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# GALLIUM NITRIDE FILM DEPOSITION BY IONIZED CLUSTER BEAM EPITAXY

ΒY

### GHULUM BAKIRI

This thesis is being submitted in partial fulfillment of the requirements, outlined by the Electrical Engineering Department, to the faculty of New Jersey Institute of Technology for the degree of Master of Science in Electrical Engineering, 1983.

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### **ABSTRACT**

Title of Thesis: Gallium Nitride Film Deposition by Ionized Cluster Beam Epitaxy

Ghulum Bakiri, Master of Science in Electrical Engineering, 1982.

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An experimental Ionized Cluster Beam System similar in design to the one developed at Kyoto University in Japan was designed and constructed in the Microelectronics Laboratory at NJIT. This involved making the working drawings, having the parts machined and then assembling the system in a vacuum system together with the necessary variable power supplies, meters, controls, gas inlets, cooling water conn-

Eleven deposition runs were then attempted to grow gallium nitride utilizing the system constructed. In a typical deposition run, gallium was vaporized from a boron nitride crucible at  $1000^{\circ}\text{C}$  and allowed to expand through a 0.5mm nozzle into a low pressure region containing nitrogen at 1-5 X  $10^{-4}$  torr in order to form clusters of about 1000 atoms each. The gallium clusters were ionized by an electron source (300 mA) and deposited on heated quartz substrates (550°C) for reactive growth with nitrogen.

For the above deposition conditions, the film growth rate was only 1 A/min and the films were highly insulating. Photoluminescence measurements and surface analysis of the grown films revealed boron contaimination which was then traced to water vapor contamination of the boron nitride crucible and heater. The water vapor, if not driven out by pre-heating the crucible in an oven, caused the boron nitride to decompose at elevated temperatures introducing reactive contaminants at the substrate surface. Further experiments showed that for GaN

deposition rates greater than 1  $^{\rm O}$  A/min (and up to 1  $^{\rm O}$ A/sec) the crucible temperature should be at least 1100 $^{\rm O}$ C but preferably around 1400 $^{\rm O}$ C.



### ACKNOWLEDGEMENT

I wish to express my sincere appreciation to all those who contributed to the work culminating in the writing of this thesis. Special thanks are due to Dr. Roy Cornely for a great deal of his time spent in reviewing my work and for his most valuable suggestions which resulted in many important improvements in this thesis. Sincere gratitude is also due to Dr. Lawrence Suchow for his advice.

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I am also grateful to my fellow students Chetan Mehta, Fred Solomon, Young Shin Pyun and many other students whose help made this work possible.

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TO MY WIFE

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#### CHAPTER I

### INTRODUCTION

Ionized cluster beam deposition (ICBD), first reported by T. Takagi et. al. of Kyoto University in 1972, is now emerging from laboratory development and is promising to become an important competing technique for the fabrication of various kinds of solid state devices and materials 936,37,51). It combines a relatively high deposition rate with several additional advantages over conventional vacuum deposition methods such as molecular beam epitaxy, thermal evaporation, the various sputtering techniques (magnetron, triode, diode) and alternative ion transport methods (40,57).

A schematic diagram of the system used in Takagi's experiments is shown in fig. 1.1. For ICB epitaxy, metal or semiconductor material to be deposited is placed in a confined crucible and heated to a high temperature so that an optimum pressure (usually between 0.01 and several torr) is maintained in the crucible. Vapour is ejected from the crucible through a small nozzle into a high vacuum chamber (pressure usually in the  $10^{-6}$  range). As the temperature of the expanding vapour is reduced due to the adiabatic expansion, a point of supersaturation is reached, condensation occurs, and clusters are formed. Clusters are groups of 500 - 2000 atoms, loosely coupled together by

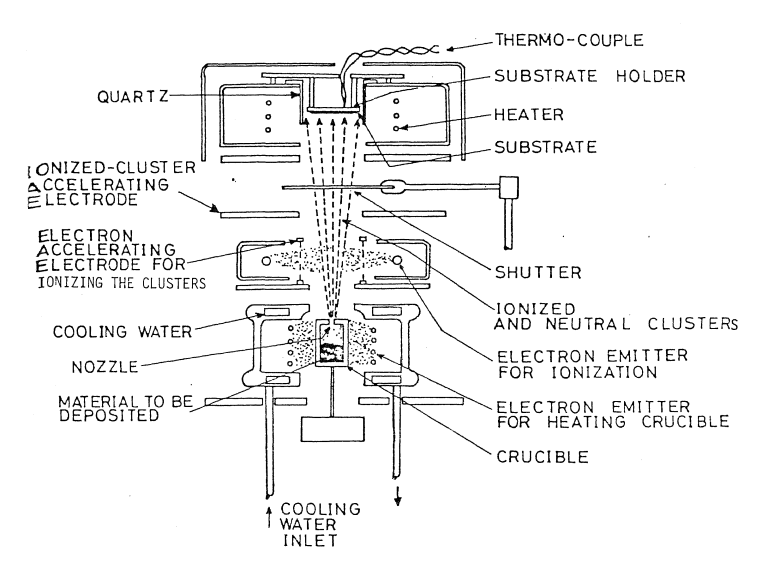


fig. 1.1 Schematic diagram of the ICB system used by Takagi Research Group. (41,46)

weak van der Waals forces. The clusters are ionized by electron impact in the ionization section located above the crucible. The cluster ions are accelerated toward the substrate by a high negative potential applied to the accelerating electrode. Ionized clusters typically comprise a few % to 40% of the total flux arriving at the substrate, the rest being neutral clusters which move toward the substrate by their ejection velocity alone.

In the reactive ion cluster beam deposition (R-ICBD), the same technique described above is used, and in addition a reactive gas is supplied from a gas nozzle through a controlled leak valve to maintain the desired gas pressure in the chamber (usually  $10^{-4}$  to  $10^{-5}$  torr). Some of the gas molecules are ionized along with the clusters, and are accelerated toward the substrate, where neutral and ionized gas atoms react with neutral and ionized particles of the vaporized metal clusters to form the desired compound film.

One important advantage of ICBD is the enhancement of surface migration of adatoms on the substrate surface during film formation, as the kinetic energy of the impinging clusters are converted to lateral movement of the individual atoms of the broken up clusters. Also, as the ionized cluster beam has an extremely small charge to mass ratio (e/m), problems due to the space charge effects in high current beam transportation (with a low voltage) and

problems encountered in depositing onto an insulating substrate can be eliminated. A third advantage is the controllability of the ICBD technique, where the dependence of the film properties on the ionization content (ratio), the acceleration voltage, source material temperature, and substrate temperature allows the optimization of deposition and enhancing the properties of the deposited film. There are also inherent advantages in the presence of even a small percentage of ions in the system, not only for accelerating these ions by the application of an electric field, but also to influence the critical parameters of initial film formation such as nucleation, coalescence, removing (scrubbing) of adsorbed impurities from the substrate surface, and chemical reactivity.

A large number of elemental and compound films deposited by the ICB process have been demonstrated. Examples have included metals such as Cu, Au, Ag, and Pb, and oxides such as BeO, PbO, and ZnO (55). Crystallographic structures and physical characteristics of these films, such as adhesion strength, density, thin layer continuity, and absence of contaminants, have been extraordinary. Very important applications of the ICB technique involve active semiconductor films. It has been shown that undoped and doped crystalline silicon films can be grown epitaxially on silicon substrates at substrate temperatures as low as 620°C.

Thermally-stable amorphous silicon-hydrogen alloy semiconductor films suitable for solar cell use have been
achieved. Compound semiconductor films such as CdTe, GaAs,
GaP, GaN, and ZnS with excellent physical, optical and
electrical characteristics have been deposited. Reference
55 includes a summary table listing the various films deposited so far by ICBD or by R-ICBD along with references
which the reader may consult for further details of the
results obtained in each case.

### 1.1 About This Work:-

The work reported in this thesis was carried out in two steps:-

1. The first step was to design and construct an experimental ICB system in the Microelectronics Laboratory at NJIT. This was done by the writer with the help of Fred Solomon and Chetan Mehta, two undergraduate E.E. students at NJIT.

Chapters II and III deal with this part of the work. Chapter II discusses the ionized cluster beam deposition method from the theoretical standpoint. The discussion includes cluster formation mechanism and the effects that varying the expansion parameters have on cluster formation. Characteristics of the ionized cluster beam, film formation mechanism, and the effects of the various control parameters on the properties of films obtained by the ICBD technique are also discussed.

Chapter III is devoted to a detailed description of the ICB system constructed at NJIT.

2. The next step was to use the ICB system thus constructed for GaN film deposition. Eleven deposition runs were attempted. The results obtained in each of these runs are described in chapter IV along with details of the experimental procedure followed in these runs.

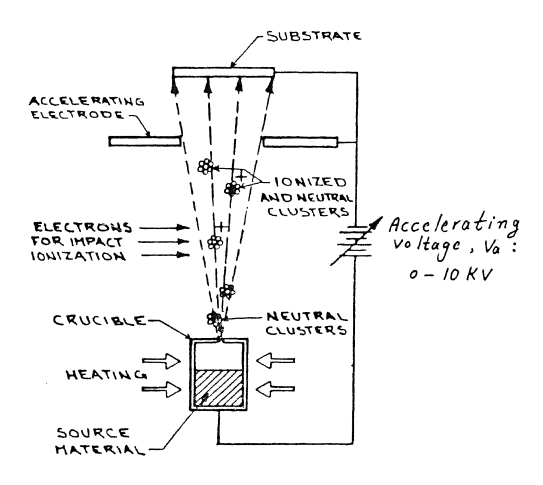
Chapter IV also deals briefly with gallium nitride material including material synthesis by chemical vapour transport methods and by the R-ICB technique, GaN optical properties and the structure of GaN light-emitting diodes. Conclusions and suggestions for future work are the subject of the last section of chapter IV (section 4.20).

#### CHAPTER II

#### I.C.B. DEPOSITION

### 2.1 General Overview

The Ion Cluster Beam (ICB) technique is a superior method for the deposition of thin films which offers controls over fundamental characteristics of film growth not possible by conventional thin-film production processes. The basic ICB deposition process is suggested in fig. 2.1 . Material to be deposited is vaporized from a confinement crucible at high temperatures into a high-vacuum chamber. Crucible nozzle configuration and operating temperatures are such that the emerging vapour undergoes supercondensation following adiabatic expansion through the nozzle, and atomic aggregate clusters, each of about 500 to 2000 loosely coupled atoms, are formed. A fraction of the clusters are then caused to become singly charged by electron impact ionization and these positively charged clusters can be accelerated toward the substrate by a high negative potential. Both neutral and ionized clusters contribute to film growth. Clusters arriving at the growth surface are broken up into individual atoms upon impact. Energy per atom is small in the case of the neutral clusters but averages  $eV_a/N$  for the ionized clusters of N atoms accelerated through potential  $\mathbf{V}_{\mathbf{a}}$ . In this chapter, cluster formation and growth will be investigated theoritically together with the characteristics of the ionized cluster beam, Film formation mechanism by the ICB deposition is then discussed taking into



account the effects of the kinetic energy and the presence of ions for film formation. Experimental results already published are cited, where appropriate, to support the theoritical discussion.

# 2.2 Cluster Formation Mechanism

# Simple View (52)

When a gas leaves a source (Temperature  $T_0$ , Pressure  $P_0$ ) and expands into a vacuum chamber (Pressure  $P_1$  in the beam), its temperature, with the assumption of adiabatic expansion, drops to a value  $T_1$  given by the relation: (3,14)

$$\frac{T_{1}}{T_{0}} = \frac{P_{1}}{P_{0}} (Y-1)/Y \qquad (1)$$

where  $\chi$  is the ratio between specific heats at constant pressure and constant volume:

f is the number of degrees of freedom of one molecule of the gas. ( )  $=\frac{5}{3}$  for monoatomic gases). From (1) it is seen that if the aperture between the source and vacuum chamber is small enough to allow a pressure ratio,  $P_1/P_0$ , of the order of  $10^{-3}$  or less, the temperature ratio,  $T_1/T_0$ , will be of the order of  $10^{-1}$  or less. If we start from the vapour of a metal just above its boiling point, i.e.  $T_0 = T_{\text{boiling}}$ , it is clear that  $T_1 = 10^{-1}$ . The jet expansion then becomes an efficient process for nucleating clusters. Once the critical nuclei have been produced, they continue to grow as long as the pressure of their surrounding

gas (vapour) remains high enough for frequent collisions to occur. In practice, this applies along the first few millimeters away from the nozzle source. Further downstream, the pressure would have dropped close to the high vacuum range  $(10^{-4} \text{ to } 10^{-7} \text{ torr})$ , at which point there are virtually no further collisions between the clusters formed and the vapour molecules, so the clusters keep their sizes constant.

A more "in depth" description of the condensation and clustering process can be given by carrying the analogy even further between metal vapour expansion into a high vacuum chamber and the expansion of a gas in a free jet. The description also requires the use of the classical theory of nucleation and condensation in the supersaturated state.

# Representative curve of the expansion

Assuming the gas(vapour) leaves the source, passing through a nozzle of diameter D, then expands as a free jet into a vacuum chamber where there is a pressure  $P_1$ , we may consider this expansion to be adiabatic and isentropic (8). For a perfect gas, it is then possible to calculate the Mach number, M, numerically for every point of the jet (33). In particular, along the axis, one obtains the function M = f(x/D), where x is the distance from the point on the axis to the orifice (21).

The temperature and pressure are then given by (2):-

$$T = T_{O} \cdot \left[ 1 + \frac{1}{2} (Y-1) M^{2} \right]^{-1}$$

$$P = P_{O} \cdot \left( T/T_{O} \right) / (Y-1)$$

$$= P_{O} \cdot \left[ 1 - \frac{1}{2} (Y-1) M^{2} \right]^{-Y/(Y-1)}$$

from which we can write

 $\log P - \log P = \left( \frac{Y}{Y} - 1 \right) \left( \log T - \log T_{0} \right),$ which is a straight line with constant slope as shown in fig. 2.2 . If the expansion starts at point A  $(P_0, T_0)$ , it will follow the line marked isentrope as far as point B where the isentrope meets the coexistence curve, xy, of the two phases. If there were equilibrium between the phases, the expansion would continue following BY. However, if the density is high enough beyond point B (as is the case normally when  $\frac{r_0}{P_1}$  is larger than  $10^3$ ), the expansion will continue along the so called "dry isentrope", BZ, into regions where the gas is in a supersaturated state. The saturation ratio,  $S = \frac{P}{P_0}$ , where  $\mathbf{P}_{\mathbf{e}}$  is the equilibrium solid vapour pressure at the temperature under consideration, rises from unity at point B, to a maximum value of  $\boldsymbol{S}_{\boldsymbol{D}}$  at point D, where the presence of clusters in the gas becomes noticeable. Further along the curve, the effect of condensation is to draw the representative point of the expansion towards the equilibrium curve.

# Simple Classical Formulation (8)

According to the classical theory of nucleation  $^{(1)}$  a cluster, which is assumed to be spherical, is in equilibrium with its vapour if it has a critical radius  $r^*$ , and contains  $n^*$  number of molecules given by:-

$$r* = 2 Q V/KT log S$$

$$n^* = 4 \pi r^*^3/3V$$

where K =the Boltzmann constant, Q =the surface tension,

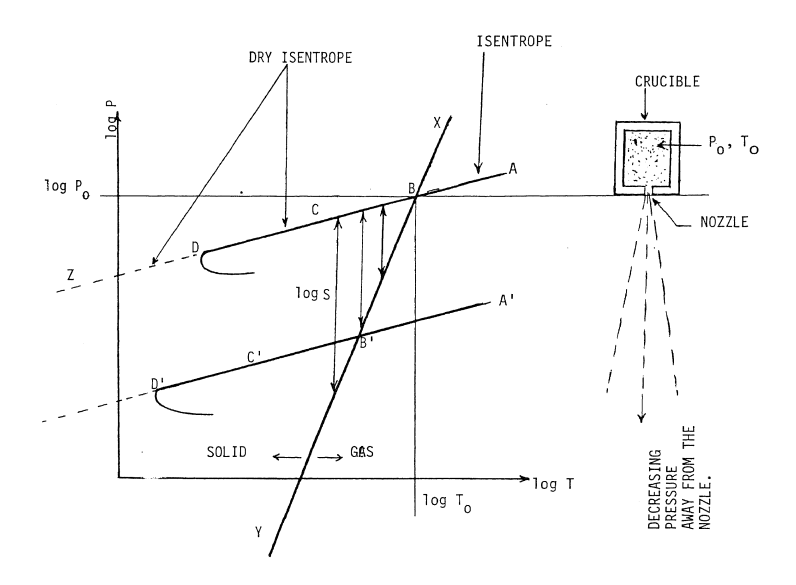


fig. 2.2 Representative curve of the "Expansion with Condensation" in the log P - log T  $\frac{1}{2} \frac{1}{2} \frac{1$ 

V = the average volume of one molecule in the cluster, T = the vapour temperature, and S = the saturation ratio defined earlier. Clusters of smaller size than critical are unstable and tend to break up and diminish in radius. On the other hand, a cluster of the critical radius will continue to grow very rapidly if it receives additional molecules through collisions.

A cluster of size n (i.e. having n molecules, where n can be less than or equal to n\*, the critical size) can come into existence simply as a result of density fluctuations in the vapour though its free energy of formation  $\Delta G_n$  is positive and rises with n. Assuming that the probability of finding a cluster of n molecules is proportional to the Boltzmann factor  $\exp(-\Delta G_n/\mathrm{KT})$ , we see that the number,  $N_{n*}$ , of clusters per cm<sup>3</sup> which reach the critical size rises rapidly as n\* gets smaller, which occurs when the saturation ratio rises:

$$(s \uparrow \longrightarrow r^* \downarrow \longrightarrow n^* \downarrow \longrightarrow \Delta_{G_{n^*}} \downarrow)$$

The nucleation rate, J, represents the number of clusters which pass the critical size per cm $^3$  per second. J is proportional to N $_{\rm n^{\star}}$ . Therefore it is apparant that the nucleation rate rises extremely rapidly with S, the saturation ratio.

### Nucleation Region

Starting from point A (fig. 2.2), the gas expands following the line BZ as described earlier. Beyond point B, S continues to rise; the critical size gets smaller and the nucleation rate starts to increase. A point is reached (C in fig. 2.2) where J is sufficiently large that it becomes

probable for a critical nucleus to be formed within a small volume (of the order of D<sup>3</sup> around the axis) during the characteristic time of the expansion. Once formed, the critical nuclei continue to grow at the expense of the surrounding vapour molecules. Point C then can be considered as the onset of nucleation. Beyond that point, the nucleation rate rises extremely rapidly. This continues until the amount of vapour condensed in the jet is no longer negligible as a fraction of the total (point D). At this point, the rate of nucleation reaches its maximum value. If it were possible to measure the number of clusters as a function of their size at point D, one would find a rapidly decreasing function as shown in fig. 2.3 . The largest clusters are less numerous since they were formed further upstream where the nucleation rate is smaller.

### Growth Region

Downstream from D, the condensation of the gas causes the pressure to drop, while the temperature remains roughly constant and then rises as result of the release of the latent heat of vaporization. The saturation ratio diminishes and with it the nucleation rate. The production of critical nuclei ceases fairly quickly but the condensation continues allowing the clusters already formed to continue to grow until they reach the high-vacuum regions where the gas density is low and collisions are minimal.

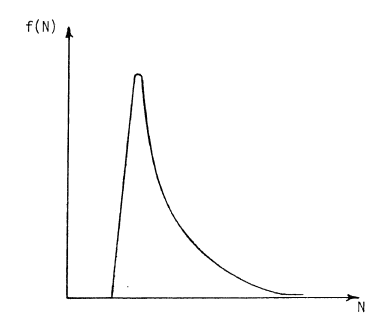


fig. 2.3 Size distribution of the clusters
 at the end of the nucleation region
 (before their growth).(8)

# 2.3 Effects of the Expansion Parameters on Cluster Formation (11)

Although the classical theory of homogeneous nucleation is not quantitatively applicable to small clusters (55), it is possible to use the theory to understand qualitatively the effects that varying the expansion parameters will have on the clusters.

#### Variation of Nozzle Diameter

A reduction in the nozzle diameter results in increasing the speed of the expanding vapour, and hence the gas stays a shorter time in the nucleation and growth regions. shorter time for growth means that the clusters formed will be smaller. Therefore a reduction of the diameter of the nozzle causes a reduction in the size of the clusters formed. In ICB experiments reported by T. Takagi group (56,52,44), nozzle diameters of up to 2 mm were shown to be adequate for cluster formation with 2 mm being the most effective. For larger nozzle sizes the intensity of the cluster beam starts to drop until it completely disappears with nozzle diameters greater than 5 mm. This is due to the pressure difference between the inside and the outside of the crucible being insufficient. Under these conditions, the vapour consists of atoms, molecules and/or small clusters comprising several atoms instead of larger clusters (55).

#### Effect of Source Pressure and Temperature

In considering the effect of source pressure and temperature on cluster formation, one cannot continue using the analogy between free gas expansion and metal vapour expansion, because in the case of the free gas both  $T_{\odot}$  and  $P_{\odot}$  are usually controlled independently, while in the case of metal evaporation, the source usually contains metal vapour at pressure  $P_{O}$  in equilibrium with the solid, so  $P_{O}$  and  $T_{\cap}$  are related. This means that in fig. 2.2, our starting point (Po, To) is not point A, but rather point B. For the expansion of gas in a free jet, the effect of an increase in  $T_{\odot}$  opposes and in effect balances the effects of an increase in  $P_{o}^{(11)}$ . Since an increase in  $P_{o}$  is almost always achieved by increasing  $T_{\circ}$  in ICB experiments, one would expect no dependence of cluster size (and hence energy) on  $P_{\Omega}$ . There is however the requirement that  $\mathbf{P}_{_{\textstyle \mathrm{O}}}$  (and hence  $\mathbf{T}_{_{\textstyle \mathrm{O}}})$  be high enough to produce a high saturation ratio as discussed earlier. In practice, it is found that  $P_0 = 10^3$ .  $P_{chamber}$  is sufficient for cluster formation. In Takagi's experiments, it was found that there is an optimum pressure Polopt for each nozzle diameter D, that resulted in the highest intensity of the cluster beam. Taking the values of Po.opt for different nozzle diameters, D, the product Po,opt. D was approximately constant. Since the mean free path  $\lambda$  is inversely proportional to P , we conclude that D/ $\lambda$  is constant.

This means that the intensity of the cluster beam is maximum for a specific ratio of (D/ $\chi$ ) which depends on the other parameters of the system.

#### 2.4 Size Distribution of the Clusters

Fig. 2.3 for the size distribution of clusters at point D (beginning of the growth region) undergoes a change as the clusters pass through the growth region. Larger clusters grow faster than smaller ones because of their larger capture cross section  $^{(8)}$ . This has the effect of broadening the distribution curve, which will then correspond more closely to the curves obtained in ICB experiments as shown in fig. 2.4 . The average diameter of clusters ranges from 50  $^{\circ}$   $^{\circ}$ 

#### 2.5 Deposition Rate:-

The deposition rate in the ICB system is primarily dependent on the source nozzle diameter, the vapour pressure inside the crucible ( $P_{\rm O}$ ) and the pressure outside the crucible ( $P_{\rm Chamber}$ ).

#### Nozzle Diameter

The deposition rate is proportional to the area of the nozzle opening, which for a circular hole is proportional to the square of the nozzle diameter. If, for example, a deposition rate of 0.4  ${\rm ^{O}}$  /sec is obtainable at certain conditions with a 1 mm diameter nozzle, the deposition rate can

<sup>\*</sup> See page 20.

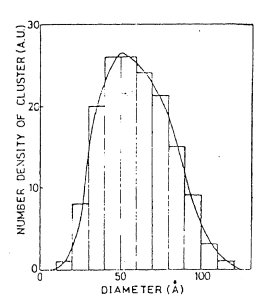


Fig. 2.4 Size distribution of Ag clusters deposited at  $P_{\circ} = 1$  torr for 30 seconds.

be increased to about  $1\text{\AA}$  /sec  $^*$  by increasing the nozzle diameter to 1.5 mm.

#### Crucible Temperature:-

This determines the vapour pressure inside the crucible,  $P_{o}$ , and must be made high enough to achieve a pressure ratio,  $P_{o}/P_{chamber}$ , of at least  $10^4$  -  $10^5$  to achieve satisfactory deposition rates.

Fig. 2.5 shows the relation between the deposition rate and the source (crucible) temperature for silicon deposition using a 1.5 mm diameter nozzle and a chamber pressure in the  $10^{-7}$  torr range. A "threshold" temperature of about 1900°C is observed below which the deposition rate is rather low, but it increases rapidly beyond this threshold temperature.

#### Chamber Pressure:-

Increasing the chamber pressure has the same effect on the deposition rate as lowering the crucible temperature, and vice versa.

<sup>\*</sup>These values are typical for silicon deposition using a crucible temperature of about 2050°C (corresponding to a vapour pressure in the crucible of 1 torr) and a chamber pressure of 10<sup>-7</sup> torr (Source: Nova, Corp. Boston, Mass. Private Communication with Mr. Allen Kirkpatrick.)

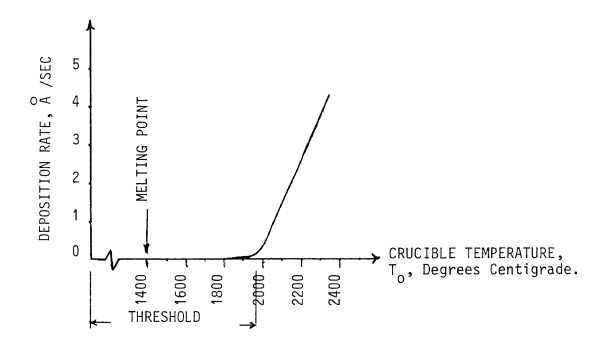


fig. 2.5 Dependence of the deposition rate of silicon on the crucible temperature.

#### 2.6 Ionizing the Clusters

A proportion of the cluster flux is ionized by crossing it with an electron beam as shown schematically in fig. 2.6. The mechanism of ionization is by electron bombardment, which knock off electrons from the cluster, rather than by electron capture, so the ionized clusters are expected to have a positive charge.

The ionization ratio (ratio of the ionized clusters' flux to the total flux) depends on: (1) the electron current for ionization,  $I_e$ ; (2) the voltage applied to accelerate the electrons,  $V_e$ ; and (3) the cluster cross section for ionization.

## The Electron Current for Ionization, Ie

The ionization ratio can be increased from zero to about 50% by increasing the electron current for ionization from zero to 500 mA. (38,55)

 $\rm I_e$  can be increased by increasing the current through the electron emitter (filament), thereby increasing its temperature. This can be achieved by increasing  $\rm V_F$ .  $\rm I_e$  can also be increased by increasing the electron accelerating voltage  $\rm V_e$ .

# Electron Accelerating Voltage, Ve

 ${\rm V}_{\rm e}$  must be high enough to accelerate the electrons to energies well above the ionization energy of the clusters. This is necessary to insure that a cluster will be ionized

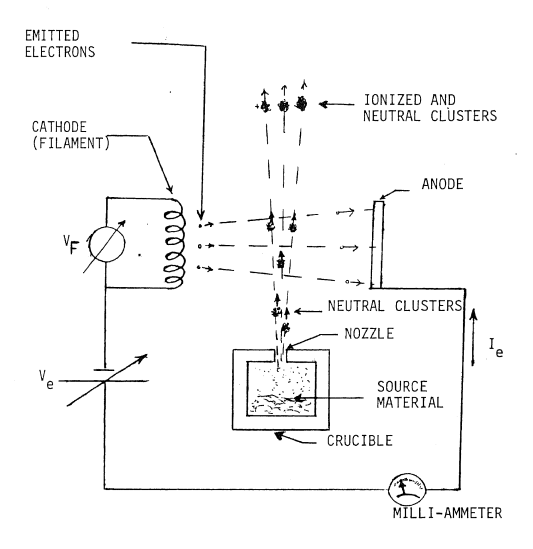


fig. 2.6 Conceptual schematic diagram of the ionization part of the ICB system.

once it is in the path of an accelerated electron.

Fig. 2.7 shows the dependence of ion current collected at the substrate as a function of ionization voltage,  $V_e$ , and ionization current,  $I_e^{\phantom{0}(49)}$ . The figure shows that a minimum voltage of approximately 50 volts is typically required for accelerating the electrons, and that an accelerating voltage of 100 volts is an optimum value at which the ion current (and therefore the ionization ratio) is maximum.

#### The Cluster Cross Section for Ionization

The cross section for ionization is a function of both cluster size (radius) and its ionization energy(i.e., the energy required to free an electron). A smaller ionization energy and a larger cluster radius result in a larger ionization cross section. The ionization energy of a cluster is somewhat lower than the ionization energy of its individual atoms and is usually in the range 5 - 10 eV (49). The radius of a cluster is obviously much larger than that of an individual atom. Therefore the clusters have a much larger cross section for ionization, which make them much easier to ionize compared to individual atoms. This is an advantage compared to ion plating deposition technology.

The ionized clusters are invariably singly charged for two reasons:-

- (1) The probability of multiple electron impact with the same cluster is very low.
- (2) The cluster atoms are only weakly bound together by

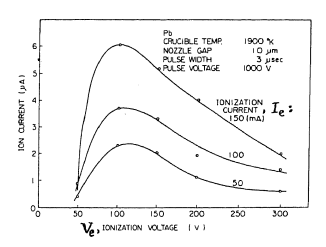


Fig. 2.7 Dependence of the ion current collected at the substrate on the ionization voltage,  $V_{\rm e}$ , and the ionization current,  $I_{\rm e}$ .

van der Waals forces; therefore even if a second charge is created, the strong electrostatic repulsion forces between the two charges will either break up the cluster or result in a charged atom leaving, and hence restoring the singly charged status of the cluster.

### 2.7 Accelerating the Clusters and Energy Distribution

The positively charged clusters can be accelerated towards the substrate by the application of a negative bias,  $V_a$ , as shown in fig. 2.1 . Although high values of several kilovolts are normally used for  $V_a$ , the energy imparted to each atom in the cluster is small and averages  $eV_a/N^{(*)}$ , which is typically in the range of a fraction of an electron volt to a few electron volts. It is within this energy range that an optimum value of energy per atom can be achieved which should be high enough for surface diffusion, but not sufficient for inducing defects in the film layer  $(E \leq 5 \text{ eV})^{(52,55)}$ .

The neutral clusters are not affected by the acceleration voltage and travel to the substrate by their ejection speed alone. It can be assumed  $^{(52,55)}$  that the expansion through the nozzle provides every atom in the vapour with a kinetic energy of the order of  $^k$   $^t$ O, where  $^k$  is the

<sup>\*</sup> e = the electron charge,  $V_a$  = the cluster accelaration voltage and N = the average Number of atoms per cluster.

Boltzmann constant and  $T_{\rm o}$  is the crucible temperature. Therefore the energy per atom in the case of neutral clusters is in the range 0.1 to 0.2 eV depending on the crucible temperature (usually  $1200^{\rm O}{\rm K}$  to  $2500^{\rm O}{\rm K}$ ).

#### 2.8 Film Growth Mechanism

Upon impact with the substrate (or film surface), the clusters break up into their constituent atoms. The velocity direction of the atoms changes from the bombardment direction to directions along the substrate surface as a result of multiple collisions as the cluster impacts the surface. The change in the direction of motion of the atoms increases the migration of atoms and their diffusion along the substrate surface. The long migration distance is one of the important characteristics of ICBD. Other effects are also present: implantation, sputtering of the substrate material and sputtering of freshly deposited material on the substrate. These effects are shown in fig. 2.8 (41).

In discussing factors that influence film growth in the ICB technique, it is convenient to consider separately the effects of:

- (1) The kinetic energy of the clusters, which is easily controlled by the acceleration voltage.
- (2) The presence of ions, which can be controlled from a few percent to about 50% by varying the electron current for ionization from zero to 500 mA.

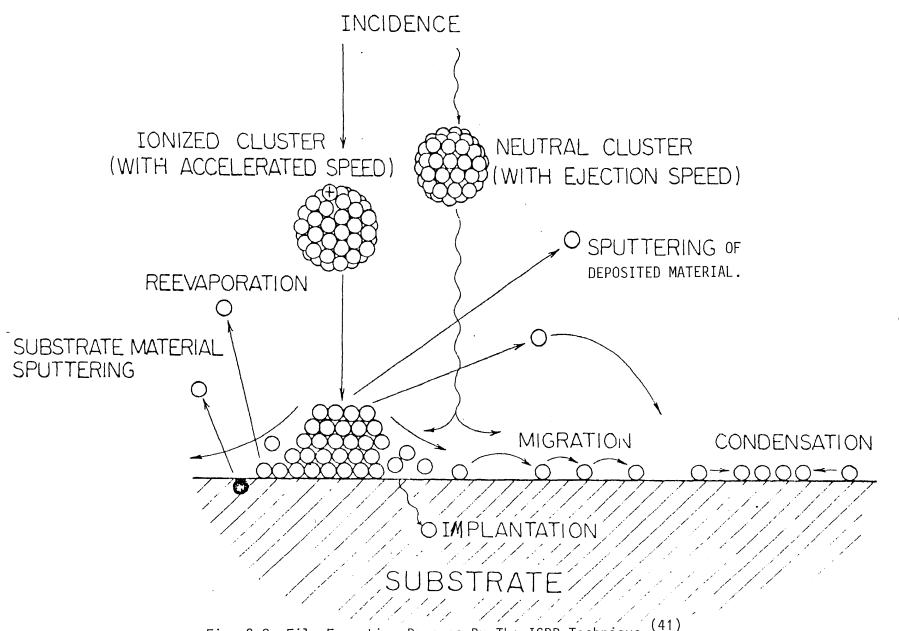


Fig. 2.8 Film Formation Process By The ICBD Technique. (41)

# 2.9 Effects of Kinetic Energy (56)

The kinetic energy of the accelerated clusters are converted upon impact to:-

- (1) sputtering energy, (2) implantation energy,
- (3) hermal energy(surface heating), and (4) migration energy of adatoms.

#### Sputtering and Implantation Effects

These are caused by the high energy tail of the energy spectrum of the flux arriving at the substrate, and result in surface cleaning and deep etching, thereby improving the adhesion strength of the deposited film. The sputtering effect can also enhance the creation of activated centers such as defects and displacement of surface atoms. These act as centers of nucleus formation during the initial stages of film formation (40). Furthermore, ion implantation and the blending of sputtered substrate material with the deposited material enhance the formation of an interfacial layer, which also contributes to strong adhesion of the deposited film.

Fig. 2.9 shows the adhesion strength of Cu films deposited on glass substrates with different values of acceleration voltage (45,46,50,57). It is clear that the adhesion strength is improved by more than an order of fmagnitude as the acceleration voltage is increased up to

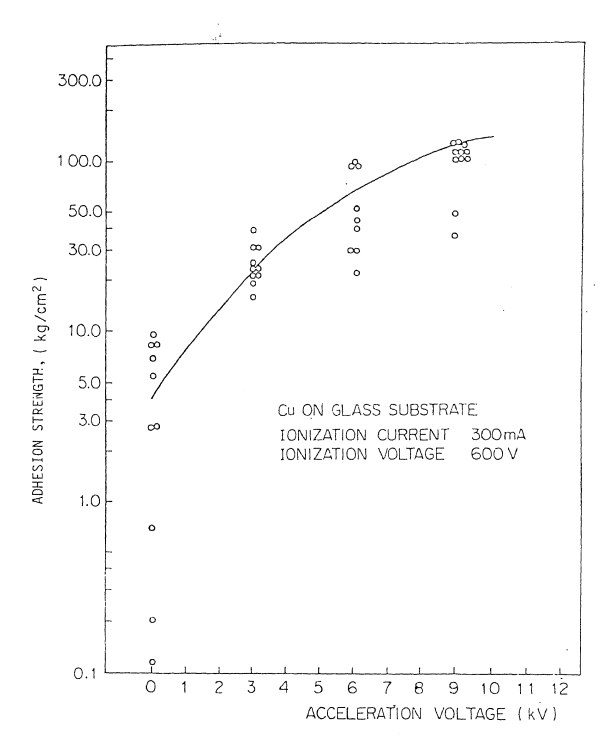


Fig. 2.9 Adhesion strength of Cu films deposited on glass as a function  $\rho f$  acceleration voltage. (45,46,50,57)

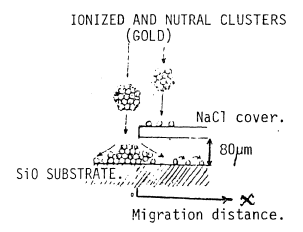
10 kv, and is improved by a factor of 100 over that obtained by conventional vacuum deposition (about 0.5  $kg/cm^2$ ).

#### Thermal Effects

Thermal energy converted from the kinetic energy of the bombarding particles is available at the surface of the substrate and the deposited layer. This allows a high surface temperature to be obtained with relatively lower bulk heating as compared to that required for conventional vacuum deposition. The increased temperature can bring about a change in morphology, and, in the case of R-ICB, an increase in the chemical reaction rates.

#### Migration Effects

The increase of migration of deposited particles with acceleration voltage was demonstrated and reported by T. Takagi's research group in 1976 (50). Gold was deposited onto silicon oxide substrates both by ICBD and by conventional vacuum deposition. The substrate surface was partially covered with a cleaved NaCl mask as shown in fig. 2.10a. The time of the deposition run was made short enough to observe the initial stages of the deposit (island formation). Electron micrographs of the resulting films near the edge of the mask are reproduced in fig. 2.10b. The results clearly demonstrate the increase of migration distance along the surface with increasing acceleration voltage. It was also found that even with zero accelerating



( a )

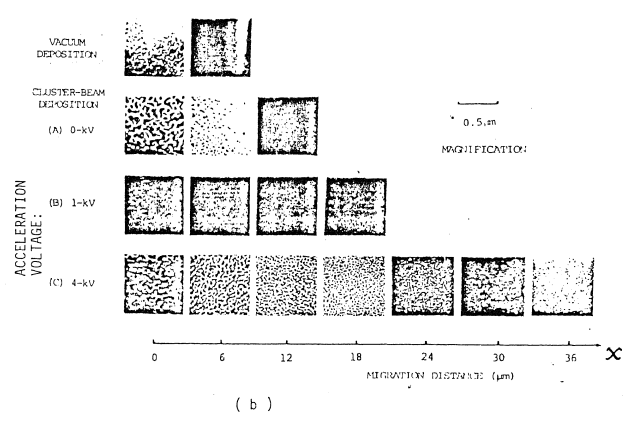


Fig. 2.10 (a) Set up for investigating the migration eafects.

(b) Electron micrographs of films formed by migrated deposits.

voltage, the migration distance is greater for ICBD than that for vacuum deposition by thermal evaporation. This can be attributed to the fact that the kinetic energy of atoms is greater in the ICB technique as a result of the ejection speed at the end of the adiabatic expansion process. The extra kinetic energy of adatoms promotes chemical reaction at the substrate surface, which is important in the case of Reactive ICB deposition.

#### 2.10 Effect of Presence of Ions

The presence of ions, even in small proportion compared to the total flux, has been found to have a profound effect on the quality of films produced by ICBD, independently of the effects of kinetic energy of the clusters. Attempts to explain how the presence of ions influence the critical condensation process in terms of the nucleation, growth of nuclei and coalescence have been made by K.L. Chopra (4,5). For our purposes, it is instructive to consider here the results reported by T.Takagi's group (43) on the deposition of ZnO films onto glass substrates using the R-ICB method. Figure 2.11 shows RHEED patterns of the films deposited at different electron currents for ionization of zero, and 300 mA. At these levels, the percentages of ionized clusters in the total flux are 0, 8 - 10%, and 25 - 30% respectively. Other operating conditions were kept constant as follows: Acceleration voltage, zero volts; Po (vapour

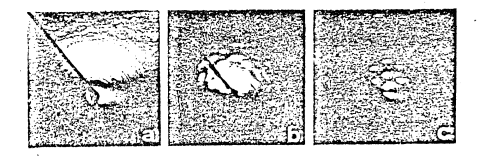


Fig. 2.11 RHEED patterns showing the influence of the electron current for ionization,  $I_{\rm e}$ , on the crystallinity of ZnO films deposited by the R-ICBD technique. (43)

(a) 
$$I_e = 0$$
, (b)  $I_e = 150 \text{ mA}$ , (c)  $I_e = 300 \text{ mA}$ 

pressure of zinc in the crucible), 0.1 torr; and P(chamber, oxygen),  $5 \times 10^{-4}$  torr. It was observed that a good quality film - having a preferred orientation, with the wurtzite c-axis perpendicular to the substrate surface - develops as the ionization current is increased.

Y. Namba and M. Toshio investigated the epitaxial growth of ZnTe on NaCl by ion beam techniques (20). They found that the epitaxial growth was notably favoured by the presence of ions in the particle flux for deposition, while no distinct dependence on the ion energy could be observed. They also found that once the first few hundred Angstroms or so are deposited, the epitaxial growth continues even when the ion content of the particles is reduced to zero. This indicates that the presence of ions plays its most important role in the initial stages of heteroepitaxial growth.

Enhanced chemical reaction activity resulting from the presence of ions is one of the principal reasons for the effectiveness of the reactive ion cluster beam method for film formation. The method has been successfully used for preparing films of compound materials such as  ${\rm ZnO}$ ,  ${\rm BeO}$ ,  ${\rm SnO}_2^{(55)}$  and nitrides such as  ${\rm GaN}^{(17,18)}$ .

# 2.11 Effect on Substrate Temperature Required for Epitaxial Growth

Epitaxial growth of films deposited by ICB has been reported to be possible at lower substrate temperatures than

that required by alternative methods  $^{(39,56)}$ . Fig. 2.12 shows the mass of Si deposited as a function of the reciprocal substrate temperature for different values of acceleration voltage and keeping the other variables constant as indicated on the graph. Superimposed on these curves are lines for  $T_{(s\leftarrow p)}$ , the transition temperature from polycrystalline to single crystal state, and  $T_{(p\leftarrow a)}$ , the transition temperature from the amorphous to the polycrystalline state. Two effects are evident in this graph:

- (1) The increase in substrate temperature with the mass deposited is greater as the voltage is increased. This is obviously due to the increased kinetic energy of the clusters, part of which is converted to thermal energy upon impact with the substrate as discussed earlier.
- (2)  $T_{(s \leftarrow p)}$  and  $T_{(p \leftarrow a)}$  decrease as the acceleration voltage is increased. This is due to the fact that the substrate temperature required for epitaxial growth is lower for higher acceleration voltages. This suggests the possibility of growing epitaxial films at low substrate temperatures, resulting in films with reduced amounts of strain, and hence, improved quality (12).

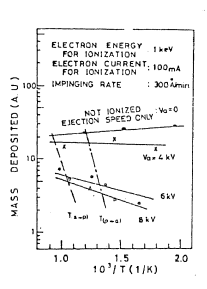


Fig. 2.12 Mass deposited vs. reciprocal substrate temperature, and the change of transition temperature for different acceleration voltages. (39,56)

#### CHAPTER III

## CONSTRUCTION OF THE ICB SYSTEM

An experimental ICB system similar to the one used by Takagi research group was designed and constructed in the Microelectronics Laboratory at NJIT. Machine shop drawings were made by Fred Solomon, an undergraduate E.E. student, and the parts were made locally in the NJIT machine shop. The complete machine shop drawings are included in Appendix A.

The system was then assembled by the writer in the chamber of a Veeco Instrument Inc. oil diffusion pump vacuum system. Variable power supplies required (minimum of 4) were arranged as well as cooling water connections, thermocouple temperature monitors, voltmeters, ammeters and nitrogen gas inlet and accessories. In the following paragraphs a detailed description of the ICB system constructed will be given.

#### 3.1 Complete System

The complete ICB system is shown in fig. 3.1. Source material vapour is generated in the nozzle assembly  $(a_1)$  and ejected into the ionization region  $(a_2)$ . The vapour condenses in the process of adiabatic expansion and clusters are formed. The clusters are ionized by electron bombardment in the ionization region. The ionized clusters are accelerated by

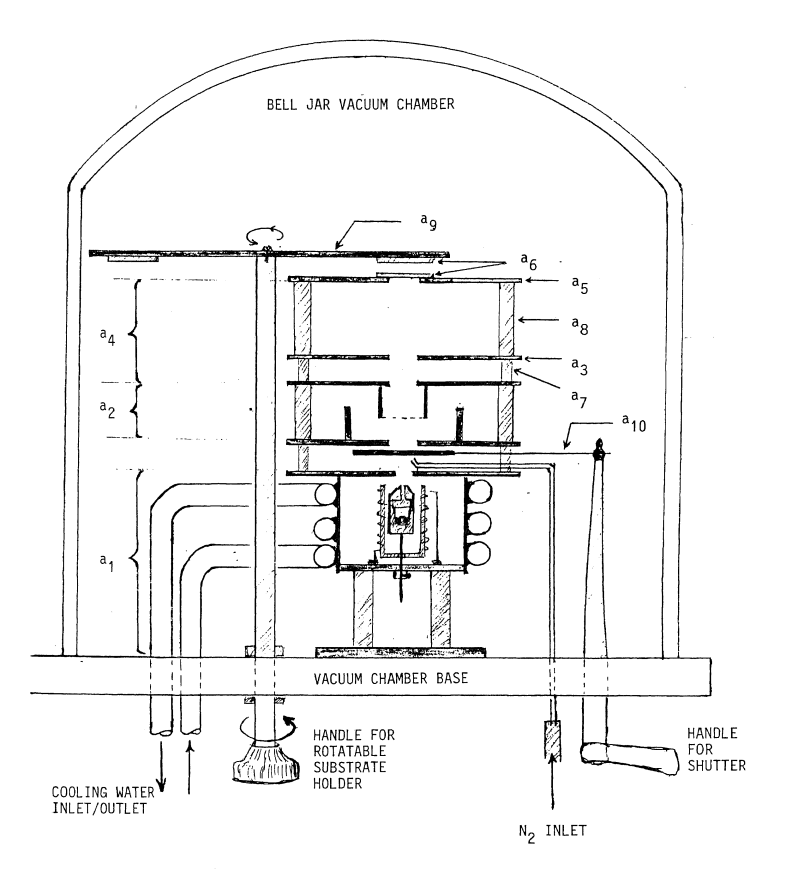


Fig. 3.1 General Assembly Drawing of the Ionized Cluster Beam (ICB) System.

the accelerating electrode  $(a_3)$  in the acceleration region  $(a_4)$  and land on the substrate  $(a_6)$  together with the neutral clusters.

#### 3.2 The Nozzle Assembly

The cluster beam generation part of the system is shown in fig. 3.2. Material to be deposited is placed in the crucible(1) and confined by the lid(2) which is designed to push-fit tightly on the crucible. The lid has a 0.5 mm diameter nozzle(3) at the center to allow source material vapour to be ejected into the high vacuum region at a high velocity, but at the same time allow the pressure in the crucible,  $P_0$ , to be at least  $10^3$  times the pressure outside the crucible. The crucible is heated by placing it in a heater (4) which in turn is directly heated by a filament(5) wrappedaround it as shown in fig. 3.3. The crucible is supported in the heater by a thin tungsten rod(6) fitted to its base. The supporting rod passes through a hole in the base of the heater and is fixed against the stainless steel baseplate by screw(7), thus allowing for the adjustment of the vertical position of the crucible inside the heater. Both the crucible and the heater are made of boron nitride material, and a 0.023" diameter tungsten wire is used as the filament material. A O - 130 volt, 45 Amps Bowerstat\*

This rating for the powerstat is used because of its availability. Actually, a 0 - 80 volts, 30 Amps powerstat would suffice.

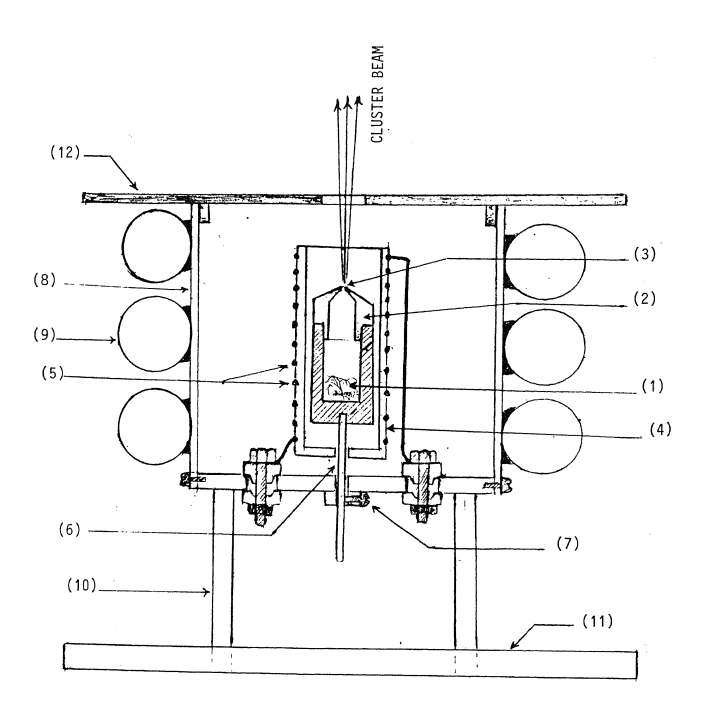


fig. 3.2 Detailed section of the Nozzle Assembly.

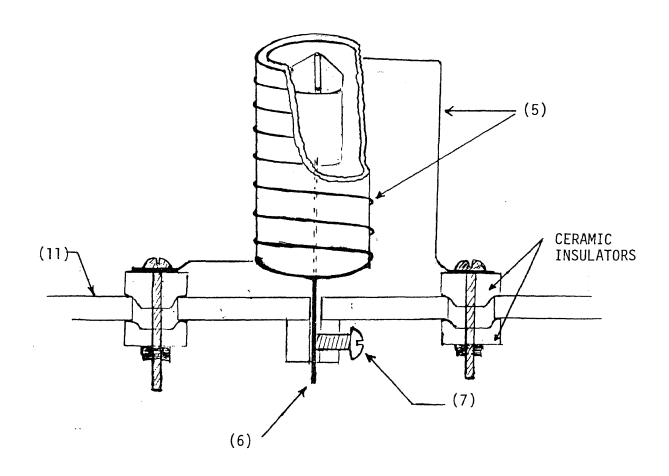


fig. 3.3 Drawing of the crucible supported inside the crucible heater.

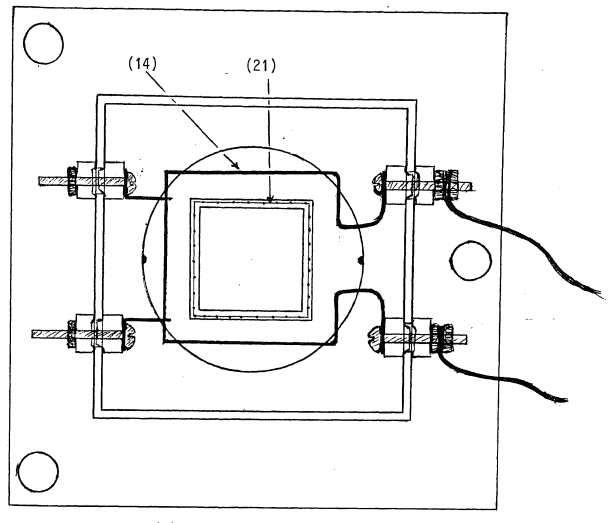
(variable autotransformer) is used to drive a high ac current through the filament, and by controlling this current, the crucible temperature can be controlled.

The assembly is surrounded by a stainless steel "box" which acts as a heat shield(8). Cooling water is circulated through copper cooling coils(9) welded on around the heat shield. The generation part of the system is supported by four 63-mm long ceramic insulators(10) fixed to the system baseplate(11). The top plate(12) acts as a baseplate to support the next stage of the ICB system, namely the ionization assembly.

#### 3.3 The Ionization Assembly

The ionizing electrode assembly is shown in fig. 3.4. It rests on the top plate of the nozzle assembly supported by three 25-mm ceramic insulators(13). The electron emitter (cathode) consists of a square loop of 0.019" tungsten wire (14) supported by four hook shaped thicker tungsten wire(15) fixed to the cathode supporting plate(16) by stainless steel screws(17) through ceramic insulators(18). The cathode leads are also terminated around stainless steel screws(19), which are brought out through ceramic insulators(20) for power connections. The cathode is heated for thermionic emission of electrons by passing a high current through it. A 0 - 130 volt, 45 Amps Fowerstat\* is used to supply this

<sup>\*</sup>This rating for the owerstat is used because of availability. Actually a O - 30 volt, 30 Amps powerstat would suffice.



(a) Top view.

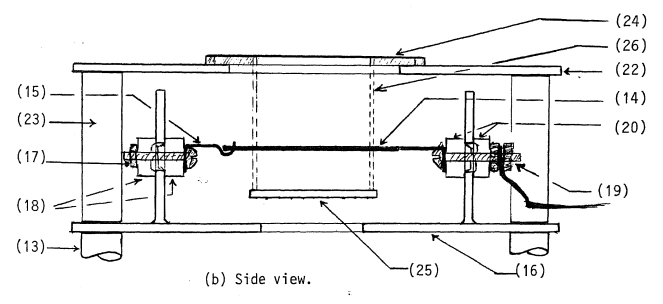


Fig. 3.4 Top and side view of the Ionization Assembly of the ICB system.

current.

The anode (fig. 3.5) is made of two stainless steel plates (24),(25) connected by 0.019" tantalum wire mesh(26) shaped like a hollow parallelopiped (21 in fig. 3.4) with a square cross section that fits at the center of the square-shaped cathode filament. The anode assembly is suspended from the anode supporting plate(22) which in turn rests on the cathode supporting plate(16), supported by three 50-mm ceramic insulators(23). The anode is biased positively with respect to the cathode to accelerate the thermally emitted electrons towards the center, where the clusters are passing. The wiring diagram for the ICB system is shown in fig. 3.6.

## 3.4 The Accelerating Electrode and the Substrate Holder

The accelerating electrode (a<sub>3</sub> in fig. 3.1) rests on the ionization assembly(a<sub>2</sub>) supported by three 25-mm ceramic insulators(a<sub>7</sub>). The fixed substrate holder(a<sub>5</sub>) rests in turn on the accelerating electrode supported by three 63-mm ceramic insulators (a<sub>8</sub>). An alternative rotatable substrate holder (a<sub>9</sub>) was also made, that could hold up to six substrates arranged in a circle, and any one substrate could be rotated externally into position to receive the cluster beam. In this manner, it would be possible (at least theoretically) to deposit on six substrates under six different sets of deposition parameters in the same pumpdown. A shutter (a<sub>10</sub>) was also included that could be controlled externally to block or allow the deposited material flux to reach the substrate.

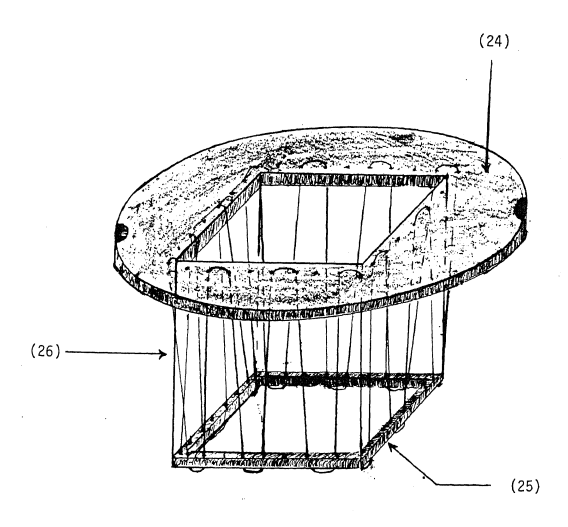


Fig 3.5 Three dimensional view of the Anode of the Ionization Assembly.

#### 3.5 Electrical (Power Supplies and Meters)

The complete wiring diagram for the system is shown in fig. 3.6. Three AC variable power supplies are used for supplying 0 -30 Amp currents through the crucible heater, the substrate heater and the ionization assembly cathode, as indicated on the diagram. Two high voltage DC supplies are also needed: V, the electron accelerating voltage (zero to 1000 volts), and  $V_a$ , the ion cluster accelerating voltage (O - 10 kV). A simple autotransformer (O - 130 volts, 45 Amps) is used for  $V_{\rm H}$ , but isolating transformers I.T.1 and I.T.2 are necessary for  $V_{\mathbf{F}}$  and  $V_{\mathbf{S}}$  respectively to provide DC isolation between the high voltage biases ( $V_{\rm p}$  and  $V_{\rm a}$ ) and the mains power supplies. The range of variation for each power supply and meter as well as ratings for the components used are all indicated on the wiring diagram(fig. 3.6). Internal wiring diagrams for the variable DC power supplies (V and V<sub>a</sub>) are included in Appendix B.

#### 3.6 Accessories

#### Substrate Heater

Originally, two tungsten filaments mounted on the substrate holder (at the back of the substrate) were used for substrate heating as shown in fig. 3.7. However, as this arrangement was not capable of raising the substrate temperature beyond 350°C, alternative substrate heaters were developed and used as described in chapter 4.

Fig. 3.6 Complete electrical wiring diagram of the ICB system. (Super-imposed on the outline of the system for easier identification).

I.T.2, 240: 6

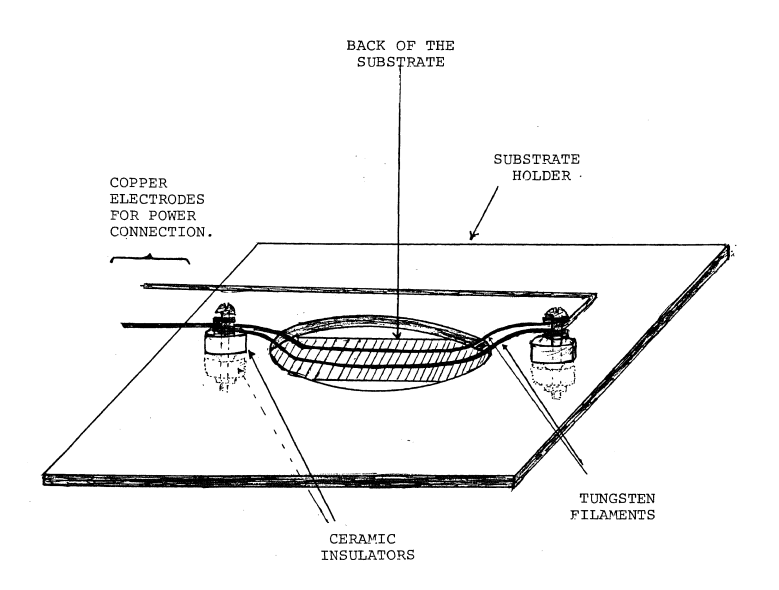


Fig. 3.7 Substrate heating arrangement used in runs No. 2 through No.6.

#### Temperature Sensors

Two type-K thermocouple sensors were included for monitoring the crucible and the substrate temperatures, respectively These were used in conjuction with a model 871 digital thermometer made Omega Engineering Inc., Stamford, Conn., which could read temperatures ranging from -40°C to 1,370°C with 1°C resolution. A battery-operated unit were used and the thermocouple leads were brought out through a high-voltage vacuum lead-through (properly insulated from the vacuum chamber base) to have the option of allowing the thermocouples near (or touching) the substrate, even when it is biased at a high potential of several kilovolts.

## Power Inlet(VacuumLead-through) Terminals

Eight vacuumlead-throughs were used for providing a vacuum-tight means of conveying the electrical power to the system through the base of the vacuum chamber. These were type 6D made by Edward Scientific (fig. 3.8a) with a rating of 10 kV, up to 150 Amps, and up to 150 C operating temperature. The diameter of thelead throughs had to be machined down to 0.992" at one point (fig. 3.8b) to fit the 1" diameter standard openings of the vacuum chamber base.

#### Nitrogen Gas Supply

A source of nitrogen supply to the vacuum chamber was needed for two reasons.

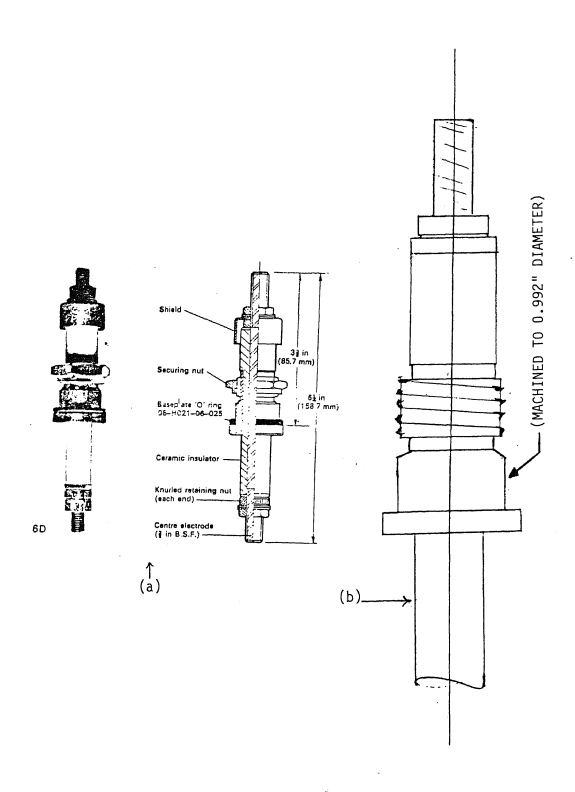


fig. 3.8 Type 6D (Edward Scientific) vacuum leadthrough used for supplying power to the ICB system.

- (1). For the system in general, to fill the vacuum chamber with  ${\rm N}_2$  after shutdown and thus allow the system to cool in an inert atmosphere.
- (2). For GaN deposition in particular, to maintain a nitrogen atmosphere of  $1-5 \times 10^{-4}$  torr during gallium deposition and thus form gallium nitride by the Reactive Ion Cluster Beam Deposition (R-ICBD) technique.

Nitrogen was piped from the  $\rm N_2$  tank through a regulator and then a leak valve to the vacuum chamber, and was terminated with a 1/8" stainless steel tube at a point (fig. 3.1) midway between the nozzle assembly and the ionization assembly. The  $\rm N_2$  outlet nozzle was made pointing upwards and as close to the crucible nozzle opening as possible without obstructing the path of the flux coming out of the crucible.

### Cooling Water Inlet/Outlet

A twin pipe, 1" vacuum feedthrough was used for providing a vacuum tight means of connecting the cooling water in and out of the vacuum chamber. The 3/8" steel tubes of the vacuum feedthrough were connected to the heat-shield cooling tubes inside the vacuum chamber using brass compression fittings. Rubber hoses were used outside the vacuum chamber for connecting the feedthrough tubes to the water supply and drain lines respectively.

# 3.7 Setting The Electron Current For Ionization

Normally, in an Ion Cluster Beam System, to set the

electron current for ionization,  $I_e$ , at a desired level (say 300 mA) at a certain value of electron accelerating voltage,  $V_e$ , (say 500 volts), one would normally increase  $V_e$  to 500 volts and then raise the cathode filament voltage,  $V_F$ , until the required  $I_e$  of 300mA is reached. However, since the DC power supply used in our system for  $V_e$  has a sharp drop in output voltage( $V_e$ ) as the output current ( $I_e$ ) is increased, one has to set  $V_e$  at a much higher value initially, so that it drops at the specified load of 300 mA to the desired value of 500 volts.

Fig. 3.9a shows the dependence of the supply voltage output ( $V_e$ ) on the output current ( $I_e$ ) for different values of no-load voltage setting. Fig. 3.9b shows typical  $I_e$  versus  $V_e$  characteristics for the ICB system for different values of filament voltage,  $V_F$ . The  $45^{\rm O}$  lines ( $V_e$  Vs.  $V_e$  and  $I_e$  vs.  $I_e$ ) are added to facilitate reflecting points between the two curves. The procedure for setting  $I_e$  at 300 mA with  $V_e$  at 500 volts is then as follows:-

- A Increase V to a higher value, say 600 volts.
- B Increase the filament voltage,  $V_F$ .  $I_e$  will start to increase and  $V_e$  decreases.
- C Repeat A and B above until the desired value of  $I_{\rm e}$  and  $V_{\rm e}$  is reached.

It should be noted that although it is possible (theor tically) to plot the values of  $I_{\rm e}$  and  $V_{\rm e}$  on fig. 3.9a and determine the no-load setting for  $V_{\rm e}$  from the same figure and

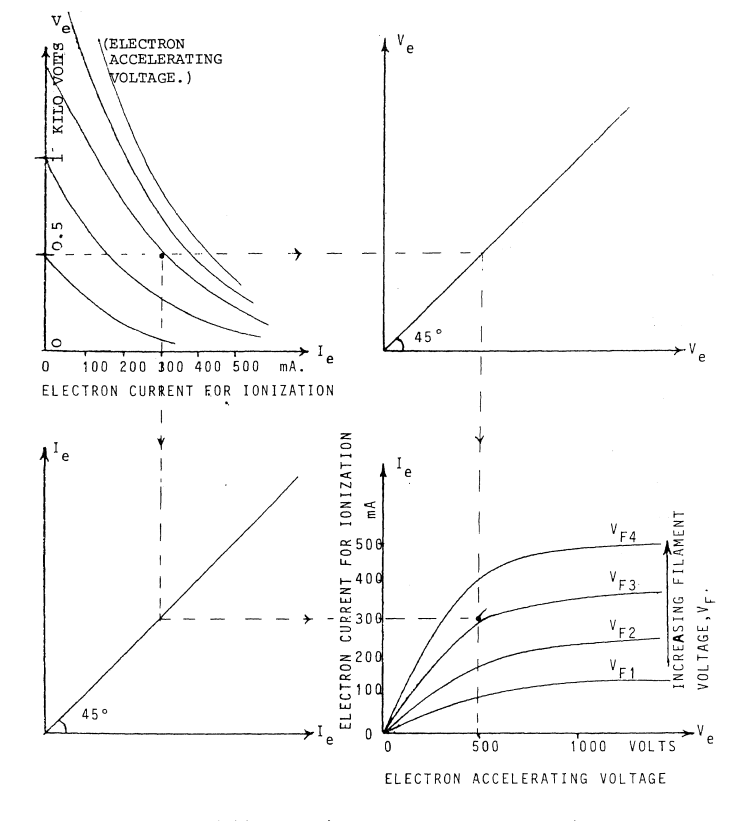


Fig. 3.9 (a)(Top left) Output characteristics (voltage vs. output current ) for the power supply used for  $V_e$ . (b)(Bottom right) Typical  $I_e$  vs.  $V_e$  characteristics for the ICB system for different values of filament voltage, $V_F$ .

then reflect the operating point onto fig. 3.9b and determine the filament voltage,  $V_{\rm F}$ , required this procedure is not followed for two reason.

- (1) The characteristics of the ICB system shown in fig. 3.9b) are only typical and depend on many additional parameters such as time, whether  $V_{\rm F}$  is being increased or decreased, and the temperature of the nearby crucible and substrate heaters which influence the cathode filament temperature by radiation.
- (2) The no-load value of  $V_e$  required for initial setting is considerably higher than the actual operating value for  $V_e$  and might be high enough to cause breakdown between the cathode and anode especially in the presence of small amounts of residual gas as is the case in the Reactive Ion Cluster Beam Deposition (R-ICBD).

These problems are avoided by following the iterative procedure described above.

### 3.8 Typical Start Up Procedure

- 1. Charge the crucible with the source material to be deposited, put the substrate(s) in, and assemble the complete ICB system in the vacuum chamber.
- 2. Turn on the cooling water lines.
- 3. Pump down for at least one hour in the 10<sup>-6</sup> torr range.
- 4. Raise the crucible temperature (with the shutter in place) up to the required value and stabilize the temperature to within a few degrees for at least 15 minutes.

- 5. Raise the substrate temperature to the desired value.
- 6. Set the ionization current at the desired level by following the procedure described in the previous section.
- 7. Apply the desired ion accelerating voltage,  $V_a$ .
- 8. Move the shutter aside to start depositing.

## 3.9 Typical Shut Down Procedure:-

- 1. Move the shutter back in place.
- 2. Reduce the ion accelerating voltage,  $V_a$ , to zero.
- 3. Reduce the electron accelerating voltage,  $V_{\rho}$ , to zero.
- 4. Reduce the cathode filament voltage,  $V_F$ , to zero.
- 5. Reduce the substrate heater and the crucible heater voltages to zero.
- 6. Introduce nitrogen in the vacuum chamber and allow the system to cool down for at least two hours before opening up the vacuum chamber.
- 7. Turn off the cooling water lines.

In addition to the above, the normal steps for start up and shut-down of standard bell jar vacuum systems should of course be followed. A detailed description of the procedure adopted in our experiments for the deposition of GaN by R-ICB is given in section 4.14 .

### 3.10 Practical Difficulties

In any experimental work - such as that reported in this thesis - many practical problems arise during the course of the experiments. In this section, a few of the more important difficulties encountered are cited, along with the solutions employed (or proposed) for each case:-

- 1. Substrate heating presented many problems. It necessitated abandoning the multiple substrate holder as discussed in section 4.7 . Modified substrate heaters had to be constructed to meet the higher temperatures ( $\Delta450^{\circ}$ C) required as discussed in sections 4.13 and 4.16 .
- 2. After every deposition run, the crucible lid would seal tightly on the crucible. Removing the lid to charge the crucible required heating the crucible e.g. using a blower or a hair dryer to melt the gallium inside it. Then the crucible would be held in a three-jaw grid(in the NJIT machine shop) and the lid turned slowly with pliers cushioned with a clean napkin or tissue paper.
- 3. After every deposition run, the top plate of the nozzle assembly (part 12 in fig. 3.2) would wrap, due to the inevitable uneven heating of the plate. This resulted in breaking of the tips of the three insulators

resting on this plate and supporting the ionization stage. The problem was partially solved by using larger diameter insulators supported by bigger screws. This eliminated the breakage problem, but the warping of the plate still resulted in tilting the top part of the ICB system slightly as the temperature was increased. To eliminate this problem in a second generation system, water cooling of this plate should be considered.

The screws feeding current to the crucible heater filament (fig. 3.3) were getting extremely hot and on occasions have been observed to melt where they touch the tungsten filament. This can be attributed to the very high temperature of the filament coupled with the relatively high thermal resistance of the stainless steel screws used. The problem could have been solved by using larger diameter screws having a lower thermal resistance. However, since larger screws required other modifications to accomodate them, it was suggested<sup>(25)</sup> doubling up the tungsten filament a short length near the screws. This lowered the equivalent resistance of this section of the filament touching the screws and hence reduced the heat generated nearer to the screws. This way the temperature of the screws was lowered and melting avoided.

- 5. After each of the first few runs, it was observed that the various tungsten filaments in the system were becoming brittle and difficult to handle. This was caused by the rapid cooling of the filaments after shutdown. To avoid this, a more elaborate shutdown procedure was adopted (section 4.15) which incorporated among other things slowly reducing the filament temperature instead of abruptly switching the supplies off.
- 6. In addition to the above, water vapour contamination presented a major difficulty. This is fully discussed in part C of section 4.20 at the end of Chapter IV.

#### CHAPTER IV

# GROWTH OF GAN FILMS BY R-ICBD

The ICB system constructed at the NJIT Micro-electronics Laboratory was used to grow GaN films. In this chapter, the details of the experiments performed are given along with discussions of the results obtained.

# A. Gallium Nitride

### 4.1 Why Choose Gan?

Gallium Nitride is a wide bandgap material which can be an efficient light emitter over a wide a spectral range including the visible (10,22,26) and the ultraviolet (24,27). It has also been made to lase in the ultraviolet region (7), and has exhibited negative electron affinity (15,23). In particular, blue-green luminescence has been observed at 5190Å with a power efficiency of 0.1% and a quantum efficiency of 1%, with no evidence of degradation over extended periods of continuous operation (28). Therefore, GaN promises a new class of numeric displays generating previously unattainable colors with highly acceptable brightness, and several additional advantages:-

- (1) The luminous area is determined by the shape of the metalization, and all the electrodes are attached at the rear surface of the crystal.
- (2) GaN is perfectly transparent throughout the visible range so that the emitted radiation can be viewed through

the crystal. Furthermore, the crystal can be a convenient support and window for LED's emitting other colors, thus permitting the fabrication of a multicolor display (29).

Thus far, the major stumbling block to the widespread use of GaN LED's has been the difficulty in growing large, homogeneous, strain free, single crystals.

# 4.2 Material Synthesis (27)

Several growth techniques have been described in the leterature for the preparation of GaN. One technique involves the flow of ammonia over gallium at a temperature greater than  $800^{\circ}$ C as follows:

$$Ga + NH_3 \longrightarrow GaN + \frac{3}{2}H_2$$

Some of the small crystallites formed by the above reaction can be made to grow at the expense of decomposing neighboring ones by heating in flowing ammonia, and thus millimeter-size crystals can be obtained.

Another technique which has enjoyed the greatest favor among researchers is a vapour transport method known as the Maruska and Tietjen method  $^{(16,28)}$ .  $GaCl_3$  is produced by passing HCl over Ga in an open flow furnace.  $GaCl_3$  (vapour) is then mixed with ammonia gas, near the surface of a sapphire substrate (at around  $1000^{\circ}$ C) on which GaN forms as a single crystal:

2 Ga(metal) + 6 HCl(gas) 
$$\longrightarrow$$
 2 GaCl<sub>3</sub>(gas) + 3H<sub>2</sub>(gas)  
GaCl<sub>3</sub>(gas) + NH<sub>3</sub>(gas)  $\longrightarrow$  3HCl(gas) + GaN(deposits on the substrate)

A third method for growing an extended area of thin GaN film is reacting Ga vapour with ionized itrogen (13). This method is the one being investigated in the present work and will be described in greater detail in the following section.

# 4.3 GaN By R-ICB Deposition

K. Matsubara and T. Takagi recently (1980) reported successful results in their attempts to grow GaN films using R-ICBD (17,18). Pure Gallium was vapourized from a crucible with a nozzle diameter of 1 mm, in a Nitrogen atmosphere. The substrate used was hexagonal ZnO film on glass. Other conditions of the deposition were reported to be as follows:

- Crucible (Gallium) Temperature: About 900°C
- Electron Current for Ionization: 300 mA.

  (At this current level, the ionization ratio of the clusters is about 25 30%).
- Chamber (Nitrogen) pressure: 5 x 10<sup>-4</sup> torr.
- Substrate temperature: 450°C.
- Acceleration voltage: Zero.

Under these conditions, it was reported that n type GaN was grown epitaxially on a ZnO-film surface.

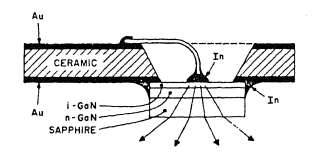
# 4.4 Important Properties of GaN Films -

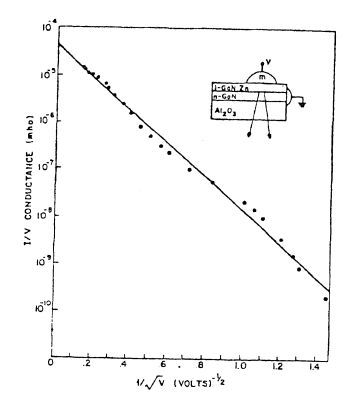
GaN is almost always n-type when undoped, with the free electron concentration usually about  $2 \times 10^{18}$  electrons/cm<sup>3</sup> and an electron carrier mobility of about 130 cm<sup>2</sup>/V.sec. The donors (which are believed to be native donors, i.e.

Nitrogen vacancies) can be compensated by the addition of Zn, Mg, Li or Be. When the material is compensated, it becomes semi-insulating, but not p-type. So far, it has not been possible to make the material p-type. The energy band gap of GaN is direct, about 3.4 eV at room temperature. Reference 26 is an excellent summary of the known properties of GaN including detailed band structure, absorption and reflectance spectra as well as their temperature-dependence.

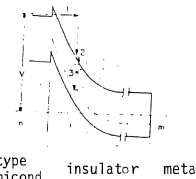
# Electroluminescence (27)

GaN light-emitting diodes have been fabricated using the MIS (metal - insulator - semiconductor) structure shown in fig. 4.1 . An undoped n type conduction layer is grown on a transparant substrate, usually sapphire. Then a Zn-doped insulating layer is grown on the n-type layer. The wafer is cut into 1-mm diameter discs which are then soldered with indium to a ceramic support. This peripheral solder makes an ohmic connection to the n-type layer. A small indium dot soldered at the center of the insulating layer forms a  $10^{-3}$ cm<sup>3</sup> area of non-ohmic connection. Such a diode can, depending on the degree of doping, emit blue, green, yellow or red light. The light is emitted at the negative electrode (cathode). I - V characteristics have been observed to exhibit a linear relationship between ln(I/V) and  $l\sqrt{V}$ . This suggests that tunneling rather than space charge limitation is the mechanism controlling the current as shown in fig. 4.2 (30). On the





(a)



n-type meta1 semicond.

Fowler - Nordheim - like plot of the I-V characteristics for a typical Zn-doped GaN diode with the metal (m) biased positively. (a) fig. 4.2

- Band diagram for the GaN diode under the m-positive bias condition. The proposed model for the process is: (1) Tunneling; (2) Impact excitation (direct case); (3) Radiative recombination.(30)

other hand, Matsubara and Takagi  $^{(18)}$  reported that I-V characteristics of GaN LED's made by ICBD follow the relation  $(I \not \sim V^n)$  with n ranging from 1 to 3. These characteristics suggested space charge limitations as the current limiting mechanism. It was also reported that the light was emitted near the positive biased contact, in direct contradiction with the results reported by J.I.Pankove  $^{(30)}$ 

# B. EXPERIMENTAL DETAILS

#### 4.5. INTRODUCTION

As a first step in exploring the prospects of growing GaN films by R-ICBD, it was decided to attempt to reproduce the results reported by the Takagi research group in 1i80 (section 4.3) (17,18). This decision might have been premature, considering that the ICB system constructed had not been tested yet with straightforward metal deposition; and that R-ICB deposition poses additional problems. These problems include among other things:

- (1) The increased possiblility of creating a plasma, and hence destroying the clusters due to plasma heating.
- (2) Sensitivity to the parameters of the deposition due to the requirement that a chemical reaction occur at the surface of the substrate in addition to the requirement that favorable conditions for film formation be attained.

Rather than discuss the best results obtained, it is instructive to scan through the runs made in sequence and

see how the results of each brought about changes in the conditions of the next run. This is necessary because the runs can be viewed as part of a learning curve.

### 4.6 Run No. 1

The first run was made using glass slides at room temperature as substrates. Deposition was made on 5 slides using the multiple substrate holder described earlier. Except for the substrate temperature, other parameters of the deposition run were adjusted to correspond to the optimum results reported the Takagi research group (17,18) as follows:

- Electron Current for Ionization: 300 mA.
- Accelerating Voltage: Zero.
- Nitrogen Pressure:  $1 2 \times 10^{-4}$  torr.

The crucible temperature was 900°C for deposition on the first three slides while the deposition time was made progressively greater for each slide. This meant that theoretically speaking, we should have obtained a thicker film on slide No. 2 than on slide No. 1. Similarly, we should have obtained a thicker film on slide No. 3 than on slide No. 2. This was not the case. The film obtained on slide No. 2 clearly appeared, upon visual examination, to be the thickest of all films, even counting the other slides (No. 4 and No. 5), which were deposited at the higher crucible temperature of 950°C instead of 900°C. It was also observed (when the system was inspected after the run) that gallium

metal had accumulated in the heater and along the rod supporting the crucible as shown in fig. 4.3. It was therefore suspected that the supporting tungsten rod might have punched through the crucible wall, and made a tiny hole in the base which allowed gallium to leak out. To avoid this problem, a new crucible was made to be used for the following runs, and extra care was taken to mount it gently on the tungsten rod without applying any unnecessary force.

### 4.7 Run No. 2

The conditions for the second run were similar to the first, except that substrate-heating was introduced. To do this, the multiple substrate-holder had to be abandoned in favor of a smaller, single substrate holder. This was necessary for several reasons. The large mass and area of the multiple substrate holder would require massive amounts of power to heat the entire plate. The large area would give rise to considerable radiation heat loss, and intolerable outgassing that might affect the performance of the small pumping system being used. There was also the practical difficulty of monitoring the temperature of movable substrates using a stationary thermocouple. For this (and subsequent) runs, quartz substrates were used. After 18 minutes from the start of the deposition, an opaque film started accumulating on the walls of the bell jar. The system was immediately shut down and allowed to cool. After the system was opened

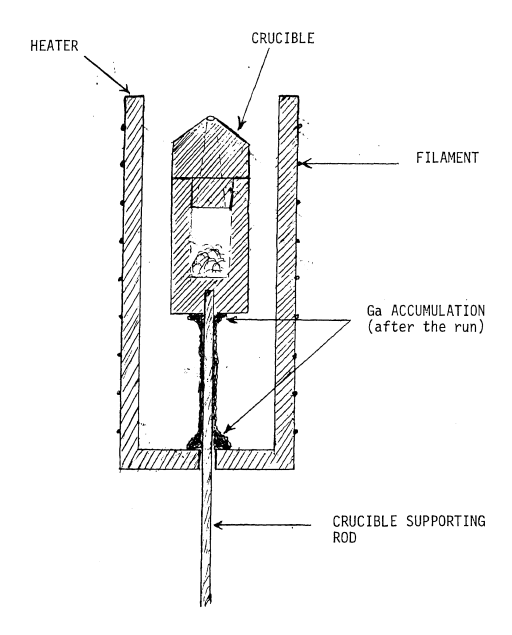


Fig. 4.3 Section of the crucible inside the crucible heater showing the accumulation of Ga after the first deposition run.

and examined, it was observed that the silver solder between the heat shield and the water cooling coils had melted. This could happen only if: (1) The cooling water was lost, e.g. accidentally shut off; or, (2) The pressure of the cooling water was so low that the cooling rate could not catch up with the rate of heat generation. The film obtained was not clean, and not uniform, and it was not carried any further to the measurements stage. The system including the bell jar and its base was thoroughly cleaned with HCl, and then trichloroethylene to remove any silver film that might have contaminated it.

### 4.8 Run No. 3:-

The conditions for this deposition run were set as follows:-

- Crucible Temperature: 1,000°C
- Electron Current for Ionization: 350 mA.
- Nitrogen Pressure in the Chamber:  $1 2 \times 10^{-4}$  torr.
- Substrate Temperature:-
  - initial: 300°C.
  - final: 330°C.

The deposition run was continued for 60 minutes after which the system was shut down and allowed to cool before it was opened. The film obtained was observed to have the following properties:-

(1) Transparent: although it could easily be distinguished from the quartz substrate by its "cloudy" appearance.

- (2) Film growth was not uniform over the substrate surface.

  There were areas of the surface where film growth appeared inhibited.
- (3) The film was soft and could be scratched off the surface of the substrate rather easily.
- (4) Film thickness was estimated to be less than 200 A , meaning that a deposition rate of only  $\approx$  3 A /min was achieved. This is rather low for ICBD.
- (5) Photoluminescence tests made at RCA Labs. in Princeton, N.J. showed no indication of the peaks normally associated with gallium nitride. (These tests will be discussed in greater detail for the film obtained in run No. 6). At this stage, the failure of the photoluminescence tests to reveal a GaN signature was attributed to any one or a combination of the following reasons:
- (a) The film was not GaN, i.e. the substrate temperature was not high enough to enhance the reaction between Ga and  $N_2$ .
- (b) The film was amorphous possibly because of the low substrate temperature.
- (c) The film was too thin to show any photoluminescence peaks.

# 4.9 Run No. 4

The conditions for this run were set similar to that for run No. 3, except that the crucible temperature was raised to 1,080°C, in an attempt to achieve a higher deposition rate. The deposition was allowed for 30 minutes, but no observable

film resulted on the substrate. At this point, it was considered that the crucible temperature was still too low to give a reasonably high deposition rate so it was decided to raise the crucible temperature even higher for the next run.

## 4.10 Run No. 5:

An attempt was made to achieve a crucible temperature of 1,300°C; but as the voltage applied to the crucible heater was increased, the power supply circuit breaker tripped although it was rated for 30 Amps. The run had to be abandoned. When the system was dismantled, it was observed that the screws carrying current to the filament of the crucible heater had melted, although stainless steel screws were used. also observed that the thermocouple tip had fused. These could have been caused by too rapid an increase in the voltage applied to the crucible heater, which did not allow sufficient time for heat conduction to establish equilibrium temperatures. Obviously, no film resulted. The system was dismantled, cleaned with HCl and then with trichloroethylene, then put together using a new filament for the crucible heater and new screws instead of the damaged ones.

### 4.11 Run No. 6

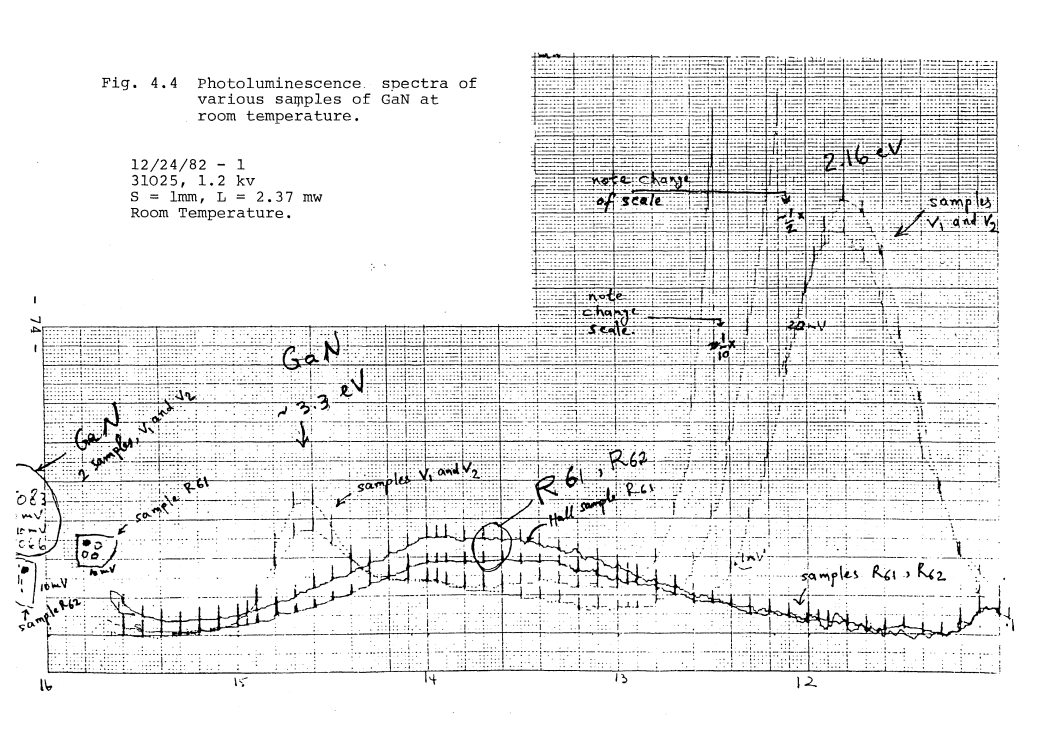
The deposition parameters for this run were set as follows:

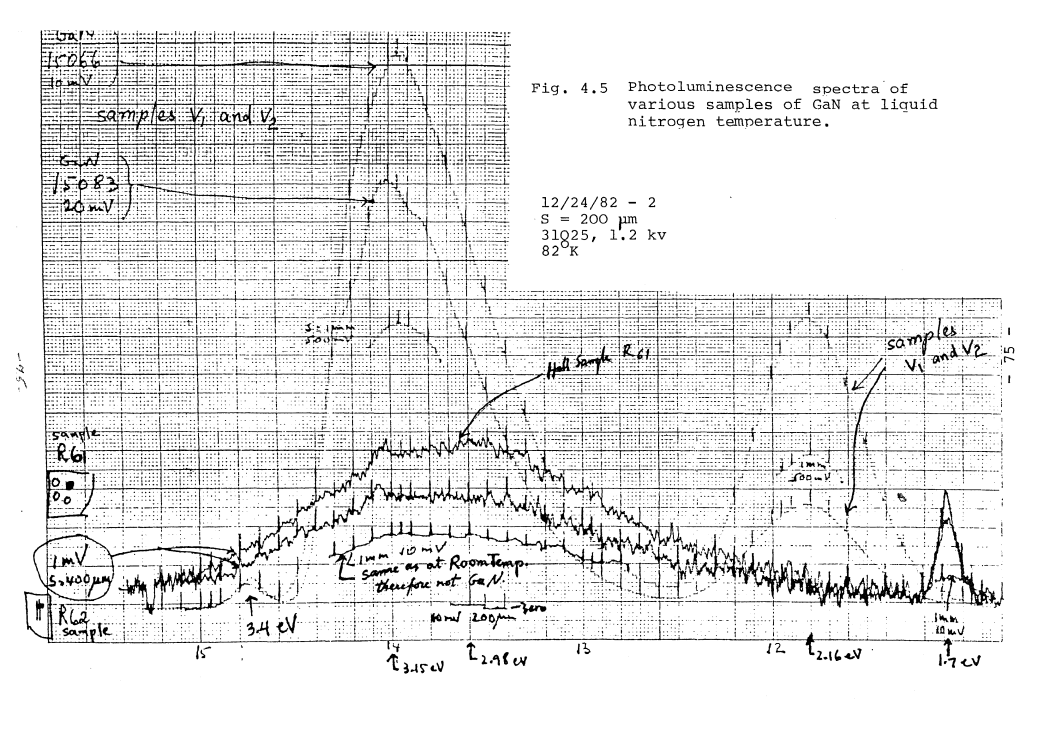
- Average crucible temperature: 1,120°C.
- Nitrogen Pressure:  $2 3 \times 10^{-4}$  torr.

- Electron Current for Ionization: 360 mA.
- Acceleration Voltage: Zero.
- Substrate Temperature:
  - initial: 340°C.
  - final: 371°C.
- Deposition Time: 90 minutes.

Two samples were obtained from this run:  $R_{61}$  and  $R_{62}$ . The films were transparent, and moderately hard to scratch, in line with the properties of known samples of GaN. ductivity measurements showed that the films were insulating. The films were then sent to RCA laboratories in Princeton for photoluminescence and absorption edge tests. In the photoluminescence tests setup of Dr. J.I.Pankove at RCA labs, the films are optically excited by a 325 nm, 2.2 mW laser beam. In optical excitation, a photon is absorbed by the semiconductor, creating an electron-hole pair which then recombines emitting one or more photons (31). Therefore by observing the emission spectra, peaks corresponding to the fundamental transition energy (band to band) as well as peaks corresponding to transitions between impurity levels, from the conduction band to impurity levels, and from impurity levels to the valence band, can be observed.

Results of such photoluminescence tests are shown in fig. 4.4 for room temperature and fig. 4.5 for liquid mitrogen temperature. These are shown for samples  $R_{61}$  and  $R_{62}$ , as



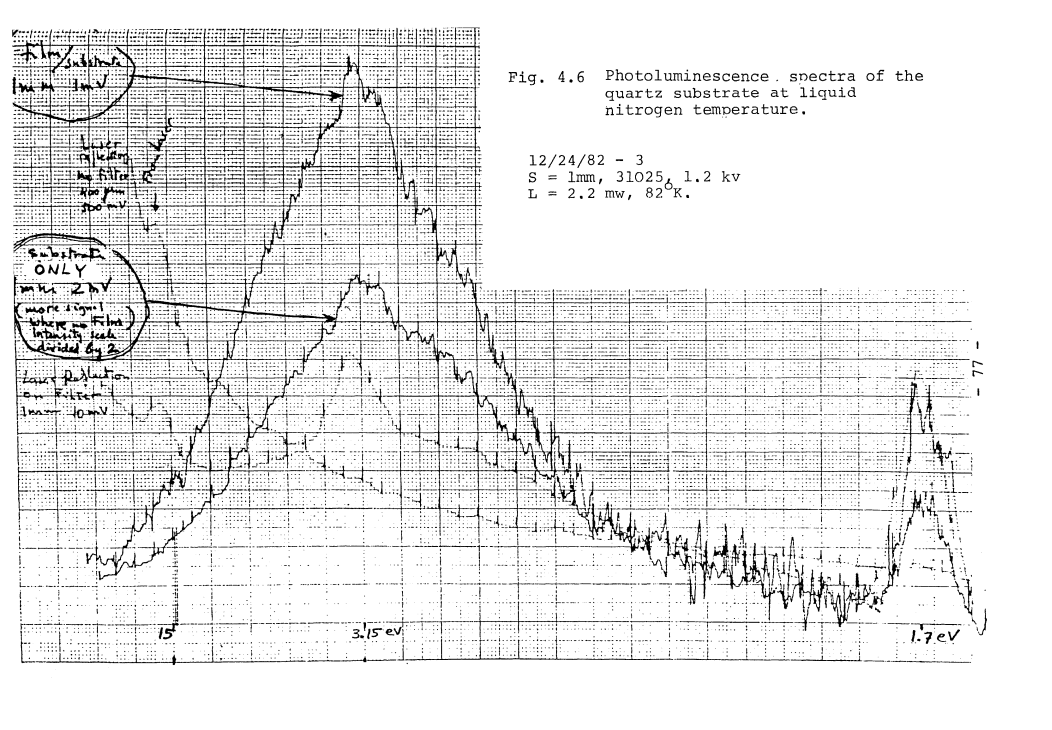


well as for two other samples of GaN,  $V_1$  and  $V_2$ , grown by the chemical vapour transport deposition method described in section 4.2 .

Examination of these two figures revealed that two peaks were present for samples  $\rm V_1$  and  $\rm V_2$  at room temperature:-

- One at a photon energy of approximately 3.3 eV.
- The other, a much higher peak, at a photon energy of approximately 2.16 eV.

At liquid nitrogen temperature (fig. 4.5), a low peak at 3.4 eV was observed, followed by a much higher peak at 3.15 eV and then the peak at 2.16eV, which was now a relatively smaller peak compared to the one at 3.15 eV. Samples R61 and R62 did not show these characteristics. Instead, a broad spectrum was obtained at room temperature. At liquid nitrogen temperature, a broad spectrum was similarly observed, but a peak at 1.7 eV was also present. This last peak was suspected to be due to the quartz substrate so another photoluminescence test was carried out on the substrate alone (at a point which was masked during the deposition of the film). The results of this test, shown in fig. 4.6, confirmed that the peak obtained at 1.7 eV could be attributed to the substrate since the peak was present in the spectra for the film/substrate as well as for the substrate alone. Since the photoluminescence spectra for the films (samples  $R_{61}$  and  $R_{62}$ ) were the same for room temperature and liquid nitrogen temperature, and they did not



correspond with the spectra of known samples of GaN, it was suspected that the films obtained in this run might have not been GaN.

### 4.12 Potential Problems

Two potential sources of trouble were identified:

- A Residual Oxygen in the vacuum chamber which, because of its high chemical activity relative to the Nitrogen, might have reacted with the Gallium to form Gallium Oxide.
- B Low substrate temperature, which could have been either too low for a reaction between Ga and  $N_2$  to occur, or, too low to produce a crystalline film resulting in the film being amorphous GaN.

## 4.13 Solutions Employed

Steps were taken to eliminate the above potential problems as follows:

<sup>\*</sup>Experimental work by C. Mehta at NJIT during the writing of this thesis has revealed that water vapour in the boron Nitride crucible can cause the boron nitride to decompose at elevated temperatures introducing additional reactive contaminants at the substrate and preventing the deposition of pure gallium. This problem will be discussed further in section 4.20 at the end of this chapter.

### Low Substrate Temperature

To increase the substrate temperature beyond  $350^{\circ}\text{C}$ , a new substrate heater was constructed as shown in fig. 4.7. The quartz substrate was directly mounted on the back side of the lower plate of the heater for efficient heat conduction. With this arrangement, substrate temperatures greater than  $500^{\circ}\text{C}$  (up to  $700^{\circ}\text{C}$ ) could be reached.

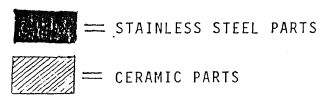
## Residual Oxygen in the Vacuum Chamber:-

Three sources of oxygen contamination to the chamber were possible:

### 1. Small Leaks

This of course was the most obvious possible source of O<sub>2</sub> in the vacuum chamber, so it was tested first. A leak detection test was carried out using a mass spectrometer leak detector, model 925-40, made by Varian Vacuum Division. By squirting Helium gas around the chamber and waiting for the mass spectrometer to respond, no leaks could be found even with the leak detector set for the highest sensitivity.

<sup>\*</sup>As outlined in the footnote on page 78 , the residual water vapour trapped in the boron nitride crucible can serve to introduce oxygen as the boron nitride reacts with  $\rm H_2O$  at elevated temperatures releasing boron oxides which act as reactive contaminants at the substrate. This is discussed further in section 4.20 at the end of this chapter.



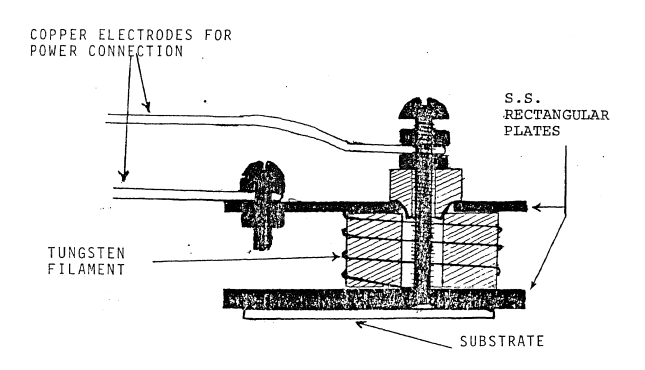


Fig. 4.7 Structure of the substrate heater used for run #7 onwards.

# 2. Oxygen Contamination from the Nitrogen Tank

To reduce the possibility of introducing small amounts of  $O_2$  present as an unavoidable impurity in the nitrogen tank; it was decided to use a <u>LIQUID</u> nitrogen reservoir as the source for nitrogen vapour. Since the boiling point of  $O_2$  is higher than that of  $N_2$ , this would insure that the  $N_2$  vapour is free of  $O_2$ , because any residual oxygen that might be present will stay as liquid in the liquid  $N_2$  reservoir. (25).

### 3. Insufficient Pumpdown and Outgassing

To eliminate any residual  $0_2$  in the vacuum chamber due to insufficient pumping down and/or outgassing of residual  $0_2$  trapped on the walls and surfaces, it was decided  $^{(6)}$  to pump down to the  $10^{-6}$  torr range overnight (over 14 hours). Then the chamber would be flushed with  $N_2$  and pumped down again to the  $10^{-6}$  torr range. This would be repeated 3 to 4 times to insure that any residual atmosphere in the vacuum chamber will be predominantly nitrogen.

Based on the above considerations, the following procedure was adopted for the runs that were attempted after run No. 6 (i.e. runs No. 7 through 11):

### 4.14 Start-up Procedure Adopted For Run No.7 Onwards

#### A . System Preparation

 The various parts of the system were cleaned with dilute HCl, then trichloroethylene, then blown dry with helium.

- 2. The crucible was filled with a fresh load of gallium then the lid was pushed in place and the crucible fitted inside the heater, making sure it is perfectly vertical, with the nozzle pointing towards the substrate. The ionization stage was fitted on top of the nozzle assembly and secured in place.
- 3. The substrate (Quartz) was cleaned in trichloroethylene for approximately two hours, blown dry with
  helium, then mounted on the back side of the substrate
  heater which in turn is secured to the substrate holder
  as shown in fig. 4.8. The thermocouple was positioned
  on one side of the substrate surface.
- 4. The nitrogen inlet was connected and the line flushed to clear any entrapped air.

### B. Start-up Procedure

- 1. The cooling water lines to the diffusion pump and to the ICB system were turned on.
- 2. The mechanical (roughing) pump was turned on and used to pump down the diffusion pump for 20 - 25 minutes. Then the diffusion pump was turned on.
- 3. The vacuum chamber was pumped down with the roughing pump to less than 50 microns of Hg.
- 4. The diffusion pump was left to warm up while continuously pumping it down with the roughing pump for about 40 45 minutes. Then the butterfly valve was

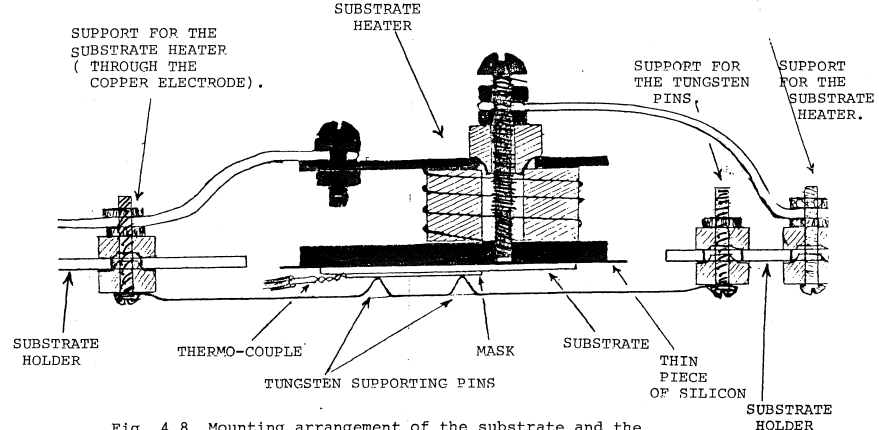


Fig. 4.8 Mounting arrangement of the substrate and the substrate heater on the substrate holder.

opened and the vacuum chamber was pumped down with the diffusion pump. Pumping down in this manner was continued for 12 - 14 hours (overnight) to a base pressure of  $1 - 3 \times 10^{-6}$  torr.

- 5. The vacuum chamber was flushed with nitrogen (obtained from a liquid nitrogen source as described earlier) to a pressure greater than 1 torr, then pumped down again to  $1-3 \times 10^{-6}$  torr. This was repeated at least 3 times.
- 6. The nitrogen leak valve was adjusted to get the desired nitrogen pressure in the chamber, i.e.,  $1 5 \times 10^{-4}$  torr.
- 7. The power supplies were set initially for zero output, then they were plugged in and switched on.
- 8. The crucible and substrate heaters' voltages were increased slowly in increments of a few volts at a time, letting the system stabilize in two ways before applying the next increment:
  - a) Re-establish its equilibrium value of pressure (because the pressure increases slightly when the temperatures of the crucible and the substrate are increased, mainly due to outgassing).
  - b) Reach a slow rate of change in both the substrate and the crucible temperatures, say  $1^{\circ}\text{C}$  rise in 15 or more seconds.

Precaution (b) above was found necessary to give enough time for heat conduction from the filaments to the

- boron nitride heater and thus avoid excessive local heating of the tungsten filaments which might lead to the breaking of the filaments.
- 9. Once the desired substrate temperature and crucible temperature were reached, the ionization filament (cathode) voltage was raised slowly up to 14 volts. Then the electron accelerating voltage was applied and both were adjusted to get an electron current for ionization ( $I_e$ ) of 300 mA with an electron accelerating voltage of at least 100 volts as described in section 3.7.
- 10. To begin the deposition, the shutter is moved aside allowing gallium flux to land on the substrate.

  After depositing for the desired period of time, the run was terminated as follows:-

### 4.15 Shut Down Procedure Adopted For Run No. 7 Onwards

- 1. The shutter was returned in place.
- 2. The electron accelerating voltage was reduced to zero.
- 3. The ionization filament (cathode) voltage was reduced until the filament was red hot instead of white hot.
- 4. The crucible and substrate heater filaments' voltages were reduced gradually and,
- 5. the mitrogen pressure was increased by allowing  $N_2$  into the chamber to a pressure of approximately 150 micron of Hg. The system was kept at this  $N_2$  pressure while continuing (4) above until the filaments were no longer visibly hot.

- 6. All power supply voltages were reduced to zero.
- 7. The butterfly valve between the chamber and the diffusion pump was closed and the diffusion pump switched off.
- 8. The chamber was filled with nitrogen and the system was allowed to cool for at least two hours before it was opened.
- 9. The cooling water to the ICBD system and to the diffusion pump were turned off.

### 4.16 Runs No. 7 and No. 8.

In both of these attempts, the procedures outlined above were followed to the point of establishing the desired substrate and crucible temperatures; however, in both runs, the substrate heaters were lost (filament broken) before the deposition could begin. In run No. 7, the carbon based ceramic used in the substrate heater disintegrated and fused tiny particles started flashing in the vacuum chamber. For run No. 8, that was replaced by a piece of boron nitride, 1" in diameter. Two turns of 0.019" diam. tungsten wire were used as the heating filament, but this broke (open-circuited at one point) before the substrate temperature could be raised above 450°C, indicating that the filament was not suitable to handle the amount of power required. Therefore it was replaced with 4 turns of 0.023" diam. tungsten wire wrapped around a 1" diam., ½" long, cylindrical boron nitride piece, grooved to accept the filament (for better heat conduction).

### 4.17 Run No. 9.

The conditions for the deposition run were set as follows:-

- Crucible temperature: 990°C ± 5°C.
- Electron current for ionization: 350 mA.
- Nitrogen pressure in the chamber:  $1 2 \times 10^{-4}$  torr.
- Acceleration Voltage: Zero.
- Substrate temperature:

initial: 618°C

final: 667°C

- Deposition time: 213 minutes.

The following observations were made on the film obtained in this run:-

- 1. The mask had moved as a result of vibration of the system during the deposition run; hence the edges were not sharp or well defined.
- 2. The stainless steel base of the substrate heater which supported the substrate had diffused into the back of the substrate due to the high temperature of the heater.
- 3. No quantitative thickness measurements were possible on the film, but it could be judged by visual inspection that the film was extremely thin.

Therefore, no further measurements were made on this film and it was decided to:

1. Use A larger nozzle diameter for the crucible (1.5 mm instead of 0.5 mm) to get a higher deposition rate.

2. Use a thin piece of silicon between the quartz substrate and the stainless steel heater base to avoid the diffusion of stainless steel into the back of the substrate. Both above modifications were made for the subsequent runs (Nos. 10 & 11).

#### 4.18 Run No. 10.

The conditions for this deposition run were set as follows:-

- Crucible temperature: 995°C ± 5°C
- Electron current for ionization: 300 mA + 20mA.
- Nitrogen pressure in the chamber:  $5 6 \times 10^{-4}$  torr.
- Acceleration voltage: Zero.
- Substrate temperature:

initial: 520°C

final: 550°C

- Deposition time: 4 hours.

After the system was allowed to cool down, the substrate/film was taken out and the following was observed/done:-

- 1. The back of the substrate was stained with impurities and various spots. These were thought to be diffused silicon at first, but further scrutiny revealed that it must have been impurities or dirt on the silicon surface rather than silicon.
- 2. The mask edges were sharp and the film was clearly visible; i.e., the constrast was very good.

- 3. The film was hard, and pressing lightly on it with a pin would not scratch or peel it.
- 4. The film was not attacked by HCl indicating that it was not gallium metal.
  - The sample was personally taken to RCA Laboratories on and the following was done/measured.\*
- The film thickness was measured to be about 250 Å.
   This indicated that the deposition rate under these conditions was approximately 1 Å per minute a rather low value for ICBD system.
- 2. Conductivity measurements were attempted, but the film proved to be highly insulating - even with 600 volts applied between 2-points only 1 mm apart.
- 3. Absorption edge measurements were made but no absorption edge was detected. This however, was thought to be an inconclusive measurement because with such a thin film, the absorption of the band gap must be very high for an absorption edge to be detected with the equipment used.
- 4. Luminescence tests were made and the results were similar to those obtained for samples  $R_{61}$  and  $R_{62}$  made in Run No.6

<sup>\*</sup>These measurements were supervised by Dr. J.I. Pankove of RCA laboratories, Princeton, N.J.

As for the previous samples, no peaks indicating the band structure of GaN were present and, this time, no peak due to the quartz substrate could be detected either. Instead, all that could be observed was some scattering effects.

#### 4.19 Run No. 11

on quartz as the substrate. The ZnO film was prepared by Dr. J.I. Pankove at RCA Labs. and photoluminescence measurements were made on the ZnO/quartz substrate for later comparison with the spectra of the sample after the deposition run. The substrate was then assembled on a thin piece of clean silicon which in turn was supported on the back of the substrate heater.

Following the procedure described in section 4.14, the deposition parameters were set as follows:-

- Crucible temperature: 1100°C
- Electron current for ionization: 300 mA.
- Nitrogen pressure in the chamber:  $1.5 \times 10^{-4}$  torr.
- Acceleration voltage: Zero.
- Substrate temperature: 590°C.
- Deposition time: 63 minutes.

The film obtained in this run was similar to the films obtained in earlier runs except for one marked difference. This film proved to be highly conductive while all previous films were highly insulating. However, the results of

photoluminescence tests made on this sample were similar to the results obtained from the substrate (ZnO on quartz) before the deposition run, and no peaks characterizing GaN could be observed. Surface analysis of the film was made at RCA labs and the results showed that the film was rich in oron, itrogen and other impurities.

#### 4.20 Conclusions and Suggestions for Future Work

#### A. With regard to the ICBD system:

Although the ICBD system construction was completed and experiments were made using the system, many of the refinements that were contemplated at the early stages were not incorporated in the system due to the extreme budget limitation at the time. Some of the more important improvements that can be suggested for the future are:

- 1. A safety interlock system is required that will automatically shut off the high voltage power supplies when the vacuum chamber is opened.
- 2. Substrate heating using a heater directly in contact with the back of the substrate is rather inefficient (because of the bulk heating) and can be a source of impurities and contaminants. Therefore surface heating of the substrate is proposed using infrared lamps focused at the substrate.
- 3. Although monitoring the crucible and substrate; temperatures

with thermocouples can be very accurate, thermocouples raised to these high temperatures can also be a source of trouble and contaminants in the vacuum chamber. Therefore monitoring the temperatures using an optical emission spectrum analyzer would be a preferable alternative.

# B. With regard to the deposition of GaN films byR - ICBD:

Our attempts to reproduce the results published to the Takagi research group, as reported in this work, were not successful. All the films obtained were very thin ( $\leq 250$  Å thickness), and the deposition rate of about 1 Å per minute was at least an order of magnitude lower than the typical deposition rates expected of ICBD systems.

The low deposition rate could be attributed to:

- either the small nozzle diameter of the crucible, or
- low crucible (gallium) source temperature.

But since the nozzle diameter was increased from 0.5 mm to 1.5 mm for runs No. 10 and 11 without much improvement in the deposition rate, we must conclude that the crucible temperature must have been lower than that required for efficient gallium vapour (or cluster) ejection. This argument can be further supported by considering the gallium vapour pressure in the crucible for the temperature range under consideration.

Referring to the vapour pressure charts (fig. 4.9), the following data are tabulated for easier reference:-

NEWTONS/METER<sup>2</sup> **+**01 -\_0 | VAPOR PRESSURE IN ATMOSPHERES έO 5000 6000 7000 - MELTING POINT 4000 000 (Rh) 000 0000 0000 TEMPERATURE DEGREES CENTIGRADE TEMPERATURE DEGREES KELVIN RULIND HR. F. W. 2 BOILING 103 , O \_0 VAPOR PRESSURE IN TORR (mm Hg) Vapour pressure charts of selected elements. 4.9

-93-

Vapour Pressure (torr)
$7 \times 10^{-4}$
$2 \times 10^{-3}$
$4 \times 10^{-3}$
$3 \times 10^{-2}$
1.5 x 10 <sup>-1</sup>
5 x 10 <sup>-1</sup>
2
4
10

Noting the above, the following observations can be made:

- 1. At  $900^{\circ}$ C, the vapour pressure of the gallium in the crucible is only marginally greater than the chamber pressure  $(1 5 \times 10^{-4} \text{ torr})$ , so only very little evaporation can be expected.
- 2. For ICB system, the vapour pressure in the crucible  $P_0$  have been reported  $^{(55)}$  to be in the range  $10^{-2}$  to 10 torr. This suggests that the crucible temperature should at least be  $1100^{\circ}$ C for gallium nitride deposition.
- 3. For ICB systems, the ratio of the pressure inside the crucible to that outside the crucible  $(p_0/p_{chamber})$  is required to be at least  $10^4$  to  $10^5$  as detailed in the earlier part of this thesis.

Since in our experiments (and those reported by Takagi and Matsubara for GaN deposition) the nitrogen pressure in the

chamber was in the 10<sup>-4</sup> torr range, the gallium vapour pressure required in the crucible would be in the 1 to 10 torr range. This corresponds to a crucible temperature of 1,400°C to 1,600°C.

Based on the above information and the very low deposition rate obtained in the runs that have been attempted so far, we can confidently conclude that a crucible temperature of  $900^{\circ}$ C is too low for GaN deposition and that future runs should be attempted at higher crucible temperatures, preferably in the range  $1,350^{\circ}$ C to  $1,600^{\circ}$ C.

#### C. The Contamination Problem

Surface analysis of the film obtained in the last run

(No. 11) was made at the RCA Laboratories in Princeton,

New Jersey. As mentioned earlier, this analysis showed that
the film was rich predominantly in boron and nitrogen. Further
expermental work by C. Mehta, a graduate NJIT student, during
the writing of this thesis revealed that the main source of
trouble was water vapour trapped in the boron nitride crucible
and heaters. Boron nitride, when exposed to the atmosphere,
quickly absorbs moisture present. This trapped water
vapour can cause the boron nitride to decompose at elevated
temperatures introducing additional reactive contaminants at
the substrate and preventing the deposition of pure gallium.
The problem can be solved by driving out the water vapour,
which can be done in two steps:

- Heating the boron nitride in an oven at 150 200°C for approximately four hours to drive most of the water vapour out.
- Then final low temperature (150 200°C) annealing in vacuum will drive out the remaining traces of water vapour.

### D. Final Remarks

Although our attempts to grow GaN films by the ICB deposition technique at the NJIT Microelectronics Lab. were not yet successful, experimental work is being continued and refinements are constantly introduced based on the wealth of experience gained through the work reported in this thesis. The main objectives of the present work were:

#### First Stage:-

The deposition of high quality GaN films using the ICB technique and the determination of the optimum values of control parameters (such as Ionization current,  $I_e$ ; Crucible temperature; Nitrogen Pressure; Substrate temperature; etc.) that will ensure consistently reproducible results.

#### Second Stage

Introducing dopants to the GaN films and the fabrication of Metal - Insulator - Semiconductor (MIS) Light Emitting Diodes (LED's).

Assuming that successful, reliably reproducible results are obtained in the first two stages, work should then be directed to investigating the possibility of commercializing the ICBD method of fabricating GaN LED's and extending the method to the fabrication of multi-colored solid state display units as suggested by Dr. J. I. Pankove (29). The technique would utilize GaN crystals both as blue light emitting diodes (LED's) and as convenient transparent supports for mounting LED's emitting other colors as discussed in section 4.1.

### APPENDIX A

MACHINE SHOP DRAWINGS FOR THE

PARTS USED IN CONSTRUCTING THE

ICB SYSTEM AT NJIT.

( Figures A.1 through A.13 attached )



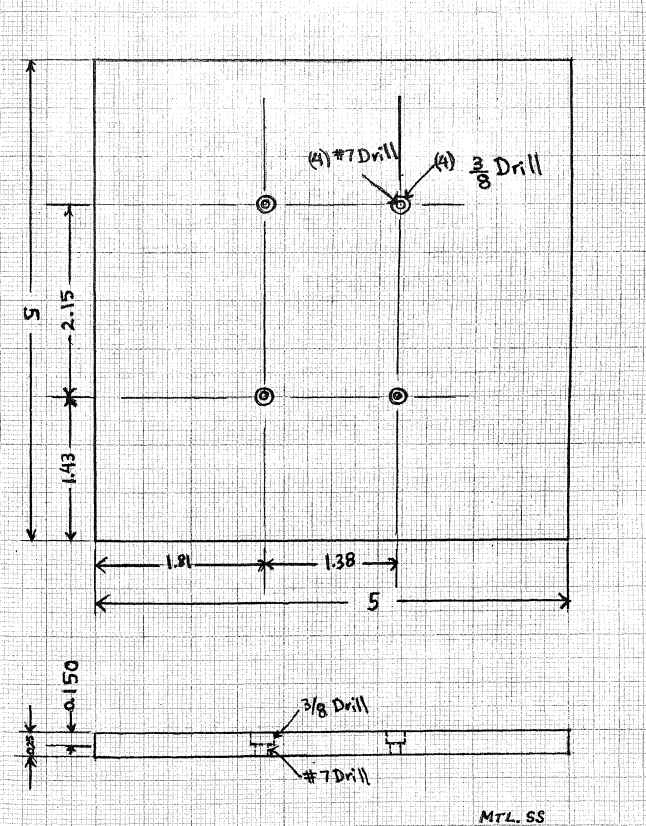
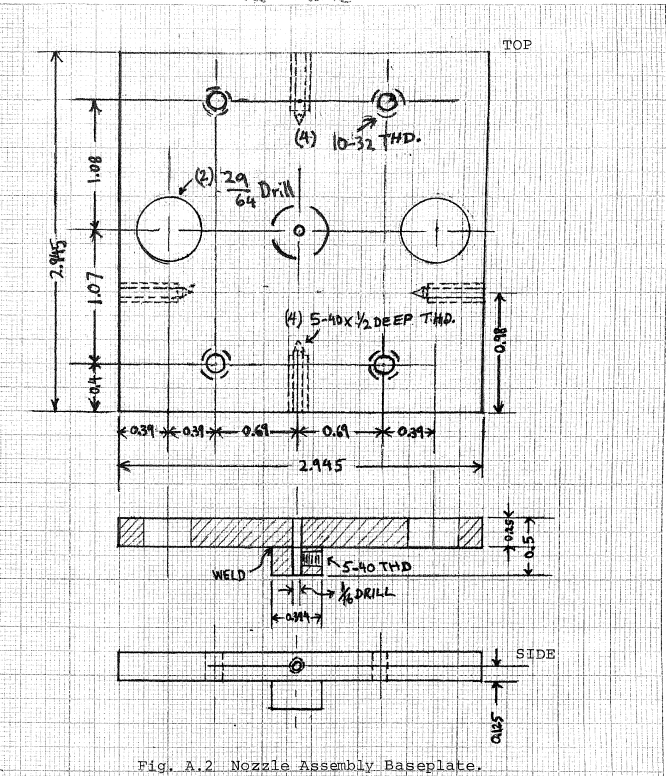
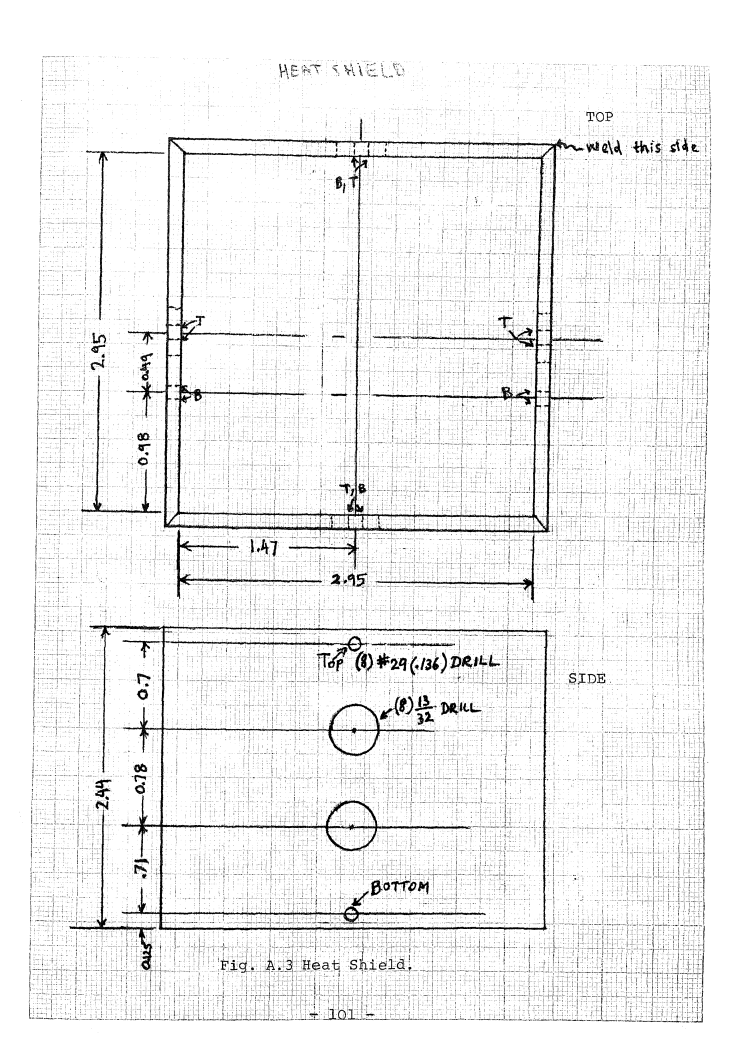


Fig. A.l System Base.



100



INVENTION BASE FLATE

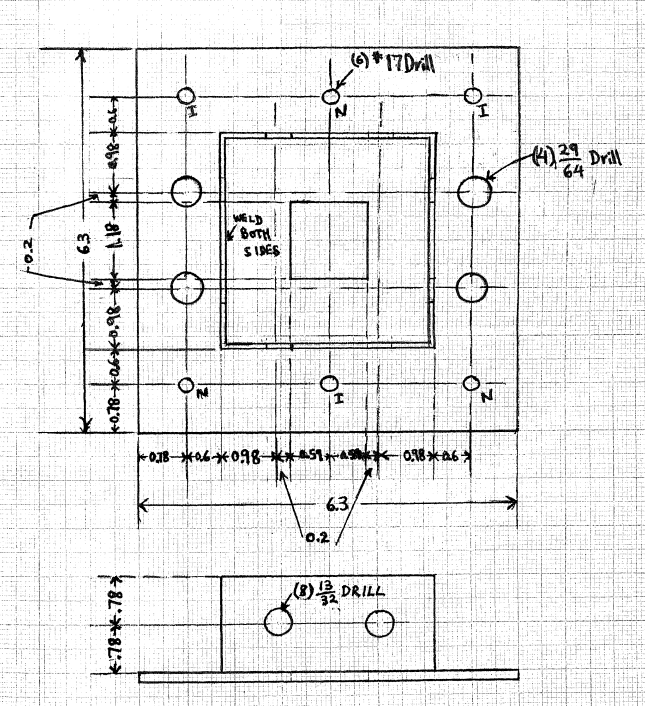
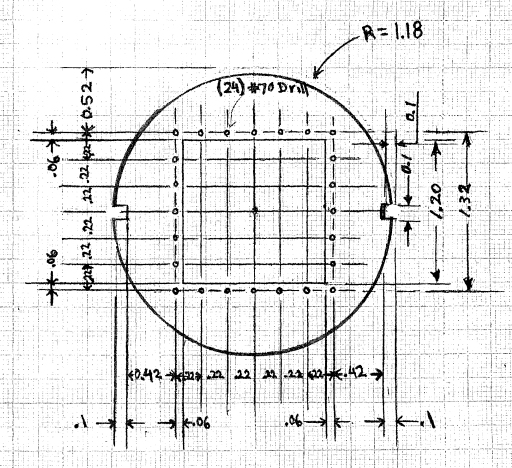


Fig. A.5 Ionization Assembly Baseplate.

4 103 -



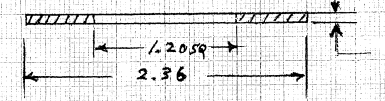
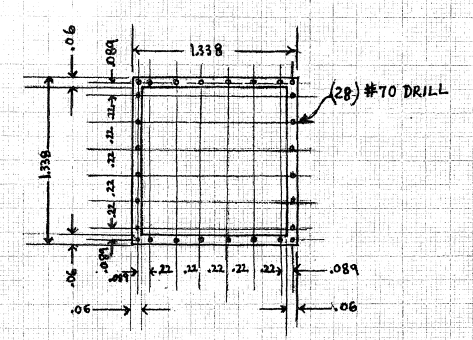


Fig. A.6 Anode upper plate.

### ANDE LOWER PLATE



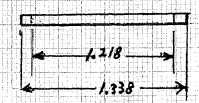


Fig. A.7 Anode Lower Plate.

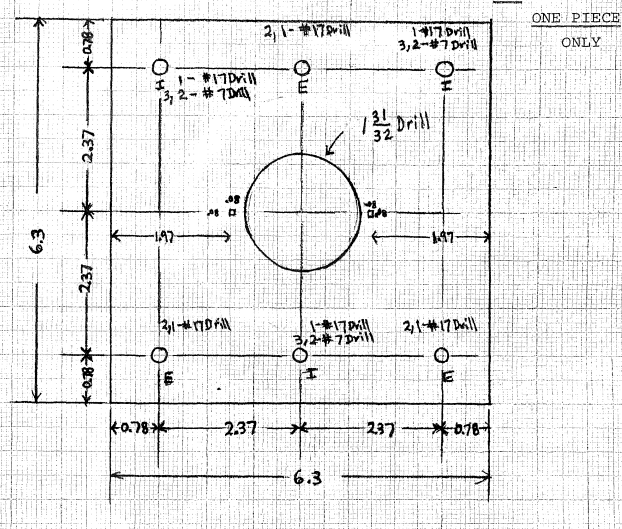


- 2-ACCELERATION ELECTRODE
- SUBSTRATE HOLDER 3-

#### (3) REQUIRED.

#### NUB ON

ONLY



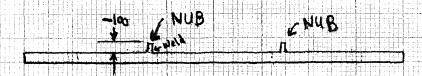
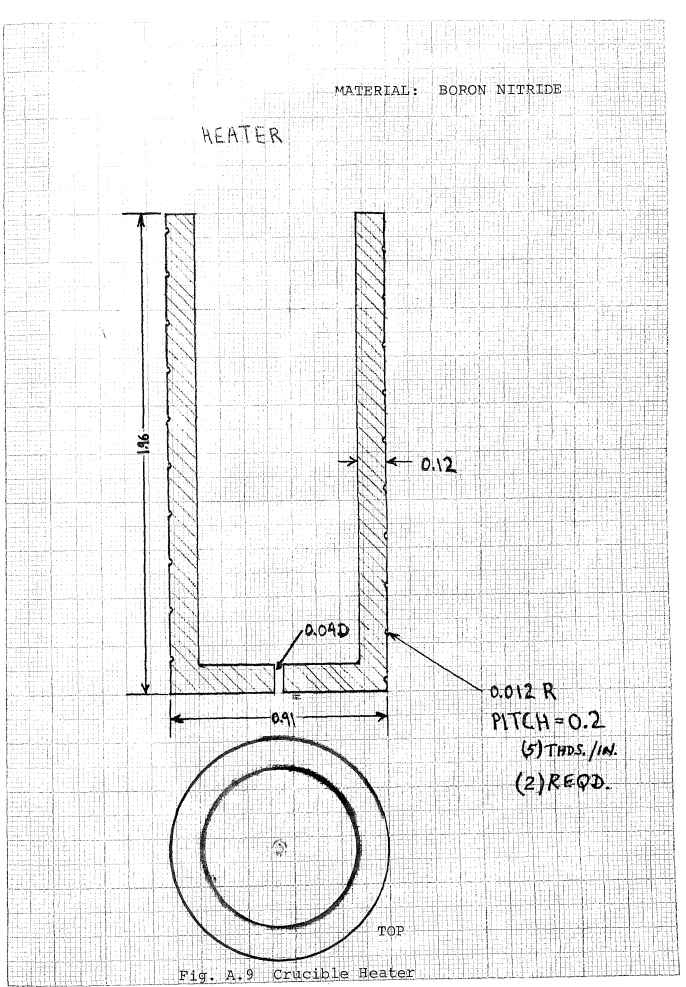
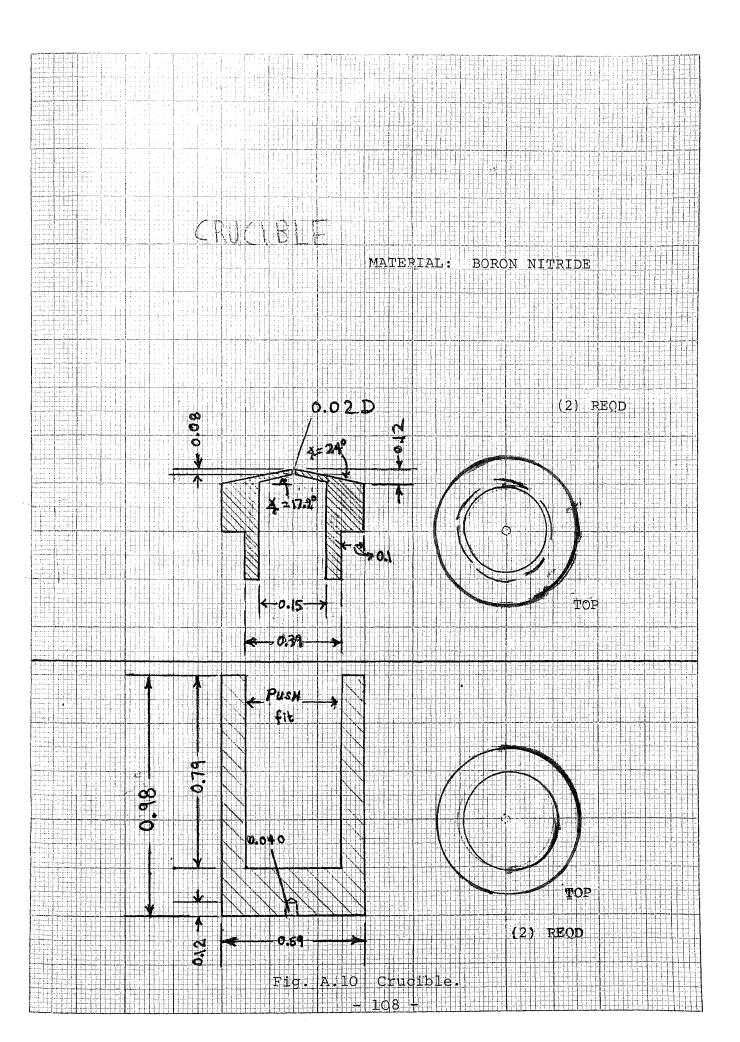


Fig. A.8 Anode Holder, Substrate holder and Acceleration Electrode.





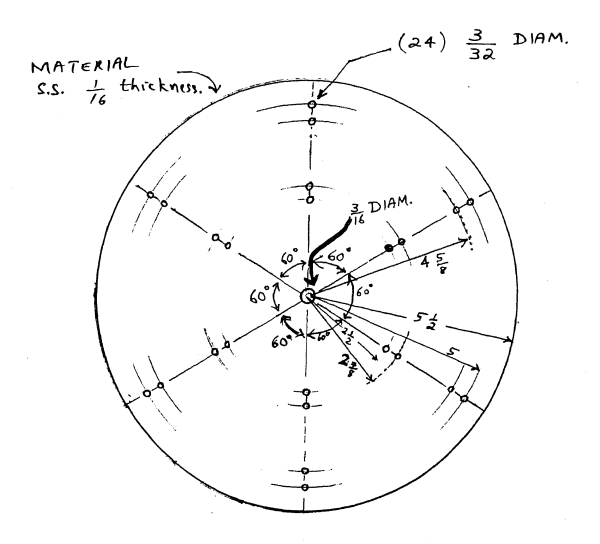


Fig. A.ll Rotating plate of the rotating substrate holder.

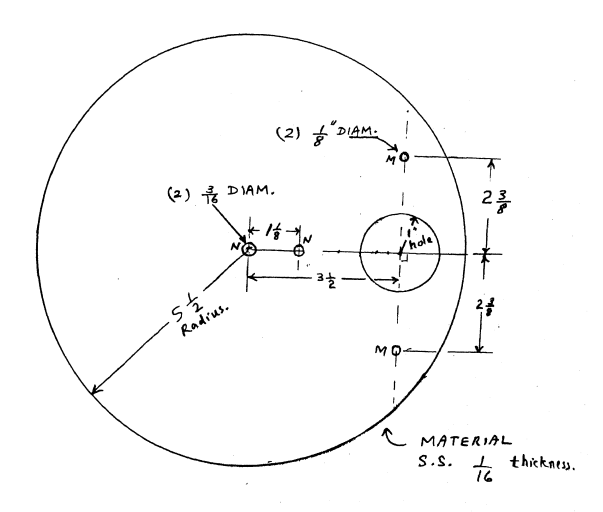


Fig. A.12 Stationary plate of the rotating substrate holder.

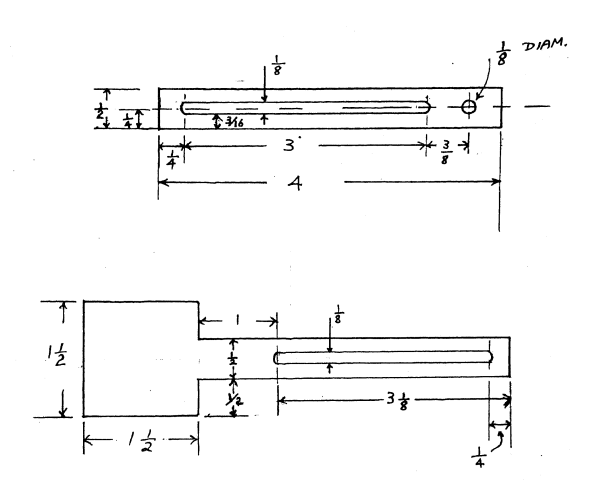


Fig. A.13 Parts for the shield (shutter).

## A P P E N D I X B

INTERNAL WIRING DIAGRAMS FOR THE DC, HIGH VOLTAGE, VARIABLE POWER SUPPLIES USED FOR  $V_{\rm e}$  AND  $V_{\rm a}$ . ( Figures B.1 and B.2 attached )

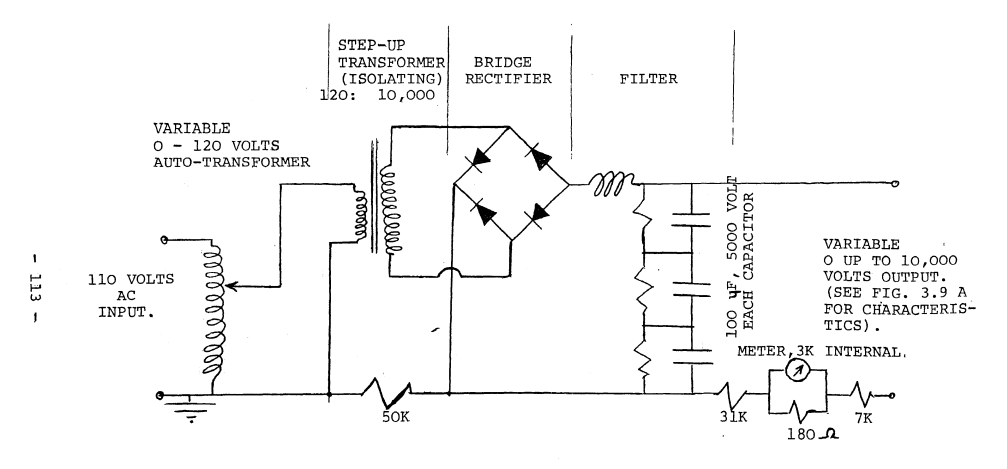


Fig. B.l Internal wiring diagram for the DC variable power supply used for  $\mathbf{V}_{\mathbf{e}}$  .

NOTES: - Existing, internally made power supply was utilized.

- Ratings not shown above were not readily available.

PART OF THE CIRCUITS OF AN ELECTRON BEAM POWER SUPPLY ( AIRCO MODEL ES6 - 110/210 ) WAS UTILIZED TO GENERATE THE DC HIGH VOLTAGE, Va, VARIABLE FROM ZERO TO 10 KV.

THE APPLICABLE WIRING DIAGRAM ( AIRCO, CALIFORNIA, DRAWING REFERENCE NO. 2F 6055 ) IS ATTACHED, AND THE FOLLOWING MODIFICATIONS WERE MADE :-

- (1) FUSES F 201, F 202 AND F 203 WERE REMOVED, ISOLATING THE ENTIRE BOTTOM SECTION OF THE DIAGRAM.
- (2) THE KEYLOCK SWITCH, S 101 WAS SET IN AN OPEN CIRCUIT POSITION.
- (3) AN EXTERNAL CONTROL WIRE WAS CONNECTED TO THE TOP SIDE OF S 101, GIVING 110 VOLT CONTROL VOLTAGE TO ACTIVATE THE STARTING MECHANISM.
- (4) TUBE V 201 WAS REMOVED AND REPLACED WITH A SIMPLE LINK.
- (5) THE POWER SUPPLY WAS ENERGISED THROUGH A THREE PHASE AUTO TRANSFORMER CAPABLE OF PROVIDING A CONTINUOUSLY VARIABLE OUTPUT ( 3 PHASE ) BETWEEN ZERO AND 208 VOLTS.
- TRANSFORMER T 204 WAS UTILIZED AS THE ISOLATING
  TRANSFORMER I. T. 2 IN FIG. 3.6 . THE INPUT TERMINALS
  H1 AND H2 WERE BROUGHT OUT AND FED THROUGH A VARIABLE
  VOLTAGE, O-240 VOLTS . THE OUTPUT TERMINALS X1 AND X2
  WERE EXTENDED OUT USING A HIGH VOLTAGE ( 15 KV PEAK),
  HIGH CURRENT ( TO 30 AMPS ) CABLE.
  - Fig. B.2. Internal wiring diagram for the high voltage, DC, power supply used for Va.

    (Above notes should be read in conjunction with the wiring diagram on page 114 A attached).

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