

Quantitative Measurement of Reduction of Phosphorus Diffusion by Substitutional Carbon Incorporation

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Abstract

Complete suppression of transient enhanced boron diffusion (TED) and oxidation enhanced boron diffusion (OED) in silicon have been achieved using substitutional carbon to reduce the excess point defect concentration in the dopant region [1]. Recent efforts have focused on removing the carbon from the active regions of the device [2,3] to avoid device degradation due to electrically active carbon defects [4] and exploring remote carbon's effect on boron TED and OED, while using the boron diffusion to probe point defect concentrations.

In this paper we measure quantitatively the effect of remotely located carbon on phosphorus and boron diffusion above a buried SiGeC layer at 850°C in oxygen or nitrogen ambients. Remote carbon, located 1250 Å below the phosphorus edge, is found to reduce the phosphorus diffusion enhancement factor due to OED from 8 to 2. The effect of the remotely located SiGeC buried layer on the excess interstitial concentration profile, which is responsible for the enhanced dopant diffusion, is probed by measuring boron and phosphorus diffusivities of in-situ doped boron and phosphorus layers above a buried SiGe(C) layer after oxidation or nitrogen anneals at 850°C. The enhanced boron diffusivity during oxidation is found to have a near linear dependence on depth ranging from 5-1.25. Finally, using x-ray diffraction and photoluminescence measurements of as-grown, buried, strained SiGe(C) structures and annealed SiGe(C) structures in oxygen or nitrogen ambient at 850°C the number of substitutional carbon atoms effectively consumed by oxidation is unambiguously correlated to the absolute number of injected interstitials using published values for the interstitial injection rate during oxidation [5].

Introduction

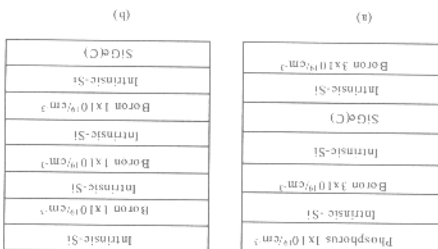
As silicon devices are continuously scaled to smaller sizes, there is an ever greater demand to reduce the diffusion of dopants. However, the enhancement of boron diffusion due to ion implant damage and oxidation, transient enhanced diffusion (TED) and oxidation enhanced diffusion (OED), poses severe challenges to this goal. It is generally accepted that boron and phosphorus diffuse primarily via an interstitial mechanism and the two dopant diffusivities are, therefore, dependent on the silicon interstitial concentration. Ion implant damage and oxidation are sources of excess interstitials which enhance the diffusivity. Therefore the reduction of TED and OED depend on the effective ability to trap or annihilate any excess interstitials that are produced during processing. Interest in the relationship between carbon and enhanced boron diffusion was recently spurred after TED was observed to strongly depend on the concentration of substitutional carbon in silicon [6], and in SiGe alloys [4,7]. Interstitial carbon can, however, introduce undesirable stable electrical defects which can degrade device quality [4,8], and therefore, the effects of remotely located substitutional carbon on boron OED and TED are being examined [2,3]. This work quantifies the dependence of phosphorus and boron diffusion above a SiGeC layer where point defects, interstitials, are injected during oxidation. Introduction of a remote, buried, SiGeC layer located below a doped surface region is found to reduce OED of both boron and phosphorus markers in the overlying region. A non-uniform interstitial profile is observed during oxidation in samples with buried SiGeC, determined using

dopant diffusivities as a probe to monitor the excess interstitial carbon concentrations in the surface region. Dopant diffusivities are found to depend on the subsurface carbon concentration in the buried SiGeC layer, and lower carbon concentrations are observed to be saturated by the dose of injected point defects. A deeper understanding of the point defect relationship with the TED both above and below the remote SiGeC layer. The subsurface carbon dependence on TED necessary carbon concentrations to optimize the reduction of boron and phosphorus OED and point defect injection is, therefore, examined using x-ray diffraction and photoluminescence measurements of the SiGeC layer. The deduced subsurface carbon concentration is found to linearly depend on time of oxidation and the dependence on oxidation time is compared with the expected net interstitial dose injected during oxidation [8] to determine a relationship between injected interstitials and subsurface carbon consumed. A ratio of one injected boron atom to 1.85 ± 0.7 substitutional carbons effectively removed from their substitutional sites is found.

Experiment

All test structures were grown using rapid thermal chemical vapor deposition (RT-CVD) [7], between 600 and 750°C using methylsilane as the carbon source. In test structure A, see figure 1 (a), two boron doped silicon layers and one phosphorus layer at the surface with and without a Si_{1-x}Ge_xCy or Si_{1-x}Ge_x layer placed between the boron

Figure 1 (a) Test structure to study boron and phosphorus oxidation enhancement
of boron nitride. This paper discusses only the top phosphorous edge and deepest
distribution. The paper discusses only the top phosphorous edge and deepest
boron nitride. (b) Test structure to examine the resulting non-uniform
interstitial profile during oxidation resulting from introduction of the boron
nitride.



(a)

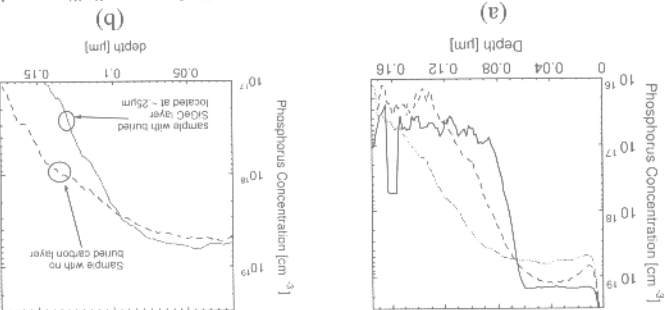
(b)

layers were used to test the effect of the alloy layers on dopant diffusivities at different locations (above and below) with respect to the SiGeC layer. The phosphorous layer was grown 650Å thick and doped to $1 \times 10^{19}/\text{cm}^3$, immediately below the surface. Both boron peaks were approximately 250Å thick and were grown to have a peak boron concentration of $5 \times 10^{19}/\text{cm}^3$ centred 2000Å and 3000Å away from the surface respectively, while the boron nitride was grown 250Å thick and doped to $1 \times 10^{19}/\text{cm}^3$, immediately below the surface. The phosphorous layer was 250Å thick and measured in nitrogen ambient atmosphere at 850°C for 30 minutes. The phosphorous and boron profiles were characterized using secondary ion mass spectrometry (SIMS) with 2 KeV Cs⁺ ions and an estimated 1–5% error in depth scales. The boron has previously been reported [2]. Boron diffusion above the SiGeC layer is effected by the substitutional carbon incorporation on the boron diffusion below the SiGeC layer.

overlying phosphorous edge in an all-silicon control sample, with a background carbon concentration below SiMS detection limits. Broadening of the phosphorous edge is observed after the nitrogen anneal, and in the oxidation case clear enhancement of the phosphorous diffusion is observed by the deeper location of the phosphorous edge relative to the average phosphorous diffusivity during the nitrogen anneal. In figure 2b, the phosphorous profiles after oxidation in the all-silicon case and that a buried $\text{Si}0.79\text{Ge}0.20\text{C}0.005$ are overall. A $4\times$ reduction of phosphorous diffusion in the sample containing the buried SiGe under the same oxidizing conditions is observed, even when the SiGe layer is buried 1500 Å below the phosphorous edge. A summary of the measured diffusivities after oxidation relative to the phosphorous edge. A summary of the measured diffusivities after the silicon case and two carbon concentrations of 0.5% and 0.5% .

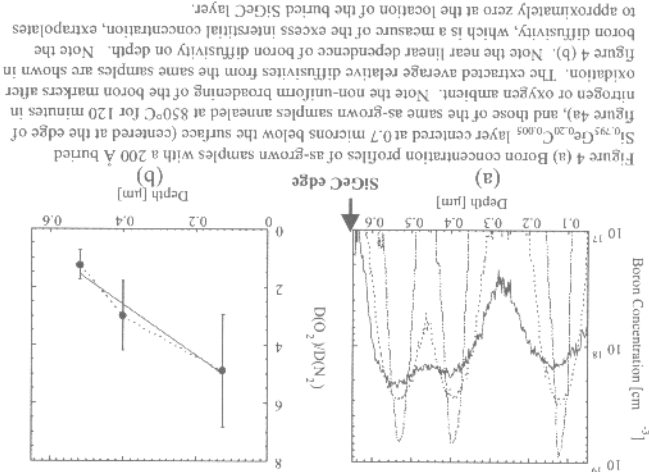
ODF of boron in the marker located below the SiGe region was observed to be completely suppressed in the 0.5% carbon case but not in the 0.5% carbon case [2], this indicates that in the 0.5% case some interstitials injected at the surface, during oxidation, pass through the SiGe blocking layer, whereas in the 0.5% case approachability all excess interstitials injected during oxidation were reacted before penetrating to the underyielding silicon region. The interstitial concentration above and below the SiGe layer is seen to depend on the reaction efficiency of the SiGe layer with the SiGe layer is diffusivity between the surface and the SiGe layer during oxidation, (fig. 1b). Three boron markers are located above a buried $\text{Si}0.79\text{Ge}0.20\text{C}0.005$ layer to probe the interface.

Figure 2 (a) Phosphorus concentration profiles (measured and calculated by SIMS) of all-silicon sample and all-Si₃N₄ sample annealed at 850°C for 30 minutes in nitrogen ambient. Phosphorus diffused into the samples from the back side of the wafer. The phosphorus profile is very similar for both samples. Note phosphorus diffuses into the silicon sample faster than phosphorus profile of phosphorus-doped silicon sample.



and quantification of its diffusion is neglected in this paper. Figure 2a shows profiles of the concentration across the alloy interface and segregation effects, so the discussion

The ability to fit the interstitial profile to near zero at the SiGeC layer depends on the Si_{0.79}Ge_{0.20} layers centred 800 Å below the surface were epitaxially grown to examine the substitutional carbon dependence on anneal times in nitrogen and oxygen ambient. Two completely undoped samples with either 400 Å thick buried Si_{0.8}Ge_{0.2} or Si_{0.79}Ge_{0.20} layers containing enough substitutional carbon to react with the interstitials injected during oxidation. To determine how much carbon is required a third set of test structures were grown. Figure 4(b) shows the average relative boron diffusivities from the same samples after oxidation. The extracted average relative boron diffusivity on depth. Note the nitrogen or oxygen ambient. Note the non-uniform broadening of the boron markers after figure 4(a), and those of the same as-grown samples below the surface (centered in the edge of Si_{0.79}Ge_{0.20}) layer centred at 0.7 microns below the surface (centered in the edge of Si_{0.79}Ge_{0.20}) and those of the same as-grown samples below the surface (centered in the edge of Si_{0.79}Ge_{0.20}) layer centred at 200 Å buried in nitrogen or oxygen ambient. Note the non-uniform broadening of the boron markers after oxidation. The extracted average relative boron diffusivities from the same samples after oxidation. Boron concentration profiles of as-grown samples with 200 Å buried in nitrogen or oxygen ambient. Note the non-uniform broadening of the boron markers after oxidation. Boron concentration profiles of as-grown samples with 200 Å buried in nitrogen or oxygen ambient. Note the non-uniform broadening of the boron markers after oxidation.



observed to have a near linear dependence on depth, see figure 4. SiGeC layer. Boron diffusivities are extracted from the boron profiles and the boron markers located closer to the clearly broader and its peak concentration is lower than those markers located closer to the depth from surface to the SiGeC alloy layer. The boron profile closer to the oxidation, the width of the three diffused boron markers is clearly not uniform over the profiles in the oxidation case are clearly broader than those in the nitrogen ambient. After growth peak is observed in both the nitrogen and the oxygen cases, however the boron minutes at 850°C in oxygen and nitrogen ambients, see figure 4. Broadening of the as-concentration in the surface region. As-grown samples were cleaved and annealed for 120 minutes at 850°C in oxygen and nitrogen ambients, see figure 4. Broadening of the as-

ambient is compared to that measured in nitrogen ambient.
SiGeC layer. Average relative boron diffusivity during anneal at 850°C for 30 min in oxygen

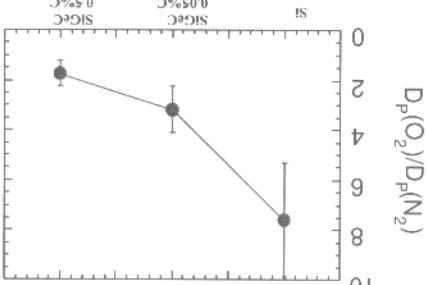
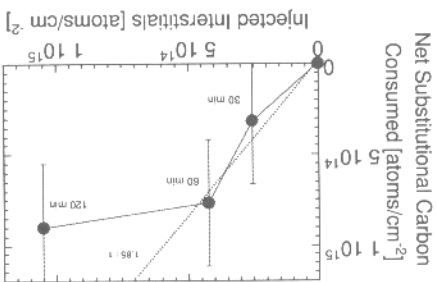


Figure 6. The injected interstitial carbon measured by type II loop defect growth [Meng et al.] is correlated to the consumed substitutional carbon measured by x-ray diffraction for oxidation times of 30, 60, and 120 minutes at 850°C.



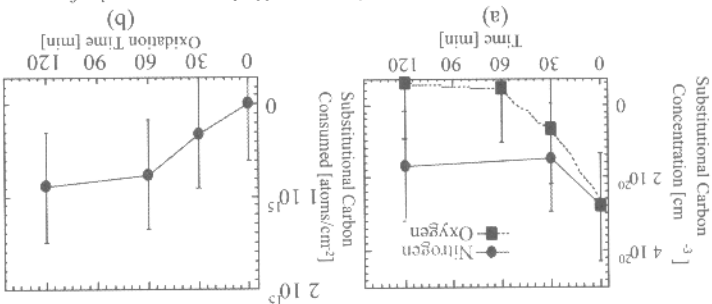
The total consumed carbon rate may be calculated from the measured substitutional carbon concentration's dependence on time. The net injected interstitials during oxidation

Discussion

oxidation time, see Figure 5.

diffusion is found to depend on annealing conditions and to decrease with increasing annealing time. The effective substitutional carbon concentration determined by x-ray diffraction measurements and found to agree well with one another, within 5% for all anneal conditions or band-gap measurement methods respectively. Using either the lattice photoluminescence and x-ray diffraction methods, the total substitutional carbon was deduced for each of the anneal conditions and found to agree well with one another, within 5% for all anneal conditions or band-gap measurement methods respectively. The net injected interstitials during oxidation time, see Figure 5.

Figure 5 (a) substitutional carbon measured in the as-grown sample of an undoped, buried, 400 Å thick $\text{Si}_{0.793}\text{Ge}_{0.206}\text{C}_{0.005}$ layer and in the same as-grown samples of the 400 Å layer measured by x-ray diffraction. Note that the lattice constant of the 400 Å SiGe layer measured by x-ray diffraction. The lattice constant of the substitutional carbon measured after 60 minutes of annealing in an oxygen ambient, but not after annealing in nitrogen ambient. Figure 5 (b) describes the difference of consumed substitutional carbon atoms after oxidation deduced from Figure 5 (a) and the as-grown substitutional carbon atoms after oxidation deduced from Figure 5 (b).



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Acknowledgements

The introduction of substitutional carbon in buried SiGeC layers is found to reduce the oxidation rate of both boron and phosphorus located above the SiGeC layer. A near linear enhanced diffusion of both boron and phosphorus is observed during oxidation of substitutional carbon in the SiGeC alloy is a buried SiGeC layer, and the diffusivity is greater than the surface. The rate of substitutional carbon consumed by oxidation was measured by x-ray diffraction and photoluminescence versus annealing times and ambient. Using published values of interstitial oxidation versus oxidation rate, the ratio of substitutional carbon atoms effectively driven from their substitutional sites to injected interstitial is correlated to be 1.85 ± 0.7 .
 The authors are grateful for enlightening discussions with H.-J. Gossman, and C. S. Rafferty. M. S. Carroll is indebted to T. Buyukimanolu and C. Magee at Evans East for assistance with SIMS analysis. License to use PROPHET from Lucent Technologies was granted from their substitutional sites to injected interstitial is correlated to be 1.85 ± 0.7 .
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Conclusion

was determined by measuring the growth of loop defects during oxidation at 900°C by Meng et al [8]. Assuming that the point defect injection during oxidation scales with oxidation rate, the injection rate of interstitials may be calculated for oxidation at 850°C from the measured interstitial injection at 900°C. In figure 6 the consumed carbon rate is shown correlated to the calculated interstitial injection rate by a ratio of 1.85 ± 0.7 to 1 which just satisfies the simple relationship of one interstitial interaction in the SiGeC alloy is more completely consumed by the reaction.