

## Reassessment of the recombination properties of aluminium—oxygen complexes in n- and p-type Czochralski-grown silicon

basic solid state physics

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The recombination parameters of aluminium—oxygen complexes in silicon have been reassessed by applying lifetime spectroscopy on several n- and p-type intentionally Al-contaminated and control samples, using a single-level defect model. The presence of the control samples has allowed greater accuracy for the extraction of the recombination lifetime. The uncertainty ranges of the parameters have been

tightened significantly by simultaneously fitting the lifetime on several samples. The electron/hole capture cross section ratio k was reassessed to be 380, in the uncertainty range of 330–460. A direct comparison of the n- and p-type samples has shown that those complexes are much more recombination-active in p-type silicon than in n-type silicon at low and intermediate injection levels.

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1 Introduction Aluminium is a common metallic impurity in solar-grade silicon materials. Concentrations in the range of  $10^{14}$ – $10^{17}$  cm<sup>-3</sup> are typical in the upgraded metallurgical silicon (UMG-Si) wafers after ingot growth [1]. Al is more difficult to remove through directional solidification than many transition metals, considering that the segregation coefficient of Al is  $k_{\rm Al} = 2 \times 10^{-3}$  [2–4], in comparison with Fe with  $k_{\rm Fe} =$  $6.4 \times 10^{-6}$  [5], Ni with  $k_{\text{Ni}} = 3.2 \times 10^{-5}$  [5, 6], and Cu with  $k_{\text{Cu}} = 8 \times 10^{-4} [5, 7]$ . Also, because of its low diffusivity [8], it is not efficient to remove Al through gettering once present in the wafers. The detrimental effects of Al on the carrier lifetime of Czochralski-grown (Cz) silicon wafers are widely recognised, and numerous authors have attributed the effects to the recombination-active aluminum-oxygen (Al-O) complexes [4, 5, 9-15].

Previous studies on the recombination properties of Al–O complexes have been based on deep-level transient spectroscopy (DLTS) [4, 5, 9, 12] or temperature- and/or injection-dependent lifetime spectroscopy (TIDLS or IDLS) [11, 12]. With DLTS, the energy level can be accurately determined, but generally only the

majority-carrier capture cross section is accessible [16]. In contrast, the lifetime spectroscopy methods are based on the Shockley-Read-Hall (SRH) model [17, 18], and can be used to determine the electron/hole capture cross section ratio k [11, 19]. However, they have a large uncertainty range for the energy level of deep levels, and it is generally not possible to separate the capture cross section from the defect concentration if both are unknown, as this technique only determines their product [11, 19]. Table 1 lists the recombination parameters of Al-O complexes from the literature. The deep level located at 0.38–0.50 eV above the valence band  $E_{\nu}$ , having a most probable value around 0.44 eV, is consistent among the different studies applying DLTS [4, 5, 9, 12]. However, k varies across orders of magnitudes in the literature. Both of the previous lifetime spectroscopy studies assumed a second shallow level to obtain a reasonable fit over the entire injection range [11, 12]. However, the shallow level has not been detected by any DLTS studies, and thus remains unconfirmed. In addition, in the absence of control samples, the authors assumed the effective lifetime to be the bulk SRH recombination lifetime [11], or obtained the SRH lifetime

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| Tahle 1  | Recombination | narameters | of the    | 11_0          | complex  | from | the liter | rature and | found in | this wor  | ŀ  |
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| authors         | $E_t$ – $E_v$ (eV)  | $\sigma_{\rm n}~({\rm cm}^2)$ | $\sigma_{\rm p}~({\rm cm}^2)$   | k              | methods            | refs.  |
|-----------------|---|-------------------------------|---|----------------|--------------------|--------|
| Marchand et al. | $0.397 \pm 0.005$<br>$0.314 \pm 0.002$<br>$0.215 \pm 0.001$ |                               | $(8.6 \pm 0.7) \times 10^{-17}$<br>$(6.6 \pm 0.2) \times 10^{-15}$<br>$(9.2 \pm 0.5) \times 10^{-22}$ |                | DLTS               | [4, 9] |
| Davis et al.    | $0.43 \pm 0.05$<br>$0.47 \pm 0.03$                          | $1.4\times10^{-14}$           | $5.7 \times 10^{-15}$   | 2.5            | DLTS               | [5]    |
| Schmidt         | 0.22–0.68<br>0.14 or 0.98                                   |                               |   | 90             | TIDLS              | [11]   |
| Rosenits et al. | $0.44 \pm 0.02$ $0.15$                                      | $3.1\times10^{-10}$           | $3.6 \times 10^{-13}$   | 870            | DLTS and IDLS IDLS | [12]   |
| this work       |   |                               |   | 380 (-50, +80) | IDLS               |        |

by correcting the measured effective lifetime with a modelled intrinsic recombination lifetime [12]. In principle, this can cause uncertainties in the fitted SRH lifetimes, and may also explain why a second shallow level was required in those studies.

In this work, we conduct a new IDLS analysis using a single-level model applied to several n- and p-type Cz wafers that have been intentionally contaminated with Al and control wafers, to reassess the recombination parameters of the defect. The control samples allow the SRH lifetime to be extracted with greater accuracy. Simultaneously, performing IDLS on several samples helps tighten the uncertainty ranges of the parameters [20], while having both n- and p-type of samples enables the recombination activity of the defect in both types of silicon to be directly compared.

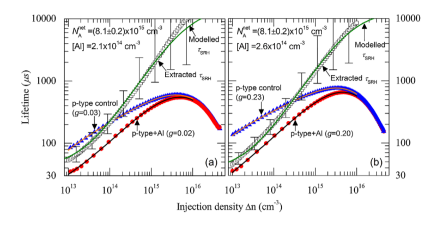
**2 Experimental** The samples investigated in this study were from four B- and P-doped compensated Czochralski (Cz) ingots supplied by Apollon Solar. The ingots were grown from 9N poly silicon feedstock that was intentionally doped with boron and phosphorus to simulate compensated UMG-Si [21]. For each type, one was grown as a control ingot and the other was intentionally contaminated with Al. The concentration of boron in the

starting melt was  $1.32 \times 10^{16} \, \text{cm}^{-3}$  in all four ingots. The concentration of Al ([Al]) in the starting melt was  $1.04 \times 10^{17} \, \mathrm{cm}^{-3}$  in the Al-doped ingots. The concentration of phosphorus in the starting melt of the n- and p-type ingots were  $3.64 \times 10^{16} \,\mathrm{cm}^{-3}$  and  $7.73 \times 10^{15} \,\mathrm{cm}^{-3}$ , respectively. The ingots were cut to wafers, and three p-type and one ntype Al-contaminated wafers and the corresponding control wafers were chosen for this study. Some key parameters of the samples are listed in Table 2. The net doping levels and the [Al] were calculated applying the Scheil's law [22]. The segregation coefficients of B, P and Al were taken as 0.8, 0.35 and 0.002, respectively [2-4]. The resistivities were measured by dark conductance, and were also predicted applying Schindler et al.'s mobility model for compensated silicon, based on the expected dopant concentrations using the Scheil's law [23]. The predicted values mostly lie within 10% of the measured values, as shown in the table. The interstitial oxygen concentrations ([O<sub>i</sub>]) in the central region of the samples were determined by the OXYMAP technique [24].

The samples were chemically etched to remove saw damage, and then went through a phosphorus gettering process at 880 °C for 0.5 h, and 600 °C for about 14 h [25]. Performing the extended heat treatment at a lower temperature (600 °C for 14 h) after the diffusion was found

**Table 2** Parameters of the samples in this study. Note that g represents the solidified fraction.

| samples                     | modelled net doping (cm <sup>-3</sup> ) | modelled [Al] (cm <sup>-3</sup> ) | $\rho$ measured with QSSPC ( $\Omega \cdot \text{cm}$ ) | $\rho$ predicted by Schindler et al.'s mobility model $(\Omega \cdot cm)$ | [O <sub>i</sub> ]<br>(cm <sup>-3</sup> ) |
|-----------------------------|---|-----------------------------------|---|---|--|
| p-type control $(g = 0.03)$ | $7.9 \times 10^{15}$                    | 0                                 | 1.96  | 2.06  | $1.3 \times 10^{18}$                     |
| p-type control $(g = 0.23)$ | $7.9 \times 10^{15}$                    | 0                                 | 1.94  | 2.05  | $1.1 \times 10^{18}$                     |
| p-type control $(g = 0.43)$ | $7.9 \times 10^{15}$                    | 0                                 | 1.99  | 2.06  | $9.9 \times 10^{17}$                     |
| p-type + Al $(g = 0.02)$    | $8.1 \times 10^{15}$                    | $2.1 \times 10^{14}$              | 2.23  | 2.00  | $1.2 \times 10^{18}$                     |
| p-type + Al $(g = 0.20)$    | $8.2 \times 10^{15}$                    | $2.6 \times 10^{14}$              | 2.20  | 1.98  | $1.1 \times 10^{18}$                     |
| p-type + Al $(g = 0.56)$    | $8.3 \times 10^{15}$                    | $4.7\times10^{14}$                | 2.25  | 1.98  | $8.4 \times 10^{17}$                     |
| n-type control $(g = 0.65)$ | $1.2 \times 10^{16}$                    | 0                                 | 0.49  | 0.58  | $8.4 \times 10^{17}$                     |
| n-type + Al ( $g = 0.64$ )  | $1.1 \times 10^{16}$                    | $5.8 \times 10^{14}$              | 0.53  | 0.63  | $8.1 \times 10^{17}$                     |



**Figure 1** Injection-dependent lifetimes of the samples (a) p-type + Al (g = 0.02) and (b) p-type + Al (g = 0.20) and the corresponding control samples. The extracted and the modelled SRH recombination lifetimes are also shown.

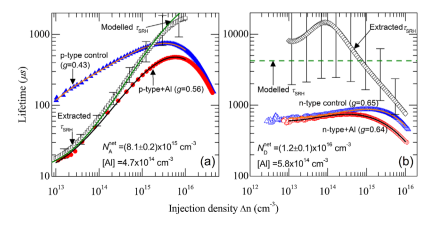
to increase the efficiency in removing fast diffusing metallic impurities [25]. Those impurities were present in all the samples before gettering, resulting in a reduced difference in the lifetimes of the control samples and the Al-contaminated samples. As confirmed by dark conductance and Fourier Transform Infrared (FT-IR) spectrophotometry measurements, the resistivity and [O<sub>i</sub>] remained unchanged, within measurement uncertainty, before and after the gettering step. After gettering, another 10 µm was etched from both sides, followed by surface passivation with plasmaenhanced chemical vapour deposited (PECVD) SiN<sub>x</sub> films. The injection-dependent lifetimes were measured with the quasi-steady-state photoluminescence (QSSPL) method, to avoid trapping effects at low injection levels [26]. The samples were annealed at 260 °C for 30 min in the dark to deactivate the boron-oxygen (BO) complexes before the measurements.

**3 Modelling procedure** To extract the SRH recombination lifetimes  $\tau_{\rm SRH}$  due to Al–O complexes, we determined polynomial fits for the measured lifetimes of the Al-contaminated and control samples, and then calculated  $\tau_{\rm SRH}$  applying  $1/\tau_{\rm SRH} = 1/\tau_{\rm eff} - 1/\tau_{\rm control}$ . The polynomial curves allowed convenient interpolation between the two sets of data points. Only lifetime data at injection levels lower than  $\Delta n = 1 \times 10^{16} \, {\rm cm}^{-3}$  were used

to extract  $\tau_{SRH}$ . At injection levels higher than this, the lifetimes of the control sample and the Al-contaminated sample are almost identical (shown in Section 4). This causes large uncertainties in the extracted  $\tau_{SRH}$ , adversely affecting the subsequent fitting of the recombination parameters.

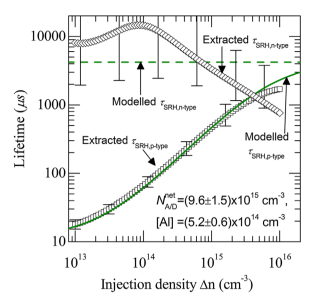
The extracted  $\tau_{\rm SRH}$  of the three p-type Alcontaminated samples were then fitted simultaneously to determine k. As will be discussed in Section 4, there is no significant difference in lifetime between the two n-type samples, which would result in large uncertainties in the extracted  $\tau_{\rm SRH,n-type}$ . So the n-type samples were not included in the fitting but were subsequently used to check for consistency with the parameters extracted from the p-type samples.

The extraction of the SRH lifetime using a control sample has the advantage of removing the effect of surface recombination velocity and intrinsic recombination on the extracted SRH lifetime. Contrary to previous studies using no control samples [11, 12], we found that a single deep level model could describe the p-type SRH lifetimes very well, and the n-type lifetime across most of the injection level range (the shallow defect affecting the high injection part of the lifetime in previous study was most likely related to intrinsic recombination or surface recombination but not bulk SRH recombination). Hence only the deep



**Figure 2** Injection-dependent lifetimes of the samples (a) p-type + Al (g=0.56) and (b) n-type + Al (g=0.64) and the corresponding control samples. The extracted and the modelled SRH recombination lifetimes are also shown. Note that the vertical axes of the two figures are not in the same scale.





**Figure 3** The extracted and modelled SRH lifetime of the Al–O complexes in the samples n-type + Al (g = 0.64) and p-type + Al (g = 0.56).

level  $E_t = E_v + (0.38-0.50) \text{eV}$  was considered in the model [4, 5, 9, 12]. When  $E_t$  lies in this range, both  $p_1$ and  $kn_1$  are negligible compared with the doping unless k > 1000, which is not true in this case as confirmed by the fitting. Thus the uncertainties in the energy level will not impact on the fitting of k. For the same reason, it was impossible to obtain the energy level via IDLS in this work. The absolute values of the capture cross sections could not be extracted either, due to the unknown concentrations of the defect complexes. Normally when fitting the lifetime,  $\sum (\tau_{\text{fitted}} - \tau_{\text{measured}})^2$  is minimized. However, when the lifetime crosses several orders of magnitudes, this method unintentionally allocates larger weights to the higher-value lifetime data. To avoid this problem,  $\sum [(\tau_{\text{fitted}} - \tau_{\text{measured}}) / \tau_{\text{measured}}]$  $\tau_{\text{measured}}$  was minimized in the fitting to determine the optimal k in this work. The uncertainty range of k was determined as follows: for every k value varied in the range of 0-10000, with the extracted lifetime data, an optimal value of  $N_t \sigma_n$  can be determined for each sample, where  $N_t$ is the defect concentration in the corresponding sample, and  $\sigma_n$  is the electron capture cross section. With the determined  $N_t \sigma_n$  and the k value, we can model the lifetime data for each sample. Then the modelled lifetime  $\tau_{\text{fitted}}$  is compared with the extracted lifetime  $\tau_{measured}$ . A feasible k is selected if  $\tau_{\text{fitted}}$  totally lies within the error bars of  $\tau_{\text{measured}}$  for every sample. The uncertain range of k is the set of all the selected feasible values.

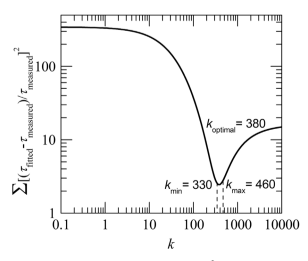
**4 Results and discussion** Figure 1 shows the lifetime measurements for the samples p-type + Al (g = 0.02) and p-type + Al (g = 0.20) and the corresponding control samples. The third group of p-type samples, p-type + Al (g = 0.56) and its control sample are shown

in Fig. 2(a). The polynomial fits are shown in solid lines in Figs. 1 and 2, agreeing very well with the measured data. The extracted SRH recombination lifetimes  $\tau_{\rm SRH}$  are also shown in the figures. The error bars were calculated assuming 12% error in the measured lifetimes [27]. As shown in the figures, the extracted  $\tau_{\rm SRH}$  is strongly injection-dependent in p-type silicon (p-Si), increasing by two orders of magnitudes from  $\Delta n = 1 \times 10^{13} \, {\rm cm}^{-3}$  to  $\Delta n = 1 \times 10^{16} \, {\rm cm}^{-3}$ .

The lifetime measurements of the n-type samples and the extracted  $\tau_{SRH}$  are shown in Fig. 2(b). Throughout the measured injection range, there is very little difference in lifetime between the two samples, indicating that Al-O complexes are weakly recombination-active in n-type silicon (n-Si). The humped shape of the extracted  $\tau_{SRH}$ can be attributed to the uncertainties, as shown by the large error bars. The n- and p-type Al-contaminated samples shown in Fig. 2 have similar net doping levels, [Al] and [O<sub>i</sub>], thus it is reasonable to assume similar concentrations of Al-O complexes in both samples. Figure 3 directly compares the extracted  $\tau_{SRH}$  in the two samples. By comparison, this defect is more recombination-active in p-Si than n-Si at low and intermediate injection levels. At highinjection level, the two lifetime curves lie within each other's error bars.

Figure 4 shows  $\sum [(\tau_{\text{fitted}} - \tau_{\text{measured}})/\tau_{\text{measured}}]^2$  as a function of k. The optimal value of k was determined to be 380, in the uncertainty range of 330–460. As shown by Table 1, this value lies between the other values in the literature. The optimal value was then used to model the  $\tau_{\text{SRH,p-type}}$  for the p-type samples, which agreed well with the extracted data from the experiments (Figs. 1–3).

The value of  $N_t\sigma_n$  could be derived for each p-type Al-contaminated sample. Assuming the same defect concentration in the samples p-type + Al (g = 0.56) and n-type + Al (g = 0.64),  $\tau_{SRH}$  could be modelled for the



**Figure 4**  $\sum [(\tau_{\text{fitted}} - \tau_{\text{measured}})/\tau_{\text{measured}}]^2$  as a function of k, when simultaneously fitting the three extracted  $\tau_{\text{SRH,p-type}}$ .

n-type sample. As shown in Fig. 2(b), the modelled  $\tau_{SRH}$ n-type lies mostly within the error bars of the extracted  $\tau_{SRH,n-type}$  from the measurements, indicating that the same deep level can explain the measured lifetimes for the n-type samples, except possibly at the highest injection levels. The modelled  $\tau_{SRH,n-type}$  is injectionindependent, because when both  $n_1$  and  $p_1$  are negligible, and  $k \gg 1$ , the lifetime is  $\tau_{SRH,n-type} = k/N_t \sigma_n v_{th}$ . Note that this is also the expression for the high-injection lifetime regardless of the doping when  $k \gg 1$ . As shown in Fig. 3, the modelled  $\tau_{SRH,p-type}$  coincides with the modelled  $\tau_{SRH,n-type}$  at high injection levels.  $\tau_{SRH,n-type}$ is injection-independent so the complexes are not very recombination-active over the entire injection range in n-Si. While at low injection levels in p-Si,  $\tau_{\text{SRH,p-type,low-injection}} = 1/N_t \sigma_n v_{\text{th}}$ , degrading by the factor k. As directly compared in Fig. 3, at low and intermediate injection levels, which are more relevant to standard solar cell operating conditions, both the experimental data and the modelling results show that the Al-O complexes are much more recombinationactive in p-Si than n-Si.

**5 Conclusions** The recombination parameters of Al–O complexes in silicon have been reassessed by applying IDLS, using n- and p-type Cz silicon wafers that have been intentionally contaminated with Al and control wafers, and assuming only a single, deep defect level. The single deep level explains the measured lifetimes in p-Si very well, and in n-Si across most of the injection level range of interest. The electron/hole capture cross section ratio k was reassessed to be 380 (-50, +80). Both experimental and modelling results show that Al–O complexes have much greater impact on the carrier lifetimes in p-Si than n-Si at low and intermediate injection levels. Concerning the potential impact of Al contamination on cell performance in low-cost solar-grade materials, n-Si is potentially a better choice.

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