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(Ph.D. Thesis)

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THIN FILM REACTIONS ON ALLOY SEMICONDUCTOR SUBSTRATES

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Ph.D. Dissertation

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by

Dale Abbott Olson

ABSTRACT

The interactions between Pt and In$_{53}$Ga$_{47}$As have been studied. In$_{53}$Ga$_{47}$As substrates with 70nm Pt films were encapsulated in SiO$_2$, and annealed up to 600°C in flowing forming gas. The composition and morphology of the reaction product phases were studied using x-ray diffraction, Auger depth profiling, and transmission electron microscopy. The reaction kinetics were examined with Rutherford Backscattering.

Results show that Pt/In$_{53}$Ga$_{47}$As reacts to form many of the reaction products encountered in the Pt/GaAs and Pt/InP reactions: PtGa, Pt$_3$Ga, and PtAs$_2$. In addition, a ternary phase, Pt(In:Ga)$_2$, develops, which is a solid solution between PtIn$_2$ and PtGa$_2$. The amount of Ga in the ternary phase increases with annealing temperature, which causes a decrease in the lattice parameter of the phase.

The reaction products show a tendency to form layered structures, especially for higher temperatures and longer annealing times. Unlike the binary case, the PtAs$_2$ phase is randomly oriented on the substrate, and is intermingle with a significant amount of Pt(In:Ga)$_2$. Following Pt/In$_{53}$Ga$_{47}$As reactions, two orientation relationships between the Pt(In:Ga)$_2$ product phase and the substrate were observed, despite the large mismatch with the substrate (~8%). At higher annealing temperatures, one of the two orientation relationships becomes dominant.
For many metal/compound semiconductor interactions, the reaction rate is diffusion limited, i.e. exhibits a parabolic dependence on time. This study shows that the early stages of the Pt/In\textsubscript{53}Ga\textsubscript{47}As reaction progress at a rate similar to that of Pt/GaAs. For longer annealing times, the reaction slows dramatically, to a rate \~one order of magnitude slower than for Pt/GaAs. This reduction in reaction rate should prove useful in producing more stable contacts to compound semiconductors by inhibiting contact/substrate reactions.

An additional result of this study was the development of an In-rich layer beneath the reacted layer. The Auger depth profile showed a substantial increase in the sample at this layer. This is a significant result for the production of ohmic contacts, as the Schottky barrier height in this system is lower for higher In concentrations.
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1.0 INTRODUCTION

1.1 Semiconductor Contacts

A well-designed metal contact is critical to the proper operation of any semiconductor device (figure 1.1). The contact provides the means for communication with the device, and therefore must exhibit very strictly defined electrical characteristics. In addition, the metallization maintains intimate contact with the semiconductor substrate, often at elevated temperatures, and consequently is prone to complex metallurgical interactions which can have a profound effect on the electrical characteristics. Despite these environmental conditions, the contact must maintain its desired characteristics for the projected lifetime of the device.

The electrical characteristics of the contact are determined by the device requirements, and can vary widely. The more basic material requirements, however, are universal: a) the electrical characteristics must be predictably and reproducibly attainable, b) the electrical characteristics of a given contact type must be uniform across each device, and across each wafer, c) once the desired characteristics have been achieved, they must not be affected by subsequent processing (e.g. annealing temperatures up to 400 °C for packaging, or 850 °C for ion implantation activation anneals), and, d) the contact must be stable; i.e., the properties of the contact should not change over the lifetime of the device.
Figure 1.1: Schematic of an FET, showing the approximate dimensions and geometry.
From the metallurgical perspective, satisfying these requirements is quite difficult, as there are many possible interactions which would interfere with the device operation. The characteristics are generally dependent on the phases and the morphology of the metal/semiconductor interface [WM79, SP80b, FR81, BI82, LL84, RO85]. If the metal reacts with the substrate, the electrical characteristics of the device may be degraded. Reaction with the metal layer can lead to the formation of electrically active defects in the substrate, or to the formation of new interfacial phases with undesirable electrical properties [WM79, SP80b, FR81]. If extensive chemical reaction occurs, definition of the contact dimensions will be difficult to maintain; the reaction may even consume critical device features. Contact areas can impinge, thereby destroying the device. During reaction, morphological changes disrupt the uniformity of the interface.

Liquid phases which may develop during processing are undesirable, as they lead to sensitivity to processing parameters (e.g. annealing temperature and time, heating and cooling rates) and rough interfaces [GR80]. Liquid phases also make control of lateral dimensions difficult, a critical parameter for VLSI devices, where control down to the submicron scale is required.

The grain boundaries of a polycrystalline contact material are also detrimental. They can provide diffusion paths for undesirable reactions, leading to the introduction of unwanted elements from metal overlayers (e.g. Au) into active regions of the device. Reactions which proceed by grain boundary (vs. bulk) diffusion will generally result in rough interfaces, and may result in localized penetration of active layers, i.e. spiking. Rough interfaces and spiking can interfere with the proper operation of the device by intruding into conducting channels. A rough interface can result in an unpredictable contact depth [BH87], while spiking can lead to short circuits and the destruction of device.

There are many other potentially damaging interactions. Excessive reaction with the substrate may consume important features, as active layers of the device may be within one micron of the surface. New phases that are the products of
contact/substrate interactions may possess undesirable electrical properties. If the contact structure is not thermodynamically stable, reactions may continue during operation of a device at an unacceptable rate, compromising the reliability of the device. This is especially serious for the specialized applications of many compound semiconductors, such as satellites or trans-oceanic optical communications devices, where replacement is expensive or impossible.

The general requirements discussed above are applicable to compound as well as elemental semiconductors, but for compound semiconductors there are two more factors which must be considered. First, the extra elements in the substrate increase the opportunity for undesirable interactions, since each additional element allows the possibility of an additional phase. Second, the stoichiometry of the semiconductor substrate must be controlled during any interactions that may occur; if the elements are removed from the compound substrate at different rates, the electrical properties of the semiconductor itself will change. It should be noted that this change in electrical behavior can be beneficial or detrimental, depending on the circumstances.

While taking all of these problems into account, the goal of contact design is the ideal contact. This contact would be thermodynamically stable, would have a smooth, abrupt interface with the substrate, would not consume excessive amounts of the substrate during fabrication, would not involve liquid phases during processing, would have no grain boundaries, and could be fabricated by conventional techniques.

1.2 Epitaxial Contacts

This is a complex set of requirements which cannot be met by simple contact structures; particularly since most single-element metallizations react with GaAs substrates [LA85]. However, more complex metallizations may result in a structure which satisfies many of the requirements of an ideal contact. For example, introduction of a thin, stable intermediate layer which inhibits diffusion between the metal contact and the semiconductor substrate would increase the lifetime of the device by reducing
further reactions. If this phase were an epitaxial single crystal, it would eliminate the grain boundaries and rough interfaces associated with a polycrystalline reaction layer. A large-grained, polycrystalline layer could be similarly advantageous, as it would reduce the number of high-diffusivity paths (i.e. grain boundaries).

Deposition

One promising technique to achieve near-ideal contact structures is direct deposition of suitable compounds on the semiconductor surface (e.g. AuGa_L/GaSb [LC85], RhGa/GaAs [GU88a], YbAs/GaAs [RI88], ErAs/GaAs [PA88b]). These compounds were chosen because they were stable in contact with the substrate, and had appropriate crystal structures and lattice parameters for epitaxial growth. In order to obtain epitaxial growth, these compounds must be directly deposited on the atomically clean semiconductor surface, and therefore require ultra-high vacuum (UHV) deposition techniques (<10^{-9} Torr), such as molecular beam epitaxy (MBE). UHV is necessary because all compound semiconductor surfaces develop a thin oxide layer (1-2nm) after even a short exposure to air [WI85]. The contacts were prepared by first treating the semiconductor surface in the MBE chamber to remove the native oxide, then depositing the desired compound. The disadvantage of this technique is the strict vacuum requirements and high cost of MBE, and consequent incompatibility with conventional patterning techniques.

Reaction

An alternate technique for producing the ideal epitaxial contact structure, without the high cost and strict vacuum requirements mentioned above, is to use a controlled metal/semiconductor interaction to form desirable structures. An elemental metallic layer can be deposited by conventional high vacuum techniques (∼10^{-6} Torr), then annealed to induce a solid phase reaction. Although it may not be possible to attain a monocrystalline, epitaxial layer, considerable improvement of the contact structure
could still be obtained by reducing interfacial roughness and polycrystallinity; such an improvement might be attained by local epitaxy of the individual grains of the polycrystal.

There are several compounds which have the potential to satisfy the requirements for an ideal contact. In many metal/A\textsuperscript{III}B\textsuperscript{V} systems, the metal and semiconductor react to form a layered structure, with a stable metal-B\textsuperscript{V} or metal-A\textsuperscript{III} phase present at the semiconductor interface. These phases provide candidates for an inert barrier layer. Many of them are cubic, and have lattice parameters within the range of important compound semiconductors. Preferred orientation relationships are often exhibited between these reaction product phases and the substrate, even some with high lattice mismatch; by reducing the lattice mismatch between the substrate and the product phase, it may be possible to create a stable layer which is effective at inhibiting contact/device metallurgical interactions.

1.3 Objective

The focus of this study is on the second method for improving the stability of a contact: controlling the metal/substrate reaction to produce desirable microstructures. In this experiment, the metallurgical effects of changing the substrate composition have been examined. This consists of varying the substrate composition and monitoring the effects on: a) phase formation, b) morphology of the reaction products, and c) kinetics of the metal/semiconductor reaction.

Ternary substrates are the focus of this study, as they can be manipulated more easily to match the required characteristics. In many of these systems, the lattice parameter follows Vegard's law, and varies linearly between the two pure binary values. This facilitates control of the lattice mismatch between the substrate and the product phases.
The Pt/In$_x$Ga$_{1-x}$As system was selected for examination in this study. As will be discussed in subsequent sections, this system satisfies many of the requirements for potential solid phase epitaxial growth. In the sections that follow, the general issues faced by designers of stable contacts will be discussed, followed by specific discussions of the Pt/In$_x$Ga$_{1-x}$As system.
2.0 CONTACTS TO SEMICONDUCTORS

This study has examined the development of stable contact structures by controlled reaction, in order to evaluate the potential of this technique for use in actual contact structures. In this chapter, current schemes for producing practical contact structures will be discussed. In order to provide a basis for understanding the motivations for the design of these structures, a summary of the electrical characteristics of metal contacts to semiconductors is presented.

2.1 Metal/Semiconductor Junctions [SZ81, RO85, RH88]

When a metal and a semiconductor are brought into contact, equilibrating processes occur that accommodate the electrical properties of the two dissimilar materials. The most significant result is the development of an electronic potential barrier at the interface, which can be utilized to control electrical conduction across the couple. Various models have been proposed to describe this equilibration, and to predict the height of the potential barrier, $\phi_B$, which develops.

Schottky Limit [SO38, MO38]

If a metal and a semiconductor are in electrical contact, the equilibrium condition must be satisfied, i.e. the Fermi levels in the metal and semiconductor must be everywhere equal ($E_{Fm}^{m} = E_{Fs}^{s}$). Under the appropriate conditions (e.g. n-type semiconductor, no surface states, flat band, $\phi_m > \phi_s$, where $\phi_m$ and $\phi_s$ are the work functions of the metal and semiconductor), electrons will flow from the semiconductor into the metal. This will result in an electric field directed from the semiconductor to the metal. Excess negative charge will accumulate on the surface of the metal, in the form of a narrow band of conduction electrons. A corresponding positive charge will build up on the surface of the n-type semiconductor in the form of uncovered positive ion cores. Because of the much lower concentration of charge carriers in semiconductor relative to the metal, the width of the charge compensating region (i.e. the depletion width; $w$)
will be much greater in the semiconductor. The charge build-up results in an upward bending of the semiconductor energy bands. When the metal and semiconductor are brought into intimate contact, a potential barrier situated in the semiconductor will be present, with a magnitude that can be modelled as:

$$\phi_B = \phi_m - \chi_s$$

i.e the Schottky barrier. \(\chi_s\) is the electron affinity of the semiconductor, and corresponds to the potential difference between the bottom of the conduction band and the vacuum level.

This model for the formation of \(\phi_B\) does not predict the barrier heights very successfully. It implies that the Schottky barrier depends on \(\phi_m\), while in practice, \(\phi_B\) for a given semiconductor is insensitive to the choice of overlaying metal.

**Bardeen Limit [BD47]**

Another model proposes that electronic states at the metal/semiconductor interface accommodate the differences in \(\phi_m\) and \(\chi_s\), producing a metal-independent barrier height for a given semiconductor. Several models have been proposed to explain the origination of these charged surface states, attributing them to intrinsic surface defects, such as \(V^{III}\) or \(V^V\) [SP80], metal-induced states [LO76], or interfacial phases [WO81].

These states may exist within the forbidden energy gap of the semiconductor, and play a role in the shape of the energy bands near the surface. If the surface is charge neutral, those states located at energies lower than \(\phi_o\) (the neutral level) will be filled, while those above \(\phi_o\) will be empty.

If the density of these states, \(N_{ss}\), is high enough, the Fermi level of the semiconductor can be pinned to the neutral level. Under these conditions, the model predicts that the barrier height of the contact will be given as:

$$\phi_B = E_g - \phi_o$$
There are approximately $10^{15}$ surface atoms per square centimeter on a typical semiconductor; $10^{13}$ surface states per square centimeter (i.e. one out of every 100 surface atoms) would be sufficient to pin the Fermi level [RO85].

**Experimental Evidence**

In reality, the behavior of the metal-semiconductor contact falls somewhere in between these two extremes, although the dependence of $\phi_B$ on the material parameters $\phi_m$, $\chi_s$, and $E_g$ has not yet been satisfactorily described.

Experimentally, some A$_{III}$-B$_{V}$ semiconductors do exhibit pinning of the Fermi level by charged surface states [BR86]. In the case of GaAs, $E_f$ is found to be pinned at $\sim 0.8$ eV below the conduction band edge. In contrast to the strict Bardeen limit, the barrier height of metals on GaAs does vary slightly with the deposited metal and surface preparation, by about 0.1 eV [BR86]. This variation is unacceptable for the production of many devices, which may require control to within 0.02 eV [BR86]. By comparison, it has been shown that the Fermi level of In$_x$Ga$_{1-x}$As does not pin for atomically clean surfaces, but does for air-exposed [BI86a,b].

### 2.2 Conduction Behavior

There are two different types of contacts to semiconductors: ohmic and rectifying. In the first case, the current-voltage behavior is linear, with the slope defined as the total contact resistance, $r_c$. This contact resistance characterizes the ohmic contact. In the second case, the current-voltage characteristic under forward-bias is exponential, while under reverse-bias, the current saturates at a very low value at low voltage. The rectifying junction is characterised by the Schottky barrier height, $\phi_B$, which appears in the argument of the exponential term. For metal-semiconductor contacts, the rectifying contact is often referred to as a Schottky contact.
Charge carriers can cross the Schottky barrier that develops at metal/semiconductor interfaces by four means: 1) by acquiring sufficient energy to jump over the barrier (thermionic emission), 2) by quantum-mechanical tunneling through the barrier (field emission), 3) by recombination in the space charge layer, and 4) by minority carrier injection from the metal into the semiconductor and recombination. The last two modes are significant only under reverse bias.

For thermionic emission, the current density between the semiconductor and the metal is determined by the concentration of carriers which have sufficient energy to jump over the intervening barrier. For carriers entering the semiconductor from the metal, the barrier is $\phi_B$. This potential is relatively independent of applied voltage, and will not change much with the bias. For carriers entering the metal from the semiconductor, the barrier is the diffusion potential, $V_d$, which represents the difference between the conduction band minimum in the bulk semiconductor and the Schottky barrier height.

$V_d$ can be affected by bias. Under forward bias, $V_d$ is diminished, allowing increased flow of electrons from the semiconductor to the metal. As $\phi_B$ is relatively independent of bias, the reverse current from the metal to semiconductor current is unaffected. Under reverse bias, $V_d$ is increased, which decreases the flow of electrons from the semiconductor to the metal.

The metal/semiconductor current under bias $V$ can be described with an exponential function:

$$J = J_s \left[ \exp\left(\frac{qV}{nkT}\right) - 1 \right]$$  \hspace{1cm} (2.3)

where

$$J_s = A^* T^2 \exp\left(-\frac{q\phi_B}{kT}\right)$$  \hspace{1cm} (2.4)

$J_s$ is the saturation current, i.e. the current under reverse bias. The constant $A^*$ is known as the Richardson constant, and depends in part on the effective mass of the
majority carrier. The \( n \) in the second exponential is known as the ideality factor, and is a measure of the deviation from an ideal, thermionic diode. For an ideal diode, \( n=1 \), for thermionic-dominant, \( n \approx 1.02 \), while in typical diodes, \( n>1 \) (\( \approx 1.2 \)).

Under reverse bias (\( V < 0 \)), the carriers in the metal must still possess energy greater than \( q\phi_B \) in order to pass into the semiconductor. Therefore, the current under reverse bias will remain at a small, constant value, thus giving the contact its rectifying characteristics. It is important to note the good rectifying contacts require a sufficiently high \( \phi_B \), otherwise an unacceptably large concentration of carriers in the metal will have sufficient energy to jump over the barrier in reverse bias. This leads to a large reverse leakage current, which may render the device unusable. The critical parameter for Schottky rectifying contacts is the contact barrier height, \( \phi_B (eV) \); good contacts generally have \( \phi_B \geq 0.7 \text{eV} \).

The second means by which carriers may cross the barrier is by field emission, also known as tunneling. This mode becomes dominant at high doping and low temperatures. Increasing the doping level in the semiconductor below the contact will decrease the width of the depletion layer, and hence the width of the Schottky barrier. If the doping is sufficiently high, the depletion layer will be thin enough to allow field emission of carriers across the barrier. At intermediate doping levels, some carriers will have sufficient energy to tunnel through the barrier near the top where it is thinner. This process is known as thermionic-field emission. Both of these processes can give rise to ohmic behavior of the contact.

The critical parameter for ohmic contacts is the specific contact resistance, \( r_c (\Omega \text{cm}^2) \). For ohmic contacts with non-zero \( \phi_B \), \( r_c \) depends on \( \phi_B \), the effective mass of the carrier, \( m^* \), the permittivity of the semiconductor, \( \epsilon_s \), and the net doping concentration, \( N(\text{cm}^{-3}) = |N_A - N_d| \). Assuming that the tunneling process dominates, the dependence of the specific resistance on the interfacial parameters is:

\[
r_c \propto \exp \left[ \frac{2\sqrt{\epsilon_s m^*}}{\hbar} \left( \frac{\phi_B n}{\sqrt{N}} \right) \right]
\]  (2.5)
Therefore, the specific contact resistance depends exponentially on the inverse square root of the doping concentration for a given barrier height.

For high doping concentrations, $r_c$ does not depend on temperature, while for low doping concentrations, the dependence is exponential. This difference in temperature dependence arises because conduction by thermionic-field emission becomes more significant as the width of the barrier increases. For field-emission ohmic contacts in III-V semiconductors, doping concentrations of $10^{19}\text{cm}^{-3}$ and above are required. In practice, however, achieving doping levels this high is quite difficult (e.g. [PA85], p. 219)

Ohmic conduction will also occur when $\phi_B = 0$. In these contacts, the current-voltage behavior is linear, i.e. ohmic. This is found, for example, in the case of InAs, where the Fermi level pins in the conduction band. Ohmic contacts on all other III-V semiconductors are of the tunneling type previously described.

$\text{In}_x\text{Ga}_{1-x}\text{As}$

The endpoints of the InAs-GaAs alloy system are very different in their contact behavior. The electronic band structures are qualitatively very similar, although the minimum energy gap between the conduction and valance bands (both are direct) are quite different [PH73]. At 300K, $E_g$ for GaAs is 1.43eV, while for InAs, $E_g$ is 0.36eV. $E_g$ for $\text{In}_x\text{Ga}_{1-x}\text{As}$ varies continuously between these endpoints (Figure 2.1), and can be expressed empirically as [MD82]:

$$E_g(\text{eV}) = 1.43 - 1.39x + 0.32x^2$$  \hspace{1cm} (2.6)

$\text{In}_x\text{Ga}_{1-x}\text{As}$ presents some difficulties to the formation of good rectifying contacts to both p- and n-type material because of its small band gap. The GaAs band gap is fairly large, and so does not present such a problem. For GaAs, $q\phi_{Bn} \approx 0.8\text{eV}$, and $q\phi_{Bp} \approx 0.5\text{eV}$ [e.g. MC87]. At the other end of the alloy system, the metal/p-InAs, the barrier height has been reported as $\approx 0.44\text{eV}$ [WA71]. This is significant, as $q\phi_{Bp}$ is
Figure 2.1: Plot of the energy gap of In$_x$Ga$_{1-x}$As vs. x. Calculated from equation 2.6. After Madelung, 1982 [MD82].
larger than \( E_g \). Assuming that the sum of the n- and p-type barrier heights is the band gap [e.g. RH88, p.25], \( \phi_{Bn} \) should be negative. In fact, the metal/n-InAs contact is ohmic.

For intermediate \( \text{In}_x\text{Ga}_{1-x}\text{As} \) compositions, the band gap is necessarily less than that of GaAs. For \( \text{In}_{0.53}\text{Ga}_{0.47}\text{As} \), which is a useful composition as it is lattice matched to InP, \( E_g (300K) = 0.75\text{eV} \). Consequently, the Schottky barriers on n-type material are quite small: \( q\phi_{Bn} \approx 0.2\text{eV} \) [KA73], \( q\phi_{Bp} \approx 0.55\text{eV} \) [VE82]. For this substrate composition, good ohmic and poor rectifying contacts are attainable on n-type, while poor ohmic and good rectifying contacts are attainable on p-type.

Kajiyama et al. [KA73] measured the Schottky barrier heights of air-exposed \( \text{Au/n-In}_x\text{Ga}_{1-x}\text{As} \) contacts as a function of \( x \). They found that \( q\phi_B \) tracked the conduction band edge, with a spacing of approximately 0.5eV, and that the barrier height relationship could be described empirically (Figure 2.2):

\[
\phi_B(\text{eV})=0.95-1.90x+0.90x^2 \tag{2.7}
\]

Combining this result with the relationship between the barrier heights and the band gaps as a function of \( x \), the largest In concentration that will allow a reasonable \( \phi_B \) for both n- and p-type material is for \( x \approx 0.25 \) (\( q\phi_{Bn} \approx 0.53\text{eV} \), \( q\phi_{Bp} \approx 0.52\text{eV} \)).

Brillson et al. [BI86a, b] measured the Fermi level stabilization energies of four different metals (Au, Al, In, and Ge) on four different compositions of \( \text{In}_x\text{Ga}_{1-x}\text{As} \). They reported that the Fermi level stabilization energies for a given substrate composition \( x \) varied considerably with the deposited metal. They attributed this variation to the formation of different interfacial phases with different metal depositions, in some cases As-rich, and in others As-deficient. In agreement with Kajiyama’s results, they found that the Fermi level tracked the conduction band edge for Au, Al, and In. However, they found considerably different values for \( \phi_{Bn,Au} \). They attributed this difference to surface preparation, i.e. air-exposed [KA73] vs. clean [BI86a, b], and suggested that air-exposure of the surfaces led to pinning.
Figure 2.2: The Schottky barrier height of Au on In$_x$Ga$_{1-x}$As vs. x. As measured by Kajiyama, 1973 [KA73].
In summary, for some metals on In$_x$Ga$_{1-x}$As, $\phi_B$ shows a definite functional relationship with $x$. For variations in the choice of overlaying metal on UHV clean surfaces, $\phi_B$ is not constant, i.e. $E_f$ is not pinned. For air-exposed surfaces, there is evidence of pinning.

2.3 Practical Contacts

For both ohmic and rectifying contacts, the requirements for contact behavior, i.e. $r_c$ and $\phi_B$, are largely determined by the specific device in question. The information provided below applies specifically to GaAs, but can be extended to other A$^{III}$-B$^{V}$ substrates.

Rectifying Contacts

The rectifying contact is characterized by the Schottky barrier height $\phi_{Bn,p}$, the ideality factor, $n$, and the sheet resistance. For conventional devices (e.g. FETs), barrier heights above 0.7eV are acceptable, although higher barriers are more desirable. The reverse saturation current, $J_s = \Lambda^* T^2 \exp(-q\phi_B/kT)$, increases with increasing $T$. Therefore, for a given application, higher operating temperatures require higher barrier heights. Conversely, for very low $T$ applications, lower barrier heights may be acceptable. Additional requirements are imposed by high radiation environments (e.g. for satellite and military applications), which can also excite electrons over the barrier.

The general requirements for rectifying contacts are as follows. $\phi_B$ must be high enough to prevent significant reverse bias current, so that as little current as possible is injected into the device through the gate (which operates under reverse bias). The barrier heights must be uniform across the device wafer. Circuits are designed to accommodate the characteristics of the rectifying contacts; if the parameters change across the device, it will not function properly. For the same reason, the barrier height must be reproducibly attainable; if the parameters change from run to run, the same
problem will arise. Higher barrier heights are better; they allow lower saturation currents, and reduce the uniformity and reproducibility requirements. The optimum contact would be single-phase, epitaxial, have an atomically abrupt interface, and have \( q\phi_B > 1.0V \).

Since most materials deposited onto GaAs form Schottky barriers, development of rectifying contacts has focussed on the high temperature stability and electrical properties of the contact material, rather than on beneficial reactions with the substrate. Materials for rectifying contacts must withstand high temperatures during processing. For self-aligned gates, the contact acts as a mask for ion-implantation, a process which requires subsequent annealing at high temperature in order to “heal” substrate lattice damage and activate the implanted species. Consequently, the contact material must be stable at high temperature, and must not react strongly with the substrate.

Elemental contacts are ineffective under these conditions, as most single-element contacts react with compound substrates [LA85]. Compound metallizations are therefore desirable, but they must not decompose or react at high temperature, and must have appropriate electrical characteristics. The refractory metal silicides and nitrides are the most promising contacts for these aspects [WL83, WU87].

The refractory contacts are deposited by chemical vapor deposition, with nitrogen present in some form (\( \text{NH}_3 \) for TiN [WL83], \( \text{N}_2 \) for NbN [WU87]) in the deposition chamber. The TiN-GaAs \( q\phi_B \) was reported as 0.83eV, the NbN-GaAs \( q\phi_B \) as 0.73eV. In the case of the NbN contact, some improvement in the ideality factor and increase in the barrier height were observed after annealing, possibly as a result of healing the damage introduced during deposition.
Ohmic contacts

In contrast to rectifying contacts, ohmic contacts require considerable interaction with the substrate in order to achieve optimum characteristics. For ohmic contacts, the basic requirement is that the voltage drop across the interface is insignificant compared to the voltage drop across the device. As described above, all contacts to $A^{\text{III}}$-$B^{\text{V}}$ semiconductors (except InAs) will exhibit a potential barrier. Examination of equation 1.1 suggests two schemes for creating low-resistance ohmic contacts: either by reducing the width of the barrier by increasing the doping concentration immediately below the contact (field emission), or by lowering the barrier height by introducing an intermediate layer of another semiconductor which has lower contact barrier heights (thermionic emission).

The requirements for the resistance of the contact are determined by the particular device. An important relation is that: $R_c = r_c/A$, where $R_c$ is the total contact resistance, and $r_c$ is the specific contact resistance ($\Omega \text{cm}^2$). This relationship allows for a large range of acceptable specific contact resistances, depending on the device application. Devices such as lasers and light emitting diodes have large contact areas (typically about $10^{-3} \Omega \text{cm}^2$), so that contact resistances in the range of $10^{-2}$-$10^{-5} \Omega \text{cm}^2$ are adequate, giving a total contact resistance on the order of 1 $\Omega$.

As minimizing the size of the device becomes more important (e.g. MESFETS and MOSFETS), the required specific contact resistance must also decrease. For typical devices, an acceptable $r_c$ is on the order of $10^{-6} \Omega \text{cm}^2$. This is determined by the limitation of the resistivity of the contact materials. This requires a doping concentration of greater than $5 \times 10^{19} \text{cm}^{-3}$, which is higher than the solid solubility limit of the dopants [BR86]. In order to achieve these doping levels, special epitaxial techniques (e.g. MBE) are required. Even when sufficiently high doping levels are achieved, making the dopant species electrically active can be challenging [PA85]. Both Ge and Si are used as amphoteric dopants in $A^{\text{III}}$-$B^{\text{V}}$ compounds. On the $A^{\text{III}}$ site, they act as shallow donors, while on the $B^{\text{V}}$ site, they act as shallow acceptors. Once the solubility
limit has been surpassed, the dopants may begin to form electrically inactive pairs [SA88].

Most conventional ohmic contacts are of the tunneling type. For these contacts, an alloy containing the contact metal and a suitable dopant are deposited onto the substrate, and then heated to incorporate the dopant into the semiconductor [PA85]. Alloyed contacts are heated sufficiently to melt some of the components to assist in the doping, while sintered contacts rely on solid-state diffusion. The most common ohmic contacts contain Ge or Si (for doping), Pd or Ni (to enhance uniformity by breaking up the native oxide) and Au or Ag (for the interconnection and low eutectic temperature) [RO85].

The Au-Ni-Ge contact is an example of this type [BR67]. The dopant is incorporated into the substrate by dissolution of the GaAs during annealing. When the substrate recrystallizes, the high solubility of Ga in the metal alloy leads to the formation of As-rich material. \( \text{Ge}_{Ga}^+ \) sites are thereby produced in the substrate. \( \text{Ge}_{As}^- \) can be produced by annealing above 500 °C to create As vacancies [SA88].

A new type of ohmic contact that is very promising is the regrown In-based contact [e.g. MU88]. In these contacts, Ni-In-W was deposited onto GaAs and annealed at high temperature (800-1000 °C). The contacts exhibit contact resistances equal to or better than the Au-Ni-Ge contact, with much better thermal stability. The contacts are believed to operate by the regrowth of a low-barrier \( \text{In}_x\text{Ga}_{1-x}\text{As} \) phase at the contact interface, with excess In forming a refractory compound.

Heteroepitaxial contacts have been made with \( r_c \approx 10^{-7}\Omega \text{cm}^2 \) [PA85]. These were achieved with highly doped (\( >10^{20}\text{cm}^{-3} \)) Ge or low \( \phi_B \text{ In}_x\text{Ga}_{1-x}\text{As} \) layers deposited epitaxially onto the substrate prior to metallization. Unfortunately, these contacts are limited by fabrication problems. They are deposited by MBE, where patterning of the contact is not practical [SA88].
Contacts to In$_x$Ga$_{1-x}$As

Excellent Schottky barrier diodes have been produced for p-In$_{53}$Ga$_{47}$As [SE85]. Barrier heights of $\approx 0.7$eV with ideality factors of $\approx 1.2$ were made by evaporating Al, Ti, Au, and Ni onto Ar$^+$ sputter-etched surfaces.

Very low contact resistance ohmic contacts have been developed to In$_{53}$Ga$_{47}$As [KR85]. The contacts were fabricated by alloying techniques, by depositing layers of Ni/Au-Sn/Ni and Ni/Au-Zn/Ni (on substrates doped to $10^{18}$cm$^{-3}$) and annealing at 400°C for 1 to 2 minutes. The best contact resistances reported were $4 \times 10^{-8}$Ωcm$^2$ for Ni/Au-Sn/Ni on n-In$_{53}$Ga$_{47}$As and $2 \times 10^{-5}$Ωcm$^2$ for Ni/Au-Zn/Ni on p-In$_{53}$Ga$_{47}$As.
3.0 Epitaxial Contacts

The most commonly used scheme for producing an ohmic contact to a III-V semiconductor is the Au-Ni-Ge type, which has been described previously. Although this contact can yield low contact resistance ($\sim 10^{-6} \Omega \text{cm}^2$), it suffers from non-uniformity (by deep diffusion of Au [GY71]) and excessive penetration into the substrate material (up to 100 nm [KU83]). As well as causing rough interfaces, Au has been well documented as a source of failure in opto-electronic devices by the formation of non-radiative recombination centers (e.g. [UE84]). In addition, the Au-Ni-Ge contact is very sensitive to annealing conditions, and not very thermally stable [KU83].

One of the most important requirements is that the contact avoids any liquid phases during processing, since these lead to sensitivity to processing parameters (e.g. temperature, time, and heating/cooling rates) and rough interfaces [GR80]. In order to improve reliability, uniformity, and dimensional control, various groups have developed methods for producing contacts with flat, abrupt interfaces. The basis of these schemes is the introduction of a stable, epitaxial contact layer onto the semiconductor substrate. In this way, atomically abrupt interfaces may be produced. Currently, there are three schemes for producing this contact layer: epitaxial deposition of suitable compounds, solid-state diffusion with epitaxial precipitation, and reaction to form epitaxial products.

It should be noted that “epitaxial” refers to overlayers which at least have a polycrystalline structure with one or two specific orientation relationships with the substrate, or at best, a monocrystalline layer with a specific orientation relationship with the substrate.

3.1 Conditions for Epitaxy [KH70, MT75, part B]

The issue of epitaxial growth between two crystals can be quantified by examining the free energy of an interface between given planes of the phases. If a well-defined
minimum in the interfacial free energy of the bicrystal vs. orientation exists, then coherent overgrowth can occur. The broader this minimum is, the greater variation in angular alignment there will be around the minimum energy configuration.

The simplest free energy approximation is based on the geometry of the couple: if a plane of crystal A can be layed on a plane of crystal B so that there is considerable overlap of A and B lattice sites (coincidence), the couple is likely to have lower energy than a random configuration. This is more likely to occur under the circumstances where the atomic configurations of the two crystal planes is similar (e.g. square lattice on square lattice), and where the dimensions are compatible (small lattice mismatch). Symmetry also plays an important role [VT74]. If the symmetry of the overlying material is less than that of the substrate, variant domains will likely result. The number of variants is determined by the number of symmetry elements present in the substrate that are absent in the overlayer. The boundaries between the domains will not in general be incoherent, but are related by the symmetry elements in the system.

In the case of identical crystal structures, the possibility of identical orientations is clear. Good matches can also be found in seemingly incompatible crystal structures: hexagonal (0001) on cubic (111) for instance, or cubic on cubic where the ratio of the lattice parameters is a suitable value (e.g. $\sqrt{2}$ or some rational number). However, consideration of the geometry of potential phases is only a rough approximation for predicting epitaxial growth.

### 3.2 Epitaxial Deposition

Epitaxial contact structures can be developed most easily by the direct deposition of a suitable compound. A candidate for the epitaxial phase must satisfy certain requirements: a) it must have a lattice parameter and crystal structure appropriate for epitaxial growth on the desired substrate, b) it must have satisfactory electrical characteristics, c) it must be stable at subsequent fabrication temperatures, d) it must not react strongly with the substrate, and e) it must be compatible with the deposition
technique.

Because of the native oxides and contamination present on air-exposed semiconductor surfaces, epitaxial deposition requires UHV techniques such as MBE. MBE allows for much more precise control of the growth conditions than is available for conventional techniques, but as a consequence leads to throughput and patterning problems. There are advantages to forming epitaxial contact layers by deposition. Most important is the potential for forming buried contact layers, by creating epitaxial semiconductor/metal/semiconductor structures. More precise control of the deposition conditions afforded by MBE allows for the subsequent deposition of device-quality semiconductor material, a requirement for 3-dimensional device structures.

Elemental Substrates

Epitaxial deposition of several compounds on (111) (CoSi$_2$ [TN82a], NiSi$_2$ [TN82b], CaF$_2$ [IS82], CrSi$_2$ [FA88], ErSi$_2$ [AV89]) and (100) (CaF$_2$ [IS82]) Si substrates has been reported. In these examples, the epitaxial layers were formed by the codeposition of the metal and Si in the MBE chamber. However, it is much more common to form epitaxial silicides by MBE deposition of the pure metal, and annealing to form the corresponding silicide.

In general, deposition of the silicide produces better crystalline quality than deposition followed by reaction, but not always (e.g. NiSi$_2$, [TN82b]). In some cases, the deposition of the silicide results in a low quality crystalline (CaF$_2$, [PF85]) or amorphous layer (ErSi$_2$, [AV89]). In these cases, improvement in the quality of the layer was obtained by annealing the specimen after deposition.

The deposited compounds are generally cubic structures, with lattice parameters close to that of silicon (e.g. CoSi$_2$ and NiSi$_2$). Hexagonal compounds have also been grown epitaxially (e.g. CrSi$_2$ and ErSi$_2$), by taking advantage of the three-fold symmetry of the (111) Si orientation.
Compound Substrates

The technique of depositing epitaxial contact structures has been applied to compound semiconductors as well. The requirements for suitable compounds are the same as those mentioned above (section 2.4.1), although the extra substrate element makes the search more difficult. Since most elemental metals react with compound semiconductors, research in this area has focussed on the deposition of stable compounds. One complication of direct deposition is the necessity for precise control over the relative flux ratio \((\alpha)\) of the elements in the compound [LC85]. Deviations from the optimum ratio can lead to undesirable reactions with the substrate; unlike the silicide/silicon case where variations in the flux ratio will result only in additional silicon or silicide.

Lince and Williams [LC85] have identified one such compound for deposition on GaSb. \(\text{AuGa}_2\) has a cubic fluorite structure, with a lattice mismatch to GaSb of only 0.3\%. According to the Au-Ga-Sb ternary phase diagram, a tie-line connects the GaSb and \(\text{AuGa}_2\) phases, which means they are stable in contact with each other. Lince and Williams were successful in forming a \(\text{AuGa}_2\) layer by codepositing Au and Ga, but found the crystallinity of the product was poor. In addition, they discovered that an excess Ga flux led to the formation of liquid Ga metal, while a Ga-deficient flux led to the formation of both AuGa and \(\text{AuGa}_2\).

\(\text{AuGa}_2\) was also deposited on GaAs substrates [LC86], but in this case the lattice mismatch was very high, \(\approx 7\%\). Lince, et al. found that a dominant orientation relationship ((001) planes of the crystallites parallel to the substrate surface) was exhibited between the deposited \(\text{AuGa}_2\) and the substrate, although other orientations were also present in the polycrystalline film. During annealing above the melting temperature of \(\text{AuGa}_2\), the deposited film broke up into islands, which exhibited the dominant orientation relationship.

Guivarc'h, et al. have experimented with rhodium compounds on GaAs [Gu88b]. Based on lattice mismatch of 0.4\%, and a cubic crystal structure, \(\alpha\text{Rh}_2\text{As}\) would be a
likely candidate for epitaxy. The lack of success of this attempt illustrates the difficulty of predicting epitaxial growth from geometric factors. They found that rather than forming an epitaxial layer with the cube axes parallel, the deposited compound was polycrystalline with four different orientation relationships present: two with (001) $\alpha$Rh$_2$As $\| (001)$ GaAs, and two with (011) $\alpha$Rh$_2$As $\| (001)$ GaAs. Most of the $\alpha$Rh$_2$As grains belonged to the latter type of orientation. They suggested that the rotated relationship was favored by the alignment of the dense-packed atomic rows in the two crystal structures.

Guivarc'h, et al. also have deposited RhGa on GaAs [GU88a]. In this case, the lattice mismatch between the two phases is quite large ($a_o$ of 0.301nm for RhGa vs. 0.565nm for GaAs), although the RhGa is much more stable in contact with GaAs than is $\alpha$Rh$_2$As. By codepositing Rh and Ga by MBE, the authors were successful in growing a highly textured film, with two orientations of the type $[100](011)_{\text{RhGa}}$ $\| [110](001)_{\text{GaAs}}$ present. Upon annealing at 550°C, however, the deposition recrystallized, resulting in a polycrystalline layer with only one orientation.

The deposition of single-crystal rare earth arsenides on GaAs has been reported (YbAs [RI88], ErAs [PA88b]). This materials have high melting points, cubic crystal structures, and acceptable electrical properties. In both cases, the rare earth and As were codeposited in MBE. Both experiments were successful in forming single crystals of the arsenide on the substrate. Palmstrom, et al. also attempted to grow a subsequent GaAs layer, but were unsuccessful due to the formation of islands. The same authors were also successful in growing high quality CoGa layers on GaAs [PA89].

Buried metallic layer heterostructures have been produced on compound substrates [HB88, TA88]. The authors utilized the techniques of epitaxial deposition and subsequent semiconductor growth by MBE. The compound they selected for their study was NiAl (grown on AlAs), which has a high melting temperature, low resistivity, and is stable in contact with the semiconductor.
3.3 Solid Phase Epitaxy

The concept of formation of an epitaxial phase utilizing only solid-state diffusion has been verified. The simplest case, Solid Phase Epitaxy (SPE), involves the diffusion of appropriate element through a solid layer, and epitaxial precipitation at the transport layer/semiconductor interface. A more difficult process requires not only diffusion, but reaction in the solid state, Solid Phase Reaction Epitaxy (SPRE).

Elemental substrates

Growth of epitaxial layers on semiconductors by solid phase epitaxy (SPE) has been demonstrated by several authors ([SA73], [CA75], [LA77]). Early examples of this technique involve the dissolution of the semiconductor substrate in some suitable material, and precipitation of an epitaxial layer upon cooling, e.g. Al-Ge [CY72] or Al-Si [SA73]. The epitaxial layer incorporates some of the solute material, which can be chosen so as to dope the substrate. This technique suffers from a lack of uniformity, because of the presence of native oxide layers, which inhibit dissolution and regrowth, and result in the formation of island structures [LA78]. The problem is especially severe with Si substrates. In addition, the semiconductor has a tendency to precipitate within the metal layer as polycrystallites when a satisfactory semiconductor interface for growth is unavailable [OT74].

The dissolution/regrowth mechanism can be extended to more complex structures. In these systems, an amorphous semiconductor layer is deposited on top of a transport layer (e.g. Al [MA72], a germanide [MJ77], or a silicide [LA77]), and then annealed. Upon annealing, the amorphous layer decomposes and recrystallizes on the crystalline semiconductor surface. The driving force for the reaction has been attributed to the reduction in free energy associated with the change in state from amorphous to crystalline [CA74].
There are two mechanisms for the transport of the amorphous material. In the case of an elemental transport layer (a-Ge/Al/c-Ge [MR72]), the mechanism is certainly diffusion of Ge through the intermediate layer. Systems with compound transport layers (e.g. a-Si/silicide/c-Si [LA77]) are more complicated, with both mechanisms possible. In the dissociation mechanism [PR76], the silicide at the crystalline semiconductor interface decomposes, with the silicon atoms growing epitaxially on the substrate. The metal atoms then diffuse through the silicide layer to the amorphous material, and react to form more silicide. Alternately, the amorphous Si can diffuse directly through polycrystalline silicide to grow epitaxially on the substrate [LA77].

The silicide layer plays an important role in the solid phase epitaxy. In order to produce this layer, the appropriate metal (e.g. Pd) is deposited on a cleaned Si substrate, commonly by electron-beam evaporation. The amorphous Si is then deposited on the Pd. In all of the SPE systems, the ratio of amorphous material to metal must be greater than one. The silicide (Pd$_2$Si) is formed by annealing the specimen (e.g. 280 °C for 20 minutes for ~50nm Pd). Studies show that the Pd consumes equal amounts of amorphous and crystalline Si in order to form the silicide ([CA75],[PR76]). By consuming both amorphous and crystalline material, the silicide formation ensures a high quality interface for the nucleation and growth of the epitaxial layer [CA75]. SPE with elemental transport layers can be hampered by the presence of native oxide layers [LA78], especially on Si. Therefore, the silicide (or compound forming) layer provides two functions. One is to provide a dissolution and transport medium for the amorphous semiconductor. The other is to provide a clean interface suitable for epitaxial growth by consumption of the original surface [MJ77].

This technique can be further refined to dope the regrown semiconductor [LA76]. Lau, et al., were able to dope the SPE Si n-type by introducing a thin Sb layer in the system: amorphous-Si/Sb/Pd/crystalline-Si (970nm/.5nm/97nm) and annealing. Upon annealing, the Sb atoms were incorporated into the epitaxial Si layer. Other studies show that during SPE, Pd is incorporated into the epitaxial layer, and Si is trapped in
the silicide [CA75].

**Compound Substrates**

The techniques of SPE also have been successfully applied to producing workable contacts on compound substrates by Marshall et al. [MA85]. They decided to introduce an epitaxial Ge layer into the contact, because of its low conduction band discontinuity with GaAs and its low barrier height with many metals. The goal was to produce ohmic contacts without the interfacial roughness of alloyed contacts [KU83], and without the stringent vacuum requirements of MBE. They developed a technique very similar to the silicide SPE described above, which had been applied to the Pd-Ge system by Majni, et al. [MJ77].

In this system, PdGe is used as the transport layer. One of the goals was to reduce the UHV requirements of MBE ($10^{-10}$Torr), so all depositions were done by electron-beam evaporation in high vacuum ($10^{-6}$Torr). After deposition of $\sim$50nm Pd, amorphous Ge was deposited without breaking vacuum. Sufficient Ge was deposited to completely react the Pd layer, with the remainder available to form the epitaxial Ge contact layer. During deposition of the Pd, the GaAs surface was prepared for epitaxial growth by a reaction which formed $\text{Pd}_x\text{Ga}_y\text{As}$. After the annealing steps, the ternary phase was no longer present.

During the first annealing stage ($325 ^\circ\text{C}, 30$ minutes), the Pd reacted with the Ge to form PdGe. After the second annealing stage, the residual amorphous Ge layer dissolved into the germanide, and diffused to the substrate, where it precipitated epitaxially at the PdGe/GaAs interface. The lattice mismatch between GaAs and Ge is $\sim$0.1%. Electrical measurements showed that the contacts were ohmic, with contact resistivities on the order of $10^{-6}$Ωcm$^2$. The authors concluded that the electrical behavior of the contacts was most likely due to tunneling. The contact characteristics degraded after annealing above $\sim$400 $^\circ\text{C}$, however.
The Ge/Pd contact has been applied to In\textsubscript{55}Ga\textsubscript{47}As with some success [CH86]. Contact resistances of $2.3 \times 10^{-6} \Omega \text{cm}^2$ to p-type material were measured, and $6 \times 10^{-7} \Omega \text{cm}^2$ to n-type. The contacts were also attractive from a processing standpoint (325 °C anneal) and thermal stability (4 hours at 300 to 500 °C).

The Au-Ni-Ge contact described in section 2.3 actually utilizes the dissolution and regrowth techniques. During the annealing, the GaAs substrate is partially dissolved in the contact material, and then regrows epitaxially on the substrate, incorporating some of the contact elements into the new material. Unfortunately, the interface is very rough and non-uniform, perhaps due to the presence of native oxides [SA88].

3.4 Solid Phase Reaction Epitaxy

Elemental Substrates

During Solid Phase Reaction Epitaxy, the products of a thin film reaction grow epitaxially on a substrate material. The growth of epitaxial silicides is an example of the simplest case of this technique. Several metals can be reacted with Si wafers to form epitaxial reaction products, with varying degrees of crystallinity (see [TN82b] for review). Epitaxial silicides include CoSi\textsubscript{2}, NiSi\textsubscript{2}, PdSi\textsubscript{2}, PtSi, CaSi\textsubscript{2} [MO88], ErSi\textsubscript{2} [AV89], and numerous refractory metal silicides [CL85].

The simplest method for producing epitaxial silicides, with the lowest vacuum requirements, involves the deposition of the metal followed by an annealing treatment to induce silicide formation. The silicide grows on the silicon substrate as an epitaxial layer. The annealing usually consists of a two stage heat treatment: a low temperature anneal (300-400 °C) to form the polycrystalline silicide, followed by a high temperature anneal (800-900 °C) to induce the preferential growth of epitaxial grains (e.g. [CJ82a]). In some cases, the epitaxial silicide can be formed in one stage.
When epitaxial silicides are formed by reaction, the overlayer is often composed of two types of grains, each with a specific orientation relationship with the Si substrate, and related to each other by some symmetry operation [CK81, TN82b]. Typically, the films have been deposited on a chemically cleaned substrate by electron beam evaporation, although some groups have deposited the pure metal by MBE. The electron-beam evaporated films are of lower quality; during annealing of the chemically cleaned substrates, nucleation of epitaxial material can only occur through holes in the contamination (oxide) layer [GI82].

Epitaxial silicides have been grown on all low-index Si planes ((100), (110), (111)). High quality, monocrystalline layers have been produced by these techniques [TN82a, AV89]. On a local scale, the silicide-substrate interface is abrupt to within 2nm, but only on (111)Si is the interface flat [CK81, TN82b]. This is due to the greater stability of the (111) interface; other orientations result in faceting of these planes, with protrusions on the order of tens of nanometers [CJ82b].

**Compound Substrates**

Solid phase reactive epitaxy on compound substrates introduces an additional problem not encountered with elemental semiconductors. In addition to the issues of lattice mismatch and interfacial compatibility, there is the possibility of the formation of multiple reaction products, one of which could impede epitaxial growth. Binary semiconductors can react with elemental metal layers to form ternary (e.g. M_{x}AB) or binary (MA_{x}, MB_{y}) phases.

The formation of a single ternary compound is not unlike the silicide case; the metal diffuses into the substrate, and reacts to form an epitaxial ternary phase. Lahav, et al., have reported that the Ni/GaAs system can undergo this behavior. In their experiment, Ni was deposited by electron-beam evaporation on chemically cleaned (111) and (100) GaAs. The two substrate orientations yielded different results. They reported that Ni_{2}GaAs phase formed epitaxially on both substrates for anneals
between 100 and 300 °C, although the product on (100) was microtwinned while the product on (111) was of higher crystalline quality. Above 350 °C, the ternary phase decomposed on the (100) substrates to epitaxial \( \beta \)NiGa and randomly oriented NiAs. Later experiments by Sands, et al. [SA86a], indicated that the ternary phase was actually Ni\(_3\)GaAs.

Similar studies have been done on the Pd/GaAs system. Oelhafen, et al. found that at lower annealing temperatures, a highly textured Pd:Ga:As ternary phase (which they identified as Pd\(_2\)GaAs) formed, while at higher temperatures (above 350 °C), epitaxial islands of PdGa were reported. Later studies by Sands, et al. [SA86b] concluded that the actual ternary composition was Pd\(_5\)(GaAs)\(_2\).

The Co/GaAs illustrates two possible interfacial epitaxial reactions, depending on annealing temperature [PA87]. At lower temperatures, a ternary Co\(_2\)Ga\(_{2-y}\)As\(_y\) (with \( y \approx 1 \)) appears at the reacted layer/GaAs interface, and exhibits a strong orientation relationship with the substrate. At higher temperatures, the ternary phase decomposes into two binaries, CoGa and CoAs, each aligned with the substrate [PA87, SA87b].

Alternatively, the product phases may segregate so that one binary phase grows with an orientation relationship with the substrate, while another grows on the surface. This has been shown in the case of Pt/GaAs, where PtAs\(_2\) forms with a strong orientation relationship on the GaAs, while Pt-Ga phases form on the surface [SI73, KM75, FO83]. To a lesser extent, the same situation develops in the Pt/InP system, where PtP\(_2\) exhibits texturing with the InP substrate, although some intermixing of the PtP\(_2\) and Pt-In phases is evident [OL89].

In summary, these metal/A\(_{\text{III}}\)-B\(_{\text{V}}\) systems exhibit three different cases for the formation of epitaxial reaction products: a ternary textured phase (Ni/GaAs, Pd/GaAs, Co/GaAs), two binary textured phases (CoGa and CoAs on GaAs), and a single textured binary phase with vertical segregation (PtAs\(_2\) and PtP\(_2\)). These systems provide the potential for the formation of epitaxial contacts by SPRE.
3.5 Phase Separation

In the Pt/In$_x$Ga$_{1-x}$As system, multiple reaction products are expected. Even if one of these phases has the inclination to form epitaxially on the semiconductor substrate, it may be impeded by the presence of other phases from forming a continuous layer. Therefore, it may be desirable to induce the segregation of the reaction products so that continuous layers can be formed. This occurs naturally in the Pt/GaAs system, although it is still unclear why the layering sequence occurs. In the next chapter, models for the occurrence of phase separation will be considered, as well as techniques for exploiting them to achieve desirable structures.
4.0 METALLURGY

This study has been motivated by the desire to develop stable contact structures on compound semiconductors. Consequently, an understanding of the possible metallurgical reactions is of profound importance. As has been mentioned previously, elemental metals in contact with compound semiconductors will generally react with the substrate, modifying the electrical and morphological characteristics of the system. The reaction products can be beneficial; therefore, inducing a desirable phase to develop at the proper interface with the appropriate morphology is very important. Some issues to consider when examining the metallurgy will be discussed in this chapter: reaction products, phase segregation, diffusion, kinetics, and diffusion barriers.

4.1 Product Phases in Pt/In$_x$Ga$_{1-x}$As

Six binary phase diagrams for the Pt:In:Ga:As quarternary system are shown in figure 4.1.a-f [MF87, HN58]. The Pt-In and Pt-Ga phase diagrams show that there are several intermetallic phases possible in these systems. However, previous studies of Pt/III-V reactions [SI73, KM75, SI78, FO83, OL89] indicate the likely product phases, which are discussed below.

The Dipnictides

In the Pt-As system, there is only one viable product phase for these reactions, PtAs$_2$. This belongs to a class of compounds, known as Pt-dipnictides, and which have the form MB$_2^V$, where B = P, As, Sb [FU65, HL65, FU67, BB74]. These compounds have the cubic FeS$_2$ (pyrite) structure, with lattice parameters within the range of many ternary substrates, which makes them nearly ideal for the purposes of this study [VI85]. In addition, these three compounds contain the group V elements commonly used in $A_a^{III}$-$A_b^{III}$-$B^V$ ternary substrates- P, As, and Sb.
Figure 4.1.a: Phase diagram of the As-Pt system, after Moffet, 1987 [MF87].
Figure 4.1.b: Phase diagram of the Ga-Pt system, after Moffet, 1987 [MF87].
Figure 4.1.c: Phase diagram of the In-Pt system, after Moffet, 1987 [MF87].
Figure 4.1.d: Phase diagram of the Ga-As system, after Hansen, 1958, p. 165 [HN58].
Figure 4.1.e: Phase diagram of the In-As system, after Hansen, 1958, p. 166 [HN58].
Figure 4.1.f: Phase diagram of the Ga-In system, after Hansen, 1958, p. 745 [HN58].
The substrate which is chosen for this study is In\(_x\)Ga\(_{1-x}\)As. This material has two attributes which make it an attractive material for device applications. First, its high electron mobility allows it to be used for the design of high speed electronic devices. Second, its bandgap energy encompasses the useful 1\(\mu\) to 2\(\mu\) range, which is desirable for low-loss optical waveguides.

The room temperature lattice parameter for In\(_x\)Ga\(_{1-x}\)As varies linearly between 0.5653\(\text{nm}\) (GaAs) and 0.6058\(\text{nm}\) (InAs). The lattice parameter for PtAs\(_2\) is 0.5967\(\text{nm}\), which corresponds to In\(_{78}\)Ga\(_{22}\)As. Unfortunately, growth of this composition is impossible without a heavily defected structure, because of the large lattice mismatch. In\(_x\)Ga\(_{1-x}\)As can be grown epitaxially on InP at a composition of In\(_{53}\)Ga\(_{47}\)As, \((a_0 = .587\text{nm}, E_g = 0.75\text{eV})\). At this composition, the lattice mismatch with PtAs\(_2\) is 1.7\%, which is low enough for epitaxial growth.

**Intermetallics**

Other compounds that are likely to form during Pt/In\(_x\)Ga\(_{1-x}\)As reactions are those that are encountered in Pt/GaAs [KM75, CO74, CC75, SI78, FO83, SA87a], Pt/GaP [SI78], and Pt/InP [BS83, OL89] reactions: PtGa and PtIn\(_2\). Both are cubic phases, with lattice parameters of .491\(\text{nm}\) and .637\(\text{nm}\), respectively [VI85]. The crystal structure of PtGa is the FeSi structure, while PtIn\(_2\) forms in the fluorite structure, which consists of a Pt FCC lattice with indium atoms filling all of the tetrahedral sites. Both lattice parameters are well outside the range of the In\(_x\)Ga\(_{1-x}\)As substrate.

Ternary phases may be encountered between the expected reaction products. In the case of PtGa, the corresponding In phase is PtIn. This phase is monoclinic, which makes extensive solid solutions between these phases unlikely. In the case of PtIn\(_2\), the corresponding Ga-based phase, PtGa\(_2\), has the identical fluorite crystal structure, with a lattice parameter of .592\(\text{nm}\), so some solubility of the two phases is expected. Introduction of Ga in PtIn\(_2\), for example, should reduce the lattice mismatch with
Morphology

During Pt/GaAs reactions, PtAs$_2$ forms with a strong orientation relationship on the GaAs substrate [SI73, KM75, FO83], although the cube axes of the two phases are not parallel. This also occurs in the Pt/InP system, where PtP$_2$ exhibits texturing with the InP substrate, [OL89]. In both Pt/GaAs and Pt/InP, the lattice mismatch between the substrate and Pt-B$^V$ phase is rather high (5.4% and 2.9%, respectively), diminishing the likelihood of epitaxial growth.

The symmetry of the dipnictides is lower than that of A$^{III}$B$^V$ substrates. As a consequence, four different variants are expected when (100) PtAs$_2$ is grown on (100) In$_x$Ga$_{1-x}$As surfaces. The crystal structure of the Pt dipnictide phases consists of an FCC Pt sublattice, with all the octahedral sites filled with pairs of B$^V$ atoms. While the FCC Pt sublattice is invariant under the symmetry operations of the substrate, the B$^V$ sublattices are not. The Pt atoms will form a continuous lattice across the domain boundaries when the dipnictide phase forms with a specific orientation relationship with the substrate (e.g. its cube axes parallel to the substrate cube axes). Thus, different domains arise from the four possible variations of B$^V$ orientations within this sublattice. The domain boundaries are likely to consist of B$^V$ point defects.

4.2 Phase Separation

Phase separation during metal/semiconductor interactions is critical for producing a desirable interface structure. The various product phases will likely have different electrical characteristics; if more than one are present at the contact interface, non-uniformity of the electrical properties will result. The product phases may have different crystal structures; if one has a tendency to form epitaxially on the substrate, another phase may intrude and disrupt the crystallinity. By understanding the reasons
for layering, better control of interfacial uniformity is possible. Vertical segregation of phases can be affected by a number of factors, including variations in solubility limits, deposition thickness, and variations in diffusivities.

**Solubility**

The existence of multiple elements in the compound semiconductor substrate provides one possibility for vertical phase separation. If the substrate elements have different solubility limits in the metal overlayer, then an excess of the lower solubility element could develop at the substrate/reacted layer interface. In the case of Pt/GaAs, Kumar [KM75] has shown that the Ga saturates the Pt, leaving an excess of As to react at the interface. Both of the group III elements in this study have substantially greater solubility in Pt than As (over the temperature range 400-800 °C): Ga ~6 atomic %, In ~3 atomic %, As negligible (figure 4.1.a, b and c, [MF87]). Consequently, more Ga and In than As can dissolve in the Pt, leaving an excess of As at the interface to react and form PtAs₂, as in the case of GaAs substrates.

**Thickness**

The thickness of the initial metal deposition can have a profound effect on the reacted morphology [SA87b, YU87a]. If the grain size of the reaction products is greater than the layer thickness, then the vertical segregation cannot occur. In this case, the reaction products would be laterally separated, i.e. intermingled. Yu, et al. demonstrated this behavior in the Rh/GaAs system by annealing thin (15nm) and thick (60nm) films [YU87a]. They found lateral segregation of the RhGa, RhAs, and RhAs₂ reaction products in the thin film, and vertical segregation (RhGa/RhAs₂/GaAs) in the thick film.
Native Oxides

Native oxide layers are present on the surface of all chemically-cleaned semiconductors [W185]. In general, these are detrimental to interfacial reactions, but some metals (e.g. Pt and Pd [SA87a]) have been shown to be effective in penetrating this layer. It is unclear what effect the presence of a native oxide layer will have on the phase segregation. There is evidence that the oxide layer acts as a diffusion filter; Auger studies of Pt/GaAs indicate that residual oxide remains at the boundaries between the segregated phases in the reacted microstructure [CC75, SA87a]. If this is indeed the case, the presence of a native oxide layer may aid phase segregation during the Pt/In$_x$Ga$_{1-x}$As reaction.

Sublimation

One technique that could eliminate some of the reaction products is anneal uncapped specimens. This allows volatile As or As compounds to sublime, resulting in the formation of only metal-III phases. Selective desorption can be useful if the As compounds in the system give rise to unwanted microstructures, or have undesirable electrical properties. Studies of uncapped Co/GaAs [PA87] and Pd/GaAs [SA87b] have demonstrated the viability of this technique. Decomposition of the substrate is undesirable, however, so it must be protected by encapsulation of the backside of the wafer, or by keeping the annealing temperature low.

Diffusivity

Probably the most important reason for phase segregation in these systems is variation in the diffusion coefficients, or diffusivities, of the elements involved in the reactions [e.g. SA87b, PA87]. The elements with the higher diffusivities will diffuse more rapidly down the chemical potential gradient, leaving an excess of the slower diffusing elements behind. This will give rise to local variations in the concentration, which will lead to variations in the phase formation. The diffusion of species in thin
films is very important to this study, and is the subject of the next section.

4.3 Diffusion [SW89, PT81, SM81]

Diffusion is the fundamental process by which a system transports material to lower its free energy. If the chemical potential of an element varies within a sample (as might occur for variations in concentration), the system is not in equilibrium. Natural processes will occur to diminish the chemical potential gradient to zero, and bring the system into a stable, low-energy configuration.

The natural process is known as diffusion. Atomistically, diffusion can be modelled by considering the local oscillations of an atom around its lattice site. When the amplitude of the vibration is sufficient, the atom can jump into an adjacent vacant site. The jump frequency, \( \Gamma \), can be described by:

\[
\Gamma = z \omega \exp \frac{-\Delta G_m}{kT}
\]

where \( z \) is the number of available vacant sites, \( \omega \) is the vibrational frequency of the atom, and \( \Delta G_m \) is the activation energy for the migration of the atom. From equation 4.1, it is evident that an increase in temperature will cause an increase in the jump frequency.

Diffusion occurs when these jumps give rise to a net flux of atoms. For the case of a one-dimensional system, the magnitude of the flux of B atoms in a medium \( \alpha \) is given by Fick's first law:

\[
J_B = -D_B \frac{\partial C_B}{\partial x}
\]

The quantity \( D \) is defined as the diffusivity, and is given by \( \Gamma \) multiplied by several factors:

\[
D = D_0 \exp \frac{-\Delta H_m}{kT}
\]
The entropy component of the activation energy, $\Delta S_m$, is brought out as a pre-exponential factor which does not depend on temperature:

$$D_0 = \lambda z \omega \exp \frac{\Delta S_m}{k}$$  \hspace{1cm} (4.4)

$\lambda$ represents geometric factors which are determined by the lattice and the diffusion mode. The diffusivity is an important measure of how easily an atom can diffuse through the matrix. Note that this quantity decreases as $\Delta H_m$ increases, thereby reducing the flux. However, $D$ increases exponentially with temperature, at a rate which is proportional to $\Delta H_m$.

The mechanism by which diffusion occurs can be divided into three regimes: bulk, grain boundary, and dislocation. In the bulk, the diffusing species moves through the crystal lattice primarily by either exchanging with vacancies, or via interstitial sites. The activation energy is rather high for this diffusion mode, hence bulk diffusion predominates at higher temperatures, where the activation energy is available. The looser atomic configuration of grain boundaries results in a lower activation energy if atoms diffuse along this path. At intermediate temperatures (below $\sim 0.75$ of the melting temperature of the medium), grain boundary diffusion can dominate. At lower temperatures (below $\sim 0.5 T_M$), diffusion along dislocation pipes can become the dominant mode.

The density of high-diffusivity paths (grain boundaries and dislocation pipes) must be considered when comparing the predominance of these three modes. Even if the activation energy for diffusion is very low, there must be a sufficient number of diffusion paths available for the mode to be significant. The total area through which diffusion can occur must be taken into account when evaluating the amount of diffusion that occurs by a given mode. For a polycrystalline sample, the cross-sectional area available in a grain for lattice diffusion is $d$, the grain size, while the cross-sectional area available for boundary diffusion is $\delta$, the width of the grain boundary ($\sim 0.5 \text{nm}$). In order to compare the relative contributions by these two mechanisms,
the diffusion coefficients must be scaled by the area available for diffusion:

\[ D_T = D_l + \frac{\delta}{d} D_{g,b}. \]  

(4.5)

As the temperature decreases the lattice diffusivity, \( D_l \), decreases more rapidly than \( D_{g,b} \); therefore, diffusion via grain boundaries will make a more significant contribution to the total flux. Similar analysis results in an expression for diffusion along dislocations, with a cross-sectional area per unit bulk area of \( g \):

\[ D_T = D_l + gD_d. \]  

(4.6)

It should be noted that all three types of diffusion will be occurring at all temperatures, but with varying prevalence, depending on their activation enthalpy, \( \Delta H_m \), and the density of the appropriate diffusion paths. Estimates of relative contributions for lattice and grain boundary diffusion for possible product phases in the Pt/In\(_x\)Ga\(_{1-x}\)As system are provided in Table I. These calculations are based on approximations for lattice vs. grain boundary diffusivities by Gupta [GP85] for pure metals.

As the complexity of a system increases, multiple species and mechanisms can contribute to the total flux. The concept of an effective diffusion constant, \( D_{eff} \), which reflects a combination of all the active diffusivities must be introduced. The active modes may include diffusion through different media, or of different species, or by different mechanisms.
Table I

Estimates of the relative contribution of lattice and grain boundary diffusivities at 400 °C. Calculated using equation 4.3, and the approximations for diffusion parameters by Gupta [GP85] for pure metals: \( \Delta H_m^{\text{l}} \) (cal/mole) = 34\( T_m \)(K), \( \Delta H_m^{\text{g.b}} \) = 25\( T_m \), \( D_0^{\text{l}} \) = 0.5(\( \text{cm}^2\text{s}^{-1} \)), \( D_{g.b}^{\text{l}} \) = 3.1, \( \delta = 0.5\text{nm} \), and \( d = 50\text{nm} \). \( R \) refers to the ratio:

\[
R = \frac{D_l}{\delta D_{g.b}}. \quad (I.1)
\]

<table>
<thead>
<tr>
<th>Phase</th>
<th>( T_m )(K)</th>
<th>( D_l )(( \text{cm}^2\text{s}^{-1} ))</th>
<th>( D_{g.b} )(( \text{cm}^2\text{s}^{-1} ))</th>
<th>( R )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt</td>
<td>2042</td>
<td>2\times10^{-23}</td>
<td>9\times10^{-17}</td>
<td>2\times10^{-5}</td>
</tr>
<tr>
<td>PtAs(_2)</td>
<td>1795*</td>
<td>8\times10^{-21}</td>
<td>9\times10^{-15}</td>
<td>9\times10^{-5}</td>
</tr>
<tr>
<td>PtIn(_2)</td>
<td>1312</td>
<td>2\times10^{-15}</td>
<td>7\times10^{-11}</td>
<td>2\times10^{-3}</td>
</tr>
<tr>
<td>PtGa(_2)</td>
<td>1205</td>
<td>3\times10^{-14}</td>
<td>5\times10^{-10}</td>
<td>5\times10^{-3}</td>
</tr>
<tr>
<td>PtGa</td>
<td>1377</td>
<td>3\times10^{-16}</td>
<td>2\times10^{-11}</td>
<td>2\times10^{-5}</td>
</tr>
</tbody>
</table>

* estimated.
The Diffusion Couple

The simplest case for the development of layering during reaction is that of the diffusion couple. In this experiment, samples of two different materials, A and B, are welded together, and allowed to interdiffuse. After a suitable amount of time, interdiffusion will give rise to a concentration gradient in the system as shown in Figure 4.2. If intermediate A:B compounds are possible in the system, they will develop in the appropriate concentration range. A layering sequence, which is a consequence of the starting geometry of the system, will result.

In a multiphase system, the reaction interface can be defined as the interface where the A and B elements meet and react to form intermediate compounds. The position of the interface is determined by the diffusivities of A and B atoms in the appropriate medium. If \( D_B^2 < D_A^2 \), for example, then the reaction will occur at the B-rich interface, and vice versa. Reactions can occur simultaneously at more than one interface.

Segregation in Pt/In\(_x\)Ga\(_{1-x}\)As

Pt/GaAs and Pt/InP have very similar reaction behavior. Both react to form layered structures, with a Pt-B\(_2^V\) phase at the substrate/reacted layer interface, and a Pt-A\(_{III}\) phase at the surface. In the Pt/GaAs case, this structure is evident even at low annealing temperatures, while the Pt, In, and P are completely intermixed for Pt/InP until higher temperature anneals. The differences between the Pt/GaAs and Pt/InP cases may be attributed to the different diffusivities of the elements involved. In the case of Pt/GaAs, the Ga diffuses out from the substrate at fairly low temperature [SI73, CC75], with the remaining As reacting with the Pt overlayer to form PtAs\(_2\). In the case of Pt/InP, the diffusion of In into the overlayer does not dominate, allowing the formation of intermixed phases until high temperature anneals [OL89].
Figure 4.2: Sequence of phase formation that results from a diffusion couple (after reference [PT81], page 104).
In the case of Pt/In$_x$Ga$_{1-x}$As, the relative diffusivities of the components becomes very important because of the number of elements involved. If a continuous PtAs$_2$ layer is to be formed at the substrate/reacted layer interface, both Ga and In must diffuse into the overlayer. Based on previous studies [SI73, OL89], the ratio of the diffusivities of Ga to Pt should be greater than the ratio of In to Pt. This indicates that the interface should be free of Pt-Ga phases, but may contain Pt-In phases. As the In concentration of the substrate increases, the lattice compatibility of PtAs$_2$ and In$_x$Ga$_{1-x}$As will improve, but the potential for intrusion of Pt-In phases will increase. There are several experimental parameters which should contribute to the segregation of the Pt-In phases, which rely on selecting reaction conditions that enhance Pt, Ga, and In diffusion, and inhibit As diffusion.

The first parameter which may induce segregation by unequal diffusivities is the annealing temperature. It may be possible to select a high enough temperature which will allow for significant Ga, In, and Pt diffusion, but minimal As outdiffusion. There is experimental evidence which supports this approach. Even when Pt/GaAs thin films are annealed over a range of temperatures (300-600°C), the layered Pt-Ga/PtAs$_2$/GaAs structure is maintained [KM75, SA87b]. This suggests that the impetus for PtAs$_2$ segregation at the interface is somewhat insensitive to temperature. In the Pt/InP system, phase segregation has been observed as low as 500°C, and becomes much more pronounced as the annealing temperature is increased to 600°C [OL89]. By annealing Pt/In$_x$Ga$_{1-x}$As at higher temperatures, it is possible that suitable phase segregation will be obtained.

Diffusion rates are affected by other factors than temperature. The mechanism (e.g. grain boundary vs. bulk diffusion) will also determine the rate of diffusion. The dominant diffusion mechanism may vary for the four elements involved. For example, changing the ratio $\delta/d$ by changing the average grain size of the polycrystalline Pt layer may alter the relative diffusion rates of the components. The grain size can be changed by varying the substrate temperature during deposition (e.g. [CP69]).
disadvantage of relying on grain boundary diffusion is that the interface will not be smooth; the reaction will proceed preferentially along the grain boundaries.

4.4 Kinetics

The kinetics of the thin film reaction is important, as the contact structure will always be thermodynamically unfavorable. This is because of the concentration gradients necessary to obtain the desired electrical properties. Consequently, a "stable" contact is really one in which the interdiffusion and interactions are much slower than the processing time and projected lifetime of the device. By slowing the reaction kinetics, the lifetime of the device can be extended.

According to the analysis done by Kidson [KD61], the growth of a $\beta$ product layer of thickness $x$ from A and B phases can be described by the following equation:

$$\frac{dx}{dt} = \frac{G\Delta C\kappa}{(1 + x\kappa/D_{\text{eff}})}$$

(4.7)

where $G$ is a constant dependent on the composition of the reactant and product phases, $\Delta C$ is the difference in composition between the A$\beta$ and $\beta$B interfaces, and $\kappa$ is the effective interfacial reaction barrier, which depends on the individual reaction constants at the two interfaces.

Equation 4.7 can be divided into two regimes by the parameter $x^* = D_{\text{eff}}/\kappa$:

$$dx/dt \sim G\Delta C\kappa, \text{ for } x << x^*$$

(4.8a)

and

$$dx/dt \sim G\Delta C\kappa/x, \text{ for } x >> x^*$$

(4.8b)

These equations can be integrated to give the dependence of the layer thickness on time:

$$x \sim t$$

(4.9a)

and
This result shows that for sufficiently small values of $x$, any system will be dominated by interface reactions, until a changeover thickness of $x^*$ (which depends on the ratio between the diffusivity and reaction rate) is reached, after which diffusion effects will dominate.

### 4.4.1 Interface Controlled

The significance of the two regimes in equations 4.9a and b can be explained as follows. When a reaction involves the growth of a phase which the reactant must diffuse through, there are two components of "resistance" that the moving species will experience. When atoms arrive at a phase boundary, they experience a resistance to movement across the interface. When diffusing through the growing medium, they also experience resistance. When the resistance to crossing the boundary dominates (as in the early stages of a reaction), the kinetics are interface controlled, and linear growth may be observed, i.e. $x = Ct$, where $x$ is the reaction variable (i.e. some measure of the progress of the reaction, such as $x = 1 - h$, where $h$ is the thickness of unreacted material).

### 4.4.2 Diffusion Controlled

When the diffusion resistance dominates, the kinetics of the metal/semiconductor reaction are parabolic, i.e. $x^2 \sim D_{eff}t$. This is the behavior exhibited by many metal/semiconductor reactions (e.g. [OT86, YU87b], including Pt/GaAs [CO74]). Plots of $x^2$ vs. $t$ will appear as straight lines under these conditions, with the slope $\sim D_{eff}$.

There are complications of this simple model which must be considered, such as the development of intermediate layers, or change in the diffusion path during the reaction.
Intermediate Layers

Sequential growth of intermediate layers could give rise to deviations from simple parabolic growth, even when the kinetics are diffusion controlled. If the reactant A is initially diffusing through the phase \( \alpha \), the diffusivity is \( D_A^\alpha \). If an intermediate layer of phase \( \beta \) develops and becomes continuous, then the diffusion should be described by \( D_A^\beta \) as well. If \( D_A^\alpha \ll D_A^\beta \), then a plot of the reaction rate would exhibit two different slopes in the two regimes. If the two diffusivities are similar, the change in slope of \( x^2 \) vs. \( t \) will be less distinct, and \( D_{\text{eff}} \) may involve a combination of \( D_A^\alpha \) and \( D_A^\beta \).
Figure 4.3: Schematic of the effect of the formation of a new phase on the diffusivity in a system. A reacts with C to form the γ phase, leaving B behind which eventually forms a continuous layer. Diffusion through the γ phase dominates the kinetics in the initial stage of the reaction. If $D_A^B < D_A^\gamma$, diffusion through the B layer will dominate at later stages.
Grain Growth

The annealing temperature of these experiments is kept at a level where bulk diffusion is unlikely to be the dominant mode (see Table I). The low temperature is necessary to avoid decomposition of the compound semiconductor substrates. Consequently, grain boundary diffusion is likely to be significant, which requires that grain growth be considered. If the dominant mode of diffusion is through grain boundaries, and the grains are growing during diffusion, the density of grain boundaries and hence the ratio $\delta/d$ will decrease. This will cause the diffusion rate to fall below $x^2 \sim Dt$.

Studies on the growth of superconducting A-15 compounds have shown a less than parabolic growth rate [FH74]. In the case of Nb$_3$Sn, the compound is grown by solid-phase reaction of pure Nb with Cu-Sn alloys. The process relies on grain boundary diffusion through the polycrystalline Nb$_3$Sn product layer. Farrell, et al., suggested that the deviation from parabolic growth could be attributed to coarsening of this product layer. As the Nb$_3$Sn grains grow, the density of diffusion paths diminishes. In a subsequent paper, Farrell, et al. [FH75], derived an expression for the reaction kinetics, which takes into account the competition between grain growth and layer growth. Assuming the following time dependence of the grain size, $L$, on time:

$$\frac{L^2}{L} = \lambda t^m$$  \hspace{1cm} (4.10)

where

$$\lambda = \lambda_0 \exp(-Q_L/kT)$$  \hspace{1cm} (4.11)

Farrell, et al. derived the following to describe the layer thickness as a function of time:

$$x \sim \left( \frac{D_{eff}}{\lambda} \right)^{0.5} t^{(1-m)/2}$$  \hspace{1cm} (4.12)

That is, the growth rate of the layer of $x \sim t^k$ can be reduced by an amount corresponding to the rate of grain growth. Evidence of this type of time dependence has been found in metal/semiconductor systems, for titanium silicide growth [CR91].
4.5 Diffusion Barriers

Many contact structures are thermodynamically unstable [NI81]. Different materials are brought into contact, each of which satisfy some requirement of the whole structure. The two materials are not generally inert with respect to each other, and can interact to degrade the contact characteristics. An intermediate layer can be introduced to the system to inhibit interactions, but because of the inherent instability of the structure, a diffusion barrier can only slow down undesirable interactions. Diffusion barriers can be introduced into the system to inhibit undesirable interactions and to extend the lifetime of the device.

There are several requirements that a material must satisfy in order to function as a diffusion barrier (see [NI81, NI78] for review): a) atomic transport across the layer should be very low. Because the thickness of contact structures is measured in tenths of microns, defect (rather than bulk) diffusion dominates the material transport. Therefore, control of the extended defects (e.g. grain boundaries) present in the contact layers can greatly reduce the diffusion through the barrier; b) the barrier should be thermodynamically stable with respect to the adjacent compounds, with low mutual solubilities; c) the barrier should adhere well to the adjacent compounds; d) the barrier should have electrical and mechanical properties which do not reduce the behavior of the system below that which necessary for proper operation. Satisfying all of these requirements simultaneously is practically impossible, so any diffusion barrier will have to be a compromise.

All of the phases which are likely to form in the Pt/In$_x$Ga$_{1-x}$As system have absolute melting temperatures above 1000K. In order to be inside the grain boundary dominated diffusion regime, the annealing temperature must be \( < .75 T_M \) [PT81]. The temperatures corresponding to this boundary are shown in Table II. The data indicates that for all of the likely product phases, grain boundary diffusion should dominate up to \( \sim 600 \)°C. Since the reaction products are stable on In$_x$Ga$_{1-x}$As, they already satisfy one of the most important requirements for a diffusion barrier. As grain
boundary diffusion is likely to dominate during fabrication, elimination of the grain boundaries could improve the capability of these phases to act as diffusion barriers.
Table II

Comparison of melting temperatures of product phases in the Pt/In$_x$Ga$_{1-x}$As system, and the temperature below which grain boundary diffusion is expected to dominate the flux.
5.0 EXPERIMENT

5.1 The Contact Metallurgy Experiment

The metallurgical behavior of a given metal/semiconductor couple can be examined with a simple experiment: the metal is deposited onto the substrate, the sample is annealed, and the metal/semiconductor interactions are evaluated. Despite its simplicity, there are some important issues which must be considered when designing the contact metallurgy experiment.

Metal contacts on semiconductors are inherently non-equilibrium structures. Large concentration gradients are necessary for two reasons. First, the properties of the structure must change quickly from a conductor to a semiconductor. Second, the presence of the contact elements in the substrate are generally undesirable. Large interfacial area to volume ratios exist within the microstructure, because of the thickness of the films (hundreds of nanometers) and the fine-grained microstructures (tens of nanometers) often encountered. Coupled with these large thermodynamic driving forces in the system are kinetically favorable reaction conditions; short diffusion distances necessary to saturate the film, and high diffusivity paths in the form of grain boundaries.

The problems associated with the non-equilibrium nature of the contact can be diminished by controlling the thermal history of the device. A thermal budget can be defined as the maximum time at temperature which a device structure can withstand without degradation of its properties below given values. Once the thermal budget of the device is set to some reasonable value, the conditions for a viable contact structure can be engineered.

When determining the conditions for a contact metallurgy experiment, the parameters must reflect the non-equilibrium nature of the contact; the stable or end phases are those which remain unchanged after exposure to typical fabrication conditions (i.e.
the thermal budget). Thermodynamic phase diagrams can be used as a guideline for predicting the stable phases, but the elemental composition of the system is artificially imposed by geometry and kinetics, and will change with time, although ideally not within the projected lifetime of the device.

Another consideration is the condition of the substrate prior to deposition. All air-exposed Al-B substrates will have 1-2nm of native oxide on the surface. The native oxides can be removed, but will quickly regrow during handling, and can be expected to be present on all semiconductor surfaces. This oxide can inhibit interactions of some metals with some semiconductors, but there are several potential contact metals which can penetrate the thin oxide layer. Alternately, the contact metallurgy experiment can be carried out entirely in vacuum, preparing the surface either by cleaving the substrate or by desorbing the native oxide, and depositing the metal without breaking vacuum. The cleaving experiment is particularly suited for studies of the electrical and chemical behavior of the interface, especially for very thin (monolayer) coverages. The experiment in this study was designed for thicker metal layers (tens of nanometers), with metal overlayers that are not impeded by native oxide.

The thickness of the deposition must be held constant when comparing specimens from different annealing conditions; very thick layers will take a long time to react, and may experience intermediate states that are not achieved with thin depositions. From a practical standpoint, the deposition thickness should be thick enough to allow a suitable range of annealing times without completely consuming the pure metal layer, but not so thick that steady state conditions are not achieved. In addition, the analytical techniques (especially plan-view TEM) put an upper limit on the deposition thickness. For this experiment, a thickness of 70nm has been selected.

When dealing with Al-B compound semiconductors, care must be taken to ensure that the system is closed during annealing. The group V element in these materials is quite volatile, and can be expected to evolve as a gas during high temperature annealing. For GaAs, preferential evaporation of the group V element can be
expected above 657 °C, for InP, above 362 °C [OB83]. In order to prevent decomposi-
tion of the substrate during annealing, either directly from the surface or through the
grain boundaries of the deposited metal film, the samples can be encapsulated in SiO₂.

5.2 InₓGa₁₋ₓAs substrates

The InAs-GaAs system was chosen for this experiment, among other things, because the substrate lattice parameter can be varied by varying the composition. Unfortunately, this makes the substrate material difficult to produce in high quality for most values x. Growth of a single crystal must be initiated by either a seed crystal (for Czochralski growth) or a substrate (for deposition techniques) with a lattice parameter very close to that of the desired InₓGa₁₋ₓAs crystal. When the lattice parameter of the seed deviates from that of the growing material, dislocations will be introduced to accommodate the strain. Due to a lack of seed crystals and substrates with suitable lattice parameters, InₓGa₁₋ₓAs is difficult to obtain for most of the composition range.

Recent developments in crystal growth by Dr. William Bonner at Bellcore, Inc. indicate that InₓGa₁₋ₓAs will eventually be produced in bulk crystalline form, but for the purposes of this experiment, InₓGa₁₋ₓAs substrates were prepared by Molecular Beam Epitaxy (MBE) on GaAs and InP substrates. In this procedure, the appropriate substrate material is loaded into a vacuum chamber equipped with sources of As, Ga, and In. The sources are known as effusion cells, and consist of a crucible containing the desired element, which can be heated to achieved a substantial vapor pressure. In the vacuum chamber (10⁻¹⁰ Torr), the vapor produce proceeds on a direct line-of-sight to the substrate, where it adsorbs onto the surface. Prior to the deposition procedure, the specimen is heated to a suitable temperature (above ~ 500 °C) to desorb any native oxides on the substrate surface. An overpressure of As is maintained to inhibit decom-
position of the substrate during the surface preparation procedure.
For this experiment, three compositions were used, all with (100) orientations: GaAs, In\textsubscript{15}Ga\textsubscript{85}As (grown on GaAs), and In\textsubscript{53}Ga\textsubscript{47}As (grown on InP). The GaAs was provided by Dr. Kin-Man Yu of Lawrence Berkeley Laboratory (Berkeley, CA 94720). The In\textsubscript{x}Ga\textsubscript{1-x}As was provided by Dr. Tim Sands and Dr. James Harbison of Bellcore, Inc (Red Bank, NJ 07701). The In\textsubscript{x}Ga\textsubscript{1-x}As was grown on a Varian Gen II MBE system, operating at a base pressure of 5\times10^{-11} Torr. The substrates were heated to 580 °C in an As overpressure to desorb surface oxides prior to deposition.

5.3 Pt deposition

Metal layers can be deposited on semiconductor substrates by a variety of means. The Pt metallizations in this experiment were deposited by plasma-enhanced sputtering, using a Perkin-Elmer Model 2400 radio-frequency plasma generator. An argon gas plasma was used to sputter the target material. The base pressure was 10^{-5} Torr, and the deposition pressure was 10^{-2} Torr.

Prior to deposition, the substrates were etched by the following procedure to remove native oxide: 2 minutes in boiling trichloroethane, 2 minutes in boiling acetone, methanol rinse, blow dry in N\textsubscript{2}, 5 minutes in de-ionized H\textsubscript{2}O:buffered HF solution (1:1), blow dry in N\textsubscript{2}. Although 1-2 nm of native oxide will regrow while the specimens are loaded into the sputtering chamber, the etching procedure will remove previous oxides, and provide the same starting surface for all specimens. On order to ensure good cooling of the substrate during the deposition procedure, the specimens were attached to a water-cooled plate with a dental wax. The maximum temperature attained by the substrate during deposition was estimated to be less than 100 °C.

The Pt was deposited using a forward power of 300 watts, and a target bias of 1350 volts; which resulted in a deposition rate of \approx 35 nm/minute. Following the Pt deposition, SiO\textsubscript{2} was deposited using a forward power of 300 watts, and a target bias of 1100 volts; the deposition rate was \approx 10 nm/minute. The final thicknesses of the
5.4 Annealing

The Pt and the \( \text{In}_x \text{Ga}_{1-x} \text{As} \) substrates were annealed in flowing forming gas (\( \text{N}_2: \text{H}_2, 95:5 \)) in a Lindburg Hevi-duty furnace, with a Eurotherm controller and an Omega type-K thermocouple. Temperatures within the furnace were controlled to within 1°C. Once the furnace and quartz holder reached the desired temperature, the specimen was placed on the holder and inserted into the furnace tube. After the designated annealing time, the specimen was air-quenched. For the kinetic studies, the samples were annealed at 400°C for times ranging from 10 minutes to 3 hours.

5.5 Analytical Techniques

In order to characterize these systems, three main features were examined, including: the progression of chemical reaction as a function of the annealing schedule, the crystal structures of the reaction products, and the morphology of the reacted layer. To determine this information, four analytical techniques were used: X-Ray Diffraction (XRD), Rutherford Backscattering (RBS), Scanning Auger Microscopy (SAM), and Transmission Electron Microscopy (TEM).

X-Ray Diffraction

The X-ray diffraction was performed as a quick, non-destructive way to examine the extent of the chemical reactions which occurred during the various annealing stages. In addition, the x-ray spectra helped identify the phases which developed by identifying the lattice spacings present in the system. Preferred orientations in the deposited and reacted material could be observed by the intensity ratios of the peaks; deviation from predicted ratios indicates an enhancement of grains with particular orientations. XRD was also very important for determining if new phases appeared for different annealing conditions. If new phases develop, the kinetic data would be
compromised, as different reactions would be occurring. This test was performed by comparing spectra from different times and temperatures to see if new d-spacings were present.

In this technique, a Siemens Kristalloflex X-Ray Diffractometer was used. The accelerating voltage was 40KeV, with a tube current of 30mA and a Cu target. A graphite crystal monochromator was used so that only peaks arising from diffraction of Cu Kα1,2 incident radiation were recorded. The x-rays were detected with a Tl-doped NaI scintillation counter.

Continuous scans were taken over $2\Theta = 15^\circ$ to $95^\circ$, with a step size of $0.1^\circ$, and a recording time of 2 seconds per step. The scans encompassed peaks down to 0.1nm d-spacings. The (100) oriented substrate material contributed two peaks in this range, corresponding to the (200) and (400) planes (GaAs: 31.6° and 66.1° and In$_{53}$Ga$_{47}$As: 30.4° and 63.3°). Based on structure factor calculations, the (200) peak will be of much lower intensity than the (400), and occasionally is not visible above the signal of the reaction products. The Pt depositions show preferred (111) or (100) orientations, which was evident by the enhancement of those peaks compared to the standard Pt powder spectrum. No specimen preparation was required for the x-ray diffraction.

Rutherford Backscattering

In order to determine the extent of the Pt/In$_x$Ga$_{1-x}$As interaction, RBS was performed. This is a non-destructive means for determining the depth distribution of the elements through the microstructure. Because of the complexity of the system, and the overlap of the signals from elements with similar atomic number (Ga and As), detailed information on the distribution of elements was impossible. However, RBS was effective in giving absolute thicknesses of the remaining unreacted Pt and In$_x$Ga$_{1-x}$As layers.
The thickness of the unreacted material was used to determine the reaction behavior of the system. The thickness of the reacted material is plotted versus the annealing time; for simple diffusion limited growth, the reaction rate should be parabolic: \(x \sim \sqrt{Dt}\). In principle, these plots can also be used to determine the activation energy of the reaction. However, in this experiment, the reaction is too complex to expect a single value.

RBS spectra were obtained by Dr. Kin Man Yu of the Lawrence Berkeley Laboratory. A 2MeV Van de Graaff accelerator was used to produce a 1.95MeV \(^4\text{He}^+\) beam. The backscattered protons were collected with a Si surface barrier detector located at an angle of 165° with respect to the incident beam. The apparatus gave a maximum depth resolution of \(\pm 5\)nm. No specimen preparation was required for RBS.

**Scanning Auger Microscopy**

Auger depth profiling is a destructive technique for determining the elemental distribution (and thereby the phase distribution) in the reacted structures. Auger spectra were obtained by bombarding the exposed specimen surface with an electron beam (10KeV), and recording the resultant Auger electron energies. Depth profiles were obtained by sputtering the specimen for a set time with an \(\text{Ar}^+\) ion beam (6 seconds at 5KeV, 1mm\(^2\) probe size), and taking a spectrum of the newly exposed surface. This procedure was repeated until the entire reacted layer was milled through (on the order of 20 minutes).

Absolute thicknesses are very difficult to determine with this technique, because of the variation in sputtering rates with changes in composition and morphology of the phases in the reacted layer. Depth profiles of elements will tend to blur the abruptness of an ideal interface, and extend the tails of the profile deeper into the specimen. The strength of the Auger signal from each of the elements cannot be easily correlated with its concentration in the layer, as the intensity depends on the element as well as the concentration. However, the sequence of elements in the microstructure can be
determined quite accurately. No specimen preparation was required for the Auger depth profiles.

**Transmission Electron Microscopy**

When several phases are present in the system, the XRD spectra are not effective in determining the phases present. For this reason, both conventional TEM and microdiffraction were used to uniquely identify the crystal structures of the product phases. TEM was also used to determine the morphology of the reaction products. Both cross-sectional and plan-view specimens were used in the TEM studies. In this study, a Philips EM 400, operating at 100KeV accelerating voltage was used. In order to resolve the microstructural features in this system, magnifications of 100,000 to 200,000x were used.

Specimen preparation is critical for obtaining good results in TEM. In order to be transparent to the electron beam, the specimen thickness must be reduced to <50nm, without introducing changes in the microstructure. Two orientations of the specimen were used to determine microstructural information. In plan-view specimens, the electron beam is normal to the original substrate surface. This orientation is useful for determining features in the plane of the reacted layer, grain size, and orientation relationships which may develop between the reacted layer and the substrate. In cross-section specimens, the electron beam is parallel to the original substrate surface. This orientation is useful for determining the depth distribution of the reaction products, and their morphology.

It should be noted that these materials are extremely fragile, so great care was taken to minimize the stress on the specimens at all stages of specimen preparation. This was especially important when handling the finished specimen.

The procedure for producing plan-view specimens in this experiment was fairly simple (figure 5.1). First, the SiO₂ encapsulation was removed by etching for two
Figure 5.1: Procedure for preparation of plan-view specimens for the TEM. a. The sample is cleaved into 2mm squares. b. The specimen is ground down to .1mm in thickness. c. A Cu grid is epoxied to the specimen to provide structural support, then a protective lacquer is applied, and the specimen is etched.
minutes in a solution of HF:de-ionized H₂O (1:1). The sample was then cut by cleaving into 2mm square specimens, which were 300 to 400 microns thick, with the deposition on one side, and the semiconductor substrate exposed on the other. The squares were then mechanically thinned down to 100 microns in thickness by removing the substrate material on a grinding wheel, with 600 grit paper. Support grids (Gilder 3mm diameter, 1x2mm slotted Cu grids) were then attached to the substrate side using Devcon 2-Ton Clear epoxy. During the grinding procedure, the specimens were fixed to a polishing stud with wax; this required that the specimens be exposed briefly to approximately 150 °C.

The specimens were then prepared for etching. The Cu support grids were protected by covering them with Tolber Miccroshield Stop-Off Lacquer. The substrate material was removed using a chlorine/methanol solution, which was prepared by bubbling chlorine through methanol until saturation. The specimens were suspended in a flow of the etching solution until the substrate was removed to perforation. At this point, most of the specimens were ready for the electron microscope. Occasionally, the etching solution did not remove enough of the reacted material; in these situations, ion milling was used to achieve the proper thickness.

The procedure for producing cross-sections was more complex (figures 5.2 and 5.3). First, the sample was cleaved into pairs of 2mm strips, which were epoxied together with the reacted layers face-to-face in the center. In order to ensure good bonding between the halves, the epoxied specimens were cured in an oven at 100 °C for 60 minutes. The specimen was then ground and polished on one side, using the following sequence of Buehler Carbimet silicon carbide grinding paper (approximate particle size in parentheses) 600 (30 micron), 1200 (18 micron), 2400 (8 micron), and 4000 (6 micron). A final polish was obtained by using a Gatan dimpler, with a felt polishing wheel and 0.05 micron Al₂O₃ slurry (3 minutes, medium speed, weight 20 grams).
Figure 5.2: Procedure for preparation of cross-section specimens for the TEM. 1mmx2mm strips of the sample are cleaved, and epoxied with the deposited layers face-to-face. Support strips of Si are epoxied on either side to provide mechanical strength.
Figure 5.3: a) The epoxied sandwich is ground down to 0.1mm in thickness. b) A shallow depression is ground in the specimen using a dimpler; the deepest part of the dimple is centered at the deposited layer interface. c) Cu support grid is epoxied to the specimen, which is then ion-milled to reduce the thickness to perforation.
At this point, the specimen was prepared for dimpling. The specimen was turned over, and thinned down to ~75 microns on 600 grit paper. A shallow depression, or dimple, centered on the two reacted layer interfaces, was formed in the specimen, using a brass wheel and 2-4 micron diamond paste (~10 minutes, medium speed, weight 20 grams), followed by a final polish with 0.05 micron Al₂O₃ slurry (3 minutes, medium speed, weight 20 grams). The thickness of the specimen at the bottom of the dimple at this point was 20 to 30 microns. A 3mm slotted Cu grid was epoxied to the specimen to provide strength. During the grinding, polishing, and dimpling procedure, the specimen was exposed to a maximum of ~150°C when attaching it to the polishing stud. Specimens in this experiment were dimpled on a Gatan model 656 Dimple Grinder.

The dimpled specimen was then thinned to the point of perforation at the bottom of the dimple. This was achieved by removing material in an ion mill, which functions by bombarding the specimen with energetic argon ions. The InP substrates used in this experiment posed a special problem for ion milling. The P atoms are much more easily removed from the substrate than are the In. Consequently, the In atoms tend to accumulate on the surface during the milling procedure, and can diffuse along the surface to agglomerate into droplets, which are detrimental to imaging in the TEM.

Indium droplet formation can be impeded by keeping the specimen surface temperature low enough to prevent diffusion of the In atoms into agglomerates. There are two measures which should be used to reduce the surface temperature: cooling the specimen with liquid nitrogen during milling, and reducing the total incident energy of the ion beam. Specimen cooling was achieved by mounting the specimen on a cold stage during milling, which was in thermal contact with a liquid nitrogen bath during milling: The total incident energy was reduced by keeping the energy of the ions low (below 3KeV), and the ion beam current low (below 0.5mA). Specimens in this experiment were milled on a Gatan model 600TMP Duomill, using guns designed for low-energy applications (Gatan model 600-37).
5.6 Summary

The analytical procedure can be summarized as follows: 1. After annealing, the specimens were examined by XRD to determine the extent of the metal/semiconductor reaction, and to identify the phases which have formed in the system (non-destructive). 2. RBS was performed to determine the thicknesses of unreacted Pt and In$_x$Ga$_{1-x}$As (non-destructive). 3. SAM depth profiling was performed to determine the depth distribution of the elements in the contact structure (this technique uses ~2mm$^2$ of material). 4. TEM was performed to determine the morphology of the reacted phases, and their orientation with respect to the substrate (this technique uses several mm$^2$ of material). The amount of material each technique requires is important because In$_x$Ga$_{1-x}$As is difficult to obtain in large quantities.
6.0 RESULTS

6.1 Pt/GaAs

In order to relate this study to the work of previous investigators (section 4.3, p. 49), annealing experiments were done on the Pt/GaAs system. The same conditions were maintained for the Pt/GaAs analysis as for the Pt/InGaAs system.

6.1.1 X-Ray Diffraction

X-ray diffraction spectra for several annealing times at 400 °C are shown in Figures 6.1 through 6.4; as deposited, 10 minutes, 15 minutes, and 40 minutes. In the as-deposited sample (Figure 6.1), Pt diffraction peaks are evident at the 2θ angles of 39.7°, 46.0°, 81.3° and 85.6°, in addition to the GaAs (200) and (400) peaks at 31.7° and 66.1°, respectively. No peaks indicative of reaction between the substrate and the Pt were observed in the as-deposited case. The intensity of the Pt peak at 39.7° peak is much higher than the Pt peak at 46.0°.

In Figure 6.2, the diffraction spectrum from the 10 minute anneal is shown. In addition to the GaAs and Pt peaks found in the as-deposited spectrum, several new peaks have appeared: 23.0°, 25.9°, 30.0°, 41.3°, 45.4°, 50.7°, and 72.3°. The Pt peaks have considerably lower intensity (e.g. the maximum counts for the peak near 40° are down by ~83%). In addition, the Pt peaks have shifted to higher angles (i.e. lower d-spacings). There is no change in the substrate peaks.

In Figure 6.3, the spectrum from the 15 minute anneal is shown. In addition to the reaction product peaks present following the 10 minute anneal, peaks have appeared at 33.7°, 36.8°, 43.1°, 62.9°, and 72.2°. The Pt peaks have decreased slightly in intensity, but have not shifted from their positions in the previous spectrum. The substrate peaks have not changed substantially, although the intensity of the (200) peak is much lower than previously.
Figure 6.1: X-ray diffraction spectrum for a Pt/GaAs specimen, as-deposited.
Figure 6.2: X-ray diffraction spectrum for a Pt/GaAs specimen annealed at 400°C for 10 minutes.
Figure 6.3: X-ray diffraction spectrum for a Pt/GaAs specimen annealed at 400°C for 15 minutes.
Figure 6.4: X-ray diffraction spectrum for a Pt/GaAs specimen annealed at 400 °C for 40 minutes.
Additional spectra were obtained for anneals of 20 and 30 minutes, but are not included here as they show no change from Figure 6.3. No new peaks developed, and no previous peaks disappeared.

In Figure 6.4, the spectrum from the 40 minute anneal is shown. At this stage in the annealing, the Pt signals have disappeared from the spectrum. The reaction product peaks are present at 25.8°, 30.0°, 33.7°, 36.8°, 41.3°, 43.0°, 45.4°, 50.7°, 52.7°, 56.5°, 63.2°, and 72.2°. All are of much lower intensity than that of the initial Pt deposition. The reaction product peak at 23.0°, which was present in all but the as-deposited spectrum, has disappeared. The GaAs substrate peaks are still present, with the (400) of much higher intensity than the (200).

6.1.2 Rutherford Backscattering

RBS spectra for various Pt/GaAs anneals are shown in Figure 6.5. The spectrum for the as-deposited specimen shows that the onset of the Pt signal occurs at the channel corresponding to surface Pt. The Pt signal increases to a broad peak, and drops off sharply at the low-energy end. There is a gap in the spectrum between the trailing edge of the Pt and the onset of the substrate Ga and As. The substrate signal increases fairly sharply, and exhibits no steps in its intensity at lower energies.

The spectrum following 10 minutes of annealing at 400°C shows a decrease in the intensity of the Pt signal, although the onset still occurs at the surface value. A step has appeared in the low-energy end of the Pt signal. The onset of the substrate signal has moved out into the gap between the Pt and GaAs, and a step has developed at the high-energies.

The intensity of the Pt signal has decreased further following the 30 minute anneal. The peak has broadened considerably, and exhibits two intensity levels. The GaAs signals have merged with the low-energy tail of the Pt peak.
Figure 6.5: Rutherford Backscattering spectra for as-deposited Pt/GaAs, and following 10 and 30 minute anneals at 400°C.
6.1.3 Auger Depth Profile

A depth profile of selected Auger transitions is shown in Figure 6.6, for a sample of Pt/GaAs annealed for 60 minutes at 400°C. In this profile, the following Auger transition energies were monitored as a function of ion milling time: As 1228 eV, Ga 1070 eV, and Pt 168 eV. The Auger computer was set to monitor the peak-to-peak height within an ~30 eV window. Because of the large number of possible Auger transitions for Pt, it was found that 12 sweeps of the energy window were necessary to get a good signal-to-noise ratio, as opposed to 6 each for As and Ga. It should be noted that the Pt signal is much lower than that for the other elements. While this may be indicative in part of low Pt concentrations, it results primarily from the larger number of Auger transitions available in the Pt atom. Consequently, each Pt Auger transition tends to have a lower intensity for a given incident radiation than do the transitions in smaller atoms.

The depth profiles in this study were taken on encapsulated specimens; this allowed for alignment of the device while sputtering the SiO₂ without affecting data collection of the reaction products.

The profile shows a very sharp increase in the Ga signal after ~5 minutes of milling. The Pt signal begins to increase soon afterwards, while the As signal remains flat until much later in the profile. After reaching a local maximum at ~10 minutes, the Ga signal begins to decrease gradually. The As signal begins to increase gradually ~2.5 minutes later, and levels off near 20 minutes. At the same time, the Ga signal reaches a local minimum and begins increasing, although at a lower rate than for the onset of the Ga signal. The Pt signal is fairly level between 7.5 minutes and 25 minutes, which is when the most Ga and As changes are occurring. Before 7.5 minutes, the Pt signal is near zero, as are the Ga and As. After 25 minutes, the Pt signal again goes to zero, while the Ga and As level off at a much higher value.
Figure 6.6: Depth profile of Auger energies for a Pt/GaAs specimen annealed at 400 °C for 60 minutes.
Features to note: First, there is no sharp transition between the various features in the As, Ga, and Pt signals. Second, the Ga signal never decreases to zero after the encapsulation is milled away. Third, the Pt signal does not vary substantially after the Ga signal first decreases, until the Ga and As level off. Fourth, the As signal does not increase until well after Ga is first encountered.
6.2 Pt/In$_{53}$Ga$_{47}$As

6.2.1 X-Ray Diffraction

Spectra of Pt/In$_{53}$Ga$_{47}$As samples annealed at several different times and temperatures are shown in Figures 6.7 through 6.13. They can be divided into two subgroups: 1. Samples annealed at 400 °C for 10 minutes through 180 minutes, and 2. Samples annealed at 350 °C to 600 °C, for long times.

as-deposited

The x-ray diffraction spectra following the deposition of Pt on In$_{53}$Ga$_{47}$As is shown in Figure 6.7. The substrate peaks are present at 30.6° and 63.4°, while the Pt deposition peaks appear at 39.7°, 45.9°, 80.8°, and 85.4°. The Pt peak at 39.7° has a much higher intensity than the other Pt peaks. No peaks are present that would indicate the interaction of the substrate and the Pt during deposition.

10 minutes

Several new peaks have appeared following this anneal (Figure 6.8), although most are of low intensity, e.g. 22.6° and 25.9°. Two of the new peaks have significant intensity; at 30.3° and 58.2°. The Pt peaks are all still present, and have shifted to larger angles (smaller d-spacings). The substrate peaks are essentially unchanged, although the intensity of the (200) is lower than was recorded in the previous spectra.

60 minutes

The spectrum obtained following the 60 minute anneal (Figure 6.9) is quite similar to that of the previous spectrum. Two peaks, at 31.5° and 41.2°, have become evident on the high angle shoulders of larger peaks. There are indications of the presence of these peaks after 20 and 30 minute anneals, but they are of too low intensity to be
convincing in those cases. Substantial reaction product peaks are present at 25.9°, 28.2°, 33.6°, 42.4°, 46.4, and 50.7°. Very high intensity peaks are present at 39.9° and 58.1°.
Figure 6.7: X-ray diffraction spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen, as-deposited.
Figure 6.8: X-ray diffraction spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 400 °C for 10 minutes.
Figure 6.9: X-ray diffraction spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 400 °C for 60 minutes.
Figure 6.10: X-ray diffraction spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 400 °C for 180 minutes.
Figure 6.11: X-ray diffraction spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 350 °C for 180 minutes.
Figure 6.12: X-ray diffraction spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 500 °C for 30 minutes.
Figure 6.13: X-ray diffraction spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 600 °C for 30 minutes.
90 minutes to 180 minutes

X-ray diffraction spectra were obtained following anneals for 90, 120, 150, and 180 minutes. The spectra differ only in the intensity of the diffraction peaks, so only the final spectrum (180 minutes) is provided here in Figure 6.10. All of the reaction product peaks are pronounced and well-defined for the longer anneals. No new peaks have appeared as compared to shorter anneals, although the peak near 41.2° mentioned previously has become much larger. There are also indications of a double peak at 25.7°.

long-term anneals

X-ray diffraction spectra were obtained for long-term anneals to determine the effect of temperature on the final reaction products. The annealing conditions for this set of data were: 350°C:180 minutes, 400°C:180 minutes, 500°C:30 minutes, and 600°C:30 minutes. The large variation in annealing time was necessary as the Pt/In$_{53}$Ga$_{47}$As reacts slowly at the lower temperatures. The spectrum for the 400°C:180 minutes anneal has been previously described.

350°C:180 minutes

Despite the long annealing time, little reaction has occurred following this anneal (Figure 6.11). The Pt peaks are still quite strong, although they have shifted from their as-deposited positions to higher angles: 40.0°, 46.7°, 81.7°, and 86.1°. One substantial reaction product peak is visible at 58.4°. In addition, a very broad signal is visible between 22° and 25°, and a narrower signal near 28°.

500°C:30 minutes

The Pt/In$_{53}$Ga$_{47}$As system has completely reacted following this anneal (Figure 6.12). All the reaction product peaks are quite pronounced, and of larger intensity than those of the lower temperature anneal. Most of the higher temperature peaks are
within 0.1° (the step size of the diffractometer) of the 400°C:180 minute anneal peaks, with a few exceptions. For example, the peak at 28.1° has shifted to 28.3°, 39.9° to 40.4° and 57.9° to 58.4°. In addition, a significant peak has appeared at 30.0°, on the low-angle shoulder of the substrate (200) peak at 30.6°.

Other features to note in this spectrum are the appearance of some small peaks, at 24.4°, 53.7°, and several between 66° and 90°. These peaks probably do not represent the appearance of new phases, as they correlate with regions of high noise in the 400°C:180 minutes anneal. In addition, a low, broad peak at the position of the Pt (200) has disappeared in going from the 400°C to the 500°C anneals.

600°C:30 minutes

This is the highest temperature anneal used in this study (Figure 6.13). This spectrum is very similar to the 500°C:30 minutes case, with the same exception noted before: some of the peaks have shifted to higher angles, the same that shifted when going from 400°C to 500°C. With respect to 500°C anneals, the peaks have shifted: from 24.4° to 24.6°, from 28.3° to 28.5°, from 40.4° to 40.7°, from 58.4° to 58.9°, and from 87.2° to 87.9°. The highest- and lowest-angle peaks in this set were too broad to be measured accurately in the 400°C:180 minutes spectrum.

The behavior of three of the peaks in the spectra these long-term anneals is of special note: those near 40°, 51°, and 58°. At 40°, two peaks nearly overlap, but by an amount that decreases as the annealing temperature increases: 1.2°, 0.8°, and 0.6°. Most of the decrease is taken up by the lower angle peak. At 58°, the peak goes through two changes as the temperature increases: it shifts to higher angles, and it develops a peak on the low angle shoulder. The peak at 51° is interesting because nothing appreciable changes about the peak as the annealing temperature increases, as will be discussed in section 7.
6.2.2 Rutherford Backscattering

Two RBS spectra are shown in Figures 6.14 and 6.15. Figure 6.14 shows the spectrum of Pt/In$_{53}$Ga$_{47}$As after deposition. The Pt on the surface of the specimen is evident from the large signal at channel 485. This signal shows a square profile, with no tail in the low-energy end of the signal. The counts decrease to zero between the end of the Pt signal, and the onset of the In signal. As in the case of the Pt, the In signal increases abruptly, without a high-energy tail. The In signal is level in intensity until the onset of the Ga and As, which increase smoothly to the continuous substrate intensity.

Figure 6.15 shows spectrum of a sample annealed for 30 minutes at 500 °C. The Pt signal in this case shows much lower intensity, and no clear dip between the Pt and the substrate Ga, As, and In signals. Both the Pt and the substrate signals have become irregular, exhibiting several broad peaks. The gap between the substrate and deposition signals no longer exists.

6.2.3 Auger Depth Profiles

Auger depth profiles have been obtained following anneals for several different annealing times at 400 °C. In addition, one profile was obtained for a sample annealed at 500 °C for 30 minutes, to examine the distribution of elements in a fully reacted sample.

Depth profiles of Auger electron energies are shown in Figures 6.16 through 6.19. The data collection conditions were the same as for those presented in Section 6.1 (page 81). In addition to the Auger energies sampled for Ga, As, and Pt mentioned previously, the In signal at 404eV was monitored. As in the case of Ga and As, 6 sweeps of the In window were taken.
Figure 6.14: Rutherford Backscattering spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen, as-deposited.
Figure 6.15: Rutherford Backscattering spectrum for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 500 °C for 30 minutes.
as-deposited

The Auger depth profile for the as-deposited specimen (not encapsulated) is shown in Figure 6.16. In this spectrum, the Pt signal is high from the start of the experiment. The signal remains constant until ~3 minutes, then decreases slowly until leveling off with a relative peak-to-peak height of ~0.5. Also near 3 minutes, the Ga, As, and In signals increase quite rapidly. The In signal rises first, followed simultaneously by the Ga and As. The Ga, As, and In signals all become roughly level after their initial increase.

Features to note: First, the Pt signal does not decrease abruptly when the Ga, As, and In signals appear. Second, the relative peak-to-peak values of Ga, As, and In change during subsequent milling. The In signal shows a slight local maximum while the Ga and As are still rising, then decreases, and gradually increases until leveling off to the substrate value. Third, the Ga signal follows a similar profile, although later in the spectrum, and the maximum intensity is not as pronounced as in the In case. The As is the slowest to reach its substrate level; not until after both the Ga and In have reached their maximum and diminished.

500 °C:30 minutes

The depth profile following the anneal at 500 °C is shown in Figure 6.17. Several new features are present in this sample. The first element to shown an increase in its Auger signal is In, which develops very rapidly after ~1.5 minutes. After reaching a maximum ~0.5 minutes layer, the In signal drops off at a similar rate. As the In signal decreases, the Ga increases very rapidly, reaching a maximum near the same milling point as the In local minimum, i.e. ~2.7 minutes. As in the case of the initial In signal, the Ga drops off rapidly after peaking. The In signal rises inversely with the Ga decrease. Simultaneous with the In increase, the As signal begins to rise, though at a slower rate. The As reaches a plateau, and then drops down to a local minimum. While As is rising to its plateau, the In signal reaches a maximum much sooner, then
begins a gradual decrease. When the As reaches its local minimum, the In has again jumped rapidly, peaked, and diminished to a constant value. Both the Ga and As reach a constant value after the In peak, in a manner similar to the as-deposited sample.
Figure 6.16: Depth profile of Auger energies for a Pt/In$_{53}$Ga$_{47}$As specimen, as-deposited.
Figure 6.17: Depth profile of Auger energies for a Pt/In₅₃Ga₄₇As specimen annealed at 500 °C for 30 minutes.
Features to note: First, both In and Ga begin the profile with peaks that develop sequentially, first In and then Ga. Second, the As profile begins with a much broader peak, that exists simultaneously with a significant In signal. Third, this In signal reaches a maximum quickly, and then decreases continuously throughout the range of the broad As peak. Fourth, at the onset of the substrate profile, the In signal peak is much more pronounced than it was in the as-deposited case. Fifth, the As signal reaches a local minimum when this In peak occurs.

400 °C: 30 minute

The Auger depth profile taken following this anneal is shown in Figure 6.18. In contrast to the fully reacted sample in Figure 6.17, the Pt signal is the first to increase in this profile. The Ga and In signals begin to rise simultaneously ~1 minute later. While the Ga signal reaches its maximum at a peak-to-peak value of ~3.5, the In continues to increase to the maximum of 10. Following its maximum, the Ga signal dips to a local minimum before rising to its substrate level. Coincident with the Ga peak at ~3.4 minutes, the As begins to rise. Coincident with the Ga dip, the As reaches a broad local maximum, before increasing to its substrate level.

Features to note: First, the Pt signal begins to rise ~1 minute before any of the other elements. Second, the In signal rises slowly, to a very high peak-to-peak value. Third, the Ga signal rises to a local maximum within the range of the In signal. Fourth, the Ga has a dip coincident with an As peak at ~4.2 minutes. Fifth, the pronounced In peak occurs just after the Pt signal disappears, and before the Ga and As have reached the substrate levels. Sixth, the Pt signal reaches its peak after its onset, then decreases continuously until it disappears at the substrate.
Figure 6.18: Depth profile of Auger energies for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 400°C for 30 minutes.
400 °C:120 minute

In Figure 6.19, the Auger depth profile following the 400 °C:120 minute anneal is shown. As in the previous sample, the Pt signal begins to rise before the other elements. In this case, the delay before the onset of other elements is much shorter. In is the first element to appear, ~0.2 minutes after the Pt. The In signal rises quickly to a short plateau within ~0.5 minutes, at a peak-to-peak value of ~3.0. The In signal rise quickly from this plateau, decreases in slope briefly, and continues on to a very high peak. The Ga signal develops later than the In, and increases very rapidly once it appears. The Ga reaches a local maximum, then drops very quickly to another minimum. At this point, it rises rapidly to the substrate value. The As signal begins to rise as the Ga signal drops. The As rises to a broad peak, diminishes slightly, and then increases to its substrate levels.

Features to note: First, the Pt appears before any of the other elements, although the milling time before the onset of the In is very short. Second, In is the second element to appear, which reaches two plateaus as it increases to a very high maximum just before the substrate. Third, the Ga is the third element to appear, and reaches a peak at the end of the first In plateau. Fourth, the Ga signal nearly disappears after the initial peak, and then rises to its substrate values. Fifth, the onset of the As signal coincides with the dip in the Ga signal, and the As reaches a broad peak at the bottom of the Ga dip. Sixth, the Pt signal peaks before the onset of the other elements, the decreases continuously until it disappears at the substrate.
Figure 6.19: Depth profile of Auger energies for a Pt/In$_{53}$Ga$_{47}$As specimen annealed at 400°C for 120 minutes.
6.2.4 Transmission Electron Microscopy

Figure 6.20 shows a Selected Area Diffraction pattern from a cross-section of a specimen annealed for 120 minutes at 400 °C. The micrograph shows In$_{53}$Ga$_{47}$As (001) pattern, superimposed with a highly textured polycrystalline pattern from the reaction products.

An SAD pattern from a plan-view specimen annealed at 400 °C for 120 minutes is shown in Figure 6.21. The SAD shows the diffraction spots from the substrate in the (011) orientation, as indicated by arrows in the figure. Well-defined diffraction spots from at least one of the reaction products are also visible.
Figure 6.20: Selected area diffraction pattern, taken from a cross-section specimen annealed at 400 °C for 120 minutes. The substrate (011) pattern is present, with a textured polycrystalline pattern overlayed which can be indexed as a Pt(In:Ga)₂ (001).
Figure 6.21: Selected area diffraction pattern, taken from a plan-view specimen annealed at 400 °C for 120 minutes. The substrate (001) pattern is visible, with highly textured polycrystalline spots present which can be indexed as a Pt(In:Ga)_2 (001) and (011).
7.0 DISCUSSION

7.1 Pt/GaAs

7.1.1 X-ray Diffraction

as-deposited

In the as-deposited sample (Figure 6.1), Pt peaks are evident at the 2θ angles of 39.7°, 46.0°, 81.3° and 85.6°, in addition to the GaAs (200) and (400) peaks at 31.7° and 66.1°, respectively. No peaks indicative of reaction between the substrate and the Pt were observed in the as-deposited case. The Pt peaks correspond to the (111), (200), (311), and (222) planes, with the intensity of the (111) peak higher than reported for a randomly oriented Pt powder. The reported relative intensities for a random sample are: (111):(200):(311):(222) 100:55:35:12 [JC88], while the data give 100:36:10:5. These relative intensities are consistent with a preference for the (111) orientation for deposited Pt, which has been previously reported for Pt/GaAs [KM75].

The preferred orientation of the initial Pt deposition can affect subsequent phases, giving them a residual preferred orientation that is not dependent on the substrate orientation. Kumar found that in the Pt/GaAs system, PtAs₂ first formed with preferred orientation attributed to the effect of the Pt preferred orientation, and subsequently developed its own relationship with the substrate.

10 minutes

The Pt/GaAs system reacts quite quickly at 400° C. In Figure 6.2, peaks from the reaction products are visible following the 10 minute anneal. These peaks indicate the development of PtAs₂, PtGa, and Pt₃Ga, with a considerable amount of residual Pt. Pt₃Ga is very difficult to distinguish from pure Pt (which is FCC). Pt₃Ga is an L1₂ ordered phase, with Pt atoms on the FCC cube corners, and Ga on the faces. It has a
lattice parameter of .3889nm, which is only 0.9% smaller than that of Pt. In order to distinguish the two phases, the superlattice reflections, which have very low intensity, must be examined. The Pt$_3$Ga (100) peak at 22.85° is the best to determine if the Pt$_3$Ga phase is present. In figure 6.2, there are peaks present at the superlattice positions, but all can be attributed to Pt or PtAs$_2$, except for the (100). Therefore, Pt$_3$Ga is a likely phase for these annealing conditions. The location of the Pt peaks has shifted to smaller d-spacings (by ~1.3%). This is consistent with the diffusion of Ga into the Pt (which reduces the lattice dimensions), and by the overlap of the Pt and Pt$_3$Ga peaks. Pt with Ga in solution will be denoted as Pt[Ga].

15 minutes

Following the 15 minute anneal (Figure 6.3), the reaction product peaks have become stronger, while the Pt signal has decreased. The relative intensities of the Pt[Ga] (111) at 40.2° and the PtGa (210) at 41.3° have reversed. The GaAs (200) peak is of considerably lower intensity than the GaAs (400) in this spectrum. This difference is to be expected in the GaAs (and In$_x$Ga$_{1-x}$As) system, where the intensity of the (200) arises from the difference in atomic scattering factor of Ga and As, as opposed to the intensity of the (400), which arises from the sum. The intensity of reflections from single crystals are much more sensitive to very small changes in orientation of the sample during XRD scans than are polycrystalline phases, so the intensity of the (200) peak is often very low.

Examination of this spectrum indicates the presence of the same phases found previously: Pt[Ga], PtAs$_2$, PtGa, and Pt$_3$Ga.

20 minutes to 30 minutes

After the 20 minute and 30 minute anneals, very little change is observed in the x-ray spectra compared to Figure 6.3, except for changes in the relative intensities of
the peaks. This indicates that no new phases have appeared, and that none of the old phases have disappeared.

40 minutes

Figure 6.4 shows the spectrum following a 40 minute anneal. There are two significant changes in this sample. First, the Pt peaks are no longer evident in the spectrum. Second, the Pt₃Ga phase is also no longer visible, as evidenced by the absence of any peak at 23.0°. This data confirms the result of other investigators (see section 4.3, p. 40) as to the final reaction products in this system: GaAs, PtAs₂, and PtGa. A summary of phase formation during the Pt/GaAs reactions is presented in Table III.

When more than one reacted phase is present, the ratios of peak intensities cannot be easily compared with the standard powder spectra. The effect of overlapping peaks from different phases on the intensities cannot be separated. However, some information can be determined once the phases have been identified. The relative intensities of the non-overlapped PtAs₂ peaks, the (200), (210), (211), (220), and (311) can be compared. The relative intensity of the PtAs₂ (200) (30.0°) peak is higher than reported for a randomly oriented powder, with the intensity of the PtAs₂ (311) (50.7°) lower than expected. The average ratio I(200)/I(311) for the data is 1.8±0.2, as compared with 0.6 for a randomly oriented powder. This change in relative intensities is consistent with texturing of the PtAs₂ product phase.
as-deposited | Pt, GaAs  
--- | ---  
10 to 30 min. | Pt[Ga], Pt$_3$Ga, PtGa, PtAs$_2$, GaAs  
40 min. | PtGa, PtAs$_2$, GaAs

**Table III**

Phases present in Pt/GaAs reactions after anneals at 400 °C, as determined from x-ray diffraction spectra.
7.1.2 Rutherford Backscattering

Rutherford Backscattering spectra from the Pt/GaAs as-deposited, 10 minute, and 40 minute anneals are shown in Figure 6.5. The spectrum for the as-deposited sample indicates that the Pt did not react significantly with the substrate during the deposition procedure. The low-energy shoulder of the peak and the high-energy shoulder of the substrate signal are sharp, with no evidence of a tail that would indicate interdiffusion is occurring. In addition, the top of the Pt and substrate peaks are fairly level, without large changes in intensity that would be indicative of changes in composition.

The spectrum from the 10 minute anneal shows an intermediate case where unreacted Pt is still present in the spectrum. The Pt peak still begins at the channel for surface Pt, but shows a step down on the low-energy side. The step indicates that the Pt has only partially reacted, and the reduced intensity of the Pt signal is consistent with a reduced thickness of unreacted Pt. The signal from the substrate elements has begun to intrude in the gap that appeared between the Pt and substrate signals in the as-deposited spectrum, indicating that at least one of these elements has diffused into the Pt.

The spectrum for the 30 minute anneal shows that there is no residual unreacted Pt, within the resolution limit of the system (~10nm). The Pt exhibits a step in its intensity, which is consistent with a change in composition across the reacted layer.

Results of the Rutherford Backscattering data are summarized in Figure 7.1. In this study, the thickness of the GaAs that has reacted was monitored in order to examine the kinetics of the Pt/GaAs reaction. The data exhibit a parabolic relationship between the amount of material reacted and the annealing time. This is consistent with a diffusion-controlled reaction, as has been previously reported for Pt/GaAs.
Figure 7.1: Plot of $x^2$ vs. annealing time, where $x$ is the thickness of GaAs that has reacted. Annealing temperature: 400°C.
The dotted line in Figure 7.1 is a linear regression of the data points. The slope of this line, which is $D_{\text{eff}}$ for this system (within a factor of 2), is $\sim 2 \times 10^{-13}$ (cm$^2$s$^{-1}$). Compared with the estimates of diffusivities for various diffusion modes and media in Table I (page 48), this value is $\sim 7$ orders of magnitude higher than for the lattice diffusivity of PtAs$_2$, and $\sim 3$ orders of magnitude for the lattice diffusivity of PtGa. If grain boundary diffusion is assumed, the measured $(d/\delta)xD_{\text{eff}}$ is $\sim 2 \times 10^{-11}$ (cm$^2$s$^{-1}$). Compared with the values for grain boundary diffusivities in Table I, this value is $\sim 3$ orders of magnitude higher than for the grain boundary diffusivity of PtAs$_2$, and on the same order of magnitude for the grain boundary diffusivity of PtGa. These data cannot conclusively determine the dominant diffusion mode in the Pt/GaAs system because of the many simplifying assumptions. However, the results indicate that the grain boundary diffusion mode is likely to have made the major contribution to the total diffusion.

7.1.3 Auger Depth Profile

Figure 6.6 shows the SAM depth profile for a Pt/GaAs sample that has been annealed for 60 minutes at 400 °C. This profile indicates the distribution of the elements in the sample, and hence the depth distribution of the phases found by XRD. It should be noted that these depth profiles tend to be less sharp than the corresponding microstructure, because of differential sputtering rates (vs. phase and morphology) and due to the dependence of the electron escape depth on the Auger energies. The ion milling procedure will also cause intermixing of the elements in the system, and tend to "push" some elements deeper into the sample. Consequently, the milling times indicated on the x-axis of the Auger depth profile is not indicative of the absolute depth in the specimen, but can only be used to identify relative positions.

In Figure 6.6, the reacted layer is encountered after $\sim 5$ minutes of milling, as indicated by the sudden increase in the Ga signal, and more gradual increase in the Pt signal. The absence of an increase in the As signal indicates that this element, and
hence the PtAs$_2$ phase is not present at the reacted layer surface. Combining the results of the x-ray diffraction studies with the Auger data, it can be concluded that the PtGa phase is located in the surface layer.

The As signal increases at $\sim$12.5 minutes, indicating the presence of PtAs$_2$. The gradual increase in As, and the simultaneous gradual decrease of Ga is consistent with a rough interface. This would be expected with grain sizes within an order of magnitude of the layer thickness, and some interpenetration of the two phases. The fact that the Ga signal never decreases to zero within this range indicates that a significant amount of Ga is present in this layer. The Ga has to be in the form of PtGa grains, as no other possibility is consistent with the x-ray data. Any other phase would give rise to unexplained peaks in the spectrum; and dissolution of Ga in PtAs$_2$ would shift the diffraction peaks of the arsenide. Close examination of Figure 6.6 shows an increase in the Ga signal and simultaneous decrease in the As signal at $\sim$21 minutes. This could arise from the presence of a particularly large PtGa grain at this point in the microstructure.

The Pt signal remains fairly flat throughout the range of the reacted layer, although it appears slightly higher in the PtGa region. This is consistent with the higher concentration of Pt in the PtGa as compared with PtAs$_2$ phases, but could also arise from different sputtering yields of Ga and As in the respective phases.

The data indicate that the interface between the reacted layer and the GaAs substrate is also rough. The Ga and As signals increase from the levels in the reacted layer to that of the substrate in $\sim$5 minutes of milling. This indicates the protrusion of PtGa and PtAs$_2$ grains into the substrate.

To summarize, the Auger data indicate that the reaction products are layered in the following sequence: GaAs/PtAs$_2$/PtGa. The data also indicate a concentration of Ga in the PtAs$_2$ layer, in the form of PtGa. In addition, the interfaces between the PtGa and PtAs$_2$ layers, and the reacted layer/GaAs are rough.
7.1.4 Summary

These experimental results are consistent with the work of previous investigators on the Pt/In$_x$Ga$_{1-x}$As system, for the case of $x = 0$:

1. The reaction kinetics are parabolic for thin films, indicating diffusion control.
2. Grain boundary diffusion is likely the dominant mode at 400°C.
3. Pt[Ga], Pt$_2$Ga, PtGa, PtAs$_2$, and GaAs are present at intermediate stages.
4. PtGa, PtAs$_2$, and GaAs are the end phases of the thin film reaction.
5. The PtAs$_2$ phase is textured.
6. The reaction products are layered in the following sequence: PtGa/PtAs$_2$/GaAs; although there is a significant amount of PtGa in the PtAs$_2$ layer.
7.2 Pt/In$_{53}$Ga$_{47}$As

In Figures 6.7 through 6.13, x-ray diffraction spectra for depositions of 70nm of Pt on In$_{53}$Ga$_{47}$As, annealed at 400°C for various times are presented. In all cases, the In$_{53}$Ga$_{47}$As substrate peaks at 30.5° (200) and 63.2° (400) are observed.

7.2.1 X-Ray Diffraction

as-deposited

In the as-deposited case (Figure 6.7), the Pt peaks are present at 39.7° (111), 45.9° (200), 80.8° (311), and 85.4° (222). These are consistent with the as-deposited Pt peaks on GaAs in Figure 6.1, with the discrepancy in the higher angle peaks due to the difficulty of identifying the peak position for very low intensity peaks. Note again the very high intensity of the Pt (111) peak, indicative of a preferred (111) orientation. The measured intensity ratios go as 100:5:4:5, as compared to 100:55:35:12 for a randomly oriented sample, and 100:36:10:5 for Pt/GaAs. This indicates an even higher preference for the (111) orientation than for Pt/GaAs. No reaction between the substrate and Pt deposition was observed by x-ray diffraction in the as-deposited samples for either case.

10 minutes

As in the case of Pt/GaAs, this annealing schedule (Figure 6.8) resulted in the reaction of the Pt with the substrate, although the generally lower intensities of the reaction product peaks in Pt/In$_{53}$Ga$_{47}$As suggest that the reaction is not as extensive at this early stage of annealing.

Comparing Figures 6.2 and 6.8, some similarities can be observed. The positions of the Pt peaks have again shifted to smaller d-spacings (by ~1.0% in Fig. 6.8), which is consistent with the diffusion of Ga into the Pt layer. The In atom is ~11% larger
than Pt (and ~14% larger than Ga), which would cause an expansion of the Pt lattice if this element were the only element to diffuse into the Pt. It is possible that both Ga and In are present, which could result in smaller shrinkage of the Pt lattice than was observed in the Pt/GaAs case (~1.3% in Figure 6.2). This possibility cannot be distinguished by x-ray diffraction.

Comparison of the non-substrate, non-Pt peaks with the PtAs$_2$ standard spectrum indicates that this phase is present. Several peaks can be attributed to PtAs$_2$: (111) at 25.9°, (200) at 30.3°, (220) at 42.6°, and (311) at 50.6°. At this early stage of annealing, it is impossible to determine if any texturing is developing in the PtAs$_2$, as only the (220) peak does not overlap with other phases. For the most part, the peak intensities are very low, with one exception. The peak at 30.3°, on the lower angle shoulder of the In$_{53}$Ga$_{47}$As (200), is consistent with PtAs$_2$ (200) and with significant intensity. For longer annealing times, additional PtAs$_2$ peaks develop that will allow for comparison of the recorded intensities with the standard, randomly oriented powder spectra.

As was discussed in section 7.1.1, the only peak available for uniquely identifying Pt$_3$Ga in these systems is the Pt$_3$Ga (100) peak at 22.85°. In Figure 6.8, a very low intensity peak has appeared at 22.6°. A peak at this angle is present in the spectrum, indicating that this phase also develops in the Pt/In$_{53}$Ga$_{47}$As reactions.

PtGa is present following the 400°C:10 minute anneal of Pt on GaAs. The existence PtGa cannot be definitely determined from the data in Figure 6.8. Any peaks which may be attributed to the PtGa phase are either too small at this stage to be convincing, or overlapping with other phases in the system. For example, the peak at 25.9° could be attributed to the PtAs$_2$ (111) or the PtGa (110).

There are peaks in this spectrum which could correspond to PtIn$_2$, although they are at slightly higher angles. These include a very strong peak at 58.2°, near the PtIn$_2$ (400) (57.9°), and a much smaller peak at 28.2°, near the PtIn$_2$ (200) (28.0°).
The peaks in the data are at smaller d-spacings than would be expected for this phase, by an average of \(~0.6\%\). A shift to smaller d-spacings could arise from the presence of Ga in this phase, i.e. Pt(In:Ga)$_2$. Indications are that at lower temperatures, nearly pure PtIn$_2$ forms, but as the annealing temperature increases, a significant solubility of Ga develops, and the lattice parameter decreases.

The possibility of a solid solution between PtGa$_2$ and PtIn$_2$, based on their identical crystal structures and the similarity of the group III elements, was discussed in section 4.1 (p. 32). The In atomic radius is 14\% larger than that of Ga. These facts suggest that it is possible for the PtIn$_2$ structure to accommodate some Ga atoms in a substitutional solid solution, with a corresponding decrease in lattice parameter.

To summarize, the probable phases present following the 10 minute anneal: Pt[Ga], PtAs$_2$, Pt(In:Ga)$_2$, and Pt$_3$Ga.

**20 minutes to 30 minutes**

The spectra taken following the 20 and 30 minute anneals (not shown) are very similar to Figure 6.8. No new phases have developed, and no previous phases have disappeared. The reaction products peaks are increasing in intensity, especially the Pt(In:Ga)$_2$ (400) at 58.1°. By extrapolating from the indexed Pt(In:Ga)$_2$ peaks, it should be possible to determine the lattice parameter, and estimate the composition of this phase. At these annealing stages, however, the presence of residual Pt peaks prevents an accurate determination of the peak positions. Peaks are developing at 31.7° and 41.1°, although their exact positions are difficult to determine because of their very low intensity and their positions on the shoulders of other peaks. They can be attributed to the PtGa (111) and (210) planes; other PtGa peaks may be present but would be overlapped by PtAs$_2$ or Pt(In:Ga)$_2$ peaks.

The probable phases present following 20 to 30 minute anneals: Pt[Ga], PtAs$_2$, Pt(In:Ga)$_2$, and Pt$_3$Ga, with PtGa possibly present.
60 minutes

The reaction products following a 60 minute anneal (Figure 6.9) at 400 °C are the same as for the previous anneals. In addition, the PtGa peaks (at 31.7 ° and 41.1 °) discussed for the 20 to 30 minute anneals are much more pronounced, indicating the presence of this phase.

Substantial PtAs$_2$ diffraction peaks are now present; the (111) at 25.9 °, the (210) at 33.6 °, the (211) at 36.8 °, the (220) at 42.9 °, and the (311) at 50.7 °. Many of these peaks overlap with PtGa and Pt(In:Ga)$_2$, so comparison of the relative intensities can be made only for the (210) and (220), which are rather low intensity peaks. The intensities of these two peaks is the same, giving $I_{(210)}/I_{(220)} \sim 1$, while the intensity ratio reported for standard powder specimens is 0.6. The standards also predict that the ratio $I_{(210)}/I_{(311)} = 0.3$, while the ratio calculated from the data in Figure 6.9 gives a value of $\sim 0.3$, despite the fact that the intensity of the (311) has contributions from the Pt(In:Ga)$_2$ phase. This result suggests that the PtAs$_2$ phase may exhibit some preferred orientation, but the effect is most likely very slight.

180 minutes

Following the 180 minute anneal (Figure 6.10), virtually all of the Pt has disappeared. There are indications of a small peak at 46 °, the Pt (200). Otherwise, the reaction product peaks are well defined. One of the reasons for performing x-ray diffraction on these samples is to determine if phase formation changes as a function of annealing time. If the phase formation is not consistent, kinetic data is difficult to interpret. The x-ray data for anneals at 400 °C indicate this is true, with the possible exception that PtGa cannot be positively identified until after the first 10 minutes. The phase formation sequence for the Pt/In$_{53}$Ga$_{47}$As system annealed at 400 °C is summarized in Table IV.
<table>
<thead>
<tr>
<th>Time Range</th>
<th>Phases Present</th>
</tr>
</thead>
<tbody>
<tr>
<td>as-deposited</td>
<td>Pt, In$<em>{53}$Ga$</em>{47}$As</td>
</tr>
<tr>
<td>10 min.</td>
<td>Pt[Ga], Pt$_3$Ga, Pt(In:Ga)$<em>2$, PtAs$<em>2$, In$</em>{53}$Ga$</em>{47}$As</td>
</tr>
<tr>
<td>20 to 180 min.</td>
<td>Pt[Ga], Pt$_3$Ga, PtGa, Pt(In:Ga)$<em>2$, PtAs$<em>2$, In$</em>{53}$Ga$</em>{47}$As</td>
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<tr>
<td>final phases</td>
<td>PtGa, Pt(In:Ga)$<em>2$, PtAs$<em>2$, In$</em>{53}$Ga$</em>{47}$As</td>
</tr>
</tbody>
</table>

**Table IV**

Phases present in Pt/In$_{53}$Ga$_{47}$As reactions after anneals at 400 °C, as determined from x-ray diffraction spectra.
Long-term Anneals

The indexed spectrum in Figure 7.2 corresponds to a fully reacted Pt/In$_{53}$Ga$_{47}$As microstructure (500 °C:30 minutes). The final product phases in this system: In$_{53}$Ga$_{47}$As, PtAs$_2$, Pt(In:Ga)$_2$, and PtGa. Only the PtAs$_2$ (210) and (220) peaks do not overlap with other phases, making determination of texturing in this phase difficult. The ratio $I_{(210)}/I_{(220)} \approx 1.6$, as compared to 0.6 for a randomly oriented sample. The intensity of the PtAs$_2$ (200) peak can be estimated from the spectrum, giving $I_{(200)}/I_{(220)} \approx 1.5$, as compared to 1.2 for a randomly oriented sample. Both of these results suggest some texturing of the PtAs$_2$ phase, but to a much smaller degree than was found in the Pt/GaAs case.

An important result of the study of long-term anneals is the existence of a solid solution between PtIn$_2$ and PtGa$_2$, or Pt(In:Ga)$_2$. Prominent peaks that can be attributed to this phase are present near 28° (200), 40° (220), 58° (400), and 86° (440) in these spectra. These data can be extrapolated to the unit cell dimension to give $a_0$. Assuming Vegard's law for small concentrations of Ga in PtIn$_2$, an expression for the lattice parameter of the ternary phase PtIn$_{2x}$Ga$_{2-2x}$ is: $a_0 = .5923\text{nm} + .0443x$. Conversely, the amount of Ga in the phase can be determined by: $x = \left( a_0 - .5923\text{nm} \right) \div .0443$.

The XRD data for these annealing conditions indicate that the lattice parameter of PtIn$_{2x}$Ga$_{2-2x}$ decreases as the annealing temperature increases, indicating a higher solubility of Ga. In Figure 7.3, the lattice parameter and Ga solubility of the Pt(In:Ga)$_2$ phase is shown as a function of annealing temperature.

A consequence of the shift in Pt(In:Ga)$_2$ peaks as a function of annealing temperature is the uncovering or covering of peaks from other phases. The peaks near 40°, 50°, and 58° are a demonstration of this behavior. The PtGa (210) peak appears at 41.1°, on the high-angle shoulder of the Pt(In:Ga)$_2$ (220). As the annealing temperature increases, the difference between these two peaks shrinks, from 1.2° to 0.8° to
0.6°. Virtually all of the decrease in spacing between these peaks comes from shifts in the location of the Pt(In:Ga)$_2$ (220).
Figure 7.2: Indexed x-ray diffraction pattern for fully reacted Pt/In$_{53}$Ga$_{47}$As specimen, 500° C:30 minute anneal.
Figure 7.3: Variation in the lattice parameter and composition of \( \text{PtIn}_{2x}\text{Ga}_{2-2x} \) with annealing temperature.
Conversely, the peak near 58° consists of the PtAs$_2$ (231) at 57.8° and the Pt(In:Ga)$_2$ (400). As the annealing temperature increases, the PtAs$_2$ peak appears first as a bulge on the low-angle shoulder of the peak at 500°C, and completely emerges at 600°C, as the Ga concentration of Pt(In:Ga)$_2$ increases.

The peak near 51° consists of the PtAs$_2$ (311) at 50.7° and the Pt(In:Ga)$_2$ (222). However, there is no significant change in this peak as the annealing temperature changes. This is probably due to the low intensity of the (222) peak; note that the Pt(In:Ga)$_2$ near 25° is very small.

The high intensity of some of the Pt(In:Ga)$_2$ peaks suggest the possibility of texturing. The (220) (40.0°) and (400) (58.2°) peaks are very strong, while the (200) (28.2°) and (440) (86.2°) are of moderate strength. No diffraction data is available for Pt(In:Ga)$_2$; by assuming that the Ga is randomly oriented on the In sites, the relative intensities should be similar to those of PtIn$_2$ and PtGa$_2$. A comparison of the relative intensities of these peaks is provided in Table V. These data indicate that there is a preference for the (400), and to a lesser degree, the (220) peaks in the 400°C and 500°C anneals when compared to PtIn$_2$ and PtGa$_2$. For the highest temperature examined in this study, 600°C, the preference is quite pronounced, with the (220) peak much more intense than expected. However, the possibility that the Ga and In are ordering on the group III sublattice can not be discounted as an explanation for the change in peak intensities.
Table V

Comparison of relative intensities of Pt(In:Ga)$_2$ for different annealing temperatures. Relative intensities for PtIn$_2$ and PtGa$_2$ are from reference [JC88].

<table>
<thead>
<tr>
<th>plane</th>
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<th>500 °C</th>
<th>600 °C</th>
<th>PtGa$_2$</th>
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</tr>
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<td>4</td>
<td>5</td>
<td>3</td>
<td>50</td>
</tr>
</tbody>
</table>
7.2.2 Rutherford Backscattering

The RBS spectrum in Figure 6.14 shows that no reaction has occurred between the Pt and the $\text{In}_{53}\text{Ga}_{47}\text{As}$ during the deposition process, within the resolution of the device ($\sim10\text{nm}$). This can be concluded by the sharp, abrupt transitions associated with the Pt low-energy and the In high-energy shoulders. In addition, the Pt, In, and GaAs signals become level, indicating no change in composition with depth with the metallization and substrate layers.

Figure 6.15 shows a fully reacted microstructure. The intensity of the Pt has diminished considerably, and the signals from the Pt and substrate peaks have merged. Unfortunately, because of the large number of elements in this system, and the overlap of the characteristic backscattering energies of the elements, it is virtually impossible to determine profile of the elements as a function of depth, and the relative compositions of the phases. Auger is much more desirable for determining this information.

A plot of the square of the thickness of $\text{In}_{53}\text{Ga}_{47}\text{As}$ reacted and annealing time is provided in Figure 7.4. This plot differs significantly from the Pt/GaAs case, which is provided in the same plot. In the initial stages of the Pt/$\text{In}_{53}\text{Ga}_{47}\text{As}$ reaction, the reaction kinetics follow a parabolic relationship with a slope of $\sim1\times10^{-13}\text{cm}^2\text{s}^{-1}$. The reaction rate in this stage is close to the Pt/GaAs case, where the slope is $\sim2\times10^{-13}\text{cm}^2\text{s}^{-1}$. For annealing times longer than $\sim60$ minutes, the reaction slows significantly, and can be characterized with a new slope of $\sim2\times10^{-14}\text{cm}^2\text{s}^{-1}$, a decrease of a factor of 5.

Compared with the estimates of diffusivities for various diffusion modes and media in Table I (page 48), both stage I and stage II are unlikely to involve lattice diffusion, especially through $\text{PtAs}_2$. The $\text{PtIn}_2$ lattice diffusion is the closest, which is still 2 orders of magnitude too slow. If grain boundary diffusion is assumed, $(d/\delta)\times D_{\text{eff}}$ is $\sim1\times10^{-11} \text{ (cm}^2\text{s}^{-1})$ in stage I, and $\sim2\times10^{-12} \text{ (cm}^2\text{s}^{-1})$ for stage II. Both of these diffusivities are about one order of magnitude slower than predicted for grain boundary
diffusion through PtGa and PtIn$_2$, and two orders of magnitude faster than estimated for PtAs$_2$.

As was discussed for Pt/GaAs, these results cannot conclusively determine the diffusion mode in Pt/In$_{53}$Ga$_{47}$As. However, the results suggest that the grain boundary diffusivity dominates the diffusion in this system. Also, a continuous PtAs$_2$ layer is unlikely, because the low diffusivities associated with this phase are inconsistent with the reaction kinetics data.
Figure 7.4: Plot of $x^2$ vs. time, where $x$ is the thickness of the substrate that has reacted.
7.2.3 Auger Depth Profile

There are some important features of the Auger profiles in Figures 6.16 through 6.19 that appear for all conditions, but to varying degrees. Once the Ga, As, and In signals indicate that the substrate has been reached, they do not level off at a constant value. In general, the In signal will reach a maximum level first before decreasing and leveling off. This could be indicative of an In rich layer at the substrate surface, or could be the result of ion milling effects. The likelihood of these two possibilities will be addressed for each profile below. Another common feature is that the Ga and As signals exchange their peak-to-peak values deep within the substrate. Due to the position of this feature in the sample, it probably arises from milling effects.

as-deposited

Although Figure 6.16 shows penetration of the Pt signal into the substrate, the RBS data indicates that there is no interaction between the substrate and the Pt following deposition. The RBS is more reliable in this case, as the ion milling during Auger profiling cause intermixing of microstructural features. Another note: the baseline (i.e. zero signal) as indicated in these profiles corresponds to a peak-to-peak value of 0.5.

The variations in the Ga, As, and In signals within the substrates are very slight in the as-deposited sample, and are most likely due to sputtering effects as discussed above. RBS does not detect any interaction for this sample, so the shape of these profiles indicate the effects of Auger data collection on the spectra. The variations of the onset of each element in Figure 6.16 should be noted. Whether as a result of differential milling rates or different electron escape depths, the In signal shows an onset much earlier than Ga and As, which appear simultaneously. It should be noted that the atomic numbers of the Ga and As are very close (31 and 33), while In is much higher (49). A similar comparison can be made of the energies of the Auger electrons: Ga 1070eV and As 1228eV vs. 404eV for In.
The depth profile in Figure 6.17 is discussed here to examine a completely reacted microstructure. Several different layers can be deduced from this figure. The first layer that is encountered (1.5 to 2 minutes) is a Pt:In phase, indicated by the peak in the In signal. The actual composition of the Pt:In phase cannot be determined from the Auger profile, as the peak-to-peak values depend on several factors other than composition.

Following the peak of the In signal is a sharp dip. The minimum of this depression in the In signal coincides with a peak of the Ga signal, indicating a layer of a Pt:Ga phase (between 2 minutes and 3 minutes). Based on the diffraction data, this phase layer most likely consists of PtGa. Although the In signal does not disappear at the minimum, Auger depth profiles are not as sharp as the microstructures they correspond to. An alternating sequence of thin layers will appear as seen in Figure 6.17; i.e., the Auger signals will not drop to zero between layers if they are thin enough. By the standards of Auger spectra, the transitions between the Pt:In and Pt:Ga signals are fairly sharp, indicating that the interfaces between these phases are not as rough as found in the Pt/GaAs system.

The Ga quickly diminishes following its peak, decreasing to a very low value in the next layer (3 minutes to 5 minutes). As the Ga decreases, the As and In signals rise, indicating a layer containing these two elements in addition to the Pt which present throughout the reacted layer. As was discussed in section 7.1.3 for Pt/GaAs, the only possible As phase consistent with the x-ray diffraction data is PtAs$_2$. Also, the Ga and In cannot be present in solution in significant amounts in this phase without affecting the PtAs$_2$ lattice parameter, and shifting the diffraction peaks.

The remaining elements in this layer indicated by the Auger (Ga, In, and Pt) are probably in the form of Pt(In:Ga)$_2$. This would be consistent with the x-ray diffraction data for two reasons. First, the diffraction peaks show texturing of the Pt(In:Ga)$_2$
product phase, which indicates that the phase develops near the $\text{In}_{53}\text{Ga}_{47}\text{As}$ substrate interface. Second, both the Ga and In signals are at a maximum at the onset of the As signal, and diminish with nearly parallel slopes as the As increase, indicating a constant composition. This would be consistent with a two-phase mixture of $\text{Pt(In:Ga)}_2:\text{PtAs}_2$, with a decreasing fraction of the ternary compound as the substrate is approached.

The final layer in the reacted microstructure cannot be identified as clearly. As was discussed at the beginning of this section, the Auger signals of the substrate do not remain constant as would be expected. However, in the case of Figure 6.17, the peak in the In signal at the onset of the substrate is quite pronounced, and unlikely to result solely from milling effects. The presence of this peak, clearly after the Pt signal has disappeared suggests that the reaction of the Pt with the $\text{In}_{53}\text{Ga}_{47}\text{As}$ has resulted in the formation of an In-rich $\text{In}_x\text{Ga}_{1-x}\text{As}$ region at the substrate/reacted layer interface. This region could result from differential diffusion rates of the Ga, As, and In from the substrate during reaction. The In-rich region would result if the diffusion of In out of substrate was slower than diffusion of Ga, which would be consistent with previous studies (see section 4.3, page 49).

400 °C: 30 minute

The layering present in the specimen annealed at 500 °C has not yet developed in Figure 6.18. In addition, a significant amount of unreacted Pt is still present, as indicated by the large Pt signal while the Ga, As, and In have not appeared. In general, the Pt signal reaches an initial peak, and then decreases continuously before disappearing at the substrate. Although it is difficult to draw conclusions about composition from Auger data, these results are consistent with an initially high Pt concentration in the reacted layer (e.g. $\text{PtGa}$), which decreases closer to the substrate (e.g. $\text{PtAs}_2$ and $\text{Pt(In:Ga)}_2$). The x-ray diffraction data for this specimen (section 7.2.1) indicated the presence of Ga in the unreacted Pt. The Auger profile is not sensitive enough to detect the presence of this element in Figure 6.18.
The depth profile of the in this figure is beginning to develop a form similar to that of the fully reacted sample, especially for Ga and As. A peak in the Ga signal is present at the reaction front (at ~2.7 minutes) followed by a dip, and finally increasing up to the substrate level. The As signal begins increasing at the same level as the Ga maximum, and reaches a broad peak just at the minimum of the Ga dip. These results are consistent with a Pt:Ga layer on top of a PtAs$_2$ layer, which is what is found in the Pt/GaAs system for the identical annealing and deposition thickness. However, the In profile has yet to develop into the profile found in the completely reacted case. The In Auger signal appears at the same point as the Ga signal. Therefore, the diffusing In atoms have reached the same point in the reacted microstructure.

It is interesting to note the amount of the Pt:Ga phase in the PtAs$_2$ layer. In the case of Pt on GaAs, the fully reacted microstructure showed a large fraction the the PtGa phase with this layer. In the case of Pt on In$_{53}$Ga$_{47}$As, however, this layer showed very little Ga, but instead a large concentration of an In-rich ternary phase related to PtIn$_2$.

As in the previous figure, there is a prominent peak in the In profile just at the reacted layer/substrate interface. Again, this could be indicative of the buildup of In at this interface, possibly in the form of In-rich In$_x$Ga$_{1-x}$As.

The x-ray diffraction data showed that the phases present in both the 500°C and 400°C anneals for 30 minutes were the same. The difference in the Auger profiles would therefore indicate only a change in the distribution of the phases in the microstructure. The distribution for this anneal would likely take the form of two- and three-phase mixtures. The phase profile for this anneal should be: Pt[Ga] / Pt:In+PtGa / PtAs$_2$+Pt(In:Ga)$_2$+PtGa / In$_x$Ga$_{1-x}$As / In$_{53}$Ga$_{47}$As.
400 °C: 120 minutes

The Auger profile shown in Figure 6.19 shows the transition between the early stages of reaction in Figure 6.18, and the final reacted structure in Figure 6.17. The onset of the Pt signal before the development of the other elements shows that the sample has not yet fully reacted. In addition, the Pt profile through the reacted layer is consistent with a gradually decreasing concentration of this element within the reacted layer as the substrate is approached.

As in the previous sample, both the Ga and As profiles show the same general features of the fully reacted profiles for Pt/GaAs and Pt/In$_{53}$Ga$_{47}$As. In this case, however, the In profile has begun to show the features of the fully reacted case. A peak is developing in the In signal at the start of the reaction front, preceding the onset of the Ga signal, and indicating the development of a Pt:In phase at the surface. Deeper in the microstructure, a plateau in the In signal is developing, coincident with the PtAs$_2$ layer. This indicates the presence of the Pt(In:Ga)$_2$ phase which was found in the fully reacted case. Finally, the prominent peak at the reacted layer/substrate interface is still present. These results show that the layering found in the final reacted layer is developing, while unreacted Pt is still present in the system.

Auger summary

The Auger profiles indicate some important similarities between the Pt/GaAs and Pt/In$_{53}$Ga$_{47}$As cases. Both show a PtAs$_2$ layer growing on the semiconductor substrate, intermixed with a significant fraction of a Pt–A$_{III}$ phase. Instead of the PtGa found in Pt/GaAs, the ternary substrate reaction results in a ternary Pt(In:Ga)$_2$ phase, and with a higher fraction of the phase present. This is not surprising, in light of the differences between Ga and In, which suggested that Ga would diffuse more readily than In, leaving more In behind near the substrate to react. It should be noted that the presence of a small amount the Ga in this layer in the form of PtGa cannot be discounted.
7.2.4 Transmission Electron Microscopy

The SAD patterns in Figures 6.21 and 6.22 confirm the texturing of the Pt(In:Ga)$_2$ phase. The reaction product diffraction spots in Figure 6.21, from a cross-section, can be indexed as Pt(In:Ga)$_2$, with [220](001)$_{\text{Pt(In:Ga)}} \parallel [020](101)$_{\text{In$_{55}$Ga$_{47}$As}}$.

The pattern in Figure 6.22 (plan view) corresponds to two different orientations of Pt(In:Ga)$_2$ overlayed on the substrate (001). The orientation relationships are:

$[220](001)_{\text{Pt(In:Ga)}} \parallel [220](001)_{\text{In$_{55}$Ga$_{47}$As}}$ and $[200](011)_{\text{Pt(In:Ga)}} \parallel [220](001)_{\text{In$_{55}$Ga$_{47}$As}}$. The latter relationship and the orientation relationship from Figure 6.21 are equivalent. The texturing of this product phase is difficult to justify on a simple geometrical model, because the lattice mismatch between the substrate and the Pt(In:Ga)$_2$ is $\sim 7.6\%$. Consequently, other factors must account for this relationship, as was described in section 3, page 22.

7.3 Summary

Kinetics

The difference in the reaction kinetics between Pt/GaAs and Pt/In$_{55}$Ga$_{47}$As is very significant. One of the purposes of this study was to determine if the system could be made more stable by changing the substrate composition and lattice parameter. The results of this study show that the kinetics are significantly different for the ternary substrate case, and can be divided into two stages. In the initial stages, the reaction rate approaches that of Pt/GaAs. In the later stages, however, the kinetics are considerably slowed. This is an important result, indicating that this system does result in slower metal/semiconductor reactions, enhancing the potential for producing a stable contact.

Because of the complexity of the system, it is difficult to determine the cause of the change in reaction rate; although the phases present in the system are essentially
the same until the unreacted Pt disappears completely, they exist in several layers. Changes in the morphology of any of these layers can result in the kinetics observed.

**Substrate Composition**

The Auger depth profiles indicate the formation on an In-rich layer at the reacted layer/substrate interface. This could take the form of a continuous In$_x$Ga$_{1-x}$As layer, or of In$_x$Ga$_{1-x}$As precipitates. The presence on an In-rich region just below the reacted layer would have profound significance. As was discussed in section 2.2, page 15, the Schottky barrier height of a contact to In$_x$Ga$_{1-x}$As decreases as the In concentration of the substrate increases. Consequently, the microstructure that these results show develops in the Pt/In$_{53}$Ga$_{47}$As system can have application in ohmic contacts, where low Schottky barrier heights are critical.

**Phase Formation**

The Pt/In$_{53}$Ga$_{47}$As reaction products are similar to those found in the Pt/GaAs and Pt/InP systems. Pt/GaAs reacts to form PtAs$_2$ and PtGa, and Pt/InP reacts to form PtP$_2$ and PtIn$_2$. The system in this study formed a combination of these phases, PtAs$_2$, PtGa, and Pt(In:Ga)$_2$. In addition, the same phases were present at the intermediate stages of the reaction, i.e. Pt$_3$Ga, and Pt[Ga].

One question that should be raised is the lack of an intermediate Pt:In compound. The x-ray diffraction data do not indicate the presence of any such phase, although the Auger depth profiles show that there is a Pt:In compound on the surface. There are several possible explanations for this result. First, neither the Auger nor the RBS is sensitive enough to detect the presence of the small amount of Ga that may exist in the surface compound. Therefore, the Pt:In compound on the surface may be Pt(In:Ga)$_2$, which would be consistent with the x-ray data. Second, the diffracting volume (i.e. thickness) of the surface Pt:In layer may be too small for its x-ray peaks
to be visible above the background noise (less than \( \sim 10\text{nm} \)). Third, the diffraction peaks of the Pt:In layer may be overlapped by the large number of peaks that already exist in these spectra. For example, Pt\(_3\)Ga has only one non-overlapped peak.

In order to resolve this issue, there are several experiments that can be recommended for future study. For example, much thicker depositions (hundreds of nanometers) could be made, and annealed to full reaction. Based on the results of this study, this would require higher reaction temperatures (e.g. \( 600\,^\circ\text{C} \)) and/or much longer annealing times (several hours). In this study, 3 hours was insufficient to completely react 70nm of Pt at \( 400\,^\circ\text{C} \). The greater thickness may result in a thicker Pt:In layer on the surface to study, although changing the conditions of the experiment to this extent may result in the formation of other phases.

**Pt(In:Ga)\(_2\)**

The Pt(In:Ga)\(_2\) product phase is certainly the most interesting phase in the system. Two orientation relationships are present for the \( 400\,^\circ\text{C} \), 120 minutes anneals:

\[
[220](001)_{\text{Pt(In:Ga)}_2} \parallel [220](001)_{\text{In}_{53}\text{Ga}_{47}\text{As}}, \quad \text{and} \quad [200](011)_{\text{Pt(In:Ga)}_2} \parallel [220](001)_{\text{In}_{53}\text{Ga}_{47}\text{As}}.
\]

These relationships develop despite the large mismatch between this phase and \( \text{In}_{53}\text{Ga}_{47}\text{As} \), about \( 7.6\% \).

The x-ray diffraction data indicate that the relative amounts of grains in these relationship changes with higher annealing temperatures. As shown in Table V, at \( 400\,^\circ\text{C} \) and \( 500\,^\circ\text{C} \) the intensities of the (220) and (400) reflections are approximately the same. At \( 600\,^\circ\text{C} \), however, the (400) becomes significantly higher, suggesting that the second orientation relationship, \( [200](011)_{\text{Pt(In:Ga)}_2} \parallel [220](001)_{\text{In}_{53}\text{Ga}_{47}\text{As}} \), becomes preferred at this temperature. This change in preferred orientation may arise from a change in lattice parameter of Pt(In:Ga)\(_2\), which decreases by \( \sim 1\% \) in going from \( 400\,^\circ\text{C} \) to \( 600\,^\circ\text{C} \) anneals. The mismatch with the substrate changes from \( 7.6\% \) to \( 6.6\% \) in going to the higher annealing temperature.
Phase Separation

The fully reacted Pt/In$_{53}$Ga$_{47}$As microstructure formed a layered sequence with many similarities to the Pt/GaAs case. For the binary substrate, the layering was PtGa / PtAs$_2$ + PtGa / GaAs. For the ternary substrate, the result was Pt:In / PtGa / PtAs$_2$ + Pt(In:Ga)$_2$ / In$_x$Ga$_{1-x}$As / In$_{53}$Ga$_{47}$As. There are three significant differences: the additional Pt:In phase on the surface, the presence of Pt(In:Ga)$_2$ as the Pt–A$^{III}$ phase in the PtAs$_2$ layer, and the change in substrate composition below the reacted layer.
7.4 Conclusions

The main conclusions of this study can be summarized as follows:

1. The reaction kinetics can be divided into two stages, where the reaction slows by a factor of 5 after \( \sim 60 \) minutes of annealing. The reduction in reaction rate has considerable potential for the formation of stable metal/semiconductor contacts.

2. An In-rich region develops at the reacted layer/substrate interface, which could be quite beneficial for the formation of ohmic contacts.

3. At \( 400 \, ^\circ \text{C} \), the \( \text{Pt(In:Ga)}_2 \) product phase exhibits two orientation relationships with the substrate: \([220] (001)_{\text{Pt(In:Ga)}_2} \parallel [220] (001)_{\text{In}_{0.5} \text{Ga}_{0.5} \text{As}}\) and \([200] (011)_{\text{Pt(In:Ga)}_2} \parallel [220] (001)_{\text{In}_{0.5} \text{Ga}_{0.5} \text{As}}\), with the latter enhanced at higher annealing temperatures.

7.5 Future Work

The results of this study suggest several interesting projects for future study. The most important would be to examine the effect of changes in the \( \text{In}_x \text{Ga}_{1-x} \text{As} \) substrate composition during these reactions. The exact nature of the In-rich region at the reacted layer/substrate interface, and its effect on electrical properties is very important. An increased In concentration in the substrate just below the metallization is likely to reduce the Schottky barrier height, which would be beneficial for the formation of ohmic contacts.

An additional study that concerns the electrical behavior of this system involves the elimination of \( \text{PtAs}_2 \) as a product phase. Due to the volatility of As, it may be possible to reduce the amount of this phase in the reacted microstructure by annealing in an open system, i.e. without encapsulation. \( \text{Pt(In:Ga)}_2 \) is likely to have better electrical properties than \( \text{PtAs}_2 \) (which is a narrow band gap semiconductor), thus the conductivity of the reacted metallization might be enhanced.
As different compositions of $\text{In}_x\text{Ga}_{1-x}\text{As}$ become available, studies of the reactions for different lattice parameters will be possible. $\text{Pt(In:Ga)}_2$ may be induced to form epitaxially for appropriate conditions, or this phase could be deposited by MBE. In this way, it may be possible to decrease the $\text{Pt/In}_x\text{Ga}_{1-x}\text{As}$ reaction kinetics even further than has been achieved in this study.
8.0 REFERENCES


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