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The formation, stability, and suitability of n-type junctions in germanium formed by solid phase epitaxial recrystallization

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Design and optimization of n-type doped regions in germanium by solid phase epitaxial recrystallization (SPER) have been studied by the authors. A systematic study is presented of process variables that influence activation and thermal stability, including preamorphization, coimplants, recrystallization temperature, and postrecrystallization thermal treatments. Unlike silicon, activation after recrystallization in germanium is not optimum where the postrecrystallization thermal budget is kept to a minimum. With the aid of modeling, a maximum peak activation of $7 \times 10^{19}$ cm$^{-3}$ was extracted. A steady increase in sheet resistance during postrecrystallization anneals confirms the formation of metastable activation by SPER. It is predicted that active concentrations of $6 - 8 \times 10^{19}$ cm$^{-3}$ are sufficient to meet targets for sub-20 nm technologies. © 2010 American Institute of Physics. [doi:10.1063/1.3452345]

As germanium provides much higher carrier mobilities than silicon, interest in this semiconductor has risen sharply in recent times, especially in the ongoing battle to boost performance and scaling.® In short, phosphorus (P) and arsenic (As) are relatively difficult to activate and diffuse quickly, leading to high resistances and limited capability to reduce the device dimensions.

Solid phase epitaxial recrystallization (SPER) is advantageous in this regard, as the low-temperature process limits dopant diffusion, while metastable solubility can be generated higher than equilibrium solubility.® A low-temperature process also reduces the risk of substrate loss and dopant out-gassing, which have been flagged as critical concerns for processing germanium substrates.® SPER has recently been explored for p-type and n-type dopant activation in germanium but a number of key issues remain. Deactivation kinetics have been studied in silicon but this aspect is largely unexplored at this point of time for n-type dopants in germanium. Moreover, optimization of process variables such as implants and recrystallization temperatures require fine tuning. Finally, it must be determined whether the dopant activation levels produced by SPER will be useful for future generation complementary metal-oxide semiconductor technologies.

Note that germanium is easier to amorphize than silicon, possibly linked to less dynamic annealing during ion implant. Amorphization occurs even during a 5 $\times 10^{13}$ cm$^{-2}$ dose P implant.® Consequently it is expected that n-type implants required for highly doped regions will amorphize the germanium substrate. In this work we focus on P, to be consistent with most experimental studies to date in this field.

Experiments were performed on (100) n-type germanium wafers, with bulk resistivity of 0.2–0.5 Ω cm. After a standard clean, wafers received a boron implant with a dose of $1 \times 10^{13}$ cm$^{-2}$ and energy of 60 keV, and a P implant with a dose of $1 \times 10^{15}$ cm$^{-2}$ and energy of 15 keV which amorphizes to a depth of 33 nm.® In some cases wafers first received a germanium preamorphizing implant (PAI) with a dose of $1 \times 10^{14}$ cm$^{-2}$ and energy of 150 keV which amorphizes to a depth of 120 nm.® One wafer received a fluorine (F) implant with a dose of $1 \times 10^{15}$ cm$^{-2}$ and energy of 10 keV. Finally, one wafer received only 5 $\times 10^{14}$ cm$^{-2}$ P dose but had an additional As implant with the same projected range, with a dose of $5 \times 10^{13}$ cm$^{-2}$ and energy of 28 keV. All implants were performed at a tilt of 7°, with a native oxide covering the substrate. Wafers were subsequently annealed for SPER in an inert ambient for 3 min at either 400 or 500 °C. Intrinsic germanium recrystallization rates are ~2 nm/s and 200 nm/s, respectively, for these temperatures, and so the SPER anneals should avoid incomplete recrystallization. All the post-SPER anneals were performed at 400 °C, in additive increments of 10, 30, 100, and 300 min. Based on the model of Ioannou et al., the 1.3 nm of substrate loss is expected for a 400 °C 443 min anneal.

Anneal temperatures were kept in the 400–500 °C range, as P diffusion is known to occur quite rapidly already at 600 °C.® Quantifying the expected diffusion is difficult as available diffusivity values were either extracted under intrinsic conditions, or at higher temperatures than studied here.® However, extrapolating diffusivity values from Chui et al., which are accurate for high concentrations, only a few nanometers of diffusion could be expected for the highest thermal budgets in this work.

Figure 1 shows experimental sheet resistance ($R_s$) data as a function of implant and anneal. The data is divided into two groups, namely, those recrystallized at 400 °C and those recrystallized at 500 °C. $R_s$ is lower when recrystallization is done at a higher temperature, which is in contrast to silicon where a low thermal budget for recrystallization is generally accepted as being the best approach. This trend was also evident in the recent results of Chao and Woo.® The gradual increase in $R_s$ in Fig. 1(b) with post-SPER annealing indicates the destruction of metastable activation. It is assumed that the lack of this trend in Fig. 1(a) is due to lower
activation created by SPER at 400 °C. The inclusion of coimplants such as PAI and F have a minor impact on the single P implant approach. Combining P with As leads to higher Rs values.

Equilibrium and metastable solubility and activation can be hindered by dopant-defect cluster formation. P tends to combine with vacancies (V) in germanium to form PnVm clusters.28,29 The initial Rs value may be influenced by the formation of PnVm or other grown-in complexes in amorphous germanium. Formation of vacancy-rich clusters is known to occur in amorphous silicon, as FmVc clusters can nucleate during SPER.30 Higher SPER temperature could limit the formation of these clusters and complexes. Using even higher temperatures (>500 °C) may not provide an added benefit, as for the same P implant (1×10^{15} cm^{-2}, 15 keV), high-temperature flash31 and laser32 anneal do not beat SPER at 500 °C in terms of the Rs versus junction depth (Xj) tradeoff.

Deactivation kinetics in post-SPER silicon have been linked to interstitials (I) emitted from the end-of-range (EOR) defect band interacting with the dopant atoms, and the formation of dopant-defect clusters, the rate of which depends on the thermal budget of the postanneal,27 the location of the EOR band with respect to the dopant,22 and the presence or not of a coimplanted nondopant species.23 In germanium, the EOR defects are relatively short lived and do not necessarily follow a classical Ostwald ripening process.27 However, due to the greater likelihood of PnVm cluster formation, rather than PnIm formation, the thermal stability of these complexes depends on the vacancy population, which often relies on thermal generation and injection from surfaces. Techniques that control these processes need future exploration.

In order to interpret the Rs data more thoroughly and evaluate the activation levels generated by SPER, modeling was undertaken using The Stopping and Range of Ions in Matter (SRIM).32 P implants with 1×10^{15} cm^{-2} dose were modeled with energies ranging from 0.5 to 20 keV, shown in the inset of Fig. 2. Rs is defined as $R_s = 1/\int_0^q \mu N_D dx$, where $X_j$ is junction depth, $q$ is electronic charge, $\mu$ is electron mobility and is a function of active donor concentration, N_D. Using the mobility data of Fistul et al.33 Rs was extracted for each implanted profile for varying N_D, assuming a diffusion-less anneal. Figure 2 shows that Rs decreases with increasing N_D and implant energy as expected. The lowest and highest experimental Rs values extracted, namely, 128.9 and 336.9 Ω/sq are plotted on the 15 keV trend line in Fig. 2. From this plot we can extract the corresponding active levels of 7×10^{19} cm^{-3} and 1.1×10^{19} cm^{-3}, respectively.

This analysis can be more useful by considering the Rs versus Xj tradeoff. Assuming an extended planar bulk architecture, channel concentration increases with MOS device and junction scaling. For the P profiles of Fig. 2, in accordance with specs in the ITRS Roadmap 2000–2009 editions,34 $X_j$ is the depth at concentrations of 5×10^{18} cm^{-3}, 2.5×10^{18} cm^{-3}, and 1×10^{18} cm^{-3}, for the 0.5–2 keV, 5–10 keV, and 15–20 keV profiles, respectively. Rs versus Xj is plotted in Fig. 3, in the form of constant N_D contour lines. The corresponding P implant energies are included on the x-axis. ITRS Roadmap target specs are added from the 2009 and 2003 editions. The 2009 edition specs are more relevant as those are the targets that industry is shooting for at this point in time. However, due to the lack of development of scaled implanted n-type junctions in germanium, to date published Rs versus Xj data is limited to $X_j$ > 60 nm.26,31,35 2003 specs are a useful guide as they concern junctions with $X_j$=28–75 nm. The afore-mentioned high-temperature flash and laser anneal data are also shown in Fig. 3.26,31 Note Heo et al.36 reported Rs versus Xj data with $X_j$<60 nm, by combining PH3 plasma doping with laser anneal.

According to the ITRS, Rs is a designated percentage of total source-drain resistance ($R_{SD}$). While Rs is not specified for different device options, $R_{SD}$ is defined with the same value for high performance (HP), low-operating power.
(LOP), and low-standby-power (LSTP) devices. Thus here $R_s$ for LOP and LSTP are taken as those listed in the ITRS Roadmap for HP devices. $X_j$ is a function of gate length ($L_{\text{gate}}$) which does change with device option. In Fig. 3 $R_s$ versus $X_j$ targets are separated for the 2003 ITRS HP, LOP, and LSTP devices. Clearly a higher active concentration is required for HP devices.

For the 2009 specs there was very little difference in targeted $L_{\text{gate}}$ and $X_j$, thus for clarity only the most aggressive target is shown in Fig. 3. $P$ implants followed by a diffusionless anneal, such as SPER, can meet the extended planar bulk ITRS specs for $L_{\text{gate}} = 17–22$ nm with an active concentration of $8 \times 10^{19}$ cm$^{-3}$. If a germanium-on-insulator architecture is used for $L_{\text{gate}} = 17–22$ nm, the resulting junction profiles can be assumed to be boxlike. The procedure of Fig. 3 was repeated for that case, and an active concentration of $6 \times 10^{19}$ cm$^{-3}$ was sufficient to hit the 2009 ITRS targets. Note, the plot is not shown here due to its similarity to Fig. 3.

In conclusion, the typical problems of n-type impurities in germanium such as diffusion, substrate loss, and outgassing can be overcome by using SPER for dopant activation. Recrystallization at 500 °C produces an active concentration of $7 \times 10^{19}$ cm$^{-3}$ which is in the order of that needed for sub-20 nm technologies.

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