3) The further slow decay after the slow rise in $I_{sc}$ of P1-P3 may result from the degradation of PECVD AlO$_x$/SiNx rear passivation [13]. However, no further decay occurs in $I_{sc}$ of P4-P5, which may be due to the higher B-O defect concentration of P4-P5 and the positive effect of B-O defect regeneration reaction on $\tau$ compensating the negative effect of back passivation degradation in P4-P5.

Figure 2: In situ variation of $I$-$V$ characteristic parameters of five groups of PERC solar cells with time during the regeneration (100°C, 1 sun, 24 h).
Interestingly, we note that at the initial stage of regeneration, the $V_{oc}$ of P1-P2 increases directly, while the $V_{oc}$ of P3-P5 first decreases and then increases, and the degradation and regeneration extents increase with increasing B and Fe contents. In order to explain the experimental phenomenon, we explore the three possibilities (BO-LID, LeTID, and dissociation of Fe-B pairs): (1) the degradation of $V_{oc}$ at the initial stage of regeneration cannot be caused by B-O defects. After the 1st LID treatment (1 sun, 45°C, and 12 h), the B-O defects in PERC solar cells exist mainly in the form of degradation state. No dark annealing treatment above 100°C was performed on the samples before the regeneration, no B-O defects in annealed state were formed [30], and no BO-LID would occur at the initial stage of regeneration. (2) LeTID (Light and elevated Temperature-Induced Degradation) was also not involved in the degradation of $V_{oc}$ at the initial stage of regeneration. Although there is still controversy about whether LeTID exists in B-doped Cz-Si [31, 32], the possibility of occurrence of LeTID in our samples has to be carefully examined. By using the method given by ref. [33] for judging whether the as-used illuminating annealing conditions can activate LeTID defects, we measured the IDLS (Injection-Dependent Lifetime Spectra) curves of five groups of lifetime samples made from same position of silicon wafers as five groups of PERC solar cells by using a WCT-120 equipment after sintering, dark annealing at 200°C for 30 min, and regeneration at 100°C and 1 sun for 24 h. It should be mentioned that the lifetime samples were prepared by the fabrication process of PERC solar cells without the steps of phosphorus diffusion, removing PSG, laser grooving on the back passivation layer, and screen printing front and back electrodes. In order to avoid the influence of Fe$^+$ on the measurement results, all the samples were kept in the dark at room temperature for 3 h before each of the IDLS measurements to make most of Fe$^+$ in the samples change into Fe-B pairs. The measurement results are shown in Figure 3. It can be seen from Figure 3 that the lifetimes of five groups of lifetime samples after the regeneration are all close to those after sintering and dark annealing at 200°C for 30 min, or even slightly higher than the latter two. This result illustrates that the LeTID defects generated in the Cz-Si samples by our regeneration treatment (100°C, 1 sun, and 24 h) can be neglected. (3) The degradation of $V_{oc}$ at the initial stage of regeneration can be attributed to the dissociation of Fe-B pairs. According to ref. [28], the dissociation of Fe-B pairs into Fe$^+$ under illumination would lead to a degradation in most $I$-$V$ characteristic parameters of crystalline silicon solar cells. In particular, $I_{sc}$ and $\eta$ always show a pronounced degradation, whereas the $V_{oc}$ either decreases or keeps constant or even increases depending on Fe contamination levels. Our observed experimental phenomenon at the initial stage of regeneration can be well explained by ref. [28]. Specifically speaking, for P1-P2 with lower Fe content, $V_{oc}$ keeps constant, while $\Delta\eta$, $I_{sc}$, and FF degrade. As a comparison, for P3-P5 with higher Fe content, all the $I$-$V$ characteristic parameters degrade. In a word, the variation of $I$-$V$ characteristic parameters of P1-P5 at the initial stage of regeneration can be well explained by the dissociation of Fe-B pairs. In addition, Fe-B pairs formed during 2.5 h LBIC scan before regeneration, together with the experimental results of the degradation extent increasing with B and Fe contents, further justify our judgement.

Figure 4 shows the in situ variation of the $I$-$V$ characteristic parameters of five groups of PERC solar cells with time during the 2nd LID (45°C, 1 sun, 12 h). As shown in Figure 4, the $\eta$ and $I_{sc}$ of all the samples suffer a certain degradation at the early stage of the 2nd LID. In contrast, $V_{oc}$ and FF of all the samples excluding the $V_{oc}$ of P2 basically remain stable throughout the 2nd LID, and the $V_{oc}$ of P5 as well as FF of P4 and P5 even show an upward trend.

At the early stage of the 2nd LID, the $I_{sc}$ and $\eta$ of all the samples suffer a degradation, while the $V_{oc}$ either decreases or keeps constant or increases. After that, the $I$-$V$ characteristic parameters of all the samples basically keep stable. Obviously, our experimental results observed at the early stage of the 2nd LID are in good accordance with the change features of the $I$-$V$ characteristic parameters induced by dissociation of Fe-B pairs [28]. Since $I_{sc}$ and $V_{oc}$ are measured at very different excess carrier concentrations, the dissociation of Fe-B pairs always results in a pronounced degradation in $I_{sc}$ and $\eta$, but different changes in $V_{oc}$, which either decreases or keeps constant or even increases, depending on the position of excess carrier concentration relative to $\Delta n_{COP}$ [28]. During the 1st LID, the degradation of all the $I$-$V$ characteristic parameters occur, which means that even the excess carrier concentration under open-circuit condition is still below the $\Delta n_{COP}$ for each sample. However, the situation for the 2nd LID totally changes. Before the 2nd LID, the minority carrier lifetimes of all the samples were improved by regeneration treatment, and Fe-B pairs were formed during 2.5 h LBIC scan. According to the calculation formula of excess carrier concentration $\Delta n = G \cdot \tau$ and constant generation rate $G$ corresponding to 1 sun light intensity for our case, the improved minority carrier lifetime $\tau$ means the increased $\Delta n$, and the increased $\Delta n$ may approach to or even surpass the $\Delta n_{COP}$, leading to a constant $V_{oc}$ or even increased $V_{oc}$ due to dissociation of Fe-B pairs. Specifically speaking, the decreased, constant, and increased $V_{oc}$ of P2, P3, and P5 indicate that the $\Delta n$ of P2, P3, and P5 are below, near, and above the $\Delta n_{COP}$, respectively. In a word, the changes of $I$-$V$ characteristic parameters of the PERC solar cells during the early stage of the 2nd LID can be given a good explanation by the dissociation of Fe-B pairs. It should be mentioned that no BO-LID would occur under the 2nd LID condition, because most of B-O defects in PERC solar cells have been converted into the stable regenerated state after the regeneration.

Figure 5 shows the relative changes of $I$-$V$ characteristic parameters of five groups of PERC solar cells before and after each process. As shown in Figure 5, for all the samples the decrease of $\eta$ during the 1st LID results from the decrease of $V_{oc}$, $I_{sc}$, and FF, while the increase of $\eta$ during the regeneration is mainly contributed by $V_{oc}$ and FF, and the decrease of $\eta$ during the 2nd LID is mainly caused by the decrease of $I_{sc}$. During the 1st LID, the PERC solar cells made from top and bottom silicon wafers have larger decay in $\eta$ and $V_{oc}$ than those made from middle silicon wafers, and the decay rate of $\eta$ is in the range of 4.43%–5.56% (relative). After the regeneration, all the samples have an increase in $\eta$, $V_{oc}$, and FF but
suffer a decay in $I_{sc}$ excluding P5. Furthermore, the rate of rise of $\eta$ increases with B content, and it ranges from 5.71% to 9.26% (relative). In addition, generally, the higher the decay rates in $V_{oc}$ and FF during the 1st LID, the higher the increase rates in $V_{oc}$ and FF during the regeneration, which can be given a good explanation from the regeneration of B-O defects. During the 2nd LID, all the samples except P2 show a good stability in $V_{oc}$ and FF but suffer a decay in $\eta$ and $I_{sc}$. Moreover, the decay rates of $\eta$ reduce to 0.33%–1.75% (relative). In addition, the relative change rates of $V_{oc}$ and FF during the 2nd LID also decrease, which are two and one order of magnitude lower than those during the 1st LID, respectively.

Because all the Cz-Si wafers were made into PERC solar cells by using the same standard industrialized process, the difference in LID and regeneration of P1-P5 should be mainly caused by the difference in impurity concentration in as-used Cz-Si wafers. In fact, it can be seen from the impurity content that B-O defects and Fe-B pairs (or Fe$_3^+$) should be responsible for the observed LID and regeneration because of significantly higher O, B, and Fe contents in the base of as-prepared PERC solar cells. The reasons are as follows: (1) for all the Cz-Si wafers we used, the contents of B and O are far larger than those of transition metal impurities. As shown in Table 1, O content is about two orders of magnitude higher than B content, while the B content is at least one order of magnitude higher than Fe content and two orders of magnitude higher than Cu and Ni contents. It should be emphasized that O and B contents listed in Table 1 just correspond to the concentrations of interstitial O and substitutional B in the silicon wafers which are much higher than those shown in as-used Cz-Si wafers, which are much higher than those of Fe, Cu, and Ni impurities with electrical activity. Therefore, the difference between the contents of B and O and those of transition metal impurities are much larger than those shown in

![Diagram showing IDLS spectra of five groups of lifetime samples corresponding to five groups of PERC solar cells measured after sintering, dark annealing at 200°C for 30 min, and regeneration at 100°C and 1 sun for 24 h.](image-url)

**Figure 3:** The IDLS spectra of five groups of lifetime samples corresponding to five groups of PERC solar cells measured after sintering, dark annealing at 200°C for 30 min, and regeneration at 100°C and 1 sun for 24 h.
After the phosphorus diffusion gettering (PDG) step included in the fabrication process of PERC solar cells, nearly all of Cu and Ni impurities can be removed from the silicon wafers excluding Fe impurities. The phosphorus diffusion condition we used is diffusion at 850°C for about 30 min, drive-in at 870°C for about 15 min, and total time of about 1.5 h for silicon wafers staying above 800°C. According to the empirical formulas given by ref. [34], we calculated the diffusion coefficients and the solubilities of Cu, Ni, and Fe impurities in silicon at 800°C, which are $4.49 \times 10^{-5}$ cm²/s, $1.24 \times 10^{-5}$ cm²/s, and $8.32 \times 10^{-7}$ cm²/s and $5.54 \times 10^{16}$ atoms/cm³, $1.58 \times 10^{16}$ atoms/cm³, and $2.84 \times 10^{12}$ atoms/cm³, respectively. As compared with the measured concentrations of Cu, Ni, and Fe (<3.94 × 10¹¹, <1.68 × 10¹³, and >3.25 × 10¹⁴ atoms/cm³ for Cu, Ni, and Fe, respectively), Cu and Ni contents are much lower than their solubilities whereas Fe content is much higher than its solubility in silicon at 800°C, which means that Cu and Ni impurities can be completely dissolved while Fe impurity can only be partially dissolved in the silicon wafers at 800°C. Thus, Cu and Ni impurities can be effectively removed by PDG step due to their high diffusion coefficients and PDG gettering efficiency [27]. In contrast, partial dissolution and lower diffusion coefficient of Fe impurity together with lower gettering efficiency caused by Coulomb attraction between Fe⁺ and B⁺ result in a part of Fe impurities remaining in the samples especially with higher B and Fe contents after PDG, which may exist in forms of Fe⁺, Fe-B pairs, Fe precipitation, etc. The recombination activity of Fe precipitation can be neglected as compared with that of Fe⁺ and Fe-B.
pairs, if they are formed by the same amount of Fe impurities [25]. (3) After the PDG and hydrogen passivation (sintering with SiNx:H and AlOx:H/SiNx:H passivation layer) steps included in the fabrication process of PERC solar cells, the base minority carrier lifetime of as-prepared PERC solar cells can be significantly improved, which means that most of the defects caused by transition metal impurities can be effectively removed by PDG or passivated by hydrogen [25, 27]. However, whether Fe⁺⁺ or Fe-B can be effectively passivated by hydrogen remains controversial up to date [25].

In conclusion, the LID and regeneration of as-prepared PERC solar cells are caused by B-O defects and dissociation of Fe-B pairs. During the 1st LID, the decay of $V_{oc}$, $I_{sc}$, FF, and $\eta$ results from B-O defects and dissociation of Fe-B pairs. The double exponential decay of $I-V$ characteristic parameters of most samples suggests the dominance of BO-LID. During the regeneration, the initial change of the $I-V$ characteristic parameters of the samples can be attributed to dissociation of Fe-B pairs, while the subsequent rise and saturation of the $I-V$ characteristic parameters are caused by the regeneration of B-O defects. The extent of rise is much larger than the extent of decay, which indicates the dominant role of B-O defects. In addition, the further slow decay after the slow rise in $I_{sc}$ of P1-P3 is ascribed to the slight rear passivation degradation. During the 2nd LID, the initial change of the $I-V$ characteristic parameters of the samples is induced by the dissociation of Fe-B pairs. By comparing the 1st LID and 2nd LID, we can see that the extent of degradation caused by B-O defects is much larger than that by dissociation of Fe-B pairs. In addition, although P5 was fabricated by using the wafer with the highest B and Fe content, it nearly shows the largest decay and rise in $\eta$, $V_{oc}$, and $I_{sc}$ during the 1st LID and the regeneration and minimum decay and highest value in $\eta$ during the 2nd LID. This result further indicates that B-O defects dominate the LID and regeneration of as-prepared PERC solar cells. In a word, the LID and regeneration of the PERC solar cells result from B-O defects playing a dominant role and dissociation of Fe-B pairs playing a secondary role.
5. Conclusion

Five groups of silicon wafers cut from a solar grade Cz-silicon ingot to the top to bottom with a certain distance were made into PERC solar cells by using the standard industrial process; then, the as-prepared PERC solar cells were treated and in situ measured on a solar cell I-V tester by 1st LID (45°C, 1 sun, 12 h), regeneration (100°C, 1 sun, 24 h), and 2nd LID (45°C, 1 sun, 12 h). The results show that the LID and regeneration of I-V characteristic parameters of as-prepared PERC solar cells are mainly caused by B-O defects playing a dominant role and dissociation of Fe-B pairs playing a secondary role. Specifically, during the 1st LID, the decay of \( V_{oc}, I_{sc}, \) FF, and \( \eta \) is induced by B-O defects and dissociation of Fe-B pairs. During the regeneration, the initial change of the electrical parameters of the PERC solar cells can be attributed to dissociation of Fe-B pairs, while the subsequent rise and saturation of the electrical parameters result from the regeneration of B-O defects. During the 2nd LID, the initial change of the electrical parameters of the PERC solar cells is ascribed to dissociation of Fe-B pairs. Moreover, the increase of \( \eta \) during the 1st LID is caused by the degradation of \( V_{oc}, I_{sc}, \) FF, and the increase of \( \eta \) during the regeneration is mainly contributed by \( V_{oc} \) and FF, and the decay of \( \eta \) during the 2nd LID is mainly induced by the decrease of \( I_{sc} \). After regeneration, the decay rate of \( \eta \) reduces from 4.43%–5.56% (relative) during the 1st LID to 0.33%–1.75% (relative) during the 2nd LID.

Data Availability

The data used to support the findings of this study are included within the paper.

Conflicts of Interest

The authors declare no conflict of interests.

Authors’ Contributions

Shuai Yuan and Siqi Ding performed experimental investigation and wrote the original draft; Bin Ai designed the experiments, analyzed the results, and revised the draft; Jiaxing Ye and Depeng Qiu discussed the experimental results; Xueqin Liang, Daming Chen, Jingsheng Jin, and Xiaopu Sun offered lots of help in preparing silicon wafers, lifetime samples, and PERC solar cells, respectively, Shuai Yuan and Siqi Ding are co-first authors.

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