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Undoped and *in-situ* B doped GeSn epitaxial growth on Ge by atmospheric pressure-chemical vapor deposition

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In this letter, we propose an atmospheric pressure-chemical vapor deposition technique to grow metastable GeSn epitaxial layers on Ge. We report the growth of defect free fully strained undoped and *in-situ* B doped GeSn layers on Ge substrates with Sn contents up to 8%. Those metastable layers stay fully strained after 30 min anneal in N₂ at 500 °C; Ge-Sn interdiffusion is seen at 500 °C but not at lower temperature. B is 100% active in the *in-situ* GeSn:B layers up to a concentration of $1.7 \times 10^{19} \text{ cm}^{-3}$. GeSn:B provides slightly lower Hall hole mobility values than in pure p-type Ge especially for low B concentrations. © 2011 American Institute of Physics. [doi:10.1063/1.3645620]

GeSn has received many interests in the last 10 years for various applications.^{1–3} An indirect to direct bandgap transition is theoretically expected for about 10% Sn in GeSn.⁴ Sn having about 13% lattice mismatch with Ge, GeSn alloys can offer interesting new routes for stress implementation: (1) tensely biaxial strained Ge layers⁵ on GeSn provides a direct bandgap and an enhanced electron mobility with respect to unstrained Ge, (2) compressively biaxial strained GeSn layers grown on Ge can provide a strained quantum well architecture for advanced metal oxide semiconductor field effect transistors,⁶ and (3) GeSn can be used moreover as a source/drain stressor materials for advanced Ge pMOSFET technology.⁷

Sn solubility in the Ge matrix is however very poor⁸ (less than 1%). Based on this thermodynamics limitation, the first monocrystalline GeSn growth techniques developed were based on out of equilibrium growth conditions using ultimately low temperatures and reduced pressures.^{9,10} Whereas the necessity of growing GeSn at low temperature is well understood (according to GeSn phase diagram), the necessity of using reduced pressure to enable metastable GeSn growth has never been confirmed experimentally. This letter shows metastable GeSn epitaxial growth on Ge substrate, with Sn content up to 8%, using an atmospheric pressure-chemical vapor deposition (AP-CVD) technique. We actually prove that reduced pressure is definitely not the key factor for growing GeSn alloys.

200 mm blanket 1 μm Ge/Si substrates were first realized with the procedure detailed in (Ref. 11). After a HF (2%) wet clean the wafers were loaded again in a 200 mm ASM Epsilon-like CVD reactor. In order to allow GeSn growth at very low temperature, Ge₂H₆ was used as Ge precursor and SnCl₄ as an innovative Sn precursor. SnCl₄ is liquid at room

temperature and was evaporated by using a vapor station connected to the CVD reactor. The main advantage of using a SnCl₄ liquid source, as compared to SnD₄ (Ref. 12) (only Sn precursor so far reported in the literature for GeSn CVD) is the total absence of instability issues (gas consumption). Moreover, SnCl₄ is a commercially available product. After a Ge pre-growth bake at 650 °C, the growth temperature was fixed to 320 °C. Fig. 1 shows a cross section transmission electron microscope (TEM) picture with associated (224) x-ray diffraction reciprocal space mapping (XRD RSM) of a 40 nm thick GeSn layer grown on a Ge substrate using atmospheric pressure. From Rutherford backscattering (RBS) measurements, a Sn content of 8% was determined in this layer. The TEM picture in Fig. 1(a) exhibits a defect free and high crystalline quality for the 40-nm-thick GeSn layer. Furthermore, the surface quality of the as-grown Ge_{0.92}Sn_{0.08}/Ge/Si heterostructure was investigated by reflection high energy electron diffraction (RHEED) analysis after *ex-situ* transfer to a MBE system. An annealing in ultrahigh vacuum up to ~420 °C resulted in an oxide-free GeSn surface showing a strong (2 × 1) surface reconstruction as seen on RHEED pattern along the [110] azimuth (Fig. 1(b)). Finally, the XRD-RSM around the (2 2 4) Bragg reflections (Fig. 1(c)) demonstrate that the grown GeSn layer is fully strained on Ge/Si (001) substrate. This result shows the fabrication of GeSn materials using an AP-CVD technique with a stable and commercially available Sn precursor.

The deposition temperature for the GeSn growth was kept low (320 °C) in order to allow Sn incorporation in Ge lattice without Sn precipitation nor agglomeration.^{10,13} Further anneals were then done after growth on the GeSn layers in order to assess their thermal stability. Anneals were performed in a rapid thermal anneal equipment at 400 °C and 500 °C for 10 and 30 min in pure N₂. Fig. 2(a) first compares (004) omega-2 theta scans on the annealed samples with the as-grown one. Although a clear GeSn XRD peak is present on all the samples, we note that the GeSn peak fringes disappear for anneals realized at 500 °C. Some interface

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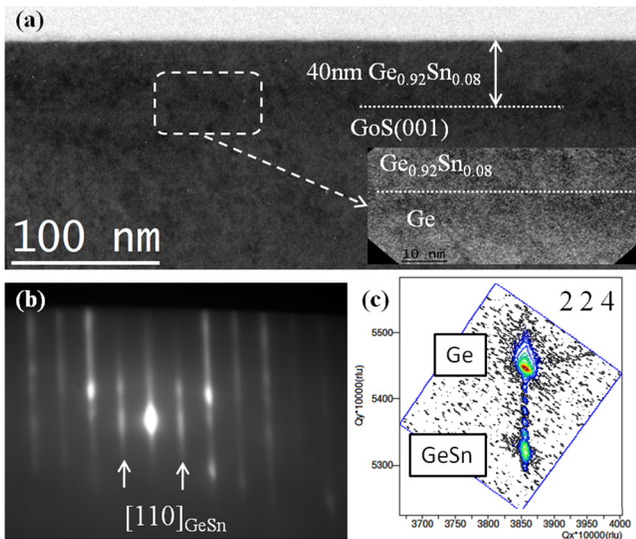


FIG. 1. (Color online) (a) Cross Section TEM of a 40 nm fully strained defect free GeSn layer on $1 \mu\text{m}$ Ge/Si buffer substrate with 8% Sn grown with AP-CVD using combination of Ge_2H_6 and SnCl_4 . (b) RHEED diagram of the $\text{Ge}_{0.92}\text{Sn}_{0.08}$ surface after deoxidation in UHV at 420°C . The pattern exhibits a strong (2×1) surface reconstruction along the $[110]_{\text{Ge}}$ direction. (c) (224) XRD-RSM of the 40 nm $\text{Ge}_{0.92}\text{Sn}_{0.08}/\text{Ge}$ bilayer showing that GeSn is fully strained on Ge.

roughening is then likely to occur for samples annealed at temperatures as high as 500°C . During optical microscope inspection (not shown here), we did not however notice any difference in surface morphology before and after anneals. The presence of clear GeSn XRD diffraction peaks confirms that the GeSn layer stays monocrystalline after anneals. Additional RSM scans (not shown here) moreover show that GeSn is also kept fully strained on Ge after anneals. We note that an important difference is observed on the XRD measurement (omega-2theta scan and also RSM) for the sample annealed at 500°C during 30 min. Both GeSn and Ge peaks are wider and a shift of the GeSn peak is observed. Being not identified, strain relaxation can then be the reason for such XRD peaks widening. Secondary ion mass spectroscopy (SIMS) was then realized on the as-grown and annealed GeSn/Ge structures to detect possible Ge-Sn interdiffusion. Fig. 2(b) compares Sn SIMS profiles in the GeSn/Ge structures before and after anneals. The Sn slope is identical in all the samples except for the one annealed at 500°C during 30 min. A clear signature of Sn diffusion in Ge is observed, responsible for the difference observed in the previous XRD measurements. We note that this diffusion signature might be caused by Sn precipitation within the GeSn layer with this anneal condition. We then conclude that the metastable 8% Sn-GeSn layer grown with our CVD approach survive to further thermal treatments at temperatures up to 500°C and duration as long as 10 min. No GeSn strain relaxation is observed after anneals but Ge-Sn interdiffusion (possibly enhanced by Sn precipitation) starts for longer anneals (30 min) at 500°C .

In-situ B doped GeSn AP-CVD growth was also investigated by combination of Ge_2H_6 , SnCl_4 , and B_2H_6 precursors. The growth of GeSn:B is of high interests for different kind of devices.^{7,14} Growth conditions were kept again at 320°C and atmospheric pressure, keeping the same $\text{Ge}_2\text{H}_6/\text{SnCl}_4$

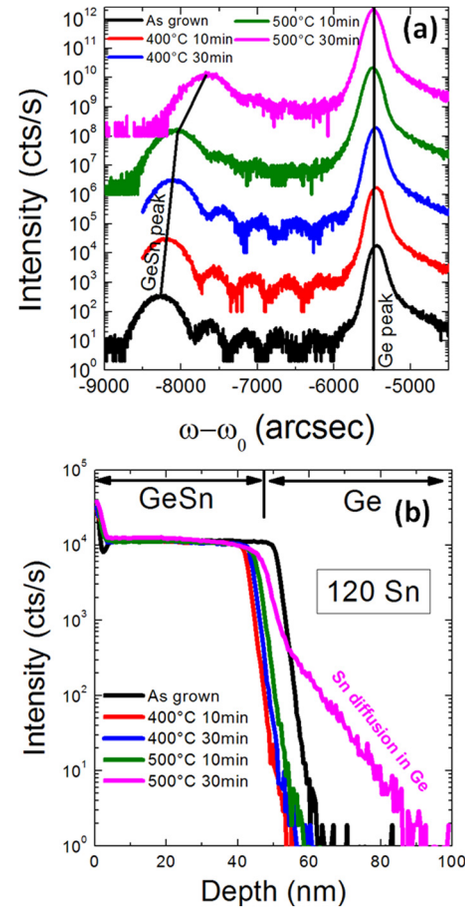


FIG. 2. (Color online) (a) Comparison of (004) omega-2 theta scan of as-grown and annealed $\text{Ge}_{0.92}\text{Sn}_{0.08}/1 \mu\text{m}$ Ge/Si with different anneal conditions: 400°C -10 min, 400°C -30 min, 500°C -10 min, and 500°C -30 min. (b) Comparison of Sn profile in the $\text{Ge}_{0.92}\text{Sn}_{0.08}/1 \mu\text{m}$ Ge/Si structure as function of anneal conditions: clear Sn diffusion in Ge is observed for an anneal performed at 500°C for 30 min.

flow ratio (giving 7% Sn in GeSn alloy for undoped layers) and adding B_2H_6 with different partial pressures during the growth. This time, 80 nm GeSn:B layers were grown on $1 \mu\text{m}$ slightly n-type doped ($1 \times 10^{17} \text{cm}^{-3}$) Ge/Si blanket 200 mm wafers. Fig. 3(a) shows the B concentration, determined by SIMS, for B_2H_6 partial pressure varying from 0.18 to 3.7 mTorr. Surface roughening and loss of crystallinity (characterized by XRD) were observed for higher B_2H_6 partial pressure. The total B concentration in the GeSn layer increases almost linearly from 5×10^{17} to $1.7 \times 10^{19} \text{cm}^{-3}$ with B_2H_6 partial pressure. The Sn content, determined by RBS (also shown in Fig. 3(a)), slightly decreases by incorporation of B (from 7.2% to 6.83%) and goes down dramatically to 4.68% for the highest B_2H_6 partial pressure. The drop from 6.83% to 4.68% is attributed to B_2H_6 surface poisoning which lowers Sn incorporation, additionally to surface degradation. For partial pressures in between 0.18 and 1.85 mTorr, a competition between B and Sn might occur during the growth of ternary GeSn:B alloys explaining a slight decrease in Sn content when B is incorporated. This competition actually occurs on the substitutional sites of the GeSn matrix. Using GeSn:B mobility values measured by micro-Hall measurements,¹⁵ reported in Fig. 3(b), and sheet resistances measured by the micro-four-point-probe (M4PP) technique,¹⁶ the active concentration of B in GeSn was

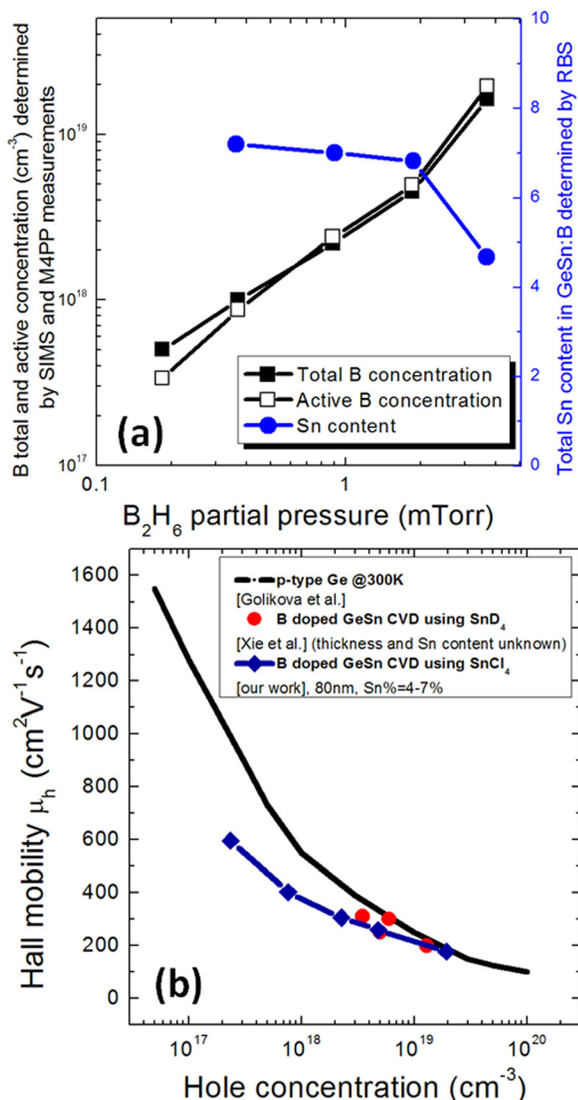


FIG. 3. (Color online) (a) Total B concentration (determined by SIMS), active B concentration (determined by M4PP measurements), and Sn content (determined by RBS) in GeSn:B layers grown on $1 \mu m$ n-type Ge/Si structures as function of B_2H_6 partial pressure: B concentration increases linearly with B_2H_6 partial pressure while Sn content is decreasing. (b) Hole Hall mobility measurements in the GeSn:B layers grown as function of the B concentration showing slightly lower mobility values than in the p-Ge bulk case.

calculated and is reported as function of the B_2H_6 partial pressure, also in Fig. 3(a). The active concentration was calculated assuming GeSn dielectric constant and intrinsic carrier concentration equal to that of Ge. We note that even for high B_2H_6 partial pressures, the B is 100% active in the *in-situ* doped GeSn:B layers grown. This means that B is incorporated in the GeSn entirely in substitutional sites taking the place of either Ge or Sn atoms. Most likely, B prefers to occupy Sn sites in order to compensate the stress within the layer (B atoms are smaller than Ge atoms which are smaller than Sn atoms) which might explain the lower Sn incorporation for highly B doped GeSn layers. We note however that the increase of B concentration in the GeSn:B layer does not compensate entirely the decrease in Sn concentration which means that a lower $SnCl_4$ partial pressure can also contribute to lower Sn incorporation. The Hall mobility values measured in the GeSn:B layers are compared with data from liter-

ature on GeSn:B layers grown by Xie *et al.*¹⁷ Our values are in good agreement with Xie's measurements and show slightly lower Hall mobility values than mobilities in pure p-type Ge bulk materials.¹⁸ This mobility degradation is likely due to alloy scattering in the Ge-Sn system. For highly doped GeSn and Ge, impurity scattering becomes the predominant cause for carrier scattering which explains that Ge and GeSn mobility values converge for B concentration higher than $3 \times 10^{18} cm^{-3}$.

In conclusion, this letter demonstrates the epitaxial growth of both undoped and *in-situ* B doped GeSn using a AP-CVD technique. We proved that reduced pressure is not mandatory and that GeSn can be made just by proper gas mixture of well chosen Ge and Sn precursors. This technique, using a production-like environment and commercially available gas precursors, opens new routes for research in group IV semiconductors. Sn can today definitely join C, Si, and Ge in the catalog of high potentials group IV materials.

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