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Breaking the Doping Limit in Silicon by Deep Impurities

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$n$-type doping in Si by shallow impurities, such as P, As, and Sb, exhibits an intrinsic limit due to the Fermi-level pinning via defect complexes at high doping concentrations. Here, we demonstrate that doping Si with the deep chalcogen donor Te by nonequilibrium processing can exceed this limit and yield higher electron concentrations. In contrast to shallow impurities, the interstitial Te fraction decreases with increasing doping concentration and substitutional Te dimers become the dominant configuration as effective donors, leading to a nonsaturating carrier concentration as well as to an insulator-to-metal transition. First-principles calculations reveal that the Te dimers possess the lowest formation energy and donate two electrons per dimer to the conduction band. These results provide an alternative insight into the physics of deep impurities and lead to a possible solution for the ultrahigh electron concentration needed in today’s Si-based nanoelectronics.

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I. INTRODUCTION

The scaling evolution for the future generations of Si-based technology, e.g., Si-based transistors in on-chip complementary metal oxide semiconductor (CMOS) devices, requires extremely high free-electron densities (around $10^{21} \text{ cm}^{-3}$) to ensure enough free carriers in each transistor [1,2]. This density is usually achieved with shallow dopants (e.g., group-V dopants such as P, As, and Sb) due to their low ionization energies (20–50 meV), low diffusivities, and suitable solid solubilities in Si. Current technologies, implantation or molecular beam epitaxy, can achieve doping concentrations greater than the solid solubility limit. However, the free-electron concentrations are still found to saturate at around $5 \times 10^{20} \text{ cm}^{-3}$ [3–5]. The electrical activation approaches a limit where the additional introduced dopant atoms cease to generate extra free carriers. During the past decades, experimental results [6–8] and first-principles calculations [9–11] were applied to investigate the microscopic nature of the electrical deactivation of donors in Si. The $\text{As}_nV$ ($n \leq 4$) model (clustering around a vacancy surrounded by As atoms) [10–17] and the dimer model (two donor atoms bound to reconfigured Si with no vacancies) [18,19] are two classical models for electrically deactivating dopant complexes. The latter, however, has been suggested to be dominant at high doping levels [19,20]. The donor dimers introduce localized deep-level states in the band gap and compensate the free electrons, preventing high electron concentrations [18,19].

Alternatively, it has been found that deep-level impurities (e.g., chalcogen dopants) with ionization energies of several hundred meV can also induce free electrons (in excess of $10^{20} \text{ cm}^{-3}$) in Si accompanied with an insulator-to-metal transition (IMT) [21,22]. This result makes chalcogens good prospects to be used for overcoming the doping limit in Si. Particularly, one puzzling observation in chalcogen-doped Si is that the electron concentration increases linearly with the dopant concentration while the substitutional fraction remains almost constant [23,24]. For traditional shallow-level dopants, interstitials, inactive clusters, and precipitates often become more energetically favorable as the dopant concentration increases, e.g., in As-doped Si [25,26], Mn-doped GaAs [27,28], and Mg-doped GaN [29,30]. Therefore, scrutinizing the atomic configuration of chalcogen impurities in Si is needed to
understand this anomalous electrical activation behavior as well as the sub-band-gap infrared photoresponse [31–33].

In this paper, we unveil the microscopic origin for the high electron density that can be achieved by chalcogen doping. We use Rutherford backscattering channeling to directly identify the lattice location of Te in Si and correlate the Te atomistic configuration with the corresponding electrical properties. It is found that, at higher Te concentration, the fraction of substitutional Te increases, contrary to the behavior of shallow dopants. By first-principles calculations, we find that substitutional Te dimers are energetically more favorable and drive the occurrence of nonsaturating free-electron densities, leading to an insulator-to-metal transition. We thus propose chalcogen dimers as effective donors in Si for ultrahigh n-type doping.

II. METHODS

Ion implantation into an intrinsic Si substrate combined with pulsed laser melting (PLM) is applied to synthesize single-crystalline and epitaxial Te-hyperdoped Si layers with a Te concentration several orders of magnitude above the solid solubility limit. Te impurities tend to diffuse toward both the surface and interior of the substrate during the PLM process [33] (see Fig. 8 in Appendix D). The samples are labeled as Te-0.25% . . . Te-2.5%, with the numbers standing for the percentage of the Te peak concentration. Rutherford backscattering spectrometry in channeling geometries (RBS-C) measurements are performed using a 1.7 MeV He\(^+\) beam. The backscattered particles are detected at an angle of 170° with respect to the incoming beam direction using silicon surface barrier detectors. Angular axial scans around different crystallographic directions are obtained using a two-axis goniometer. The electrical properties of the PLM-treated Te-hyperdoped layers are measured using a commercial Lake Shore Hall Measurement System in a van der Pauw configuration [34]. The structural and electronic calculations in the framework of the density functional theory (DFT) are performed by solving the Kohn-Sham equations through the plane-wave pseudopotential approach implemented in the QUANTUM ESPRESSO (QE) open-source code [35]. Further details about experiments and computations can be found in Appendices A and B.

III. RESULTS AND DISCUSSION

A. Electrical properties of Te-hyperdoped Si layers

Figure 1(a) shows the carrier concentration of Te-hyperdoped Si samples measured by the Hall effect at 300 K as a function of doping concentration. The carrier concentration shown here is in the range of 2.0 \(\times\) 10\(^{19}\) to 8.3 \(\times\) 10\(^{20}\) cm\(^{-3}\). Being completely different from previously published results for As-, P-, and Sb-doped Si...
[19,20,36–43], the electron concentration increases monotonically as the doping concentration increases and does not show any sign of saturation even at an atomic dopant concentration as high as 2.5% (exceeding $10^{21}$ cm$^{-3}$). The unsaturated carrier concentration in the Te-hyperdoped Si layers is approaching $10^{21}$ cm$^{-3}$, which is the value required for the next generation of Si technology [1,4,44]. This shows an outstanding behavior of deep impurities as compared with traditional $n$-type dopants in Si (e.g., As and Sb) at very high doping levels, where the reported carrier concentration is never found to be larger than $6.5 \times 10^{20}$ cm$^{-3}$ [45–47]. Figure 1(b) shows the room-temperature resistivity of Te-hyperdoped layers as the function of dopant concentration. Previous results for group-V dopants (i.e., As, P) doped Si [36,38–40,48] are also included. The smallest room-temperature resistivity of Te-hyperdoped Si samples is as low as $7.5 \times 10^{-4}$ $\Omega$ cm and is comparable to the smallest resistivity value for As-doped Si. In addition, the carrier mobility has also been obtained from the Hall effect measurements. As shown in Fig. 1(c), the mobility is in the range of 10–40 cm$^2$/V s, which lies in the same order of the reported values for As-doped Si at a high doping regime ($10^{20–21}$ cm$^{-3}$) [39] (see Appendix C for further details about the electrical transport measurements). IMT is also observed in Te-hyperdoped samples as shown by their temperature-dependent conductivity (see Fig. 5 in Appendix C). Samples Te-0.25% and Te-0.50% with donor concentrations below the Mott limit (see Appendix C) show an insulating behavior with a strong temperature-dependent conductivity. However, samples with the Te peak concentration exceeding 1.0% present a radically different behavior, exhibiting a conductivity with negligible variation with temperature and a finite value of more than $10^2$ S/cm as the temperature tends to zero. It is therefore crucial to understand this anomalous dependence of the electrical activation on the dopant concentration in Te-hyperdoped Si by scrutinizing the dopant lattice configuration at the atomic scale.

### B. Lattice location of Te dopants in Si matrix

RBS-C is a unique technique to measure the lattice location of the dopants [49] (see Appendix D for further details). Figure 2 shows the angular distributions of He ions backscattered by Te and Si as a function of the tilt angle around the Si (100) axes in the selected samples. More angular scans in the vicinity of the three main crystallographic directions (⟨100⟩, ⟨110⟩, and ⟨111⟩) are displayed in Appendix E. Two typical parameters are used to characterize the angular distributions, $\chi_{\text{min}}$ and $\psi_{1/2}$. The minimum yield in the angular scan, $\chi_{\text{min}}$, qualitatively describes the ability of dopants to block the channels. The critical angle $\psi_{1/2}$ represents the half width at half maximum of the angular scans between $\chi_{\text{min}}$ and 1. It is related
to the displacement of substitutional atoms [50–52]. Both parameters are calculated for Si and Te at the same depth. The main results and their implications are as follows:

(i) For sample Te-0.25%, there is a peak in the middle of the angular scan. Some Te ions are located at low-symmetry interstitial sites lying around the axes of the crystal lattice. This peak decreases with increasing Te concentration. At low concentrations, Te ions occupy both substitutional and interstitial sites, whereas at higher concentration they preferentially go to substitutional sites.

(ii) The interstitial fraction estimated from the angular scan drastically decreases at Te concentrations up to 1.0% and tends to a small constant value at higher concentrations [Fig. 2(d)]. It is worth noting that 1.0% is also the critical concentration where the IMT takes place in Te-doped Si [see Fig. 1(a)]. This result indicates a strong correlation between the atomic configuration of Te ions and the electronic properties of the samples.

(iii) At higher concentrations, $\psi_{1/2}$-Te is narrower compared to $\psi_{1/2}$-Si. This suggests that the Te dopants do not perfectly substitute the Si lattice sites. There exists a displacement ($r_0$) from the ideal substitutional site [49]. Based on the experimental data, the projections of the displacement $r_0$ of Te atoms perpendicular to the three main axes can be estimated (see Appendix F). The average value $r$ obtained from the angular scan is found to be around $0.42 \pm 0.03$ Å. This result is in agreement with the average displacement ($r_0 = 0.46$ Å, for Te concentration of 1.36%) obtained from the first-principles calculations shown later.

To further prove that the majority of Te ions are located at the substitutional lattice sites at high Te concentrations, we perform angular-scan mapping around three major crystalline axes. Figure 3 shows representative angular-dependent two-dimensional backscattering-yield patterns of the sample with the highest Te concentration (Te-2.5%). The backscattering yield patterns are extracted from the integral backscattering signals of Si and Te in the vicinity of $\{100\}$, $\{110\}$, and $\{111\}$ in sample Te-2.5%. Prominent channeling effects are observed for both the Si and the Te signals and their patterns show a high similarity along $\{100\}$, $\{110\}$, and $\{111\}$ orientations and the closest-packed planes $\{100\}$, $\{110\}$, and $\{111\}$). These angular-scan mappings unambiguously demonstrate that the Te-hyperdoped Si layer is epitaxially regrown during the PLM treatment and that the majority of Te atoms occupy the substitutional sites of the Si matrix at such high Te concentrations.

C. Ab initio calculations

We directly probe the lattice location of Te ions in hyperdoped Si by RBS-C. These results suggest that the decrease of Te interstitials is accompanied by an increase of displacements of Te atoms from the ideal substitutional site. To understand this, first-principles calculations (see Appendix B for further computational details) have been carried out to compute the formation energy for three types of defects as a function of Te doping concentration. We define the formation energy of a defect ($D$) as

$$\Delta E_F \equiv E_D^{\text{SC}} - n_{\text{Si}}^{\text{SC}} \mu_{\text{Si}} - n_{\text{Te}}^{\text{SC}} \mu_{\text{Te}},$$

where $E_D^{\text{SC}}$ is the total energy of the supercell (SC) with defect $D$, $n_{\text{Si}}^{\text{SC}}$ and $n_{\text{Te}}^{\text{SC}}$ are the numbers of Si and Te atoms in the supercell, and the chemical potentials $\mu_{\text{Si}}$ and $\mu_{\text{Te}}$ correspond to pure Si and SiTe$_2$ in equilibrium with each other, respectively. In Fig. 4(a), we summarize our results for the most relevant types of defects: substitutional single Te (Te$_{\text{Si}}$) and substitutional Te dimer (Te$_{\text{Si}}$-Te$_{\text{Si}}$). Moreover, the results for interstitial defects (Te$_{\text{interstitial}}$ at hexagonal $H$ and tetrahedral $T$ symmetry sites) at the selected concentrations are displayed in the inset of Fig. 4(a) for comparison. Generally, the formation energy of the interstitial sites ($T$ site and $H$ site) is much larger (several eV per Te atom) than that of the substitutional site in the Te-hyperdoped Si system, which results in an energetically unfavorable formation of interstitial Te. Surprisingly, the Te$_{\text{Si}}$-Te$_{\text{Si}}$ dimer presents the lowest formation energy among all types of defects considered, and the difference in the formation energy (per Te atom) between a single Te$_{\text{Si}}$ dopant and a Te$_{\text{Si}}$-Te$_{\text{Si}}$ dimer is around 0.3 eV. This result suggests that Te$_{\text{Si}}$-Te$_{\text{Si}}$ dimers are more energetically favorable. Note that, at low concentrations, the average distance between Te atoms might be too large to promote the formation of dimers. As shown in Fig. 4(a), the Te$_{\text{Si}}$ formation energy increases sharply with increasing Te concentration on the insulating side, which is consistent with the previously reported results about Se-hyperdoped Si [22]. However, in the regime exceeding the threshold for the IMT, the defect formation energy is relatively stable; it only slightly increases with the Te concentration. Therefore, as the doping Te concentration increases, the penalty for assigning additional Te atoms becomes less costly and ultimately renders the Te$_{\text{Si}}$-Te$_{\text{Si}}$ dimer configuration stable. If Te$_{\text{Si}}$-Te$_{\text{Si}}$ dimers form, the lattice positions of Te are slightly displaced from the ideal substitutional sites. From the ab initio relaxed positions of Te dimers, the displacement is estimated to be around 0.46 Å for samples with a Te concentration of 1.36% (see Appendix F). The Te$_{\text{Si}}$-Te$_{\text{Si}}$ dimer formation and the displacement are in excellent agreement with the findings obtained from the experimental results shown in Figs. 2 and 3.

In Figs. 4(b) and 4(c) we show the density of states (DOS) for our calculations of Te$_{\text{Si}}$-Te$_{\text{Si}}$ and single Te$_{\text{Si}}$ at a Te concentration of 1.56%, i.e., in the metallic regime. The projected DOS on the Te atomic
Fig. 3. Experimental two-dimensional backscattered-yield patterns of the sample Te-2.5% obtained from RBS-C spectra. The patterns are extracted from the integral backscattering signal from Te [(a), (c), and (e)] and Si [(b), (d), and (f)] in the vicinity of the ⟨100⟩, ⟨110⟩, and ⟨111⟩ directions. The patterns for the Si and the Te signals match well with each other, indicating that the majority of Te impurities are substitutional.

Orbital has been summed up and displayed for Te single (red solid line) and Te dimer (black solid line); the Te projected DOS represents the contribution of electronic states localized at Te atoms to the total DOS. Figures 4(b) and 4(c) show a small gap approximately centered at −0.7 eV for Te_{Si} and −0.4 eV for Te_{Si-Te_{Si}}, respectively, that separates the conduction band from the valence band. The modification of the bulk Si band gap is caused by doping. The Fermi energy (the zero of the horizontal energy scale) is located in the conduction band. By comparing the contribution of projected DOS on Te species with the total DOS, we notice that for the conduction band the Te projected DOS is small, denoting that the conduction electrons are delocalized on the whole crystal, thus producing a metallic phase. By performing the same comparison for the valence band, we can notice that for the dimer case the gap between the valence and conduction band is significantly reduced with respect to the single Te case, because of the presence of electronic states (of s and p character) at about −0.7 eV that are localized at the Te atoms, as can be seen in Fig. 4(c) by noticing the significant contribution of Te projected DOS to the total DOS at this energy. In both panels, the position of the Fermi energy denotes a metallic phase in agreement with the experimental data in Fig. 1(a). By integration of the local DOS from the conduction band minimum (CBM) up to the Fermi energy, we compute the number of carriers provided by each type of defect. From our ab initio simulations, we find that a single Te_{Si} acts as a double donor, which is consistent with other reports [53,54]. At variance, one Te_{Si-Te_{Si}} dimer provides only two electrons to the conduction band; i.e., each Te composing the dimer provides only one electron to the conduction band. From Fig. 1 we can notice that at high Te concentration, the electron concentration...
FIG. 4. (a) Normalized (per atom) formation energy ($\Delta E_f/n^{SC}_{Te}$) of an isolated substitutional Te (circles) and a Te-Te dimer (diamonds) as a function of Te concentration, computed via ab initio calculations. Inset: Formation energy of high-symmetry interstitial sites ($T$ and $H$ denoted with squares and stars, respectively) as a function of Te concentration; lines are a guide for the eye. Ab initio calculations of density of states (DOS) at a Te concentration $x = 1.56\%$ for a single Te dopant (b) and a Te$_{Si}$-Te$_{Si}$ dimer (c). The sum of the DOS projections on the Te atomic orbitals is also displayed. The zero of the energy scales (dashed lines) corresponds to the Fermi energy.

(8.1 × 10$^{20}$ cm$^{-3}$) is approaching the dopant concentration (1.25 × 10$^{21}$ cm$^{-3}$) due to the formation of Te$_{Si}$-Te$_{Si}$ dimers. The lowest resistivity in our films is also comparable with As- and P-doped Si for similar doping levels [as shown in Fig. 1(b)]. We anticipate that the electron concentration can be further increased by optimizing the Te implantation fluence and annealing parameters, therefore opening a new avenue toward ultrahigh $n$-type doping for the Si-based new-generation microelectronics. As a final remark, we cannot exclude the appearance of Te$_{nV}$ complexes from RBS-C [55,56]. However, the measured high carrier concentration supports the idea that Te$_{nV}$ complexes cannot be the major states since they are electrically inactive centers [7,12,14,16,20].

IV. CONCLUSION

In conclusion, we demonstrate that the long-standing $n$-type doping limit in Si can be overcome by using deep-level donors (e.g., Te) instead of traditional shallow-level donors (e.g., P, As, Sb). We have provided direct experimental and theoretical evidence for the formation of Te$_{Si}$-Te$_{Si}$ dimers with increasing Te concentration that are found to be responsible for the nonsaturating electron concentrations and the insulator-to-metal transition. This work provides insight toward understanding the lattice configurations and electronic structures of deep-level dopants in semiconductors.

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APPENDIX A: EXPERIMENTAL DETAILS

The single-side polished Si (100) substrates (intrinsic, $\rho \geq 10^4 \, \Omega \, cm$) are implanted with Te ions at six different fluences at room temperature with implantation energies of 150 and 50 keV (see Table I). The combined implantation energies are used to obtain a uniform distribution of Te in the implanted layer. The peak Te concentrations ($c_{pk}$) are firstly calculated using SRIM code [57] and then verified by RBS-C measurements and SIMNRA [58,59] simulation. The ion-implanted samples are exposed to a single laser pulse with the energy density of 1.2 J/cm$^2$. The whole amorphous implanted region is melted and then recrystallized with a solidification speed of the order of 10 m/s while cooling down [60] during the annealing process. After
TABLE I. Sample definition and related parameters. The sample names refer to the peak Te concentration of the as-implanted layer.

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Implantation energy (keV)</th>
<th>Implantation doses (cm(^{-2}))</th>
<th>Tellurium peak concentration (cm(^{-3}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Te-0.25%</td>
<td>150</td>
<td>7.8 \times 10^{14}</td>
<td>1.25 \times 10^{20}</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>3.1 \times 10^{14}</td>
<td></td>
</tr>
<tr>
<td>Te-0.50%</td>
<td>150</td>
<td>1.6 \times 10^{15}</td>
<td>2.50 \times 10^{20}</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>6.2 \times 10^{14}</td>
<td></td>
</tr>
<tr>
<td>Te-1.0%</td>
<td>150</td>
<td>3.1 \times 10^{15}</td>
<td>5.00 \times 10^{20}</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>1.2 \times 10^{15}</td>
<td></td>
</tr>
<tr>
<td>Te-1.5%</td>
<td>150</td>
<td>4.7 \times 10^{15}</td>
<td>7.50 \times 10^{20}</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>1.9 \times 10^{15}</td>
<td></td>
</tr>
<tr>
<td>Te-2.0%</td>
<td>150</td>
<td>6.2 \times 10^{15}</td>
<td>1.00 \times 10^{21}</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>2.5 \times 10^{15}</td>
<td></td>
</tr>
<tr>
<td>Te-2.5%</td>
<td>150</td>
<td>7.8 \times 10^{15}</td>
<td>1.25 \times 10^{21}</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>3.1 \times 10^{15}</td>
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</tr>
</tbody>
</table>

that, the Te-rich layer is recrystallized and its thickness is approximately 120 nm [33].

The RBS-C measurements are carried out using a 1-mm-diameter collimated 1.7-MeV He\(^+\) beam of the Rossendorf van de Graaff accelerator with a 10–20-nA beam current. Energy spectra of backscattered He are detected at an angle of 170° with respect to the incoming beam direction using silicon surface barrier detectors with an energy resolution of 15 keV. Because of the so-called channeling effect, the backscattering probability sensitively depends on the direction of the incident beam with respect to the crystallographic orientation of the sample and is significantly reduced around the crystal axes and planes [61]. Angular axial scans are performed using a two-axis goniometer by recording integral backscattering yields in energy windows corresponding to scattering from Si and Te atoms (see Fig. 10) as a function of the angle of incidence around different crystallographic axial directions (⟨100⟩, ⟨110⟩, and ⟨111⟩). In addition, two-dimensional angular backscattering patterns are obtained.

The electrical properties of the Te-hyperdoped Si samples are examined using a commercial Lake Shore Hall system in van der Pauw geometry [34]. Samples are measured in the temperature range from 2 to 300 K and a magnetic field perpendicular to the sample plane is swept from −6 to 6 T. The gold electrodes are sputtered onto the four corners of the squarelike samples. The native SiO\(_2\) layer is removed by hydrofluoric acid (HF) etching prior to the sputtering process. Silver glue is used to contact the wires to the gold electrodes.

**APPENDIX B: COMPUTATIONAL DETAILS**

We perform structural and electronic calculations in the framework of the DFT, solving the Kohn-Sham equations through the plane-wave pseudopotential approach, as implemented in the QE open-source code [62]. We use ultrasoft pseudopotentials [63–66] in the separable form introduced by Kleinmann and Bylander [67], generated with a Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional and smearing techniques with a Gaussian broadening of 5 mRy. We choose a 50-Ry cutoff radius for the electronic valence wave function and 500-Ry cutoff radius for the charge density. The formation energy is computed according to Eq. (1) of the main text; the more natural choice for \(\mu_{Si}\) corresponds to the chemical potential of bulk silicon (usually referred as the Si-rich chemical potential), evaluated according to the standard procedure (see, e.g., Refs. [68,69]) by taking the total energy of the unit cell computed by DFT divided by the number of atoms contained in the unit cell. Our choice for \(\mu_{Se}\) corresponds to the Te chemical potential of bulk SiTe\(_2\) [18] in equilibrium with bulk silicon. So, \(\mu_{Se}\) is evaluated by subtracting the chemical potential of bulk Si from the total energy of the SiTe\(_2\) unit cell computed by DFT and dividing the result by 2. Note that, for the present case, the choice of the chemical potential of Te simply shifts the zero of the energy, i.e., the vertical axis in Fig. 4(a) in the main text.

The \textit{ab initio} simulations of doped silicon are performed by using the supercell method. In a similar way as done, e.g., in Ref. [22], we vary the size of the supercell to simulate different concentrations of impurities (in our case, the Te concentration ranges from approximately 0.4% to 3.1%). Periodic boundary conditions are used. So, in the hyperdoping regime considered in the present work, each impurity interacts with its own images, simulating a uniform distribution of the same defect type. This approximation is expected to provide qualitative estimation of average properties like the defect formation energy of defect types representing a fraction of the total defect population, while it is expected to provide a reliable quantitative estimation of
average properties of defect types representing the large majority of defects present in the sample. In the case in which only one type of defect is present or the concentration of the other types of defects is negligible, the present method would provide exact results within the density functional approximation technique used. In general, the atomic positions and the lattice constants are fully relaxed by the Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm [21] with a threshold on the forces of 1 mRy/\(a_0\). However, since we find that lattice constants satisfy Vegard’s law (i.e., the lattice parameter varies linearly as a function of doping concentration) also for the higher doping regime we consider, we fix the lattice parameter for a few structures at a diluted concentration (thus having the largest supercell size considered in the present simulations) by linear interpolation of structurally relaxed results and relax only the atomic positions by the BFGS algorithm.

**APPENDIX C: ELECTRICAL TRANSPORT MEASUREMENTS**

The Te-hyperdoped Si samples with the single-side polished intrinsic Si substrate (\(\rho > 10^4 \Omega \text{cm}\)) are used for the electrical characterization. The Hall measurements and resistivity measurements of samples are measured using the van der Pauw technique [34]. Based on the basic physical principle of the Hall effect, the carrier density of the samples can be estimated using standard techniques and a magnetic field \(B\) ranging from -4 to 4 T. In some cases, it is convenient to use a layer or sheet density \(n_s\) instead of bulk density by using \(n_s = IB/eV_H\), where \(I\) is the current, \(B\) is the magnetic field, \(V_H\) is the Hall voltage, \(e\) (1.602 \times 10^{-19} \text{C}) is the elementary charge, and \(\gamma\) is the Hall scattering factor that is generally assumed to be 1 heavily doped Si. If the conducting layer thickness \(d\) is known, the bulk density \(N\) can be determined as \(N = n_s/d\) [34]. Here, the carrier concentration \(n\) ranges from 2.0 \times 10^{19} to 8.3 \times 10^{20} \text{cm}^{-3}, which is calculated by assuming the effective thickness of the tellurium-doped layer as 125 nm [33]. This relatively large magnitude of the donor concentration is comparable to the shallow-level impurity doped Si with just-metallic concentrations [70] and is consistent with the metallic-like samples in S- and Se-doped Si [21,22]. Moreover, the carrier mobility can be obtained via \(\mu = |V_H|/IB_{Rs} = 1/qnR_s\), where \(R_s\) is the sheet resistance. The obtained carrier mobility is in the range of 10–40 cm²/Vs and decreases with increasing Te doping concentration. These values are almost of the same order as in Si heavily doped with As in the high doping regime (\(10^{20–21} \text{cm}^{-3}\)) [39].

The sheet resistance \(R_s\) of samples over the temperature range 2–300 K is measured. The bulk electrical resistivity \(\rho\) is calculated using \(\rho = R_s \times d\) and the conductivity \(\sigma = 1/\rho\). The effective layer depth \(d\) is 125 nm, which is verified by the RBS results (see Fig. 7). Figure 5 shows the temperature-dependent conductivity with the Te peak concentration varied from \(1.25 \times 10^{20} \text{cm}^{-3}\) (0.25%) to \(1.25 \times 10^{21} \text{cm}^{-3}\) (2.5%) in a temperature range of 2–300 K. There, the room-temperature conductivity of Te-hyperdoped Si samples is higher than \(10^2\) S/cm, which confirms that the intrinsic Si substrate has no influence on the transport properties of the Te-hyperdoped Si layer considering the respective thickness. As shown in Fig. 5, a rigorous experimental evidence of an IMT lies in the measurement of nonzero conductivity as the temperature

![FIG. 5. The temperature-dependent conductivity of Te-hyperdoped Si in the temperature range of 2–300 K. Samples with lower Te concentrations (Te-0.25% and Te-0.50%) show a strong temperature-dependent conductivity, indicative of the insulating state. Samples with higher concentrations (Te-2.0% and Te-2.5%) exhibit a conductivity comparatively insensitive to temperature down to 2 K, indicating an impurity-mediated transition to the metallic-like state.](image)

![FIG. 6. The conductivity of sample Te-0.25% as a function of inverse temperature. The red solid lines show the fits of data to Eq. (C3), yielding an activation energy of 46.5 meV.](image)
tends to zero (described previously [33]). Samples Te-0.25% and Te-0.50% show insulating behavior with a strong temperature-dependent conductivity. Samples with the Te peak concentration exceeding 1.0% present a dramatically different behavior, exhibiting a conductivity with negligible variation with temperature and a finite value of more than 10^2 S/cm as the temperature tends to zero. At 2 K, the conductivity of the samples with 1.0% and 0.25% Te concentration differs by nearly 4 orders of magnitude in spite of the difference of just four times in the concentration. Samples with Te concentration exceeding 1.0% show a quasimetallic behavior. Moreover, the room-temperature conductivity of Te-hyperdoped Si is around 600 S/cm, while the conductivity of S- and Se-hyperdoped Si is about 110 and 300 S/cm respectively at similar doping levels (approximately 4.2–4.9 × 10^20 cm^-3) [21,22].

As Mott [71,72] originally derived, the insulator-to-metal transition in semiconductors can be described as

\[ n_c^{1/3} a_H = 0.25, \]  
(C1)

where \( a_H \) is the first Bohr radius of the donor electrons. The first Bohr radius can be scaled as

\[ a_H = \frac{e^2}{8\pi \epsilon_0 \epsilon_r E}, \]  
(C2)

where \( E \) is the activation energy of the localized states, and \( \epsilon_0 \) and \( \epsilon_r \) are the permittivity of free space and the high-frequency dielectric constant, respectively. Thus, by taking into account the activation energy as 199 meV for a single Te atom [53,54] and 158 meV [73] for a Te dimer and by assuming that the Mott criterion is approximately valid also for the dimer configuration, the isotropic Bohr radius is calculated as 3.1 and 3.9 Å, respectively. Therefore, the critical electron concentration for the Mott transition of a single Te atom in hyperdoped Si is approximately 5.24 × 10^20 cm^-3, corresponding to a Te impurity concentration of 2.65 × 10^20 cm^-3 since single substitutional Te atoms are double donors, while for a Te dimer in hyperdoped Si, the critical electron concentration for the Mott transition is approximately 2.65 × 10^20 cm^-3, corresponding to a Te impurity concentration of 2.65 × 10^20 cm^-3 since one substitutional Te dimer donates two electrons. This rough estimation suggests that the onset for metallicity is almost the same for single Te or Te dimers.

Sample Te-0.25% is away from the transition point. However, the Te atoms can be “ionized” even in sample Te-0.25% at room temperature because the large Te concentration gives rise to the formation of an impurity band that provides weakly localized or free electrons. To describe a thermally activated conduction process at high and low temperatures, we use

\[ \sigma = \frac{\sigma_0}{\exp[E_0/k_B T]}. \]  
(C3)

where the pre-exponential constant \( \sigma_0 \) and the activation energy \( E_0 \) are the free parameters. Over a limited temperature range, the measured conductivity data fit to this model quite nicely (as shown in Fig. 6). The fitting results indicate the thermally activated contribution to the conductivity of Te-hyperdoped Si. For sample Te-0.25%, a high-temperature activation region (300–200 K) with an activation energy of approximately 46.5 meV (\( E_0 \)) is observed, which is comparable to the thermal energy of room temperature (around 26 meV). Therefore, in sample Te-0.25%, it is possible to have a thermally activated conductivity. At low temperature, this sample shows variable-range hopping conductivity with a Coulomb gap. It can be fitted with a \( 1/T^{1/2} \) dependence (not shown here).

APPENDIX D: RBS-C MEASUREMENTS

Figure 7 shows the representative RBS-C spectra of a PLM-treated Te-hyperdoped Si layer with different Te concentrations, also including the virgin Si and the as-implanted wafers. The random spectrum of the as-implanted layer reveals a thickness of about 120 nm for the Te profile, in an approximately Gaussian distribution. No channeling effect is observed in the implanted layer because of the amorphization caused by ion implantation. From the RBS-C signal, the PLM-treated layer shows a minimum backscattered yield \( \chi_{\text{min}} \) of 4% (defined as the ratio of the aligned to the random yields near the sample surface from 740 to 920 keV), which is comparable to that of virgin Si substrate. This result indicates a full recrystallization and an epitaxial growth of the PLM-treated layers.

Moreover, a channeling behavior of Te (from 1380 to 1510 keV) is observed in the PLM-treated samples, which proves the incorporation of Te into the Si lattice sites. The
FIG. 8. The RBS random spectra and the fitting by the SIMNRA code for sample Te-1.5% before and after the PLM treatment.

depth profile of the Te distribution before and after PLM treatment can be obtained by fitting RBS spectra using the SIMNRA code [58, 59]. Figure 8 shows the Te signal taking from the random spectrum for sample Te-1.5% before and after PLM treatment. It is evidenced in the Te signal that Te atoms tend to diffuse toward both the surface and the substrate side during the PLM process. The sample name is indicative of the peak Te concentration of the as-implanted layer. This has been verified in our previous work by transmission electron microscopy [33].

APPENDIX E: DETAILED ANGULAR SCANS

Figure 9(a) shows the schematic of the Si matrix with the implanted impurities that can take place at different lattice (I, II, III, and IV) sites. Figure 9(b) shows the angular scans for the respective lattice sites: the dashed line corresponds to the impurity, while the solid one to Si. If both angular scans match well with each other, impurities are located at the substitutional sites with high symmetry. The angular scan for impurities occupying interstitial sites is displayed in Fig. 9(b). Normally, two typical parameters are used to characterize the angular distributions, $\chi_{\text{min}}$ and $\psi_{1/2}$. In detail, $\chi_{\text{min}}$ corresponds to the minimum value of the backscattered ion angular spectrum with respect to the random direction normalized to 1 [52]. Qualitatively, it presents the ability of dopants blocking the channels. The critical angle $\psi_{1/2}$ represents the half width of the angular scans at half maximum between $\chi_{\text{min}}$ and 1 (full width at half maximum of the angular scan spectrum), and it is related to the displacement vector of substitutional atoms [50].

Figure 10 shows the angular distributions for Te and Si at the same depth in the vicinity of the three main crystallographic directions, namely ⟨100⟩, ⟨110⟩, and ⟨111⟩, recorded off a major crystallographic plane. Two typical parameters $\chi_{\text{min}}$ and $\psi_{1/2}$ are simultaneously calculated for Si and Te at the same depth. The three axial angular scans for Te follow those of the Si host in all the Te-hyperdoped Si samples, indicating that the dominant fraction of the Te atoms is located at substitutional [50, 75] sites. For samples Te-0.25% and Te-0.50%, the $\chi_{\text{min}}$-Te...
FIG. 10. (a)–(r) The angular scans about the \(\langle100\rangle\) [(a)–(f)], \(\langle110\rangle\) [(g)–(l)], and \(\langle111\rangle\) [(m)–(r)] axes for all the Te-hyperdoped Si samples. Angular distributions have been normalized in a random direction. The red circles correspond to Te atoms and black squares to Si; smooth curves are drawn through the symbols.
is significantly higher than that of Si. This suggests that a non-negligible fraction of the Te atoms does not occupy substitutional sites and those atoms are located at the low-symmetric interstitial sites lying around the axes of the crystal lattice. As depicted in Figs. 10(a), 10(b), 10(g), 10(h), 10(m), and 10(n), the interstitial site impurities are clearly evidenced. For samples with higher doping concentrations, a puzzling and interesting behavior is observed from the angular scans. The fraction of the interstitial Te clearly evidenced. For samples with higher doping concentrations, the general expectation, the interstitial fraction decreases from the angular scans. The fraction of the interstitial Te dominates, and the sample 0.25% is a mixed case. However, a quantitative estimation is difficult from the current data.

**APPENDIX F: TELLURIUM DISPLACEMENT ESTIMATION**

The RBS/C angular scans shown in Fig. 10 for Si and Te suggest that the Te atoms may be displaced from the ideal substitutional Si lattice sites. To obtain quantitative information about the Te location, the angular distributions as a function of the equilibrium displacement of an atom from a lattice row [51] are calculated. The parameter $\psi_{1/2}$ can be expressed as follows for displacement calculation [52]:

$$\psi_{1/2} = \sqrt{\left[ \frac{1}{2} \ln \left( \frac{C^2 a^2}{r_{\min}^2} + 1 \right) \right]} \times \sqrt{\frac{2Z_1 Z_2 e^2}{Ed}}, \quad (F1)$$

where constant $C^2$ is around 3; $Z_1$ and $Z_2$ are the atomic numbers of the incident particle and of the considered atom, respectively; $e^2$ is equal to 14.4 eV Å; $E = 1.7$ MeV is the incident ion energy, and $d$ is the average atomic distance in the considered atomic row ($d_{\text{aver}}(110) = 5.43$ Å, $d_{\text{aver}}(110) = 3.84$ Å, $d_{\text{aver}}(111) = 4.70$ Å) [52]; and

$$r_{\min}^2 = \rho^2 \ln 2 + r_0^2. \quad (F2)$$

Here, $\rho$ is the thermal vibration amplitude and $r_0$ is the projection (on a plane perpendicular to the channeling direction) of the equilibrium displacement of the Te atoms with respect to the lattice sites. $a$ is the screening distance, which is related to the Bohr radius ($a_0 = 0.528$ Å).

**TABLE II.** The experimental values of the displacement are obtained from formula F1, which corresponds to the lattice sites projected perpendicularly to the channeling direction. The (100) direction is perpendicular to the Si substrate. Moreover, the computational values of the displacement are from the first-principles calculations, where an average displacement of 0.46 Å is obtained.

<table>
<thead>
<tr>
<th>$r_0$ (computational values)</th>
<th>$r_{(100)}^0$</th>
<th>$r_{(110)}^0$</th>
<th>$r_{(111)}^0$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Te-0.25%</td>
<td>0.36 ± 0.03</td>
<td>0.40 ± 0.03</td>
<td>0.34 ± 0.03</td>
</tr>
<tr>
<td>Te-0.50%</td>
<td>0.37 ± 0.03</td>
<td>0.40 ± 0.03</td>
<td>0.36 ± 0.03</td>
</tr>
<tr>
<td>Te-1.0%</td>
<td>0.42 ± 0.03</td>
<td>0.40 ± 0.03</td>
<td>0.38 ± 0.03</td>
</tr>
<tr>
<td>Te-1.5%</td>
<td>0.43 ± 0.03</td>
<td>0.41 ± 0.03</td>
<td>0.45 ± 0.03</td>
</tr>
<tr>
<td>Te-2.0%</td>
<td>0.46 ± 0.03</td>
<td>0.42 ± 0.03</td>
<td>0.46 ± 0.03</td>
</tr>
</tbody>
</table>

as follows:

$$a = 0.8853 \times a_0 \times \frac{1}{\sqrt{(Z_1^{1/2} + Z_2^{1/2})^2}} = 0.11114. \quad (F3)$$

Here, the range for the average error of the projection displacement is determined by the dispersion of the parameter $\psi_{1/2}$.

For comparison, we computed the displacements obtained in our *ab initio* simulations. To minimize the numerical error, which in our simulation is comparable to the variation of the displacement as a function of the Te concentration, we average the values of displacements of Te impurities obtained from first-principles calculations for TeSi over the whole range of the investigated concentration. The average results, corresponding to the average Te concentration of 1.36%, are displayed in Table II (bottom panel) in good agreement with the experimental data for 1.0% and 1.5% Te concentrations.

The corresponding first principles average displacement is equal to 0.46 Å.


