Growth of Silicon-Germanium Alloy Layers

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1. INTRODUCTION

The advancements in crystal growth technologies, such as molecular beam epitaxy (MBE), gas source molecular beam epitaxy (GSMBE), organometallic vapour phase epitaxy (OMVPE), and chemical vapour deposition (CVD) have enabled ultra thin epitaxial semiconductor layers to be routinely grown with monolayer precision in thickness and composition control to about one atomic per cent. In the past, main focus was on homoepitaxial films and closely lattice-matched combination of materials such as $Al_{2}Ga_{1-x}As/GaAs$ and $In_{0.53}Ga_{0.47}P/InP$. The possibilities for lattice-matched heteroepitaxial systems are relatively limited as shown in Fig. 1. In the last decade, focus was placed on lattice mismatched combination of materials as these material systems also offer unique new prospects, such as the use of strain to modify electronic and optical properties. Recently, alloys of group IV elements, including carbon, silicon, germanium and tin have been actively pursued for use in heterojunction devices compatible with conventional silicon processing technology. Major advancements have been made in binary $SiGe/Si$ alloys where the built-in strain and the composition of a pseudomorphic $Si_{1-x}Ge_{x}$ layer on silicon substrate affect the band structure, energy gap as well as the band offset significantly. High performance devices and circuits based on $Si/Si_{1-x}Ge_{x}$ heterostructures have been demonstrated. There is now considerable literature on various aspects of $Si_{1-x}Ge_{x}$ epitaxial growth and devices.

Strain-induced modification of $Si/Ge$ films have a significant impact on the band structure and carrier transport. When a thin film with a larger lattice constant (e.g. $Si_{1-x}Ge_{x}$) is grown on a smaller lattice constant substrate (e.g. silicon), the film maintains in-plane lattice constant of the substrate and is under a biaxially compressive strain. Figure 2 shows the band offset between a strained $Si_{0.5}Ge_{0.5}$ film grown on silicon. This is known as the type-I band alignment where the entire band offset occurs in valence band [Fig. 2(a)] while band offset in conduction band is very small. This type of structure is favourable for hole confinement.
Table I. Summary of deposition systems

<table>
<thead>
<tr>
<th></th>
<th>MBE</th>
<th>Gas source MBE</th>
<th>UHVCVD</th>
<th>LRPCVD/RTCVD</th>
<th>VLPCVD</th>
<th>Atmospheric CVD</th>
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<td>10^{-10}</td>
<td>10^{-10}</td>
<td>10^{-3} to 10^{-5}</td>
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<td>(In situ) cleaning</td>
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<td>Yes</td>
<td>No</td>
<td>Yes</td>
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<td>Dopant gas</td>
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<td>B_{2}H_{6}, AsH_{3}, PH_{3}</td>
<td>B_{2}H_{6}, AsH_{3}, PH_{3}</td>
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<td>550-625</td>
<td>550-850</td>
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<td>Si_{1-x}Ge_{x}(x=per cent)</td>
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<td>0.1-100</td>
<td>&lt;40</td>
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<td>SiC, SiGe, SiGe, GeC, GeSn</td>
<td>SiC, SiGe, SiGe, GeC, GeSn</td>
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and has been exploited in several novel heterostructure devices, viz., buried channel p-MOSFETs, p-MODFETs and HBTs.

Similarly, a smaller lattice constant silicon epilayer will be under biaxial tension when grown on a larger lattice constant relaxed Si_{1-x}Ge_{x} substrate. Figure 2(b) shows the band offset for a strained silicon epilayer grown on a relaxed Si_{1-x}Ge_{x} substrate. In this case, type-II band offset occurs and the structure has several advantages over the more common type-I band alignment, as a large band offset is obtained in both the conduction and valence bands, relative to the relaxed Si_{1-x}Ge_{x} layer. This allows both electron and hole confinements in strained silicon layer making it useful for both n- and p-type devices for strained Si/SiGe-based CMOS technology. Since strained silicon provides both larger conduction and valence band offsets and does not suffer from alloy scattering (hence mobility degradation), a significant improvement in carrier mobility can be achieved. Strained silicon is more difficult to grow compared to strained Si_{1-x}Ge_{x}, since bulk Si_{1-x}Ge_{x} substrate is currently not available and until recently, growth of relaxed Si_{1-x}Ge without forming large concentration of defects due to dislocation was difficult. However, the ability to achieve both n-MOS and p-MOS devices using strained silicon provides a promising alternative for next-generation high performance SiGe-CMOS technology.

In this paper the growth of group IV binary alloy films has been reviewed. The deposition of heteroepitaxial films using various reactors has been critically examined. As the reactor configurations differ substantially, the advantages and disadvantages
of each system are compared. The deposition and characterisation of binary \( Si_{1-x}Ge_x \), \( Si_xGe_{1-x} \), \( Ge_{1-y}C_y \), and alloy layers containing tin are discussed. Growth of strained silicon on relaxed \( SiGe \) buffer layers, poly-SiGe film and hydrogenated amorphous SiGe (a-SiGe:H) film is also considered.

2. GROWTH OF BINARY ALLOY LAYER FILMS

\( SiGe \) alloys have attracted considerable attention during the last decade, and it is now recognised that the \( Si/Si_{1-x}Ge_x \) material system is the most practical route to heterojunction devices on silicon substrates. The present review emphasises \( Si/SiGe \) technology that can be integrated into silicon processes. The growth of other binary alloys, such as silicon-carbon, silicon-tin, and germanium-carbon for each of these material systems has its own merit. In applications of strained silicon layers, it is important to grow thick relaxed \( SiGe \) layers as buffer. Poly-Si\(_{1-x}Ge_x\) and hydrogenated amorphous SiGe (a-SiGe:H) films have applications as gate electrodes and interconnects in integrated circuits, thin film transistors (TFTs) for flat panel displays and high efficiency multijunction solar cells, respectively.

Most of the early work on binary \( Si_{1-x}Ge_x \) alloy films was carried out using MBE whereas growth using CVD systems started much later. A comparison of different deposition systems is given in Table 1.

2.1 Growth of \( Si_{1-x}Ge_x \) Films

In case of strained layer epitaxy, lattice mismatched strained layers can be grown without misfit dislocations if the thickness is kept below the critical thickness. The stability of strained layers was first studied by Frank and Van der Merwe\(^{12}\).

The term critical thickness was initially defined to denote the transition from a strained to relaxed layer. Using a continuum model, authors calculated the critical layer thickness, \( h_c \), below which the strained layer is expected to be in the thermodynamically stable state. Above \( h_c \), it is energetically favourable to relieve the strain via dislocations. Since then, there have been a number of reports on calculations of \( h_c \) differing from each other in the assumed energy stored in a dislocation. Van der Merwe\(^{17,18}\) was the first to calculate the critical thickness as a function of increased lattice mismatch by minimising

![Figure 2. Band alignments between silicon and \( Si_{1-x}Ge_x \) on two substrates: (a) silicon and (b) \( Si_{1-x}Ge_x \).](image-url)
fits their experimental data. Many other researchers have contributed for critical thickness theories based on energy, mechanical equilibrium, and kinetics of dislocations.22-24

The first trial of \( \text{Si}_{1-x}\text{Ge}_x \) growth on silicon was attempted in 1977 by Kasper25, et al. at 750 °C using MBE. As the temperature was high, three-dimensional growth took place and islanding was observed. The first commensurate growth of \( \text{Si}_{1-x}\text{Ge}_x \) layer on silicon was reported by Bean20, et al., who used a low temperature (550 °C) growth technique. At this low temperature, it was found that the critical thickness of the epilayer was several times higher than the value that can be obtained from the mechanical equilibrium theory. The key feature of this low temperature growth is that two-dimensional growth of strained \( \text{Si}_{1-x}\text{Ge}_x \) containing a high concentration of germanium is possible. For example, for a pseudomorphic growth of \( \text{Si}_{0.6}\text{Ge}_{0.4} \) on silicon, the growth temperature must be lower than 550 °C. Bean20, et al. found that the maximum germanium incorporation before the occurrence of non-planar growth depends on the deposition temperature, as shown in Fig. 4. At 750 °C, the maximum germanium mole fraction is 10 per cent.

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**Figure 3.** Critical thickness of \( \text{Si}_{1-x}\text{Ge}_x \) layers as a function of germanium mole fraction. Lines show theoretical kinetic model for various growth temperature.

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**Figure 4.** Plot of growth regimes vs growth temperature and mole fraction \( x \) for \( \text{Si}_{1-x}\text{Ge}_x \) grown on silicon substrates. The solid line and the open circles refer to MBE, while the solid circles refer to LRP samples which exhibit planar growth.
Figure 5. Dopant accuracy for two dopant source concentrations. Limitations are flow controller absolute accuracy and offset, as well as temperature-induced fluctuations.

whereas at 550 °C, 100 per cent germanium mole fraction is possible. It is speculated that higher deposition temperatures resulted in surface mobility, high enough to cause surface tension problems and islanded growth. Deposition rates of up to 600 nm/min for silicon are possible. However, typical $Si_{1-x}Ge_x$ deposition rates are in the 30 nm/min range for greater profile control. Also, extremely abrupt compositional profile control is possible by the use of mechanical shutters. At 630 °C, the $Si_{1-x}Ge_x$ growth rate decreases with increasing germane ($GeH_4$).

Ultra high vacuum chemical vapour deposition (UHVCVD) system developed by Meyerson$^{27}$, et al, uses base pressure of 10$^{-9}$ Torr, growth pressure of 10$^{-6}$ Torr, and a growth temperature as low as 550 °C. This system, however, has few limitations. As the source gas flow is increased (at constant temperature) to increase the germanium concentration during a $Si_{1-x}Ge_x$ layer growth, the growth rate increases.

The deposition pressure is about 1-2 mTorr with deposition rates around 1-2 nm/min. Because of the low deposition pressure, gas depletion does not occur like in standard low pressure chemical vapour deposition (LPCVD) poly silicon reactor at 0.4 Torr. In situ doping of both types is possible and boron dopant content in the film is linear as shown in Fig. 5. The addition of $PH_3$ to silane ($SiH_4$) produces $n$-type doping. Although dopant control is similar to that of boron, the dynamic range over which $P$ content remains stable is far more limited as shown in Fig. 6 and allows the incorporation of up to $1 \times 10^{20} \text{ cm}^{-3}$ (Note the discontinuous jump in, and subsequent stability of, phosphorus incorporation in the presence of germanium). Apparently no problems with islanded growth at high germanium concentrations appear, because the deposition temperature is below 550 °C. The addition of $GeH_4$ enhances the growth rate of $Si_{1-x}Ge_x$ over silicon homoepitaxy at 550 °C. The catalytic effect of $GeH_4$ in removing hydrogen from the surface is cited as the reason for the increased $Si_{1-x}Ge_x$ growth rate. Germanium incorporation in alloy layers is roughly linear in source germanium content, while the behaviour of growth rate is somewhat more complex, as shown in Fig. 7. Although blanket depositions are trivial, problems appear with patterned oxide depositions because the HF-dip cleaning technique is hydrophilic on oxide regions. Polysilicon layers on top of the oxide may eliminate this problem at the expense of greater process complexity. This problem may ultimately limit the manufacturability of the UHVCVD
technique. Selective silicon epitaxy was examined at Tohoku University and selective Si$_{1-x}$Ge$_x$ deposition was studied by Greve and Racanelli, who observed that GeH$_4$ enhanced the nucleation time of polycrystalline material on oxide regions of patterned wafers.

Using rapid thermal chemical vapour deposition (RTCVD) growth technique, at a base pressure of 10$^{-7}$ Torr, which is much lower than that used in limited reaction processing (LRP), SiGe films have been grown at 900 °C. Si$_{1-x}$Ge$_x$ layers need to be deposited at lower temperatures to avoid relaxation and the three-dimensional growth problems. So the deposition temperature was reduced to 625 °C for Si$_{1-x}$Ge$_x$ and increased to 850 °C for silicon cap layers. Fortunately, the Si$_{1-x}$Ge$_x$ growth rate increases with the addition of GeH$_4$. The Si$_{1-x}$Ge$_x$ growth rate enhancement with increasing GeH$_4$ is similar to the UHVCVD result. The maximum incorporation before three-dimensional growth by MBE does not seem to apply to limited reaction processing chemical vapour deposition/rapid thermal chemical vapour deposition (LRPCVD/RTCVD). Layers having up to 45 per cent germanium were deposited at 630 °C. One of the major problems with reducing the temperature is the increased oxygen incorporation in the Si$_{1-x}$Ge$_x$ layers. The contamination source was traced to high oxygen content present in SiH$_2$Cl$_2$. SiH$_4$ does not have the same problem because the gas with much higher purity is available. The oxygen incorporation problem diminishes with the use of a load-lock and a point-of-use filtration of SiH$_2$Cl$_2$.

Atmospheric pressure chemical vapour deposition (APCVD) has been used to grow good quality epitaxial silicon and SiGe films. This technique offers temperature flexibility over a wide range, excellent temperature uniformity in the commercial process, and a simple conventional flowthrough design which uses no vacuum pumps. Ultra clean APCVD process has been used to grow films at low temperatures, 550-850 °C, by rigorously minimising oxygen and moisture in the growth environment using load-lock, point-of-use gas purification and ultra clean gas handling. Si$_{1-x}$Ge$_x$ layers were deposited over the temperature range from 600-900 °C. At 625 °C, the deposition rate of Si$_{1-x}$Ge$_x$ alloy was found to increase with GeH$_4$ partial pressure; the amount of germanium in the alloy increased rapidly at low GeH$_4$ partial pressures and appears to approach saturation at higher GeH$_4$ partial pressures.

The deposition rate depends slightly on the SiH$_2$Cl$_2$ partial pressure, with a slight decrease in deposition rate with increasing SiH$_2$Cl$_2$ partial pressure at higher GeH$_4$ partial pressures. The deposition kinetics appear similar to the LRP/RTCVD system since SiH$_2$Cl$_2$ and GeH$_4$ are used. When the deposition rate of the silicon component over the temperature range from 600-900 °C was normalised by the germanium content in the alloy, an Arrhenius-type behaviour was observed with an apparent activation energy of about 1.9-2.0 eV suggesting that the deposition process for silicon is limited by reaction kinetics, rather than by mass transport. The germanium component of the deposition rate behaves quite differently as shown in Fig. 8. In low temperature range (600-650 °C), the deposition rate displays an Arrhenius-type behaviour with an apparent activation energy of 0.7 eV, much lower, than that of the silicon component. At higher temperatures, the deposition rate of the germanium component depends only slightly on temperature. Remote plasma-enhanced chemical vapour deposition (RPCVD) has also been used, for silicon and Si$_{1-x}$Ge$_x$ epitaxy.
The researchers demonstrated that good quality layers with a few atomic per cent of carbon ($y \leq 0.05$) can be grown by MBE if a low growth temperature (500-600 °C), and a growth rate of 0.2 nms$^{-1}$ are used. Amorphous growth occurs for lower substrate temperatures or higher carbon concentration. The thermal stability of Si$_{1-y}$Cy/Si strained-layer superlattices was studied by Goorsky et al. The superlattices were grown by MBE as described above, with three concentrations of carbon: 0.003, 0.008 and 0.013. The superlattices were stable on annealing for 2 hr at 800 °C. Between 800 °C and 900 °C, strain relaxation occurred by interdiffusion of carbon and silicon at the interfaces. At 1000 °C and above, precipitation of SiC$_2$ was observed. An X-ray diffraction rocking curve for a ten-period Si$_{1-y}$Cy superlattice, taken at glancing angle of incidence, is shown in Fig. 9. Each period of the superlattice consisted of a 320 Å silicon layer and a 110 Å C$_{0.003}$Si$_{0.997}$ layer. The alloy peaks were quite sharp. TEM studies showed no evidence of dislocations or other defects. For 5 per cent carbon concentration, the layers grown at 450 °C were amorphous. At higher temperatures, precipitates of SiC$_2$ were formed.

One possible application of Si$_{1-y}$Cy alloy is the fabrication of symmetrically-strained SiC/SiGe layers.
superlattices on a silicon substrate. Eberl, et al. reported fabrication of $C_{0.01}Si_{0.99}Ge$ superlattices on silicon. The strain in the superlattices alternates between tensile and compressive in the individual $Si_{1-x}Ge$ and $Si_{1-x}C$, alloy layers, respectively. The individual layer thickness were 10 nm and 13.5 nm, respectively. X-ray rocking curves were measured and average strain in the superlattice was estimated. The strain was considerably less than that expected in a superlattice in which $C_{0.01}Si_{0.99}$ layer is replaced by an silicon layer. It is clear from these results that perfect symmetrically-strained superlattices can be fabricated by adjusting the relative germanium and carbon content and/or by changing the thickness of the individual layers.

Faschinger, et al. reported the electrical properties of undoped and antimony-doped $Si_{1-x}C_x$ alloys on high resistivity (1000 $\Omega$-cm) silicon grown by MBE at 500 °C. The layers could be doped with antimony at low growth temperatures (at 350 °C). However, a significant antimony segregation takes place on the surface and leads to doping of the subsequently deposited layers. Modulation-doped $Si/Si_{1-x}C_x/Si$ structures with the modulation doping in the silicon layer (to avoid unintentional doping of $Si_{1-x}C_x$ channel and silicon spacer layer) exhibited enhanced electron mobilities with peak values of about 10,000 cm$^2$/V.s at 100 K, indicating that $Si_{1-x}C_x$ is strained on silicon and forms an electron channel.

Growth of dilute SiC epitaxial layers on silicon using RTCVD have been reported. Ray, et al. reported the growth of polycrystalline and epitaxial layers of metastable $Si_{1-x}C_x$ alloys using silane and propane at different temperatures with varying amount of carbon ($x = 0.01, 0.02$ and $0.05$) using RTCVD. Stoichiometric SiC$_2$ layers were also grown for comparison by increasing propane to silane flow ratio. Following usual cleaning schedules, $p$-type Silicon < 100 > wafers were pretreated with HF (2:1) diluted in DI water vapour just before loading into the chamber. They have also reported solid phase epitaxy (SPE) growth of $Si_{1-x}C_x$. Epitaxial regrowth by rapid thermal annealing of C-implanted silicon and SiGe layers was used in the study for the formation of metastable phases. CO was used as the source gas for C-implantation. Ion state was chosen to incorporate 1, 2 and 5 per cent carbon into silicon with two different energies. All the samples were rapid thermally annealed in nitrogen (1.2 l/min) at 1046 °C for 30 s.

Fourier transform infrared (FTIR) spectroscopic measurements were carried out on all RTCVD and SPE grown layers to study the nature of carbon incorporation. The absorption peaks due to the localised vibrational mode (LVM) of substitutional carbon in silicon at 16.7 μm (wave number 607.5 cm$^{-1}$) and the strong phonon absorption mode in SiC$_2$ at 12.6 μm (wave number 794 cm$^{-1}$) were monitored. Strane, et al. reported the synthesis and detailed structural characteristics of $Si_{1-x}C_x$ strained layers on silicon by SPE. Carbon was introduced into pre-amorphised silicon by multiple energy implantation to form uniform carbon concentration. Pre-amorphisation was done using 60, 30 and 10 keV silicon ions with doses of $3 \times 10^{15}$, $6 \times 10^{14}$ and $5 \times 10^{14}$ cm$^{-2}$. This was followed by carbon implantation with doses of $2.34 \times 10^{14}$, $9.78 \times 10^{14}$ and $4.44 \times 10^{14}$ cm$^{-2}$, respectively, with corresponding energies of 25, 12.5 and 5 keV. Simulations showed that this procedure yielded a flat carbon profile. A 30 min, 450 °C anneal was used to remove end of the range damage. A 30 min anneal at 700 °C was used for the SPE growth.

High quality layers could be fabricated with carbon concentrations up to 1 atomic per cent by this technique. FTIR spectroscopy measurements showed that the carbon occupies substitutional lattice sites. Film stability was studied in the temperature range 810-925 °C and the changes in strain were measured by double crystal ray rocking curves. Carbon and oxygen concentrations were measured by SIMS and microstructure by TEM. On annealing the layers at 875 °C for 240 min, the 607 cm$^{-1}$ local mode due to substitutional carbon decreased and a broad peak at 810 cm$^{-1}$ was observed. On annealing the layers for 1200 min at the same temperature, the local mode disappeared and the broad peak became stronger indicating the precipitation of SiC$_2$.

2.3 Growth of Ge$_{1-x}$C$_x$ Strained Layers

Metastable Ge$_{1-x}$C$_x$ alloys have been prepared by MBE under nonequilibrium conditions at relatively low growth temperatures with significant carbon concentrations of $\gamma = 0.01$. The pure germanium
layer's growth on silicon shows a typical Stranski-Krastanov (SK) mode, i.e. the growth starts in a two-dimensional mode up to a certain thickness and transition occurs to a three-dimensional island mode. The use of surfactant can inhibit three-dimensional island formation. Osten et al. applied the concept of antimony-mediated growth for the preparation of two-dimensionally grown Ge$_{1-y}$C$_y$ layers on silicon. First a silicon buffer layer was grown followed by deposition of one monolayer of antimony. A 300 Å Ge$_{1-y}$C$_y$ layer was then deposited with a growth rate of 0.125 Å s$^{-1}$. The substrate temperature was 500 °C during the whole growth process. The amount of carbon co-evaporated with germanium was adjusted to attain a total carbon concentration of 1 per cent. The reflection high energy electron diffraction (RHEED) technique was used to monitor the in-plane lattice spacing of the layer during the growth. Lattice constants perpendicular to the interface ($a_{\perp}$) and parallel to the interface ($a_{\parallel}$) were also measured using X-ray techniques and strain in the layers was determined. Kolodzey et al. have grown crystalline germanium-carbon alloys having a cubic diamond lattice on silicon (100) using MBE at 600 °C. Measurements on thick relaxed alloy layers showed that up to 3 atomic per cent carbon was incorporated, which reduced the lattice constant and increased the energy gap (0.875 eV) compared to bulk germanium.

### 2.4 Growth of Alloy Layers Containing Tin

One approach for realisation of a direct energy gap group IV alloy system involves alloying tin with silicon or germanium to form epitaxially stabilised diamond cubic Sn$_{x}$Ge$_{1-x}$/Ge and Sn$_{x}$Si$_{1-x}$/Si heterostructures. Growth of tin-based heterostructures is challenged by the large lattice mismatch between α-tin and silicon (19.5 per cent), very low solid solubility of tin in crystalline silicon ($5 \times 10^{19}$ cm$^{-3}$), and pronounced tin segregation to the surface during growth at ordinary silicon epitaxy temperatures (T > 400 °C). Growth conditions similar to MBE on antimony delta-doped layers in silicon had to be used for the growth incorporating tin at high fractions while maintaining a high epitaxial quality. Min et al. have fabricated ultra thin, coherently strained Sn$_{x}$/Si and Sn$_{x}$Si$_{1-x}$/Si alloy quantum well structures with substitutional tin incorporation far in excess of the equilibrium solubility limit via substrate temperature and growth flux modulations in MBE. Tin/silicon single and multiple quantum wells with tin coverage up to 1.3 monolayer, Sn$_{0.02}$Si$_{0.98}$/Si multiple quantum wells of up to 2 nm and Sn$_{0.16}$Si$_{0.84}$/Si multiple quantum wells up to 1.1 nm were determined to be pseudomorphic.

He et al. grew strain-compensated epitaxial layer of Sn$_{x}$Ge$_{1-x}$/Sn$_{y}$Si$_{1-y}$ alloy films on silicon (100) with composition of tin and carbon greatly exceeding their normal equilibrium solubility in silicon. Amorphous Sn$_{x}$Si$_{y}$C alloys were deposited by molecular beam deposition from solid sources followed by thermal annealing. In situ monitoring of crystallisation rate was done using time-resolved reflectivity. SPE for Sn$_{0.94}$Sn$_{0.01}$C$_{0.01}$ occurs at a rate about 20 times slower than that of pure silicon. The film was found to be dislocation free with good substitutionality of tin and carbon.

### 2.5 Growth of Strained-Silicon on Relaxed Buffer Layers

The present status of growth of strained silicon on relaxed SiGe buffer layers on silicon using various techniques has been reviewed. Experimental studies for the last few years on strained SiGe materials have resulted in the understanding of strain relaxation kinetics and optimisation of graded buffer layers wrt relaxation and surface morphology. These parameters are of crucial importance as they are interdependent and are affected by growth temperature, grading rate and composition.

It is now known that the problem of high threading dislocation densities in relaxed layers may be avoided using a series of low mismatched interfaces and increasing the germanium concentration in steps (step grading) or linearly with a relatively high growth temperature. Because of gradual increase of the lattice mismatch in such a buffer, the misfit dislocation network is distributed over the range of compositional grading rather than being concentrated at the interface to the silicon substrate. The greatly improved buffer quality via the compositional grading lowered the threading dislocation density by three
orders of magnitude and resulted in a much improved electron mobility at low temperatures. Strained layer epitaxial growth on patterned substrates has been attempted\(^7\) which can reduce epitaxial threading dislocation densities by up to two orders of magnitude.

It appears that the competition between dislocation nucleation and propagation determines the final threading dislocation density in the film. The compositional grading is believed to promote propagation while suppressing nucleation of dislocations and to lead to reduced amounts of surface strain, and thus allowing higher growth temperatures\(^8,9\). In fact, the use of a compositionally graded, relaxed, \(\text{Si}_{1-x}\text{Ge}_x\) buffer layer has been advocated as virtual substrate and allows the strain in the film to be tailored at will\(^9\).

Powell\(^8,\) et al. proposed a method for producing an almost dislocation-free relaxed \(\text{SiGe}\) buffer layer on silicon on insulator (SOI) substrates. In this process, the top silicon layer on a wafer-bonded or oxygen-implanted SOI substrate is etched back to an ultra thin layer (50 nm or less). The chemical bond at the \(\text{Si}/\text{SiO}_2\) interface is weak enough so that silicon layer can be considered as quasi-free-standing. When this is used as a substrate for the growth of a thick \(\text{SiGe}\) epitaxial film, it will be energetically favourable for the ultra thin silicon layer to relax rather than the \(\text{SiGe}\) layer itself. If this happens, the threading ends are terminated at the weakly bonded \(\text{Si}/\text{SiO}_2\) interface and the \(\text{SiGe}\) epitaxial layer relaxes without the generation of threading dislocations within the \(\text{SiGe}\) layer.

Many methods exist for deposition of strained silicon on thick relaxed \(\text{Si}_{1-x}\text{Ge}_x\) films on silicon. Gibbons\(^9,\) et al. at Stanford were one of the first groups to demonstrate high quality relaxed thick \(\text{Si}_{1-x}\text{Ge}_x\) film on silicon. The lamp-heated limited reaction processing reactor (LRPCVD)\(^9,10\) was used to grow linearly-graded \(\text{SiGe}\) buffer layers and strained silicon. High quality, epitaxial, relaxed \(\text{Si}_{1-x}\text{Ge}_x\) layers have been grown by RTCVD by Jung\(^10,\) et al. Further improvements of the relaxed buffer (step-graded) layer formation using APCVD with intermediate \textit{in situ} annealing at high temperature have been reported by Kissinger\(^11,\) et al. Threading dislocation densities as low as 100 cm\(^{-2}\) were found indicating that most of the misfit dislocations really extended throughout the wafer.

High quality, completely lattice-relaxed \(\text{SiGe}\) buffer layers have been grown on silicon (100) using MBE in the range 750-900 °C and compositional grading of the order of 10 per cent m\(^{-1}\) or less with final germanium concentrations of about 30 per cent. Xie\(^12,\) et al. have grown compositionally graded relaxed \(\text{Si}_{1-x}\text{Ge}_x\) buffer layers on silicon with various composition gradients and temperatures. The authors reported a threading dislocation density in fully relaxed \(\text{SiGe}\) buffer layers grown using both\(^13\) MBE and RTCVD in the range 10\(^5\)-10\(^6\) cm\(^{-2}\). GSMBE has also been successfully employed\(^14,15\) for the growth of high quality completely lattice-relaxed step-graded, \(\text{SiGe}\) buffer layers on silicon (100) in the range 750-800 °C. A more abrupt compositional \textit{transience} of the \(\text{SiGe}/\text{Si}\) interface is expected in GSMBE grown quantum wells, owing to reduced germanium segregation at the heterointerface\(^16\) than in those grown by solid-source MBE, where germanium segregation has been recognised as an important issue\(^17\). A method of controlling threading dislocation density in relaxed graded buffers containing 50 to 100 per cent germanium has been developed by Cutric\(^18,\) et al. using chemical-mechanical polishing (CMP) though the release of immobile dislocations located in dislocation pile ups. The characteristic cross-hatch surface roughness and the underlying strain fields of the misfit array can overlap, blocking threading-dislocation glide leading to dislocation pile ups during the growth of thick and high germanium content buffer layers.

The growth of only a micron thick relaxed buffer layer is possible\(^19,20\) using \textit{stepwise}-graded buffer based on a combination of \(\text{Si}_{1-x}\text{Ge}_x\) and \(\text{Si}_{1-x}\text{Ge}_x\text{C}_y\). The buffer concept is based on the fact that the addition of carbon to a \(\text{SiGe}\) layer not only reduces the strain, but also stabilises the layer. Due to a very strong local strain field around the individual carbon atoms, dislocation glide requires a higher energy in \(\text{SiGeC}\) than in unperturbed, strain equivalent \(\text{SiGe}\) on silicon. Using this method, a threading dislocation density below 10\(^5\) cm\(^{-2}\) was obtained for 73 per cent relaxed homogeneous \(\text{Si}_{0.7}\text{Ge}_{0.3}\) layer on top of the buffer structure. A stepped \(\text{Si}_{1-x}\text{Ge}_x\) buffer with the identical thickness
and strain profile grown at the same temperature shows a threading dislocation density above $10^7$ cm$^{-2}$. The ternary SiGeC material therefore should be considered like a new material with its own strain degree and relaxation behaviour rather than like a SiGe film with artificially reduced strain.

Structural characterisation and film quality of strained silicon layers are usually carried out by TEM (both plan view and cross-section) for the determination of defects/dislocations within the layers and measure the thickness of the layers, and energy dispersive spectroscopy (EDS) utilising a scanning TEM for film composition. Rutherford backscattering spectroscopy (RBS) also yields similar information.

High resolution X-ray diffraction (HRXRD) is typically used to determine strain. In this technique, the lattice constant, perpendicular to the sample surface, $a_{\perp}$ is determined. To extract germanium content and strain state of the film, it is also necessary to find the $a_{\parallel}$. This is achieved either by measuring the germanium content by some other method (e.g. RBS) and $a_{\perp}$ can be measured by grazing incidence X-ray technique. An alternative technique that measures the strain in the film directly is Raman spectroscopy. Sputter-depth profiling by SIMS is used to study the chemical composition, e.g. germanium profile. However, this method has a limiting depth resolution of about 3 nm and cannot give a reasonable profile for ultra thin films.

Fitzgerald et al. used triple-crystal X-ray diffraction and conventional plan-view and cross-sectional TEM for the determination of strain relaxation. Compositionally graded (10 per cent germanium/μm) $Si_{1-x}Ge_x$ films grown at both MBE and RTCVD at 900°C reveal that for $0.10 < x < 0.53$ the layers are totally relaxed. $Si_{1-x}Ge_x$ cap layers grown on these graded layers are threading-dislocation-free when examined with plan-view and cross-sectional TEM, Electron beam induced current (EBIC) images were used to count the low threading dislocation densities. The dislocation densities measured by EBIC from MBE grown samples with final germanium concentrations of 23, 32, and 50 per cent were found to be $4.4 \times 10^5 \pm 5 \times 10^4$, $1.7 \times 10^6 \pm 1.5 \times 10^5$, and $3.0 \times 10^6 \pm 2 \times 10^6$ cm$^{-2}$, respectively.

### 2.6 Growth of Poly-Silicon-Germanium Films

Doped polycrystalline silicon (poly-silicon) is commonly used as gate material in MOS structures and as contact material for bipolar transistors. Poly-SiGe film has emerged as a new material for advanced CMOS technology and low temperature TFT fabrication for large area display electronics. Commonly used source gases for deposition of poly-SiGe are $SiH_4$ and $GeH_4$ in the range 400-600 °C.

RTCVD has been employed to grow poly-SiGe films. However, these films suffer from higher oxygen content and difficulty for nucleation on oxide. Very low pressure plasma CVD (VLPCVD) has been found to yield high quality films on oxidised silicon substrates at a lower temperature (400-600 °C) and pressures < 4 mTorr using an RF power of 4 W only. Compared to LPCVD, the RPCVD gives higher growth rate, smaller grain sizes, direct deposition on oxide, improved structural properties, such as smoother surface and a more columnar grain structure.

King et al. also used the conventional LPCVD technique to deposit polycrystalline $Si_{1-x}Ge_x$ films of different compositions at 625 °C and at pressures between 0.1-0.2 Torr using $SiH_4$ and $GeH_4$, as the precursor gas sources. The films were heavily doped with boron and phosphorus by ion implantation at a dose of $4 \times 10^{15}$ cm$^{-2}$ and energy 20 keV (for boron) and 60 keV (for phosphorus) and annealed at 900 °C for 40 min in argon. The resistivity of boron-doped films was substantially less than that of poly-silicon films with the same doping level, the resistivity decreasing with germanium mole fraction up to $x = 0.6$. On the other hand, the resistivity of phosphorus-doped films decreased slightly up to a germanium mole fraction of $x = 0.45$ and increased considerably for higher germanium mole fractions. The rapid thermal annealing led to a further reduction of resistivity at a higher germanium mole fraction even at a low peak anneal temperature (500-700 °C). It was thus evident that the temperature required to activate boron decreases dramatically with germanium content. For example, as reported, the resistivity of $Si_{1-x}Ge_x$ of thickness...
25 nm annealed for 30 s at 500 °C is about 10 Ω-cm which is half of that of the silicon film annealed for 30 s at 900-1000 °C. The mobility also increases significantly with germanium content. Heavily boron-doped poly-Si$_{1-x}$Ge$_x$ film with $x = 0.5$ may be a superior gate-electrode material for MOSFETs and as a good interconnect layer in VLSI. On the other hand, heavily phosphorus-doped poly-Si$_{1-x}$Ge$_x$ films degrade with increasing germanium content above 35 per cent. Since the electron affinity does not change appreciably with the mixing of germanium in silicon, the work function of p-type Si$_{1-x}$Ge$_x$ can be more effectively controlled by changing the germanium content as compared to the N-type material. This was experimentally confirmed by King, et al.

Recently, Bang, et al. reported sheet resistance, Hall mobility, and effective carrier concentration as a function of annealing parameters for boron and phosphorus ion implanted films of poly-silicon, Si$_{0.75}$Ge$_{0.25}$ and Si$_{0.5}$Ge$_{0.5}$ films. The films were ion implanted with boron or phosphorus at dosages between $5 \times 10^{14}$ and $4 \times 10^{15}$ cm$^{-2}$, and then thermally annealed between 550-650 °C for 0.25-120 min. Boron-doped films showed decreasing minimum sheet resistance with increasing germanium mole fraction (up to $x = 0.51$) with extended annealing, while phosphorus-doped films exhibited the reverse trend as discussed earlier. Figure 10 shows the minimum measured sheet resistances of boron and phosphorus-doped films versus anneal conditions, implant dose, and germanium mole fraction.

### 2.7 Growth of Hydrogenated Amorphous Silicon-Germanium Alloys

Hydrogenated amorphous silicon-germanium alloys are being developed for the narrow bandgap material in tandem solar cells, for optical detection and image sensing. The bandgap of a-SiGe:H can be varied from 1.75-1.0 eV by changing the content of germanium and make the material suitable for detection of light emitted from commercial laser diodes or LED. The thin film can be deposited at a low temperature of about (250 °C) on glass substrates.

CVD methods, such as RPECVD, microwave plasma CVD and photo CVD have been used for growing a-SiGe:H thin films. The films are deposited by mixing hydrogen with source gases: SiH$_4$ and GeH$_4$. The bandgap is primarily controlled by the germane fraction $f = GeH_4/(GeH_4 + SiH_4)$. Typical conditions of rf plasma-assisted deposition are: rf power 30-60 mW/cm$^2$, pressure 0.2 - 1.0 Torr, flow rates of GeH$_4$-SiH$_4$ mixture 2-20 sccm, hydrogen flow rate of 5-50 sccm. Dilution with hydrogen causes a small decrease in the bandgap and improves the structural and electronic properties. In the presence of diluent hydrogen gas, the dissociation of GeH$_4$ is enhanced relative to SiH$_4$ and the incorporation of germanium is thereby increased.

Evidence from small angle ray scattering confirms that the degradation of electronic properties of a-SiGe:H films with germanium content $> 0.2$ is due to microstructural growth. Microstructural defects can be removed to some extent by proper use of hydrogen dilution of source gases and by controlling the rf power density.

### 3. CONCLUSION

A critical review on growth of group IV binary alloy films, in particular, SiGe, SiC, GeC and SiSn has been made. Various growth techniques, including MBE, CVD and SPE using implantation are compared.
MBE offers perhaps the broadest range of growth conditions and mostly used as a research tool, but MBE cannot be regarded as production-friendly. UHV-CVD offers the possibility of multiwafer growth, which is certainly attractive for volume production. APCVD, using the dichlorosilane chemistry, represents revolutionary development. It can be stated that insight into the detailed growth mechanism is rapidly progressing since a few years now, based on investigations of the fundamental surface science studies. There now exists a detailed, and mostly quantitative understanding of the epitaxial growth and cleaning procedures. However, growth rate data in the low temperature regime as a function of pressure over a broader pressure range (10^{-3} - 100 Torr) is needed. Additionally, fundamental surface studies using modern analytical techniques, preferably in real CVD conditions in view of the important role played by hydrogen, would be necessary to remove much of the uncertainty from the development of epitaxial growth processes.

Compatibility of binary alloy films with standard silicon IC technology requires that high temperature step avoided after the growth of the films. Since standard silicon processing steps, such as implantation annealing, typically exceed the strained layer deposition temperature, thermal stability of strained layers is of utmost importance. The Matthews-Blakeslee curve imposes severe limitations on stable-strained layer thickness and germanium concentration. Understanding the relaxation processes of metastable group IV alloy layers is imperative if thicknesses and germanium or other constituent concentrations greater than the equilibrium curve are needed.

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