Understanding the distribution of iron in multicrystalline silicon after emitter formation: Theoretical model and experiments

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We studied the behavior of iron in multicrystalline silicon during phosphorus diffusion by spatially resolved measurements and physical modeling. We present improvements to the previously used models for internal gettering in multicrystalline silicon and phosphorus diffusion gettering. 2-dimensional simulations are used for optimization of the phosphorus diffusion processes for intentionally contaminated wafers regarding the iron distribution, without changing the emitter characteristics. Simulations and experimental results show a reduced interstitial iron concentration after an additional low temperature step at the end of the phosphorus diffusion. The concentration of iron precipitates was reduced by a short annealing at 900°C before the phosphorus diffusion, leading to a carrier lifetime three times higher than compared to the standard process.

I. INTRODUCTION

The development of industrial solar cell concepts with high efficiency potential increases the demands on the quality of multicrystalline (mc) silicon. At the same time, selective and lowly doped emitters with lower gettering efficiencies have become more commonly used. Therefore a crucial factor is metal contaminations, which often limits the recombination lifetime in mc silicon. An effective gettering of the metals during solar cell processing could be an alternative to the cost-intensive purification of the feedstock materials. A concentration of metals in the emitter or in a few large precipitates1 drastically reduces the recombination activity of the impurities. If the processes are optimized by enhanced temperature ramps to gain a higher carrier lifetime, the risk of shunting due to large metal precipitates2 can not be disregarded. It has already been found that an extended low temperature gettering after solidification3 or at the end of the phosphorus diffusion process,4 reduces the interstitial iron (Fei) concentration and could increase the solar cell performance5 for highly contaminated silicon. Rinio et al.6 showed that the low temperature step is mainly effective at the edge of a mc ingot, where the metal contamination is high. We used simulations to find an optimized phosphorus gettering process for moderate iron concentrations and tried (experimentally) a pre-anneal at 900°C besides a post low temperature step. An emitter diffusion with an additional peak before the phosphorus diffusion aims for a reduction of the iron precipitate density. Plekhanov et al.7 suggested such a short pre-anneal for aluminum gettering from theoretical considerations. In this paper the results of theoretical and experimental investigations of the iron behavior during the phosphorus diffusion processes are presented. The comparisons of experimental and theoretical results test our iron models and improve the theoretical understanding.

II. MODEL FOR IRON IN MC SILICON

The differential equations for dissolved iron, iron precipitates, and phosphorus (for the phosphorus diffusion process) are solved, including their interaction terms. The main differences of the precipitation model used here which is based on the Fokker-Planck Equation,8,9 to the common model based on Ham’s law,10,11 are: i) that the size distribution of the precipitates is modeled, giving more information and more accurate results in low supersaturated samples, and ii) that instead of the precipitate density itself, the density of sites at which a nucleation of precipitates is possible is used as an input parameter. This allows for the nucleation and complete dissolution of the precipitates during the simulation. The nucleation site density in mc silicon is assumed to be proportional to structural defects. Due to the high nucleation site density in mc silicon, precipitates grow only at a small portion of the nucleation sites. As was experimentally observed in Ref. 12 the resulting precipitation rates are not proportional to the density of structural defects and therefore are less sensitive to uncertainty in the input parameter, i.e., the proportionality factor between the nucleation site density and the density of structural defects. Additional trapping of iron at structural defects due to the local distortion of the lattice13 is considered by an additional term in the solubility. The predictive model for the phosphorus in-diffusion from a POCl3 source distinguishes between the inactive concentration and the active phosphorus concentration, which is responsible for the higher solubility of iron in the emitter and determines the sheet resistance. The simultaneous interaction of iron with phosphorus in the emitter and with the unevenly distributed structural defects via precipitates is taken into account by using a two dimensional structure.

A. Trapping of iron at extended defects

Multicrystalline (mc) silicon is an inhomogeneous material with dislocation densities ranging from $10^2$ cm$^{-2}$ in good grains to more than $10^3$ cm$^{-2}$ in dislocation clusters.
and grain sizes varying from microns to several centimeters. Mobile impurities are trapped at these extended defects by two mechanisms: i) precipitation of supersaturated impurities, and ii) segregation due to higher solubility in the strain field. Iron concentrations 3 orders of magnitude higher than in undisturbed areas can be found in the vicinity of the defects.\(^{14}\)

In the case of mc silicon the density of possible precipitation sites is the product of the dislocation density and the linear density of precipitation sites along a dislocation. The linear density of precipitation sites was chosen to be \(l_p = 3.3 \times 10^5 \text{ cm}^{-1}\). For simplification, grain boundaries are interpreted as lines of dislocations. It is important to note that although the density of possible precipitation is proportional to the structural defects the density of precipitates is not.

Increased equilibrium iron solubility due to local distortion of the lattice in the vicinity of extended defects (SD) is considered by adding an additional term to the iron solubility \(C_{\text{Sol}}\):

\[
C_{\text{Sol}}(\text{SD}) = C_{\text{Sol}}(\text{mono}) (1 + C_1 \times \text{SD}),
\]

\[
C_1 = 0.5 \pi r_{SD}^2 \exp\left(\frac{E_i}{k_B T}\right),
\]  

We decided to modify our first approach\(^{15}\) for the segregation to extended defects and reduce the influence of structural defects on the solubility at high temperature, which is not consistent with theoretical analyses of the interaction of metals and dislocations.\(^{16}\) The area of the strain in which the iron is preferentially located is experimentally difficult to measure and estimated with the radius \(r_{SD} = 15 \text{ nm}\). In this area the interaction energy between dislocations and iron point defects \(E_i\) is inhomogeneous, reaching more than 1 eV in a small core region and less than 0.8 eV in the vicinity of the dislocation.\(^{16,17}\) The factor 0.5 in \(C_1\) is derived from the side with negative \(E_i\). We use an interaction energy between dislocations and iron point defects \(E_i\) of 0.6 eV.

The solubility in the vicinity of extended defects used in this model influences the iron distribution only at lower temperatures and has almost no influence at temperatures above 800°C. Due to the higher proportion of large grains on the overall wafer, the higher solubility in the vicinity of extended defects often has only a minor effect on the mean Fe\(_i\) concentrations. In most cases a precipitation model alone should be able to explain mean values as was found in Ref. 18.

### B. Phosphorus diffusion gettering of iron in mc silicon

Based on the standard diffusion models from the Sen-taurus Process\(^{8}\) platform,\(^{19}\) the diffusion parameters and the boundary conditions for phosphorus are improved.\(^{20}\) During the POCl\(_3\) flux the phosphorus concentration at the surface of the phosphorus silicate glass (PSG) remained constant, while segregation of the phosphorus at the silicon PSG interface is simulated, taking the PSG growth into account.

The redistribution of iron during phosphorus diffusion is modeled with a segregation mechanism, caused by the high active phosphorus concentration in the emitter. A higher solubility of iron in the emitter is modeled by assuming double negatively charged substitutional iron and a pairing of this substitutional iron with positively charged phosphorus. A substitutional state can be expected from a first principle analysis\(^{21}\) and a double negatively charged state in combination with a phosphorus pairing from solubility measurements.\(^{22}\) The solubility of the neutral iron is independent from the net doping in silicon, whereas the equilibrium concentration of negatively charged states and pairs increases in the emitter with the concentration of active phosphorus. We use the same effective segregation coefficient \(k_{Fe}(P^+)\) between emitter and base as in Refs. 20 and 23:

\[
k_{Fe}(P^+) = 1 + 1.5 \times 10^{-27} \exp\left(\frac{1359}{T}\right) \times \left(\frac{p_i}{n}\right)^2 \times P^+, \tag{2}
\]

with the parameters achieved from the experimental data. The additional term in the segregation coefficient proportional to the active phosphorus concentration and the square of the electron concentration \(n\) is the contribution of the iron phosphorus pairs to the solubility. Gilles et al.\(^{22}\) found a decreased diffusivity of iron in highly phosphorus doped silicon. Therefore substitutional iron and iron-phosphorus pairs are assumed to be immobile and only interstitial iron contributes to the diffusivity.

### III. EXPERIMENT

Parallel wafers from the middle and top regions of a block intentionally contaminated with 20 ppma iron in the melt and a reference block are used for the experimental study of internal and external gettering. The 190 \(\mu\)m thick p-doped wafers were chemically polished in order to remove the saw damage and flatten the surface. They were divided into groups: as-cut, phosphorus diffusion (PD) 1–3, and oxidation. For each process variation three \(5 \times 5 \text{ cm}^2\) wafers are processed. Three different temperature profiles were used in the phosphorus diffusion processes (Fig. 1): a standard process (PD 1), a process with a slow cooling ramp and one hour at 550°C after the phosphorus diffusion (PD 2), and a process with a 900°C peak before the phosphorus diffusion (PD 3). After the emitter formation in a tube furnace with the POCl\(_3\) source, the wafers were cooled in air to room temperature. For a better comparison, the sheet resistances should be the same after the three phosphorus diffusions. After stages PD 1 and PD 3, a sheet resistance of 90 \(\Omega\)/sq (a possible industrial standard in the near future), was measured on both sides of FZ reference wafers. Due to the lower solubility of phosphorus at 550°C and the resulting deactivation, the sheet resistance after PD 2 is 20% higher (in the experiment and the simulation). Afterward, the PSG layers as well as 5 \(\mu\)m silicon (including the emitter) at both sides of the wafer are removed and silicon nitride layers are deposited at 350°C for surface passivation. Oxidation processes with equal temperature profiles enable us to distinguish between the redistribution of iron due to the high temperature and due to phosphorus diffusion gettering.

Spatially resolved lifetime measurements captured by quasi-steady-state photoconductance calibrated photoluminescence imaging were performed. One measurement was
obtained after keeping the wafer in the dark where FeB pairs were formed, and the other measurement was obtained after the splitting of the FeB pairs. FeB pairs have been broken by 2 min laser illumination (790 nm, photon density \(6.375 \times 10^{17} \text{cm}^{-2} \text{s}^{-1}\)). From the two measurements and the different Shockley-Read-Hall (SRH) lifetime dependencies of Fei and FeB pairs, an image of the interstitial iron concentration can be calculated. With a Si-CCD camera a resolution of \(50 \times 50 \text{ lm}^2\) for \(50 \times 50 \text{ mm}^2\) wafers are achieved. Wafers are excited by a 790 nm laser with a photon density of \(8.67 \times 10^{16} \text{cm}^{-2} \text{s}^{-1}\) (equivalent to 0.34 suns), generating carriers near the front surface. The low laser intensity ensures that only an insignificant fraction of the FeB pairs break during the 0.5 s of lifetime measurement. In samples with low carrier lifetime the FeB pairs split more slowly under illumination (see Ref. 25 for a discussion). This allows us to use integration times up to 3 s for as-cut samples in order to reduce noise. For the reference samples with a high lifetime the excess carrier density is nearly homogeneous over the whole sample thickness, due to the electron diffusion. For as-cut samples, the underestimation of the Fei concentration due to the inhomogeneous carrier depth profiles is corrected as proposed in Ref. 26. The interstitial iron concentrations are calculated, using the parameters from Istratov et al. for Fei and from Macdonald et al. for FeB.

IV. RESULTS

The phosphorus diffusion gettering model for iron as well as the process variations have been tested at: i) reference wafers with a low total iron concentration which is considerably reduced during phosphorus diffusion, ii) wafers with a medium total iron concentration which is slightly reduced during phosphorus diffusion, and iii) wafers with a high total iron concentration which stays almost constant during phosphorus diffusion. The simulated Fei concentrations in a model structure with areas of different crystal quality are compared with measured Fei concentrations. However, first the results without an external gettering sink are presented.

A. Internal gettering for wafers with an iron concentration of \(3.5 \times 10^{13} \text{cm}^{-3}\)

The phosphorus diffusion and the iron gettering are simulated for a 2-dimensional cross section of a wafer through a grain boundary, a grain 3 mm in width and 210 \(\mu\text{m}\) at a highly dislocated area [see Fig. 2(a)]. The simulations of the precipitation during the cooling process of the block start directly after solidification, using the iron concentration measured by neutron activating analyses (NAA) and the temperature profiles of the crystallization process. The simulated Fei concentrations for wafers in the as-cut state with a total iron concentration of \(3.5 \times 10^{13} \text{cm}^{-3}\) are shown in Figs. 2(b) and 2(c). The simulated mean Fei concentration of \(3.5 \times 10^{12} \text{cm}^{-3}\) is consistent with the measured \(2.1 \times 10^{12} \text{cm}^{-3}\) ([see Fig. 3(a)]. The precipitation sites are located along the dislocation and therefore not evenly distributed in the grains. The simulated Fei concentration varies around the dislocations in the grain. A similar effect is not expected in the measurements, due to the pixel size of \(50 \times 50 \text{ lm}^2\) and the carrier diffusion length of 50–500 \(\mu\text{m}\). If we consider blurring due to carrier diffusion in our simulation and average over \(50 \mu\text{m}\), the variation around a dislocation is less than 5%, which is in the range of background noise. The simulated Fei concentration in the grain is around six times higher than in the vicinity of the grain boundary and eight times higher than in the vicinity of the dislocation cluster, which is in good agreement with the measurement. Comparing the measured and simulated Fei concentration, one has to keep in mind the carrier diffusion during the measurements, which is responsible for smearing around structural defects and explains the smaller contrast in the Fei images. The denuded zones appearing in the Fei imaging can be found with the same widths in the simulated results.
The simulated as-cut state is used as an input file for the simulation of further processing steps. After an oxidation with the standard temperature profile a mean Fe\textsubscript{i} concentration of 2 \times 10^{12} cm\textsuperscript{-3} is measured [see Fig. 3(b)], whereas 4.4 \times 10^{12} cm\textsuperscript{-3} was expected from the simulation [Fig. 2(b)]. In the simulation as well as in the measured Fe\textsubscript{i} images, the contrast between the grain and the grain boundary becomes smaller after oxidation. The denuded zones get thinner, due to out-diffusion from the grain boundaries\textsuperscript{30,31} and less iron is precipitated during the fast cooling.

B. Emitter process variation for wafers with an iron concentration of 3.5 \times 10^{13} cm\textsuperscript{-3}

The simulated Fe\textsubscript{i} concentrations after the three phosphorus diffusions are shown in Fig. 4 together with the Fe\textsubscript{i} image after PD 3. After the standard phosphorus diffusion the Fe\textsubscript{i} concentration is reduced by 1 order of magnitude within the grains, but remains almost constant at the dislocation cluster and the grain boundary, due to dissolution of the precipitates. The additional peak (PD 3) has only a small influence on the concentration of dissolved iron, whereas the additional plateau at 550°C (PD 2) is quite effective. The reasons for this are: i) the higher segregation coefficient [Eq. (2)] between emitter and base, leading to an enhanced external gettering, and ii) precipitation of highly supersaturated iron during the long low temperature step. The simulated total iron concentration after phosphorus diffusion is shown in Fig. 4(a) for the cross section shown in Fig. 2(a). The Fe\textsubscript{i} concentration in the bulk decreases toward the surface in the simulation, due to the gettering effect of the emitter. (b) The simulated Fe\textsubscript{i} concentrations (depth-averaged) after the three process variations compared to the as-cut state. The Fe\textsubscript{i} concentration in the vicinity of the grain boundary and the vicinity of the dislocation cluster is higher than within the grain, but the contrast is rather small for the three process variations. (c) Measured Fe\textsubscript{i} concentration after phosphorus diffusion with an additional peak (PD 3). A higher Fe\textsubscript{i} concentration was measured at pixels with grain boundaries and at dislocation clusters, e.g., in the lower left side of (c).

Table I summarizes the measured mean carrier lifetimes at parallel wafers (Fe\textsubscript{i} state/excess carrier density \( D_n = 10^{14} \text{cm}^{-3} \)).

![FIG. 3. (Color online) Measured Fe\textsubscript{i} concentration in (a) the as-cut state and (b) after oxidation. The mean Fe\textsubscript{i} concentration in both cases is 2 \times 10^{12} cm\textsuperscript{-3}, but after oxidation the Fe\textsubscript{i} concentration in the vicinity of structural defects is higher and the denuded zones are smaller.](image)

![FIG. 4. (Color online) (a) Simulated Fe\textsubscript{i} concentration after PD 3 for the cross section shown in Fig. 2(a). The Fe\textsubscript{i} concentration in the bulk decreases toward the surface in the simulation, due to the gettering effect of the emitter. (b) The simulated Fe\textsubscript{i} concentrations (depth-averaged) after the three process variations compared to the as-cut state. The Fe\textsubscript{i} concentration in the vicinity of the grain boundary and the vicinity of the dislocation cluster is higher than within the grain, but the contrast is rather small for the three process variations. (c) Measured Fe\textsubscript{i} concentration after phosphorus diffusion with an additional peak (PD 3). A higher Fe\textsubscript{i} concentration was measured at pixels with grain boundaries and at dislocation clusters, e.g., in the lower left side of (c).](image)
Figure 5 summarizes the simulated and measured results for the wafers with an iron contamination of $3.5 \times 10^{13}$ cm$^{-3}$. The simulated Fe$_i$ concentration is higher than the Fe$_i$ imaging of the wafers after phosphorus diffusion with a preceding temperature peak (PD 3) (see Fig. 4) and standard phosphorus diffusion (PD 1). The deviation of the Fe$_i$ concentration could be due to uncertainties in the input parameters in the simulations, i.e., total iron concentrations that are too high, incorrect temperature ramps, and a segregation coefficient that is too low in the model, or additional gettering effects. Systematic errors in Fe$_i$ imaging must also be taken into account. The Fe$_i$ concentration in the bulk decreases toward the surface in the simulation, due to the gettering effect of the emitter (see Fig. 4). This depth dependence Fe$_i$ concentration and the low lifetime lead to an underestimation of the Fe$_i$ concentration up to a factor of 2 in the measurements. Considering the number of uncertainties in the experiment and the simulations the correlation is satisfying.

C. Reference wafers

For the as-cut wafers from the reference ingot, a mean Fe$_i$ concentration of $1.6 \times 10^{11}$ cm$^{-3}$ and $6.8 \times 10^{11}$ cm$^{-3}$ were measured. The total iron concentration was below the NAA detection limit of $7.2 \times 10^{12}$ cm$^{-3}$. The Fe$_i$ concentration slightly increases after oxidation, but is still smaller than the solubility during oxidation and the concentration in the contaminated wafers. This allows the assumption that the iron in the reference wafers is completely dissolved during oxidation. In the contaminated samples the Fe$_i$ concentration after the oxidation is nearly 50% of the solubility at the oxidation temperature. Due to the smaller precipitation rates during the cooling process in the lower contaminated samples and the small differences between the Fe$_i$ concentrations before and after oxidation, more than 50% of the iron is in the interstitial state or paired with boron atoms after oxidation and is already in the as-cut state. After the three phosphorus diffusion processes, the Fe$_i$ concentrations are below the detection limit and the carrier lifetimes are at the same level (see Table I).

D. Emitter process variation for wafers with an iron concentration of $1 \times 10^{16}$ cm$^{-3}$

For the wafers from the top of the contaminated ingot, a total iron concentration of $1 \times 10^{16}$ cm$^{-3}$ was measured with NAA. The Fe$_i$ images before and after the standard phosphorus diffusion along with the corresponding simulations are shown in Fig. 6. Table II summarizes the measured carrier lifetimes and mean Fe$_i$ concentrations in areas of different crystal quality. Areas with high dislocation density and grain boundaries have lower Fe$_i$ concentration and higher lifetime before phosphorus diffusion. In the area with large grains the Fe$_i$ concentration and the carrier lifetime is as high as in the wafers contaminated with $3.5 \times 10^{13}$ cm$^{-3}$. The wafers from the top region have a higher proportion of areas with high crystal defect density than the wafers from medium ingot height, resulting in an overestimation of the mean Fe$_i$ concentration in the simulation. A mean Fe$_i$ concentration of $3.3 \times 10^{12}$ cm$^{-3}$ was simulated for the as-cut state, whereas $1.5 \times 10^{12}$ cm$^{-3}$ was measured for the wafer shown in Fig. 6(a). Again, the contrast between the grain and areas with high density of structural defects in the simulation is in consistency with the measurement [see Figs. 6(a) and 6(c)].

After PD 1, the iron concentration is reduced and the carrier lifetime is increased from 0.8 µs to 9.8 µs in the large grains [rectangle in upper left part of Fig. 6(a)], indicating that the dissolved iron was limiting the carrier lifetime.

FIG. 5. (Color online) Comparison of simulated and measured mean Fe$_i$ concentration in the as-cut state and after different processing steps (see Fig. 1).

FIG. 6. (Color online) Measured Fe$_i$ concentration (a) before and (b) after phosphorus diffusion in comparison with the simulations (linescan) (c) for an initial iron concentration of $1 \times 10^{16}$ cm$^{-3}$. After phosphorus diffusion with an additional low temperature step the measured Fe$_i$ concentration is below the detection limit, therefore only the simulation is shown. The rectangles in the upper left part and in the lower right part of (a) are used to represent areas with high and low crystal quality.
before the phosphorus diffusion. A further reduction of the Fe concentration below the detection limit by an additional 550°C anneal improves the carrier lifetime slightly to 11 μs which is comparable with good grains in medium contaminated wafers. In areas with a high density of crystal defects [rectangle in the lower right part of Fig. 6(a)], the reduction of the Fe concentration to 2.8 × 10^{11} cm^{-3} (PD 1) increases the carrier lifetime by a factor of three. An additional low temperature ramp (PD 2) reduces the Fe concentration below the detection limit and increases the carrier lifetime in large grains.

After phosphorus diffusion the simulated Fe concentration in areas with high dislocation density is around 1.7 × 10^{11} cm^{-3} and almost equal to the measured concentration, whereas the Fe concentration in the grain is higher than in the measurements. This leads to higher values in the grain than in areas with a high density of structural defects, although the contrast between areas of different crystal quality is lower than in the as-cut state. The simulation for a further low temperature step shows an almost homogenous Fe concentration of 9 × 10^{9} cm^{-3}.

V. DISCUSSION

A. Interstitial and total iron concentration

The simulated and measured mean Fe concentrations as well as the simulated total iron concentrations after PD 1 for various initial iron concentrations are summarized in Fig. 7. Phosphorus diffusion reduces the total and Fe concentration by 2 orders of magnitude up to an initial iron concentration of 1 × 10^{13} cm^{-3}. Above an initial iron concentration of 5 × 10^{13} cm^{-3} only the mean Fe concentration is significantly reduced.

Although the differences in the Fe concentrations of a series of measurements are reliable, the use of different SRH-parameters for calculating the Fe concentrations from the carrier lifetimes in the two states could result in an overall shift of the Fe concentrations. The published SRH-parameters for Fe and FeB reveal considerable as shown in differences.28 Thus the consistency of the shapes of the measured and simulated curves is more important than an exact match of the values. The Fe concentration in the as-cut state increases rapidly for low initial iron concentrations and has a maximum determined by the cooling ramp after crystallization, whereas the proportion of precipitated iron increases rapidly for iron concentrations above 1 × 10^{12} cm^{-3}. The differences in the crystal defect density of the measured samples are not considered in the simulated mean values in Fig. 7, leading to larger deviations especially for the highly contaminated top wafers (see Table II). As in the simulations the measured Fe concentrations after phosphorus diffusion also increase for an increasing initial iron concentration for high initial concentrations. However the number of measured points is too small to confirm the two different regimes of the simulated curve, i.e., the slower rise for high initial iron concentrations is not discernible.

Kvande et al.34 and Coletti et al.35 found higher Fe concentrations after crystallization and phosphorus diffusion in experiments with intentionally contaminated mc silicon. The qualitative shape of the measured curves is comparable with our simulations, although an emitter with lower sheet resistance and probably different temperature ramps after solidification were used.

B. Carrier lifetime and precipitates

The Shockley-Read-Hall lifetime for an Fe concentration of 7 × 10^{10} cm^{-3} is 80 μs, indicating that Fe is not responsible for the differences in the carrier lifetimes between the iron contaminated wafers (7 μs) and reference wafers (32 μs) after standard phosphorus diffusion. Consequently a further reduction of the Fe concentration (PD 2) increases the carrier lifetime only for the highly contaminated wafers from the top of the ingot. If the iron point defects are not responsible for the reduced carrier lifetimes after phosphorus diffusion, the latter could be due to crystal defects caused by the higher metal concentration or other iron defects, which could be, in principle, gettered. The large increase in the carrier lifetime after PD 3 (with 900°C peak) is evidence that iron precipitates are detrimental after phosphorus diffusion.

![Image](image_url)

**FIG. 7.** (Color online) Comparison of simulated and measured mean Fe concentration in the as-cut state and after standard phosphorus diffusion for various total iron concentrations.

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**TABLE II.** Measured carrier lifetimes τ (Fe state/excess carrier density Δn = 10^{14} cm^{-3}) and mean Fe concentrations in the marked areas of Fig. 6.

<table>
<thead>
<tr>
<th>Crystal quality</th>
<th>τ [μs]</th>
<th>Fe conc. [cm^{-3}]</th>
<th>τ [μs]</th>
<th>Fe conc. [cm^{-3}]</th>
<th>τ [μs]</th>
<th>Fe conc. [cm^{-3}]</th>
</tr>
</thead>
<tbody>
<tr>
<td>High [see Fig. 6(a)]</td>
<td>0.8</td>
<td>2.2 × 10^{12}</td>
<td>9.8</td>
<td>1.0 × 10^{11}</td>
<td>11</td>
<td>&lt;2 × 10^{10}</td>
</tr>
<tr>
<td>Low [see Fig. 6(a)]</td>
<td>1.6</td>
<td>9.3 × 10^{11}</td>
<td>4.8</td>
<td>2.8 × 10^{11}</td>
<td>5.0</td>
<td>&lt;1 × 10^{11}</td>
</tr>
</tbody>
</table>
For this reason we look more closely at the simulated concentrations of precipitates after the different high temperature steps and the recombination activity without Fe\textsubscript{i} and FeB recombination (Fig. 8). The recombination activity without recombination due to Fe\textsubscript{i} point defects can be extracted from the measured lifetime of parallel wafers:

$$R_{\text{woFe}} = \frac{1}{\tau_{\text{other}}} = \frac{1}{\tau_{\text{measured}}} - \frac{1}{\tau_{\text{Fei}}}.$$ (3)

The precipitates have been sorted into three classes (small, medium, and large) in order to show the redistribution of iron atoms during the high temperature process steps.

Without phosphorus gettering, a small shift toward larger precipitates for oxidized wafers show dissolution of small precipitates during the high temperature step and a ripening of large precipitates. An additional external gettering reduces the total iron concentration and thus the number of precipitates. The equilibrium concentration of iron around large precipitates at 900°C is $4 \times 10^{13}$ cm$^{-2}$ \cite{36}, leading to a complete dissolution of small and medium sized iron precipitates and a shrinking of larger precipitates during the short 900°C peak. Coming from higher temperatures (PD 3), the concentration of mobile iron atoms is higher at the start of the phosphorus diffusion and iron can be quickly and effectively gettering. Another advantage of the temperature peak can be seen in Fig. 8. Compared to the cooling process after solidification, the supersaturation after the 900°C peak is much lower, leading to a growth of the existing large precipitates with almost no nucleation of new precipitates. Therefore the densities of small and especially medium sized iron precipitates are lower after the process with the 900°C peak. Small precipitates nucleate only at the temperature ramp to room temperature and the subsequent SiN$_x$ deposition process. The recombination activity of the precipitates depends on the size of the precipitates,\textsuperscript{37} thus small precipitates have only a minor effect on the carrier lifetime. Buonassisi et al.\textsuperscript{1} showed that a concentration of iron in a few large precipitates results in higher carrier lifetimes. The same trend could be observed in the sum of medium and large precipitates and the recombination activity without Fe\textsubscript{i} and FeB recombination, allowing for the assumption that the reduction of precipitates is responsible for the higher lifetimes after phosphorus diffusion with an additional temperature peak. One has to keep in mind that a linear dependency between precipitation density and recombination activity cannot be expected, due to the extremely inhomogeneous distribution of the dislocations and precipitates. The additional 900°C peak halves the number of precipitates, but their concentration is high enough to explain the differences in carrier lifetime between the contaminated wafers and the reference wafers. The pre-anneal would have almost no influence on the simulation results, using a model with a mean radius for all precipitates.

The redistribution of the impurities due to the additional 900°C peak and the resulting gain in lifetime depends strongly on the level and the solubility of the impurities. Using higher peak temperatures with rates of solubility well above the iron concentration could result in high Fe\textsubscript{i} concentrations, if the precipitate density is too low to trap the remaining Fe\textsubscript{i} atoms after phosphorus diffusion.

VI. CONCLUSIONS

A 2-dimensional model for precipitation and phosphorus diffusion gettering of iron in multicrystalline silicon is proposed, which explains the experimental results for emitter process variations. The simulations have been realized for different iron contamination levels and are consistent with the spatially resolved Fe\textsubscript{i} measurements and carrier lifetimes at all process stages. The dissolved iron around dislocations and grain boundaries in higher contaminated as-cut samples is captured by precipitates, leading to lower concentrations of dissolved iron at the structural defects as shown in the simulations and the experiments. We observe and explain the strong reduction of the Fe\textsubscript{i} concentration within the grains and minor changes in areas with high defect densities during standard phosphorus diffusion.

Iron point defects, which are detrimental in the as-cut state, are reduced by more than 1 order of magnitude during standard phosphorus diffusion and can further be reduced to concentrations below $10^{10}$ cm$^{-3}$ with a 90 min low temperature anneal at 550°C. Strong indications were found that iron precipitates and not interstitial iron are limiting the carrier lifetime after phosphorus diffusion in the contaminated wafers. We propose a short temperature peak directly before the phosphorus diffusion in order to significantly improve the gettering effect. We experimentally observed an increase in the carrier lifetime for wafers contaminated with $3.5 \times 10^{13}$ cm$^{-3}$ iron after emitter diffusion by a factor of 3. Our simulations indicate that the large increase in lifetime is due to a reduced density of small and medium precipitates. The additional temperature peak before the phosphorus diffusion improves the carrier lifetime not only inside large grains but also in areas with high crystal defect density. We expect that a 900°C pre-annealing is not always ideal and the optimal peak temperature depends on the level of impurities.
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