Theoretical study of Mbe growth of indium gallium arsenide semiconductor compound on gallium arsenide substrate

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THEORETICAL STUDY OF MBE GROWTH OF InGaAs SEMICONDUCTOR
COMPOUND ON GaAs SUBSTRATE

by

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ABSTRACT

Theoretical Study of MBE Growth of InGaAs Semiconductor Compound on GaAs Substrate

by

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The theoretical study of Molecular Beam Epitaxy allows us to model and construct an experiment with the same conditions. Growth modeling investigates compound semiconductor characteristics during the MBE growth which can achieve the best results to control the quality of growth. Growth modeling also is less expensive and faster than experiments. The wide variation in the band-gap and lattice constants between $\text{InAs}$ and $\text{GaAs}$ is a subject for a variety of optical and electronic device applications involving $\text{InGaAs/GaAs}$ systems. In this material system, the perfection is intrinsically controlled by the surface segregation of $\text{In}$ due to its larger atomic size compared to $\text{Ga}$. In this work, a rate equation model is developed including several surface processes such as segregation from the crystalline layer to a surface riding $\text{In}$ segregated layer and incorporation from the segregated $\text{In}$ layer to crystalline layer and gallium desorption to surface layer. The rate of the processes are assumed Arrhenius type with concentration dependent activation energies. The simulated $\text{In}$ incorporation coefficient versus substrate temperature is in excellent agreement with the experimental data [1] for various $\text{As}$ overpressure. For a constant $\text{As}$ overpressure, $\text{In}$ incorporation decreases with increasing temperature. For a constant temperature,
In incorporation increases with increasing As overpressure. The In desorption versus time results from experiments and our simulation match very well. The desorption process has two components, one arising from the physisorbed layer of In and the other from the surface of the crystal. The activation energy for these processes for an isolated adatom are 0.18 eV and 2.6 eV, respectively. These observations are explained based on the interplay of competing surface processes such as segregation and incorporation.
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CHAPTER 1

INTRODUCTION

1.1 Objective of the Thesis

Today, the increasing demand for advance semiconductor device technology has attracted researchers to study a wide variety of issues in semiconductor fabrication. Molecular beam epitaxy (MBE) is one of the most interesting growth techniques to produce fast opto-electronic devices with unique optical and electronic properties. Molecular beam epitaxy is an epitaxial process that employs evaporated material on heated substrate under ultra vacuum of $10^{-8}$ to $10^{-10}$ torr. MBE technique has a number of advantages over the traditional techniques such as chemical vapor deposition (CVD). The main advantage is low-temperature processing. Low-temperature processing minimizes out-diffusion and out-doping. Another advantage is the precise control of doping, and thickness control to atomic dimensions that MBE technique allows. MBE technique also allows in-situ monitoring of growth control during the growth processes through several analytical tools. Reflective high-energy electron diffraction (RHEED) is one of the experimental tools to monitor cation and anion incorporation rates, alloy composition, and oxide desorption. It is also an important tool to measure growth rate and thickness [21]. The in-situ reflection mass spectroscopy (REMS) is one of the analytical tools in MBE growth for measuring of native oxide desorption, sticking coefficients of group III, desorption rates, incorporation rates and fluxes [8]. Effective segregation during InGaAs/GaAs growth can be observed by using both in-situ RHEED, ex-situ secondary ion mass spectrometry
(SIMS) and photoluminescence (PL)[8].

The $\text{InGaAs/GaAs}$ is one of compound semiconductors which has a variety application in solid state microwave, optoelectronic technologies, quantum well devices, and semiconductor lasers. It has been reported that device performance can be improved if the base of the transistor is fabricated from $\text{InGaAs}$ [65]. Electron transport in $\text{InGaAs}$ has some very remarkable features for microwave applications. The great interest in this system for device application is due to the electronic and optical properties of band gap and lattice constant differences between $\text{GaAs}$ and $\text{InAs}$ [1]. Particularly, the $\text{InGaAs/GaAs}$ heteroepitaxial system can be used in monolithic integration of optoelectronic devices for near infrared and for visible light regions [21].

The growth dynamics of the $\text{InGaAs/GaAs}$ system have been investigated for understanding of important surface processes. Investigation of the molecular beam epitaxy (MBE) dynamic processes is for controlling growth processes and improving reliability and productivity. Several studies [1-19] have shown that the dynamic of processes during the growth of $\text{InGaAs}$ are complex, particularly at temperatures where indium desorption, segregation, and incorporation are all important.

In spite of the interest and experimental work on the growth processes and device applications, the understanding of the growth processes for the reliable control of the growth is limited. In this thesis, a rate equation is employed to model the $\text{InGaAs}$ alloy semiconductor MBE growth. In incorporation coefficient versus substrate temperature and $\text{In}$ and $\text{Ga}$ desorption versus growth time are investigated and compared with experimental results using numerical simulation of the model to validate the model.
1.2 Organization of the Thesis

The thesis is organized into five chapters. The overview of MBE growth, properties and application of InGaAs devices, experimental dynamic studies of InGaAs MBE growth and theoretical models employed for growth studies relevant to this thesis are discussed in chapter 2. The physical and mathematical basis of the theoretical model along with all equations related to the model are described in chapter 3. Chapter 4 is devoted to a discussion of fitting procedure to obtain the theoretical model parameters, results and the limitation of the model. Finally the conclusions and recommendations regarded to this study are given in Chapter 5.
CHAPTER 2

LITERATURE SURVEY OVERVIEW

2.1 Molecular Beam Epitaxy

The word “epitaxy” is originally from the Greek words “epi” meaning “upon”, and “taxis” meaning “to arrange”. Thus, epitaxy means the ability to add and to arrange atoms upon a single crystalline surface of a substrate. Epitaxial growth is a growing process of one material on a single crystal substrate of the same material (homoepitaxy) or of a different material and chemical composition (heteroepitaxy). Epitaxial films of semiconductors can be grown from vapor phase epitaxy (VPE) using chemical vapor growth epitaxy (CVE) or physical vapor growth such as molecular beam epitaxy (MBE), and liquid phase epitaxy (LPE). Each technique has its own advantages and disadvantages. MBE technique is one of the sophisticated techniques which requires high vacuum equipment that can produce vacuum in the order of $10^{-8}$ torr. This technique has the capability to allow abrupt changes in doping or in composition, to control precisely of very thin films monolayers, to grow high quality multilayer structures in relatively low temperatures and minimizing out-diffusion. Due to precise control of the beam fluxes, highly order crystalline films of one or more material layers can be deposited.

In MBE technique, the molecular beams are generated under UHV conditions normally from Knudsen-effusion-cells containing the constituent elements whose temperatures are accurately controlled to achieve a good flux stability. Computer controlled temperatures of the substrate and each of the sources and the operation of shutters,
dictate the desired chemical composition and the doping of the epitaxial films. The molecules of different species of beams have negligible collisions or interactions before reaching the surface of the substrate as their mean free paths are very long. Epitaxial growth occurring on the substrate surface involves a series of surface processes like adsorption of the atoms on the substrate surface, surface migration of the adsorbed atoms, incorporation of the atoms into the crystal lattice and thermal desorption of the species. The crystal surface has crystal lattice sites created by the surface dangling bonds and are characterized by their individual chemical activity.

The growing surface is accessible for observation atom using powerful real-time surface-science diagnostics. The ultra-high vacuum allows monitoring of the growth with \textit{in – situ} tools like reflection high-energy electron diffraction (RHEED). RHEED is routinely used to monitor the crystal structure and microstructure of growing surfaces. Reflection mass spectrometry (REMS) and modulated beam mass spectrometry (MBMS) are used to monitor the chemistry of growing surfaces, and reflectance difference spectroscopy (RDS) is used to monitor the composition and optical properties of the growing surfaces. Surface segregation in the growth of alloy materials can be studied via surface sensitive techniques such as auger electron spectroscopy (AES), x-ray photoemission spectroscopy (XPS) and combination of secondary ion mass spectroscopy (SIMS) and photoluminescence (PL)\cite{9}. Temperature-programmed desorption (TPD) \cite{6} is also employed to investigate the tendency of indium to segregate at the surface of \textit{InGaAs} films during molecular beam epitaxy. TPD analysis is used for quantitative assessment of surface populations and binding energies of adsorbed species present on solid surfaces. The TPD analysis of the surface indium population can be carried out as a function of \textit{InGaAs} thickness, \textit{GaAs} cap thickness, growth temperature and incident arsenic dimer flux.

In a nutshell, the device engineer can control and produce the state of the surface
including composition, crystal structure and smoothness and subsequently, the quality of the material very precisely. The surface scientist can study, directly, the real-time evolution of surface structure, micro structure and composition.

2.2 Difference Between VI Elemental and III-V Compound Semiconductors

Semiconductors such as \( Si, Ge, C \), etc have a diamond structure with the same atom on both sublattices. If the atoms are different in the two sublattices, the structure is called zinc blende structures. Semiconductors such as \( GaAs, AlAs, CdS \) fall in to this category. Some semiconductors crystallize in the “wurtzite” structure. Wurtzite is similar to the zinc blende structure. Both structures are fourfold coordinated, except that the former has a hexagonal close packed (hcp) Bravis lattice rather than a face centered cubic (fcc). Semiconductors such as \( InAs, ZnO, GaN \) have a wurtzite structure.

Semiconductors such as \( GaAs, InP \) and \( InGaAs \) have their bottom of the conduction band and top of the valance band at the same \( K \) (momentum vector) location in the energy (E) versus momentum vector (K) plot. Such materials are optically active and are called direct band gap semiconductors. \( Si, Ge \) are indirect band gap materials and have very weak interactions with light and can not be used for efficient optical devices.

One of the other advantages, of the III-V compound semiconductors is the high mobility of these materials compared to elemental semiconductors such as \( Si, Ge \), resulting in devices with reduced parasitics and improved frequency response. Room temperature mobilities in high quality \( GaAs \) samples are \( 8500 \ cm^2/V^{-1}s^{-1} \) compared to only \( 1500 \ cm^2/V^{-1}s^{-1} \) for \( Si \). One of the semiconductors which is important for high speed devices is \( In_{0.53}Ga_{0.47}As \). This material has a very high room temperature mobility of about \( 11000 \ cm^2/V^{-1}s^{-1} \). At low temperature, the mobility is dominated
by alloy scattering effects, and is less than, the mobility in pure GaAs low field [69].

The greatest impact of compound semiconductors has been in areas where their unique properties allow functions that can not be performed by silicon. These include transferred electron devices, light-emitting diodes, lasers and infrared photodetectors. Of the many compound semiconductors currently under investigation, gallium arsenide and related compounds are the most technologically advanced.

2.3 Application of InGaAs Compound Semiconductor Devices

The importance of ternary compound semiconductor devices and their application form the foundation of solid-state microwave and optoelectronic technologies used in many modern communication systems. It has been reported that device performance can be improved if the base of the transistor is fabricated from InGaAs for following reasons:

- The reduction of the band-gap in InGaAs compared with GaAs gives a greater emitter-base valance band offset. This helps to ensure that holes are not reverse injected into the emitter.

- The higher electron mobility in the base allows a shorter base transit time and a consequent improvement in device speed.

- The hole mobility is dependent on the direction of the strain, being higher in growth plane with consequent lower base sheet resistance, and lower perpendicular to the growth plane and hence limiting the movement of the holes in that direction.

- The lower growth temperature for InGaAs allows more p-type dopant to be incorporated. High doping levels can be achieved, reducing further the sheet resistance.
• There is the possibility of grading the InGaAs alloy composition allowing the electrons to be swept out of the base by a high electric field in the conduction band [65].

InGaAs is an excellent material for long haul communications (at 1.55 micrometer). One of the application of InGaAs devices is in optical communication system as a p-i-n photo detectors. They provide switching circuitry in optical communication system which use optical fiber instead of metallic cable. Another possible application of compound semiconductor is in semiconductor lasers using quantum wells and quantum dots. Quantum dot (QD) as an active region in a semiconductor injection laser enhances ultra low threshold current density and provides high thermal stability in the semiconductor lasers.

2.4 Incorporation of In in InGaAs MBE Growth

Indium incorporation coefficient as a function of substrate temperature [1,5,9,10,15] and growth time [2,11] have been studied experimentally for various V/III ratios, In compositions and As molecular specious, i.e. As$_2$ and As$_4$. Indium incorporation is enhanced by using high V/III ratio and arsenic dimer, As$_2$. It is observed that there is In enrichment at the surface of the interface in heterointerfaces involving InGaAs.

Woodbridge et al [10] studied indium incorporation in InGaAs grown by MBE, at a growth rate of 1 monolayer/second (ML/s), temperature of 550°C and various As overpressures. They found that the free In surface population can be suppressed by increasing the As flux. The growth of InGaAs/GaAs quantum well structures at temperatures above about 550°C may therefore be improved using high V/III ratios or by the use of a growth interruption at the InGaAs/GaAs interface.
2.5 Indium Segregation Studies

Muraki et al [9] investigated the surface segregation of In atoms during MBE growth and its influence on the energy levels in InGaAs/GaAs quantum wells (QWs), by combining secondary-ion mass spectroscopy (SIMS) and photoluminescence (PL). The growth rate was about 0.7 μm/h for InGaAs. They developed a model for interpreting their experimental results. In their model, the segregation probability \( R \) is calculated from the segregation length (\( \lambda \)) obtained from secondary ion mass spectroscopy (SIMS) measurement and using the relation \( R = \exp(-d/\lambda) \), where \( d \) is half the lattice constant of GaAs (2.83Å). They observed that when growth temperature is raised from 370°C to 520°C, the segregation length increases from 0.8 to 2.9 nm. They showed that the indium surface segregation length is highly dependent on growth temperature for low V/III flux ratios of 4 and 12.

Kao et al [8] studied the segregation ratio \( R \) under various deposition conditions in real-time. Kao performed experiments similar to those of Muraki et al [9]. In-situ reflection mass spectroscopy (REMS) was used for real-time monitoring of surface composition during epitaxial layer growth to improve control of InGaAs compositions. They found that throughout the temperature range 470-560°C, the REMS signal dependence yields an activation energy (4.08 eV) which agrees well with that of Evans et al [5]. The high values of \( R \) (0.7-0.8) were observed under normal device layer growth produces a surfaces layer with high In content (i.e., InAs). They suggested a model for segregation process where the segregation ratio \( R \) is defined as the fraction of In atoms on the top layer which segregate to the next layer. Only \( (1 - R) \) of the total surface In incorporates into the growing film. The In composition in the \( n^{th} \) monolayer is then:

\[
x_n = x_0(1 - R^n), (0 < n < N : \text{in the InGaAs layer})
\] (2.1)
\[ x_n = x_0(1 - R^n)R^n\,^n-N, (n > N: \text{in the GaAs overlayer}) \] (2.2)

where \( x_0 \) and \( N \) are the nominal \( In \) mole fraction and the QW width in monolayer, respectively.

Woodbridge et al [10] found that indium segregates to the surface during growth above 550°C and a constant surface concentration results at low indium flux. They found that up to two mono-layers of indium may have segregated onto the surface during the growth of 200 Å of \( \text{In}_{0.25}\text{Ga}_{0.75}\text{As} \) at 560°C. They also observed the indium segregation at higher substrate temperature under the effects of indium flux and for V/III flux ratios of 20 and 30.

Hazay et al [17] studied the segregation to the surface of various third-column atoms in ternary arsenide (\( \text{GaAlAs}, \text{InAlAs}, \text{InGaAs} \)) grown by molecular beam epitaxy. The surface segregation of third-column atoms used ternary alloys (\( \text{Ga}_{0.7}\text{Al}_{0.3}\text{As} \), \( \text{In}_{0.52}\text{Al}_{0.48}\text{As} \), \( \text{In}_{0.53}\text{Ga}_{0.47}\text{As} \), \( \text{In}_{0.2}\text{Ga}_{0.8}\text{As} \)) was studied by \textit{in-situ} auger electron spectroscopy (AES) and x-ray photo-emission spectroscopy (XPS). The peak intensities due to the different alloys were considered. Hazay et al suggested that an elementary model involving an exponential law with absorption length \( L \) provides an excellent match with experimental data. The reduced segregation rate, \( R \), relevant to elements \( A \) and \( B \) in the alloy \( A_xB_{1-x}\text{As} \) was found using the follow equations:

\[
R_A = x_b + (x_s - x_b)(1 - \exp\left(-\frac{a^*}{L_A}\right)) \quad (2.3)
\]

and

\[
R_B = 1 - x_b + (x_b - x_s)(1 - \exp\left(-\frac{a^*}{L_B}\right)) \quad (2.4)
\]

where \( a^* \) is half the lattice constant and \( x_s \) and \( x_b \) are, the surface and bulk compositions of the \( A \) element, respectively. It has been suggested that segregation found at the interfaces could be explained as the difference between 'normal' and 'inverted' interfaces of \( \text{InGaAs}/\text{GaAs} \) system. The tendency of third-column atoms to sur-
face segregation in ternary alloys is summarized by the relation $In > Ga > Al$ as in heterostructure between binary arsenide. The relation shows that indium atoms have the highest tendency to surface segregation. Hazay et al showed that during the growth of $InGaAs$ on $GaAs$ by MBE, the indium atoms segregate at a ratio of more than 0.8 under the conventional growth conditions for $InGaAs$. The transition from two-dimensional to three-dimensional growth occurs when the amount of indium reaches to about 1.7 monolayers, with indium composition greater than 0.25.

Kaspi et al [7] suggested the use of low substrate temperature and higher V/III ratios as a means of reducing indium surface segregation to improve both normal and inverted interfaces. They found that the surface segregation of indium atoms during MBE of $InGaAs$ layers highly influences the composition in the vicinity of both the normal and the inverted $Ga(Al)As/InGaAs$ interface. They observed that the intended alloy composition in $In_{0.22}Ga_{0.78}As$ is not reached to the actual composition unless it grows until nearly 35Å from the $InGaAs$ on $GaAs$ interface for growth temperature 500°C. The compositionally graded region in $InGaAs$ can be eliminated by pre-adsorbing a fixed amount of $In$ onto the $GaAs$ surface to match the surface segregated layer during steady state, before depositing the $InGaAs$ layer. The amount of indium in the floating layer at steady state is observed to more than double from 0.63 monolayer (ML) at 425°C to 1.3 ML at 520°C.

2.6 Indium Desorption

The desorption of group III elements during MBE of III-V semiconductors is of great importance in the control of the thickness and the composition of structures grown [11]. There are two main techniques used in the study of this phenomenon. One way is to observe the temperature dependence of the growth rate and infer the desorption rate [2]. The other is to measure the desorption flux directly by using
modulated beam mass spectroscopy (MBMS) [5,8]. Measurement of the growth rate can be achieved \textit{in-situ} by the reflection high energy electron diffraction (RHEED) intensity oscillation technique or \textit{ex-situ} by layer thickness measurements. One of the distinguishable indium desorption dependences is on the substrate temperature during the growth. This study can be possible by using MBMS [5].

Zhang \textit{et al} [3] investigated \textit{Ga} and \textit{In} desorption in the \textit{InGaAs} and \textit{InAs} layers grown via MBE at relatively high substrate temperatures. For \textit{Ga} desorption studies, \textit{InGaAs} and \textit{InAs} were grown for 10 min at substrate temperatures range of 560°C to 630°C. Indium and gallium molecular beam fluxes were given including \textit{As}_2 and \textit{As}_4 were measured using RHEED intensity oscillation and maintained constant. The logarithmic desorption rate of \textit{Ga} from \textit{GaAs} surfaces was plotted against inverse substrate temperature. Two distinct temperature dependences for indium desorption from \textit{InAs} were reported. One is shown to be independent of surface indium adatom population, the other is shown to be dependent on indium adatom population. The two rate limiting processes operate at two different temperature regions and are independent of one another. It is found that there is a small difference in the desorption rate between the two cases Langmuir evaporation and growing conditions. The activation energy for desorption is approximately 4.0 eV[3].

Mozume \textit{et al} [19] studied the indium desorption during the molecular beam epitaxy of \textit{InGaAs/GaAs} growth using RHEED intensity oscillations. The \textit{InAs} mole-fraction was varied from 0.07 to 0.25. \textit{Ga}, \textit{In} and \textit{As} fluxes were monitored by the ion gauge at the substrate position. The V/III flux ratios were varied from 8 to 20. The growth rate was about 0.5 \textmu m/h for \textit{GaAs}. The flux evaporating from the surface was completed by the Hertz-Knudsen-Langmuir equation:

\begin{equation}
J_{In} = 3.51 \times 10^{22} \times \left(\frac{M}{T}\right)^{\frac{1}{2}} \times P_{torr} (cm^{-2}.s^{-1})
\end{equation}

(2.5)
where $M$ is the atomic mass in grams, and $P_{\text{torr}}$ is the partial pressure of the evaporating species at $T^\circ K$. The time for In evaporation, $t$, from the $In_xGa_{1-x}As$ surface is determined:

$$t = \frac{3.1 \times 10^{14} X}{J_{In}(s^{-1})}$$

(2.6)

where $X$ is the indium composition and $J_{In}$ is the indium flux. The indium desorption activation energy obtained from the temperature dependence of $InAs$ growth rate agree well with the enthalpy of $InAs$ decomposition. The RHEED pattern transition time, $t_1$, for switching from $InGaAs$ to $GaAs$ reconstructions during heteroepitaxial growth of $GaAs$ on $InGaAs$ was found to be independent of $InGaAs$ layer thickness.

Incorporation/desorption rate variation at heterointerfaces in III-V molecular beam epitaxy were studied by Evans et al [2,5]. Three heterointerfaces, $AlGaAs$ on $GaAs$, $InGaAs$ on $GaAs$ and $SbGaAs$ on $GaAs$, were investigated. The V/III flux ratios were varied from 8 to 20. The growth rate was about 0.5 $\mu$m/h for $GaAs$. A mass spectrometer was used in the experimental method to monitor the desorption rate of partially incorporation species such as $In$, $Al$ and $Sb$. The incorporation coefficient $\alpha_M$ of species $M$ is defined by:

$$\alpha_M = 1 - \frac{F_d(M)}{F_i(M)}$$

(2.7)

where $F_d(M)$ and $F_i(M)$ are $M$'s desorb and incident fluxes, respectively. The $In_yGa_{1-y}As/GaAs$ system with $y=0.15$ was studied for substrate temperature between 547 and 637°C. Constant incident fluxes were used (1 monolayer (ML)=2.83Å for $GaAs$): $F_i(\text{In})=0.09$ ML/s, $F_i(\text{Ga})=0.5$ ML/s. The arsenic tetrameric beam flux pressure was $1.38 \times 10^{-5}$torr. It was observed that incorporation rates vary with time during the growth and it significantly influences the compositional profiles and layer thickness. The indium desorption flux versus time shows that the time required to
reach steady state is about 8 second, or 4 ML growth. As long as indium desorption flux is not varied in a step-like manner and indium desorption is less than expected, the indium composition is not constant during the growth. The variation of the In composition, y's values with position from the InGaAs/GaAs for different substrate temperatures were investigated. The relation between In incorporation coefficient, $\alpha_{In}$, and y for In$_y$Ga$_{1-y}$As was suggested to be:

$$ y^{-1} = 1 + \frac{F_i(Ga)}{\alpha_{In}F_i(In)} $$

(2.8)

The value of $F_d(In)$ for steady state "bulk" InGaAs growth is found to vary with calculated $y$ value, i.e. $F_d(In) = K_d \times y$, $K_d$ is a first order In desorption rate constant. The variation of $K_d$ with temperature is expected to be Arrhenius, i.e.

$$ K_d = \nu \times \exp\left(-\frac{E_a}{kT}\right) $$

(2.9)

where $\nu$ is a frequency factor and $E_a$ is the activation energy associated with In desorption. An Arrhenius plot of $F_d(In)/y$ gives a value of $E_a = 3.4$ eV[2].

2.7 Strain

There have been several experimental studies reported in the literature to study the effect strain layer on the electrical and optical quality of compound semiconductors [22,23,24,25,26]. InGaAs strained layers on GaAs show useful characteristics, e.g. higher carrier mobility and the splitting of heavy and light holes in the valance band. It has been shown that the InGaAs grown in the substrate temperature range 400-420°C has a mobility more than 6200 cm$^2$ V$^{-1}$ s$^{-1}$. In this study, highly strained layers grew without misfit dislocations for the In composition range of 0.3 to 0.4. To achieve 2-dimensional electron gas and electron mobility, In$_x$Ga$_{1-x}$As layers must be grown 6 to 10 nm thick. Misfit dislocation [23] occurrence during growth has been studied by RHEED of In$_x$Ga$_{1-x}$As on gallium arsenide substrate where indium
composition is $x=0.3, 0.38$ and 0.5. For highly strained layers, it was observed that GaAs must be preserved carefully at 590°C before InGaAs growth. It was observed that the surface energy has a greater effect than the details of dislocation for very thin films. The results are compared with various models of dislocation nucleation and good agreement is found for heterogeneous misfit accommodation by 60 degree dislocations.

Coman et al [24] found that in order to reduce the strain in lattice-mismatches during epitaxial growth, one must use thin compliant growth substrates. They also observed that strain can be used to modify molecular beam epitaxy growth kinetics, such as cation desorption and migration. They introduced a new method for producing laterally confined structures which is extremely flexible and can be applied to any material system. Such a method, called strain-modulated epitaxy, utilizes thin, compliant substrates which are patterned on the bottom, bonded surface. The strain produced during growth will be partitioned between the substrate and the lattice-mismatched epitaxial layer according to:

$$\epsilon_f = \frac{h_s}{h_f + h_s} \epsilon_0$$  \hspace{1cm} (2.10)

where $\epsilon_f$ is the new strain film, $\epsilon_0$ is the total strain of the system, $h_s$ is the thickness of the substrate, and $h_f$ is the thickness of the grown film.

2.8 RHEED

Reflection high-energy electron diffraction (RHEED) is an integral part of the molecular beam epitaxy chamber to study the initial static semiconductor surface geometries (e.g. surface smoothness, disorder, steps etc.) and stoichiometric phases (e.g. changes of surface reconstruction pattern with respect to substrate temperature). In RHEED, a high energy beam of electrons in the range of 5-40 kV is directed
towards the surface at a grazing angle of about 1° to 3°. This is ideal for MBE where the molecular beams impinge on the surface at near-normal incidence.

Fujita *et al* [27] studied grown $In_xGa_{1-x}As$ layers with various alloy compositions ($0 \leq x \leq 1$) on GaAs substrates. The grown layers and heterointerfaces were characterized by RHEED and X-ray diffraction. The RHEED patterns indicated that the crystal nucleation is two-dimensional throughout the epitaxial growth for $0 \leq x \leq 0.4$, but for $0.5 \leq x \leq 1$, it is three-dimensional at the early stage of the growth followed by two-dimensional growth. All the experimental results consistently showed that the crystalline quality degrades with the increase of $x$ from 0 to about 0.5, but beyond 0.5, it tends to become rather improved with $x$.

### 2.9 Comparison of Growth Modeling Techniques

One aim of the numerical simulation is to compare the results with experimental results to validate the physical model. Detailed information calculated by simulation can be achieved faster with less expense. Another feature of the calculation by simulation is to study phenomena which is not amenable to analytical theories. There are several theoretical tools available to study the growth process MBE and metal organic chemical vapor deposition (MOCVD) such as Monte Carlo simulations (MC), molecular dynamics simulation (MD), stochastic models, and rate equation models. The theoretical studies of MBE must be able to probe the surface growth kinetics to predict the macroscopic growth rate as a function of substrate temperature, composition, surface and interface roughness, and the partial pressure of the reactants.

The term Monte Carlo refers to the random simulation by calculating the probability distribution of a physical event or series of events based on the expression for the rates. Monte Carlo simulations are based on the rigid lattice of finite size [59,60]. The surface kinetic processes, typically considered in the MC simulation, are
incorporation, surface migration and reevaporation. The rate, $K$, corresponding to individual processes at the (local) temperatures, $T$, are usually taken to be of the Arrhenius form, given by:

$$K(T) = K_0 e^{\frac{-E}{k_B T}}$$

where $K_0$ represents an attempt rate for the process, $E$ is the kinetic energy barrier for the process, $k_B$ is the Boltzmann's constant, and $T$ is the temperature [60]. MC simulation provides macroscopic growth in formation such as growth rate, surface and interface and roughness compositional guiding. It allows RHEED simulation based on kinematic theory of electron diffraction.

Molecular Dynamics (MD) is growth simulation method based on actual atomic dynamics. In this model, no rigid lattice is assumed unlike the MC simulation. In this model, particle experiences and reacts to the potential of the rest of the particles which are held fixed in their position. The potential is usually obtained from a definite physical model. In order to follow the dynamics of a many particles system, one could calculate the force of each particle by considering the influence of each of its neighbors. In the dynamical calculation, with all the given initial velocities and positions, the future behavior of the system is determined using the potential at a point and Newton's second law. The MD model is used for studying the surface kinetics during the epitaxial growth, the dynamics of strained layer epitaxy and the evolution of the surface diffusion coefficient [61]. In the dynamical method, it is practical to follow a small system of molecules for only about one microsecond at low temperature. Another limitation of the method is to consider the smallness of the system studied. Only short range interaction between particles can be considered. The solution of the coupled equations of motion for any particle of the system in MD restricts the number of particles and also the range of real time simulation because of limitations in CPU time. The specific advantage of MD simulations is that the
surface kinetics can be studied to get atomic details [62].

Ashu et al [30] performed molecular dynamics simulation of \( In_xGa_{1-x}As \) on substrate \( GaAs(100) \) to show the dynamics of threading dislocations in the over layers and the formation of misfit dislocations at the heterojunction interface. They developed a code by using a modified Tersoff potential [31,32], which simulates the threading dislocation dynamics in the \( InGaAs \) over layer, and also the formation of interface misfit dislocations. To completely simulate the \( InGaAs \) structure, potential energy function PEF’s for the \( In-In, Ga-Ga, As-As, In-As, Ga-As \) and \( In-Ga \) interactions are required. Following Smith [34], who used the functional form due to Tersoff [31,32], Ashu et al fitted PEF’s for four of these types of interactions (\( In-In, Ga-As, In-As \) and \( In-Ga \)), using Smith’s [34] parameters for the remaining two. The interaction energy between each bond is defined as \( V_{ij} \) for the \( i_{th} \) atom to the \( j_{th} \) atom. The functional form of the PEF’s used is:

\[
E = \sum_{i,j} V_{ij}
\]  

\[
E = \left( \frac{1}{2} \right) \sum_{i,j} f_c(r_{ij})(A_{ij}e^{-\lambda_{ij}r_{ij}} - b_{ij}B_{ij}e^{-\mu_{ij}r_{ij}})
\]

The definition of \( b_{ij} \) and \( f_c(r_{ij}) \) have been described in literatures [30-34]. These relations are described as functions of \( r_{ij} \) and \( \theta_{ijk} \). \( r_{ij} \) are bond lengths of the \( i_{th} \) atom to the \( j_{th} \) atom and \( \theta_{ijk} \) are the angles of three atoms \( i, j \) and \( k \) and each index runs from 1 to 3 for \( As, Ga \) and \( In \), respectively. The constants \( A_{ij}, B_{ij}, \lambda_{ij} \) and \( \mu_{ij} \) for the various interactions are shown in table by Ashu [30]. Ashu showed the formation of misfit dislocation for \( In_xGa_{1-x}As/GaAs(100) \) systems. It was observed that continued deposition results in the excess strain being relieved by the formation and migration of defects. In the simulation of dislocations, the inner and boundary region atoms are placed according to the strain fields predicted by elasticity theory. In this simulation, the inner and boundary region atoms are placed according to the
strain fields predicted by elasticity theory. By using this method, a 45° dislocation strain field was imposed on the substrate and a 60° in the over layer [30].

The Stochastic model is based on the master equation approach with quasi-chemical approximation and the solid on solid restriction [48-49]. The time evolution of the epilayer is described by the rate of change of macrovariables of growth such as concentration of atoms, the atom-atom pair concentration of atom-vacancy pair concentrations. The model involves solving simultaneous non-linear differential equations and hence computationally less intensive but does not provide microscopic details of the atoms. The surface kinetic processes considered in this method are the relaxation processes such as the adsorption, the evaporation also the surface diffusion processes such as the intralayer and the interlayer diffusion. These models are studied with either pure relaxation kinetics or pure diffusion kinetics. Saito and Krumbhar (SK model) [48] studied the combined influence of the relaxation and the surface diffusion processes.

Venkatasubramanian [49] developed a stochastic model for the MBE growth kinetic studies of compound semiconductors based on the work of Saito et al [48]. The model developed at first for diamond cubic lattice and later for the two-sublattices zinc blende structure based on the master equation approach and modified solid-solid restriction. Therefore the atom is not absorbed exactly on top of another atom but in a vacant site whose projection falls in between a pair of nearest neighbor atoms. The time evolution of the epilayer is described by the rate of change of a complete set of macro variables such as coverage of atoms in a layer, atom-atom pair concentration etc. The kinetics of the low temperature GaAs were studied using the modified model [51] which in addition to the surface processes like incorporation, evaporation and migration, included the kinetics of the physisorbed layer of As, loosely bound to the surface of the growing crystal by Van der Waals type binding. The thermally
activated surface processes are considered rate limiting to dictate the growth of the film.

The modified stochastic model developed by Muthuvenkatraman et al [52] considered the surface kinetics such as incorporation of arsenic from the physisorbed layer onto the substrate, the intra-layer migration, inter-layer migration and evaporation processes of the gallium arsenide. The antisites incorporation from the physisorbed arsenic (PA) layer and the evaporation of the antisites were studied and fitted to Arrhenius form of equations with incorporation lifetime $\tau_{in}$ and evaporation lifetime $\tau_{ev}$ factors and activation energies for incorporation and evaporation.

A rate equation model proposed by Krishnan et al [70] included the presence and dynamics of physisorbed arsenic (PA) riding the growth surface in the low temperature molecular beam epitaxy (MBE) of (100) gallium arsenide (GaAs). The model results for the dependence of $As^+_{Ga}$ and $As^0_{Ga}$ concentrations on beam equivalent pressure (BEP) and growth temperature agreed well with experiments [41]. Using the same kinetic model for the temporal behavior of the surface, the contribution of the PA layer to the reflection high energy electron diffraction (RHEED) intensity was computed based on kinematic theory of electron diffraction [51]. The results were in agreement with experimental results [40].
CHAPTER 3

THE THEORETICAL MODEL FOR MBE GROWTH DYNAMICS STUDIES

3.1 Introduction

In this chapter, a rate equation model was used in the MBE growth dynamics studies of compound semiconductors. The physical basis of the model includes several plausible surface kinetic processes and its mathematical formulation is in terms of first order non-linear differential equations. The numerical method to solve the first order differential equations, the fourth-order Runge-Kutta method, is also briefly described.

3.2 The Kinetic Rate Equation Model

The MBE growth simulation of InGaAs on GaAs [100] substrate was considered. Growth of compound semiconductors is a result of dynamic processes occurring on a surface riding physisorbed material layer (PM) and the surface of the crystalline epilayer. The PM layers may contain any or all of the species that are used in the growth. The atoms/molecules in this layer are physisorbed on to the surface by Van der Waals type binding. The PM layer undergoes several dynamic processes such as the adsorption of atom onto crystal, the evaporation of atom out of it into vacuum and the segregation of atoms from the crystal into the PM layer. The rates of these processes are assumed to be Arrhenius type with the form:

\[ \tau_i = \tau_{o,i,e} e^{\frac{E_i}{k_B T}} \]  

(3.1)

where \( \tau_{o,i,e} \) is the time constant for the process 'i', \( E_i \) is the activation energy for the processes, \( k_B \) is Boltzmann constant, and \( T \) is temperature in Kelvin. Figure 4.1
depicts a schematic picture of the surface processes of the PM layer. The surface dynamic processes considered for the epilayer in the model are adsorption, evaporation, interlayer and intralayer migration, and segregation. The rate of adsorption depends on the flux rate and the availability of proper surface site with surface covalent bonds satisfaction. The segregation from the crystal layer is allowed only for In. The rate of evaporation and migration of atoms are modeled based on Arrhenius type rate equations with frequency factors and activation energies:

\[ R = R_0 e^{-\frac{E_{act}}{k_B T}} \]  

where \( R_0 \) is the frequency prefactor, \( E_{act} \) is the activation energy, \( k_B \) is the Boltzmann constant and \( T \) is the temperature in Kelvin. The atom interactions are assumed to be pairwise and only up to second nearest neighbor interactions are considered. The activation energy for the segregation process, i.e., from the crystal to the physisorbed state, is assumed to be smaller than that for the evaporation process, but larger than that of the surface migration process.

The time evolution of the growing epilayer is described through the change of macro variables resulting from the surface processes. The macro variables of growth are normalized with respect to the maximum number of possible atoms in the layer. The macro variables considered are the layer coverages of \( Ga \), \( As \), and \( In \) in their respective layers and are given as:

\[ C_{Ga}(2n) : \text{layer coverage of Ga in the } 2n^{th} \text{layer} \]
\[ C_{As}(2n + 1) : \text{layer coverage of As in the } 2n+1^{th} \text{layer} \]
\[ C_{In}(2n) : \text{layer coverage of In in the } 2n^{th} \text{layer} \]

PM layer coverage \(, C_{Ga}^{PM}, C_{As}^{PM} \text{ and } C_{In}^{PM} \)

where \( n \) is the layer index, with the regular \( Ga \) and \( In \) belonging to even numbered layers, and the regular \( As \) belonging to the odd numbered layers. The layer coverage
of atoms is 1 when the layer is completely full and 0 when the layer is completely empty. Only a monolayer of the PM layer will be exposed to that epilayer surface, and hence is dynamically active. Therefore, a constraint on the PM layer coverage is $C_{Ga}^{PM} + C_{As}^{PM} + C_{In}^{PM} \leq 1.0$. The time evolution of the layer coverage of Ga in the $2n^{th}$ layer due to the various surface processes is given by:

$$\frac{dC_{Ga}(2n)}{dt} = \left( [C_{As}(2n - 1) - C(2n)] \left( J_{Ga} + \frac{C_{Ga,phy}}{T_{Ga}} \right) \right) (A1) + [C_{As}(2n - 1) - C(2n)]$$

$$\times \left( e^{-\frac{E_d}{kT}} \left( \frac{C_{Ga}(2n + 2)}{C(2n + 2)} \right) [C(2n + 2) - C_{As}(2n + 3)] \right)$$

$$+ \left( e^{-\frac{E_d}{kT}} \left( \frac{C_{Ga}(2n - 2)}{C(2n - 2)} \right) [C(2n - 2) - C_{As}(2n - 1)] \right) (B1)$$

$$- \left( e^{-\frac{E_d}{kT}} \left( \frac{C_{Ga}(2n)}{C(2n)} \right) [C(2n) - C_{As}(2n + 1)] \right)$$

$$\times \left( [C_{As}(2n + 1) - C(2n + 2)] + [C_{As}(2n - 3) - C(2n - 2)] \right) (C1)$$

$$- \left( e^{-\frac{E_d}{kT}} \left( \frac{C_{Ga}(2n)}{C(2n)} \right) [C(2n) - C_{As}(2n + 1)] \right) (D1)$$

(3.4)

where the term $A1$ denotes the increase in $C_{Ga}(2n)$, due to adsorption of Ga from the incoming molecular beam. The rate of adsorption is the product of the available sites for Ga incorporation on the surface, $[C_{As}(2n - 1) - C(2n)]$, and the fluxes of Ga, $J_{Ga}$ from the molecular beam and $\frac{C_{Ga,phy}S_{In}}{T_{In}}$ from the PM layer. The sticking coefficient of Ga is taken as unity. The term $B1$ describes the increase in $C_{Ga}(2n)$ due to migration into the $2n^{th}$ layer from adjacent Ga layers indexed $(2n + 2)$ and $(2n - 2)$ and fraction of available sites for Ga in the $2n^{th}$ layer is $[C(2n - 2) - C_{As}(2n - 1)]$. The rate of migration is described by Arrhenius type rate equations with frequency factor, $R_0$, and activation energy, $E_d$. The cation sublattice contains two possible elements, Ga and In. Thus, the layer coverages satisfy:

$$C(2n + 2) = C_{Ga}(2n + 2) + C_{In}(2n + 2)$$

of the fraction of the $(2n+2)^{th}$ layer exposed, only a portion of it belongs to Ga. Thus, the fraction $\frac{C_{Ga}(2n+2)}{C(2n+2)}$ is used to make sure that only the Ga portion is considered for
migration. Similar arguments hold for the \((2n-2)^{th}\) layer also. The activation energy for a particular layer is a function of layer coverage of that layer, the activation energies of isolated atoms, \(E_{d,iso}\), and the second neighbor atom-atom pair interaction energy, \(E_{GaGa}\) and \(E_{InGa}\). The factor of four is used to allow for four possible neighboring atoms. In the mathematical form, the activation energy for \(Ga\) diffusion for the \((2n-2)^{th}\) layer is given as:

\[
E_{d,Ga}(2n-2) = E_{d,Ga,iso} + 4E_{GaGa}C_{Ga}(2n-2) + 4E_{GaIn}C_{In}(2n-2)
\]

Thus \(E_{d,Ga}(2n-2)\) is equal to \(E_{d,Ga,iso}\), when the coverage is very small. Its value is given by above equation with \(C_{In}(2n-2) + C_{Ga}(2n-2) = 1.0\), when the layer is full.

The term \(C1\) denotes the decrease in \(C_{Ga}(2n)\) due to migration of \(Ga\) out of the \(2n^{th}\) layer to the adjacent layers, \((2n+1)\) and \((2n-3)\). The description of the rate of this process is similar to term \(B1\), with \(E_d(2n)\) being the activation energy for migration from the \(2n^{th}\) layer, \([C_{As}(2n+1) - C(2n+2)]\) and \([C_{As}(2n-3) - C(2n-2)]\) being the fractions available \(Ga\) atoms for migration into the adjacent layers \((2n+1)\) and \((2n-3)\), respectively. \(\frac{C_{In}(2n)}{C_{Ga}(2n)}[C_{As}(2n) - C(2n+1)]\) is the fraction of \(Ga\) atoms in the \(2n^{th}\) layer. The value of \(E_{d,In,iso}\), \(E_{InIn}\) and \(E_{GaIn}\) are in Table 4.1. The term \(D1\) describes the evaporation of \(Ga\) atoms from the \(2n^{th}\) layer resulting in the decrease in \(C_{Ga}(2n)\) with activation energy for evaporation, \(E_{e,Ga}(2n)\) and the fraction of the \(2n^{th}\) layer exposed, \(\frac{C_{Ga}(2n)}{C_{Ga}(2n)}[C_{As}(2n) - C(2n+1)]\).

The description of the activation energy for evaporation, \(E_{e,Ga}\), is similar to that of \(E_{d,Ga}\) and is written as:

\[
E_{e,Ga}(2n) = E_{e,Ga,iso} + 4E_{GaGa}C_{Ga}(2n) + 4E_{GaIn}C_{In}(2n)
\]

with \(E_{e,Ga,iso}\) is the evaporation energy for the isolated \(Ga\) atom. The description of the activation energy for segregation, \(E_{s,In}\) is similar to that of \(E_{d,Ga}\) and is written
\( E_{a, In}(2n) = E_{a, In, iso} + 4E_{In}C_{In}(2n) + 4E_{Ga}C_{Ga}(2n) \) \( (3.5) \)

with \( E_{a, In, iso} \) is the segregation activation energy for the isolated In atom. Equations similar to Eqn. (3.4) are written for Ga in the \( 2n^{th} \) layer and As in the \( (2n + 1)^{th} \) layer. In our model, all the parameters including the incorporation rate from the physisorbed As state are kept identical for both As\(_4\) and As\(_2\) fluxes to make the model simple. The time evolution of the layer coverage of In in the \( 2n^{th} \) layer is written as:

\[
\frac{dC_{In}(2n)}{dt} = \left( \left[ C_{As}(2n - 1) - C(2n) \right] \left( J_{In} + \frac{C_{In, phy}}{R_{In}} \right) \right) (A2) + \left[ C_{As}(2n - 1) - C(2n) \right] \left( \left[ C_{In}(2n + 2) \right] \left( C(2n + 2) - C_{As}(2n + 3) \right) \right) (B2) \\
+ R_0e^{-E_{d,In}(2n)} \left( \frac{C_{In}(2n)}{C(2n - 2)} \right) \left[ C(2n - 2) - C_{As}(2n - 1) \right] (C2) \\
- R_0e^{-E_{e,In}(2n)} \left( \frac{C_{In}(2n)}{C(2n)} \right) \left[ C(2n) - C_{As}(2n - 1) \right] (D2) \\
- R_0e^{-E_{d,In}(2n)} \left( \frac{C_{In}(2n)}{C(2n)} \right) \left[ C(2n) - C_{As}(2n - 1) \right] (E2) \]
\( (3.6) \)

Note that Eqn. (3.6) is similar to (3.4) except for the substitution of Ga with In.

The activation energies \( E_{d, In} \) and \( E_{e, In} \) are given by:

\[
E_{d, In} = E_{d, In, iso} + 4E_{In}C_{In} + 4E_{Ga}C_{Ga}
\]

and

\[
E_{e, In} = E_{e, In, iso} + 4E_{In}C_{In} + 4E_{Ga}C_{Ga},
\]

respectively. The description of terms \( A2, B2, C2 \) and \( D2 \) are similar to that of \( A1, B1, C1 \) and \( D1 \) except the substitution of Ga with In and the last term \( E2 \) related to In segregation to the PM layer.
The time evolution of the layer coverage of As in the \((2n + 1)\)th layer, \(C_{As}(2n + 1)\), is written as:

\[
\frac{dC_{As}(2n+1)}{dt} = \left( [C(2n) - C(2n + 1)] \left( J_{As} + \frac{C_{As,phy}}{\tau_{in}^{As}} \right) \right) (A3) + [C(2n) - C(2n + 1)]
\]

\[
\times \left( R_0 e^{-\frac{E_{d,As}(2n+3)}{kT}} \left( \frac{C_{As}(2n + 3)}{C(2n + 3)} \right) \right) [C(2n + 3) - C(2n + 4)]
\]

\[
+ R_0 e^{-\frac{E_{d,As}(2n-1)}{kT}} \left( \frac{C_{As}(2n - 1)}{C(2n - 1)} \right) [C(2n - 1) - C(2n)] \right) (B3)
\]

\[
- R_0 e^{-\frac{E_{d,As}(2n+1)}{kT}} \left( \frac{C_{As}(2n+1)}{C(2n+1)} \right) [C(2n + 1) - C(2n + 2)]
\]

\[
\times \left( [C(2n + 2) - C_{Ga}(2n + 3)] + [C(2n - 2) - C(2n - 1)] \right) (C3)
\]

\[
- R_0 e^{-\frac{E_{d,As}(2n+1)}{kT}} \left( \frac{C_{As}(2n+1)}{C(2n+1)} \right) [C(2n + 1) - C(2n + 2)] \right) (D3) (3.7)
\]

Terms A3, B3, C3 and D3 are similar to those of Eqn. (3.4).

Thus, coupled nonlinear first order differential equations given by Eqns. (3.4), (3.6) and (3.7) are obtained from the time evolution of all the macro variables for every layer. These equations are simulated with additional equations for describing the dynamics of Ga, In and As in the PM layer. The time evolution of the PM layer coverage of \(i\)th species, \(\frac{dC_i}{dt}\), is given by:

\[
\frac{dC_{i,phy}}{dt} = \left( J_i(1 - S_i) \right) - \frac{C_{i,phy}}{\tau_{ev}^i} - \frac{C_{i,phy}S_i}{\tau_{in}^i}
\]

\[
+ R_0 e^{-\frac{E_{d,i}(2n)}{kT}} \sum_{\text{all layers}} \left( \frac{C_i(2n)}{C(2n)} \right) [C(2n) - C_{As}(2n + 1)] (3.8)
\]

where \(i\) represents either Ga or In except that there is no segregation of Ga from the crystal which is given by the last term on the RHS. A similar equation is written for physisorbed As without the segregation term and the As getting incorporated in to the anion sublattice. The sum of the coverage of Ga, In and As are not exceeded than 1 as only a monolayer coverage of the PM layer is effective in the surface dynamics. \(J_i\) is the molecular flux of \(i\)th coming onto the substrate and its unit here is in atom/sec. The unit of flux is usually in atoms/cm².sec. and it can be
converted to atom/site.sec. which is simply written as atom/sec. The conversion is performed using the effective area per crystalline site which in case of GaAs substrate with lattice constant $a=5.6533\text{Å}$, is given by $a^2/2$ and is equal to $15.97\text{Å}^2$ in the (100) growth direction. In the above equation, the first term denotes the increase in PM coverage due to arrival of $i^{th}$ species flux into the PM layer. The next two terms denote the net loss of the PM layer coverage due to evaporation and chemisorption into the appropriate site, respectively. The last term is gain of the PM layer coverage due to the segregation of atoms from the crystal which applies only to In. The various $\tau$'s are the time constants representing the respective surface processes. $S_1$ in Eqn. (3.8) represents the total fraction of the appropriate surface available for the incorporation of atoms.

3.3 Runge-Kutta Method

Runge-Kutta method [68] is one of the Euler methods to solve differential equations. It is a numerical method that requires only the initial points in order to begin the algorithm and improve the solutions. Consider the simple case of a single first-order differential equation, $\frac{dy}{dt} = f(y, t)$. The values at time step 'i', $y_i$ and $t_i$, are given. The goal is to extrapolate across the time interval $\Delta t$ to estimate the values at step $i + 1$. In order to carry out this extrapolation, the Runge-Kutta scheme first estimates where the center of the interval is located. The value of $t$ at the center is, $t_i + \Delta/2$. The Runge-Kutta algorithm, then, evaluates the slope of the function at the mid point of the interval and uses this slope to extrapolate all the way across the interval. The corresponding equations are:

$$k_1 = f(y_i, t_i)\Delta t$$

$$k_2 = f(y_i + k_1/2, t_i + \Delta t/2)\Delta t$$

$$y_{i+1} = y_i + k_2$$
In other words, \( y_t + k_1 / 2 \) is the Euler estimate for \( y \) at the center of the interval and \( k_2 / \Delta t \) is the slope at the center. The error in this estimate for \( y_{t+1} \) is proportional to \((\Delta t)^3\), in contrast to the Euler method for which it is on the order of \((\Delta t)^2\). Hence, in this sense the Runge-Kutta method more accurate than the Euler method. The most commonly used Runge-Kutta method is the one comes from the Taylor expansion of \( y(t + \Delta t) \). For simplicity let \( \Delta t = \tau \). The accuracy of the method is higher with considering higher degree of \( \tau \). The Taylor expansion is given as:

\[
y(t + \tau) = y(t) + \tau \dot{y} + \frac{\tau^2}{2} \ddot{y} + \frac{\tau^3}{3!} y^{(3)} + \cdots
\]

\[
y(t + \tau) = y(t) + \tau g + \frac{\tau^2}{2} (g_t + g_y) + \frac{\tau^3}{6} (g_{tt} + 2g_{gy} + g_{yy}) + \cdots
\]

where the indices are for partial derivatives. For example, \( g_{yt} = \frac{\partial^2 g}{\partial y \partial t} \). one can also write the solution at \( t + \tau \) as:

\[
y(t + \tau) = y(t) + \alpha_1 k_1 + \alpha_2 k_2 + \alpha_3 k_3
\]

with

\[
k_1 = \tau g(y, t)
\]

\[
k_2 = \tau g(y + \beta_{21} k_1, t + \beta_{21} \tau)
\]

\[
k_3 = \tau g(y, t \beta_{31} k_1 + \beta_{32} k_2, t + \beta_{31} \tau + \beta_{32} \tau)
\]

for the expansion of the term \( O(\tau^3) \). The flexibility in choosing the parameters \((\beta_{21}, \beta_{31}, ...)\) provides one more way to increase the numerical accuracy in practice. The most common fourth-order Runge-Kutta algorithm which truncated Taylor expansion at the term \( O(\tau^4) \), is given by:

\[
y(t + \tau) = y(t) + \frac{1}{6} (k_1 + 2k_2 + 2k_3 + k_4)
\]

where

\[
k_1 = \tau g(y, t)
\]
\[ k_2 = \tau g(y + \frac{k_1}{2}, t + \frac{\tau}{2}) \]
\[ k_3 = \tau g(y + \frac{k_2}{2}, t + \frac{\tau}{2}) \]
\[ k_4 = \tau g(y + k_3, t + \tau) \]

where \( y = g(y, t) \). The above solution of parameters satisfies the required equations. This algorithm can be modified according to the problem under study.

3.4 Computational Details

Description of evolution of each bilayer of InGaAs requires three first order nonlinear differential equations, one of which describing the time evolution of each of the normalized macro variables. In this work, simultaneous growth of 80 bilayers and the PA layer are considered, requiring a total of 243 \((= 80 \times 3 + 3)\) coupled nonlinear first order differential equations. The system of equations were integrated using the Fourth-order Runge Kutta method described in section (3.3) with time steps of less than \(10^{-6}\) s to get the values of each of the macro variables as a function of time, for a growth time of 10 seconds. The growths were simulated on the Silicon Graphics supercomputer ORIGIN-2000. The average coverage of Ga, As and In in individual layers and the PM layers at the end of growth are obtained from the solution of the differential equations. A fraction of layer coverage of the particular species is obtained by this method.

3.5 Conversion of \(J_{As}\) to BEP

Experimentally, the As flux is described in terms of BEP for a given Ga flux, whereas our model requires the flux in number of monomer atoms per site per second. The conversion between the two flux definitions is accomplished using the following
equation:
\[
\frac{J_{As_4}}{J_{Ga}} = \frac{P_{As_4}}{P_{Ga}} \frac{\eta_{Ga}}{\eta_{As_4}} \sqrt{\frac{T_{As_4}}{T_{Ga}}} \times \frac{M_{Ga}}{M_{As_4}}
\]  
(3.9)

where \( \frac{P_{As_4}}{P_{Ga}} \) is the BEP, \( J \) is the flux and \( T \) is the absolute temperature and \( M \) is the molecular weight. \( \eta \) is the ionization efficiency for the respective species relative to nitrogen and is given by:
\[
\frac{\eta}{\eta_{N_2}} = \left[ \left( \frac{0.4Z}{14} \right) + 0.6 \right]
\]  
(3.10)

where \( \eta_{N_2} \) is the ionization efficiency of diatomic nitrogen and \( Z \) is the atomic number. In Eqn. (3.9), the \( As \) is assumed to be a tetramer. The values used for MBE growth of \( GaAs \) are: \( Z_{Ga}=31; Z_{As_4}=4\times33; T_{As_4}=1173^\circ K; T_{Ga}=573^\circ K; M_{Ga}=69.72 \) and \( M_{As_4}=4\times74.92 \). The number of sites per cm\(^2\) in case of \( (100) \) \( GaAs \) is obtained as \( 1\mu m/hr. = 2.77 \text{ Å/sec.} \), since one bilayer of \( GaAs \) is half of the cubic lattice constant equal to \( 2.82 \text{ Å} \), then \( 1\mu m/hr. = 0.98 \text{ atoms/site.sec.} \). The equivalent surface area for a \( (100) \) site is \( 6 \times 10^{-16} \text{ cm}^2 \) and hence, the number of sites per cm\(^2\) is obtained as \( 6.26\times10^{14} \). Using the conversion factors described in the above paragraph along with Eqn. (3.10), Eqn. (3.11) can be rewritten as:
\[
J_{As}(\text{monomer/cm}^2\text{.sec.}) = 4.0 \times 1.46 \times 10^{14} \times \left( \frac{P_{As_4}}{P_{Ga}}(\text{BEP}) \right) J_{Ga}(\mu m/hr.)
\]  
(3.11)

where \( 4 \) is used for converting the tetramer to monomer. In the growth simulations, composition, indium flux \( J_{In} \) and gallium flux \( J_{Ga} \) were kept the same as the experimental data. To convert the \( As_4 \) and \( As_2 \) BEP's to fluxes (monomer/sec.), the following equations were used:
\[
J_{As_4} = 0.2359 \times (\text{BEP}) \times J_{Ga}
\]  
(3.12)

and in the case of \( As_2 \):
\[
J_{As_2} = 0.58526 \times (\text{BEP}) \times J_{Ga}
\]  
(3.13)

where \( \text{BEP} \) is the V/III ratio.
CHAPTER 4

RESULTS AND DISCUSSION

4.1 Model Parameter Fitting Procedure

The model involves several parameters such as time constants and activation energies, which are initially unknown. These unknown parameters are established according to the following procedure. Experimental conditions employed by Fournier et al. [1] were simulated and the model parameters were adjusted systematically until In incorporation coefficient values for substrate temperature 803 and 903°K fitted well with the experimental data for a As$_4$ BEP of 36. Once the parameters were established, these parameters were used for the rest of the simulations with As$_4$ and As$_2$ at growth conditions employed by several independent research groups [1,3,7,8,17]. Detailed descriptions of these parameters and their values are discussed below.

The activation energy for incorporation processes of Ga and As from the PM layer to the crystal surface are assumed to be independent of temperature (i.e., $E_{in}^{Ga}=0.0$ eV and $E_{in}^{As}=1.0$ eV). The activation energy for the incorporation of In from the PM layer onto the crystal, $E_{in}^{In}$ is assumed to be linearly dependent on the In coverage in the physisorbed layer and is given by:

$$E_{in}^{In} = 0.5C_{In,phy}$$  \hspace{1cm} (4.1) 

where $C_{In,phy}$ is the coverage of In in the PM layer. Similarly the activation energies for the In, Ga and As evaporation process from the PM layer, $E_{ev}^{In}$, $E_{ev}^{Ga}$ and $E_{ev}^{As}$ are assumed to be linearly dependent on their own coverage in the physisorbed layer and are given by:
The prefactor of time constants for incorporation and evaporation processes are obtained according to the Arrhenius equation and related to the activation energies which were described earlier in Eqn. (3.1).

The evaporation, segregation and diffusion processes in the surface of the epilayer are assumed to be thermally activated and are modeled with the frequency factor, $R_0$ and activation energy given by Eqn. (3.2). $R_0$ is also linearly dependent on the substrate temperature, and is given by:

$$R_0 = 2.08 \times 10^{10} \times T$$

This is based on the phonon frequency obtained using the equi-partition energy principle. The frequency prefactor of diffusion processes are assumed constants. The frequency prefactor of In segregation is considered to be linearly dependent on the substrate temperature, and is given by:

$$R_{0,s} = 1.743 \times 10^{10} \times T$$

The segregation process from the PM layer is allowed only for In. It is noted that $R_{0,s}$ is smaller than the $R_0$ of evaporation and diffusion. All the model parameters and their dependences on the surface coverage are summarized in Table 4.1.

4.2 InGaAs Segregation and Desorption Studies

For this study, the growth conditions of Fournier et al [1] were used. The fluxes were: $J_{Ga}=0.714 \mu m/h; J_{In}=0.192 \mu m/h; J_{As1}$ and $J_{As2}$ BEP in the range of 17 to
36. The substrate temperature was in the range of 500-700°C. The $In$ incorporation coefficient, which is defined as a ratio of the total $In$ incorporated to the total $In$ deposited, was obtained for various growth temperature for both $As_4$ and $As_2$ for a BEP of 36. Plots of $In$ incorporation coefficient versus substrate temperature obtained from simulation are shown in Figure 4.2 along with the experimental results of Fournier et al [1]. The agreement is excellent for $As_4$ and fair for $As_2$ for entire temperature range. It is noted that there is no difference between the model parameters for $As_4$ and $As_2$. The $In$ incorporation decreases with temperature for both $As_4$ and $As_2$ due to increased segregation of $In$ to the PM layer and evaporation of $In$ to the vacuum. The $In$ incorporation coefficient is larger for $As_2$ than $As_4$ at the same BEP. The primary reason for this is that the actual flux of $As$ monomer/site/sec. for $As_2$ is more than that of $As_4$ given by Eqns.(3.12) and (3.14). Thus, in our model, no difference in reactivity between $As_4$ and $As_2$ is considered which makes the model simple to use.

Plots of $In$ incorporation coefficient versus temperature for $As_4$ BEP of 17 along with the experimental results of Fournier et al [1] are shown in Figure 4.3. The agreement between the results is excellent below 570°C. Above 570°C, simulation results are lower than the experimental values but agree well with the values for BEP of 36. The experimental values saturate at 0.2, even though the physical reasons suggest that at 630°C, it should be close to zero, especially since the incorporation coefficient is close to zero for BEP 36.

Plots of $In$ incorporation coefficient versus temperature are shown for various $As_4$ BEPs in Figure 4.4. As BEP increases, the incorporation coefficient increases due to reduced lifetime for $In$ surface atoms for the evaporation and segregation processes. It is observed that to achieve a high $In$ incorporation a low substrate temperature below 570°C and high BEP of $As_4$ above 17 are needed.
Desorption parameter of the \(i^{th}\) species, \(D_i\), was found as the difference between the arriving atoms and the change in the total atom concentration in the crystal and the PM layer in a preset short period of time. Mathematically, it can be written as:

\[
D_i = J_i(\Delta t) - \sum_{\text{all grown layers}} \left[ C_i(2n)(t + \Delta t) - C_i(2n)(t) \right] - \left[ C_{i,phy}(t+\Delta t) - C_{i,phy}(t) \right]
\]

(4.2)

where \(\Delta t\) was arbitrarily chosen as 0.1 s for the simulation. The \(In\) and \(Ga\) fluxes were on from 0 to 5 s and at 5 s, the \(In\) flux was terminated. Plots of \(In\) desorption parameters, \(D_{In}\), (computed using Eqn. (4.2)) versus time for a \(As_4\) BEP of 36 is shown in Figure 4.5. These results agree qualitatively well with the experimental results of Ref.[1]. A quantitative comparison cannot be made due to the arbitrary nature of the experimental results. The \(In\) desorption rate initially increases rapidly and reaches a steady state within 2 s. After the \(In\) flux is terminated, the \(In\) desorption flux decreases exponentially to zero. As expected, the desorption rate is larger for higher temperatures. Additionally, \(D_{In}\) shows periodic oscillations in the desorption flux which is related to the periodic exposure of the cation and anion layers due to layer-by-layer evaporation from the crystal. Even though there are noticeable oscillations in the experimental data [1], it is not as periodic as our results.

Indium desorption parameters versus time is shown in Figure 4.6 for 903°K for \(As_4\) BEPs of 36 and 17. The indium desorption for higher BEP is larger which agrees qualitatively with Fourier et al. The reason for this behavior is that at high BEP, the coverage of \(As\) in the PM layer increases which causes decreasing the coverage of \(In\) in the PM layer. Thus less \(In\) incorporates into the crystal. In other words more \(In\) adsorbs.

Relative desorption parameter (RDP) is defined as the ratio of steady state the desorption parameter \(D_{In}(T)\) to \(D_{In}(803°K)\) where 803°K is the lowest temperature in our study. RDP was obtained for several temperatures from Figure 4.5 for a \(As_4\)
BEP of 36. A plot of RDP versus substrate temperature is shown that along with the experimental data of Fournier et al is shown in Figure 4.7. The agreement between the results is excellent for most of the temperature range. Experimental [1] as simulation plots of RDP versus temperature for a $As_4$ BEP of 17 shown in Figure 4.8 also shows good agreement.

The MBE growth simulation was also performed for GaAs growth experiments of Kao et al [8]. Simulated gallium desorption parameters, $D_{Ga}$ given by equation similar to Eqn. (4.2) in the presence of $As_4$ flux is shown in Figure 4.9. The activation energy for desorption was found 2.92 eV from Figure 4.9. This value of 2.92 eV is smaller than that obtained by Kao et al [8]. But our value is reasonable, considering the fact that a surface Ga surrounded by four in-plane Ga neighbors will have an activation energy of about 3.5 eV and a Ga is surrounded by by two in-plane Ga atoms will have about 3.1 eV.

Our investigations strengthened the previous suggestions in the literature [1] that there are two components to the desorption process, one from the surface riding In and the other from the crystal. The activation energy for these processes for an isolated adatom are 0.18 eV and 2.6 eV, respectively. Plots of In layer composition versus layer number is shown in Figure 4.10 for various substrate temperature at a BEP of 36. The growth simulation were performed for 10 s at a growth rate of 0.912 ML/s. These results agree fairly well with the experimental results of Ref. [8]. At lower temperatures the In composition uniform over 10 layers for most temperatures. The segregation of In spreads over at least 10 layers which suggests that these will considerable roughness of alloy mixing at heterointerfaces.

Segregation coefficient, $R$, can be obtained using Eqn. (2.1) and data Figure 4.10 following equation:
\[ \log R = \frac{1}{n} \log \left( 1 - \frac{x_n}{x_o} \right) \]  

(4.3)

where \( n \) is the number of the layer and \( x_o \) and \( x_n \) are the nominal composition and the composition of the \( n^{th} \) layer, respectively. Plots of \( R \) versus temperature obtained for several BEPs of \( As_2 \) and \( As_4 \) is shown along with the experimental data of Kao et al. [8] for \( As_4 \) BEP of 6 in Figure 4.11. Qualitatively, the results are in good agreement. In general, the segregation coefficient, \( R \) increases non-linearly with temperature and attains a maximum value of 1.0 at 850°K for a As \( As_4 \) BEP of 17. The temperature at which the maximum \( R \) is attained increases with BEP as segregation rate decreases with BEP.

4.3 General Observations and Growth Implications

- The \( In \) incorporation to the surface layer decreases in substrate temperature higher than 540°C, more indium atoms incorporate to the the growth surface when the BEP ratio is higher than 20.

- The \( In \) segregation rate is large for higher temperatures and low As over-pressure. Thus, to minimize the \( In \) segregation, one should adopt lower temperatures and high As over-pressures. But, the temperature should not be set so low that the other thermally activated surface processes such as migration and As molecular adsorption by reaction are suppressed.

- The \( In \) desorption for BEP of 36 has a higher rate compare to 17, the actual desorption ratio shows the same behavior in experiment and simulation, it goes up more rapidly by increasing substrate temperature when BEP is higher.

- \( In \) desorption has two components, one from the surface riding \( In \) layer and the other from the surface of the crystal itself. The former component is smaller
than the latter.

- $\text{As}_2$ limits the $\text{In}$ segregation rate more $\text{As}_4$ of the same BEP and it appears that $\text{As}_2$ is a better choice for limiting $\text{In}$ segregation. Therefore, cracked $\text{As}_4$ should be employed.

- The simulated $\text{In}$ composition versus growth monolayer number shows that the $\text{In}$ segregation for substrate temperature range 803-903$^\circ$K, starts at the 4th monolayer and increases by increasing the number of layers.

- For lower $\text{As}$ BEIP, $\text{In}$ segregation occurs at lower temperatures.

4.4 Advantages and Limitations of the Model

The kinetic rate equation model developed calculates the change in concentration of elements in each epilayer grown at each interval of time. Since the model is described by a system of differential equations, the calculations can be performed easily with less computational time. The model considers surface kinetic processes like incorporation, evaporation, migration, deposition, nucleation, growth of islands and interlayer and intralayer migration of atoms from the islands. The model is simple and not limited by crystal size. The doping kinetics in the crystal growth can be performed with ease. Any number of elemental sources can be considered with all surface processes applicable. The main disadvantage of the model is that the microscopic details of the atoms, such as size and shape of the islands, cannot be obtained. The position of atoms or the energy cannot be determined and hence the sites available for antisites are considered only from the total number of atoms in the layer. The activation energies for evaporation, $E_e$ and migration, $E_d$ considered with four neighbor atoms is only approximate and the energies may be a different function of the coverage of atoms.
Table 4.1: Fitted model parameters such as energies, time constants and frequency factors and their dependences

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Model Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Phi_{Phy,Ga}$</td>
<td>prefactor for physisorbed Ga incorporation</td>
<td>$10^{-3}$ s</td>
</tr>
<tr>
<td>$\Phi_{Phy,As}$</td>
<td>prefactor for physisorbed As incorporation</td>
<td>10.0 s</td>
</tr>
<tr>
<td>$\Phi_{Phy,In}$</td>
<td>prefactor for physisorbed In incorporation</td>
<td>0.2 s</td>
</tr>
<tr>
<td>$\Phi_{Phy,Ga}$</td>
<td>prefactor for physisorbed Ga evaporation</td>
<td>100.0 s</td>
</tr>
<tr>
<td>$\Phi_{Phy,As}$</td>
<td>prefactor for physisorbed As evaporation</td>
<td>$10^{-4}$ s</td>
</tr>
<tr>
<td>$\Phi_{Phy,In}$</td>
<td>prefactor for physisorbed In evaporation</td>
<td>$10^3$ s</td>
</tr>
<tr>
<td>$R_0^{d,Ga}$</td>
<td>frequency factor for Ga for diffusion</td>
<td>$2.4 \times 10^8$/s.</td>
</tr>
<tr>
<td>$R_0^{d,As}$</td>
<td>frequency factor for As for diffusion</td>
<td>$4.38 \times 10^7$/s.</td>
</tr>
<tr>
<td>$R_0^{d,In}$</td>
<td>frequency factor for In for diffusion</td>
<td>$4.38 \times 10^5$/s.</td>
</tr>
<tr>
<td>$E_{Ga}$</td>
<td>activation energy for incorporation of Ga</td>
<td>1.0 eV</td>
</tr>
<tr>
<td>$E_{As}$</td>
<td>activation energy for incorporation of As</td>
<td>0.5 $C_{phy}^{In}$ eV</td>
</tr>
<tr>
<td>$E_{In}$</td>
<td>activation energy for incorporation of In</td>
<td>$0.188$ eV</td>
</tr>
<tr>
<td>$E_{Gu-Ga}$</td>
<td>2nd neighbor atom-atom pair interaction energy for Ga</td>
<td>0.0 eV</td>
</tr>
<tr>
<td>$E_{As-As}$</td>
<td>2nd neighbor atom-atom pair interaction energy for As</td>
<td>0.188 eV</td>
</tr>
<tr>
<td>$E_{Gu-In}$</td>
<td>1st neighbor atom-atom pair interaction energy for Ga-In</td>
<td>0.0 eV</td>
</tr>
<tr>
<td>$E_{In-In}$</td>
<td>2nd neighbor atom-atom pair interaction energy for In-In</td>
<td>0.173 eV</td>
</tr>
<tr>
<td>$E_{Gu-As}$</td>
<td>1st neighbor atom-atom pair interaction energy for Ga-As</td>
<td>0.94 eV</td>
</tr>
<tr>
<td>$E_{In-As}$</td>
<td>1st neighbor atom-atom pair interaction energy for In-As</td>
<td>0.86 eV</td>
</tr>
<tr>
<td>$E_{Ga}$</td>
<td>activation energy for diffusion for isolated Ga atom</td>
<td>1.2 eV</td>
</tr>
<tr>
<td>$E_{d,iso}$</td>
<td>activation energy for diffusion for isolated In atom</td>
<td>1.3 eV</td>
</tr>
<tr>
<td>$E_{Ga}$</td>
<td>activation energy for diffusion for isolated As atom</td>
<td>1.2 eV</td>
</tr>
<tr>
<td>$E_{Gu}$</td>
<td>evaporation activation energy for isolated Ga atom</td>
<td>2.63 eV</td>
</tr>
<tr>
<td>$E_{e,iso}$</td>
<td>evaporation activation energy for isolated In atom</td>
<td>2.13 eV</td>
</tr>
<tr>
<td>$E_{As}$</td>
<td>evaporation activation energy for isolated As atom</td>
<td>2.63 eV</td>
</tr>
<tr>
<td>$E_{Gu}$</td>
<td>activation energy for the Ga evaporation</td>
<td>$0.18$ eV</td>
</tr>
<tr>
<td>$E_{In}$</td>
<td>activation energy for the In evaporation</td>
<td>$0.18$ eV</td>
</tr>
<tr>
<td>$E_{As}$</td>
<td>activation energy for the As evaporation</td>
<td>$0.18$ eV</td>
</tr>
<tr>
<td>$E_{Gu}$</td>
<td>segregation activation energy for the isolated In atom</td>
<td>$2.1$ eV</td>
</tr>
</tbody>
</table>

* $C_{Ga}^{phy}$ - Ga coverage in the physisorbed layer

** $C_{In}^{phy}$ - In coverage in the physisorbed layer

*** $C_{As}^{phy}$ - As coverage in the physisorbed layer
Figure 4.1: A schematic picture of the surface processes in MBE growth of InGaAs.
Figure 4.2: Comparison between experimental [1] and simulated results for \(\text{In}\) incorporation coefficient versus substrate temperature for a BEP of 36 with \(A_s^2\) and \(A_s^4\) fluxes.

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Figure 4.3: Comparison between experimental [1] and simulated results for In incorporation versus substrate temperature for a As$_4$ BEP of 17.
Figure 4.4: Simulation of In incorporation versus substrate temperatures for various BEPs of As$_4$. 

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Figure 4.5: Simulation of Indium desorption rate versus time for various substrate temperatures for an As$_4$ BEP of 36.
Figure 4.6: Simulation of Indium desorption rate versus time substrate temperatures 903°K for $A_s$ BEPs of 36 and 17.
Figure 4.7: Comparison of simulation and experimental results [1] for relative desorption parameters of Indium versus substrate temperature for a $As_4$ BEP of 36.

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Figure 4.8: Comparison of simulation and experimental results [1] for relative desorption parameters of Indium versus substrate temperature for a $As_4$ BEP of 17.
Figure 4.9: Simulation of Ga desorption rate versus inverse of substrate temperatures.
Figure 4.10: Simulation of Indium composition versus monolayer number for various substrate temperatures for a $\text{As}_4$ BEP of 36, for 10 sec. growth.
Figure 4.11: Simulation of Indium segregation versus substrate temperatures for $As_4$ BEPs of 36, 17 and a $As_2$ BEP of 36 along with the experimental data of Kao et al [8] for a BEP of 6.
Figure 4.12: Simulation of Indium segregation versus substrate temperatures for $As_4$ BEPs of 36, 20, 17, 12, and 6 and a $As_2$ BEP of 36.
The wide variation in the bandgap and lattice constants between InAs and GaAs allows for a variety of optical and electronic device applications. Many of the possible devices involve heterostructure in the active portions of the device and hence, the compositional and structural perfection of the interfaces is of paramount importance. In this materials system, the perfection is intrinsically controlled by the surface segregation of In due to its larger atomic size compared to Ga. In spite of several experimental investigations, there is a lack of a thorough understanding of the underlying surface dynamic processes and their interplay. In this work, a rate equation model is developed including several physically sound surface processes such as segregation from the crystalline layer to a surface riding In segregated layer and incorporation from the segregated In layer to crystalline layer. The rate of the processes are assumed Arrhenius type with concentration dependent activation energies. The simulated In incorporation coefficient versus substrate temperature is in excellent agreement with experimental data [1] for various As overpressure. For a constant As overpressure, In incorporation decreases with increasing temperature. For a constant temperature, In incorporation increases with increasing As overpressure. The In desorption versus time results from experiments and our simulation match very well. The desorption process has two components, one arising from the physisorbed layer of In and the other from the surface of the crystal. The activation energy for
these processes for an isolated \textit{In} adatom are 0.18 eV and 2.6 eV, respectively. These observations are explained based on the interplay of competing surface processes such as segregation and incorporation.

5.2 Future Recommendations

- Simulate \textit{GaAs} and \textit{InAs} desorption versus temperatures and compare results to Zhang \textit{et al} [3]. If results of activation energy do not agree, adjust model parameters

- Simulate pre-deposition of \textit{In} to achieve abrupt \textit{InGaAs/GaAs} heterostructure as suggested by Kaspi \textit{et al} [7].

- Explore independent way of obtaining bond parameters to use in the simulation instead of fitting.
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