Diffusion and Phase Change Characterization by Mass Spectrometry

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Diffusion and Phase Change Characterization by Mass Spectrometry

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Part I

INTRODUCTION

The mass spectrometer has long been used as an analytical tool in many disciplines. Relative to the space program, the mass spectrometer was one of the first major analytical instruments to be utilized from the very inception of the nation's program.

Virtually every major rocket was equipped with a mass spectrometer to measure the atmosphere either on ascent or descent and in all recent missions the mass spectrometer has provided a wealth of information which cannot be obtained with any other type of instrumentation. Many types of mass spectrometers have monitored the upper atmosphere of the earth and are continuing to do so. Additionally, deep space probes can be expected to continue to utilize mass spectrometry, and the surfaces of planets are amenable to analysis by mass spectrometry via telemetering of their chemical composition.

Another major area in which mass spectrometry has made a contribution to NASA sponsored programs relates to the high purity materials that have been developed for solid state electronics. Without mass spectrometric analysis to develop a solid state technology, the rapid advance of transistors and other microcircuitry, computers, etc. would not have been possible, inasmuch as the mass spectrometer has a sensitivity for analyzing trace elements that generally exceeds all
other analytical methodologies.

The present work, however, relates to a more generic type of materials research that has applicability to high strength metals and alloys for advanced space vehicles, and even to the ultimate performance of composites in supersonic transports. Hence, this research represents an even further extension of the manifold applications for mass spectrometry that have occurred during the last two decades.

Specifically, this paper addresses the problem of the high temperature diffusion in metals, and it does so in a very unique manner. Until the last few years, there was rather little interest in the trace elements of metals as opposed to some of the major constituents that are generally recognized as alloying elements. However, we are now approaching an era in which the efficiency and reliability of many systems, e.g., freedom from corrosion, the attainment of theoretical yield strengths, and the macroscopic properties of materials generally are dependent upon trace metals and their transport to grain boundaries. Hence, while this present research effort is focused on a few specific materials, the technology that it utilizes should have a very broad scale application, and it is in this context that we are presenting results of a preliminary nature which will hopefully lead to an extended future study of diffusion in many engineering materials.

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The first basic theory for calculating diffusion coefficients was presented by Adolf Fick in 1855. Fick who modified Fourier's heat conduction equations, hypothesized that in an isotropic medium the quantity of diffusing substance $Q$, which passes in a unit of time through a unit of transverse cross sectional area, is proportional to the concentration gradient measured along the normal to the section:

$$Q = -D \frac{dC}{dx}. \quad (1)$$

The above equation is called Fick's first law for equilibrium flow, $C$ is the concentration of the diffusion substance in units of mass/cm$^3$, $x$ is the spatial coordinate in units of cm., and $D$ is the diffusion coefficient with units of cm$^2$/sec. In the case of non-equilibrium flow, Fick's second law can be derived from the first law by considering the rate of accumulation of the diffusing substance in a given element of volume as the difference between the incoming and outgoing fluxes per unit time. This yields

$$\frac{dC}{dt} = D \frac{d^2C}{dx^2} + D \frac{d^2C}{dy^2} + D \frac{d^2C}{dz^2}. \quad (2)$$

Fick's laws were shown to be approximations to more general transport equations, and they may apply only to several atomic distances from solute sources or sinks, when the solvent is a homogeneous medium and when the solute concentrations are small. Boltzmann extended the application of Fick's equations to large solute concentrations, and these modified equations were graphically solved by Matano in 1936 for metallic...
diffusion. To date, experimental determinations of the diffusion coefficient have always followed a prescribed procedure. The solute material was allowed to diffuse at a known temperature and for a known time into a block of solvent material of such a shape that Fick's laws yielded an exact mathematical solution for the diffusion process in the system. Next, the solute concentration was measured at known points throughout the solvent. These values of solute concentration, and the diffusion coefficient was found algebraically or graphically. The problem of experimentally determining the diffusion coefficient thus became one of accurately determining the solute concentrations throughout the solvent, and of fulfilling the initial and boundary conditions applicable to a particular form of Fick's laws.

The methods which various experimenters have used to determine the solute concentrations and to satisfy the Fick's law boundary conditions in their experiments are quite diverse. An excellent summary of these different methods have been compiled by Gertsriken and Dekhtyar. 6

The introduction of radioisotopes in the 1940's enabled the concentration of a radioactive solute in a solvent specimen to be determined quite readily by nuclear radiation detectors. It is now advantageous to illustrate these standard methods.

**Serial Sectioning:** This method involving radioactive tracers is most frequently used in the study of diffusion. The sample after being held at a given temperature for a known amount of time (annealing time) to allow the radioactive tracer to diffuse into the sample, is subjected to a series of sectioning
operations. The sample is weighed before and after each cut to determine the thickness of the material removed. The cuttings are collected and their activity measured with a Geiger or scintillation counter. From the weights and the density, the coordinates of the midpoints of each slice, \((x_l, x_r)\), are calculated. A plot of the natural logarithm of the activity versus \(x^2\) has a slope of \(-1/4Dt\) for volume diffusion. Hence, if the time of the anneal, \(t\), is known, the diffusion coefficient \(D\) may be calculated.

The main advantage of the serial sectioning method is that it is simple, direct, and doesn't depend on the properties of the radiation from the radioactive material used, assuming no damage to substrate by ionizing radiation from the radioisotope.

**Residual Activity Method:** This method is similar to the serial sectioning analysis in that layers of thickness \(x\) are removed, but the total remaining activity, \(I\), of the sample is measured in this case.

In the case of weakly absorbed radiation (strong gamma rays) a plot of \(\ln(-dI/dx)\) versus \(x^2\) should be a straight line of slope \(-1/4Dt\); and in the case of strongly absorbed radiation (weak beta-rays) a plot of \(\ln(I)\) versus \(x^2\) should also be a straight line of slope \(-1/4Dt\).

**Surface Decrease Method:** In this method, the total activity of the specimen is measured as a function of time. No sectioning of the sample is necessary.
Autoradiography: The principle of this method is to determine the photographic density of the blackening of an exposed emulsion as a function of the distance from the interface of the specimen with the initial radioactive deposit. The specimen is usually cut at a measured angle $\alpha$ (usually $90^\circ$) to the initial face, and the cut face is placed in contact with a piece of appropriate X-ray film. The blackening of the emulsion (which is usually found to be directly proportional to the concentration of activity) is measured with a micro-densitometer. A plot of $\ln$ (photographic density) versus $x^2$ has a slope of $-1/4Dt$ from which $D$ may be calculated if the anneal time, $t$, is known. This method offered two distinct advantages; 1) since the filament was small, the times involved in heating and cooling it were small, and 2) because of the source mounting and the high vacua of the mass spectrometer the diffusing system was negligibly affected by any reaction between it and the surrounding environment at the experimental operating temperatures.

With the advent of sophisticated electronics, new devices have been developed for studying solid state phenomena. The electron-beam microprobe is one of the most important as far as diffusion studies are concerned. The device collimates a beam of electrons into a 1 $\mu$ diameter "pencil" which is directed onto the specimen surface at the spot to be analyzed. The electron bombardment causes characteristic X-rays to be emitted from the sample. These X-rays are then detected and analyzed to determine composition of the bombarded surface.
The limit of detectability is about \(10^{-14}\) grams for elements of high atomic number.\(^7\)

Existing procedures and techniques used for material characterization are adequate for materials which are to be operated in a normal temperature environment. However, at elevated temperatures (above 800°C), there is a need for improved measuring techniques. It is very important to understand the effects of high temperature operation on the materials, i.e., to determine whether or not the elevated temperature will cause matter from the environment to be absorbed into the material and then be transported through it, or whether some constituent of the material will be removed. The transport of constituents either into or out of the material can change the structural integrity of the material. As the high temperatures diffusing system tends to react with its environment. Also rapidly heating and cooling of the solute-solvent system will often adversely affect the diffusion measurements. In 1960, McCracken and Love\(^8\) developed a system for determining diffusion coefficients utilizing a mass spectrometer, and a thermal ionization source.

Using a mass spectrometer with a thermal ionization source, only solute atoms of a single isotope would be observed, and diffusion rates in a specified temperature range could be obtained. The assumption is made that Fick's solution for one dimensional diffusion in an outgasing thin plate can be applied. The solute ion current out of the filament as a function of time is thus:
\[ J_{\text{ion}} = A \exp \left( -\frac{\pi^2 D t}{x^2} \right) \]

where \( A \) is a constant of proportionality, \( D \) is the diffusion coefficient, \( x \) is the filament thickness, and \( t \) is the measurement time. Hence, the slope of a plot of the natural logarithm of the solute ion current versus time will yield the desired diffusion coefficient, provided the thickness of the filament is known.\(^9\)

**General Advantages of the Mass Spectrometric Technique:** There were many reasons for choosing the mass spectrometric approach for determining diffusion characteristics. As the mass spectrometer analyzes one mass at a time, there is no ambiguity caused by interfering masses. Since the filament is small, the times involved in heating and cooling it are also small. The probability of a reaction occurring between the high temperature specimen and the surrounding environment is negligible due to the mounting and the high vacua of the mass spectrometer. Finally, the sensitivity of the instrument permits the impurity level to be less than a part per billion, thus no doping of the sample is necessary and commercial grade material can be utilized.

**Importance of Sample Materials:** The materials studied in this work were tantalum, zircaloy-2, and 304 stainless steel. These materials were chosen for two basic reasons. First, they represent a family of materials which go from being ultra pure (99.996\% Ta) to a 1.5\% alloy with zirconium, to a 70\% Fe,
18.5% Cr, 9% Ni, alloy (304 stainless steel). Hopefully, this will enable one to draw conclusions about diffusion rates in alloys rather than ultra-pure single crystals. Secondly, these materials are of current interest to the nuclear industry where higher operating temperatures mean higher efficiencies.

The alkali metals were chosen as the elements of interest primarily due to their relatively low ionization potential (making them more suitable for thermal ionization) and due to the fact that they probably appear in most materials as naturally occurring impurities - thus eliminating doping of the host matrix. It was decided that concentrating these experiments on an entire chemical family might yield an overall diffusion mechanism which would allow diffusion rates of other impurities and sample materials to be approximated.

Use of trade names or names of manufacturers in this report does not constitute official endorsement of such products or manufacturers, either expressed or implied, by the National Aeronautics and Space Administration.
A. General Considerations

Diffusion is one of the mechanisms by which matter is transported through matter. From the theory of specific heats, it is known that atoms in a crystal oscillate around their equilibrium position. Occasionally these oscillations become large enough to allow an atom to change sites. The net result of many such random movements of a large number of atoms is actual displacement of matter, the movement being activated by the thermal energy of the crystal.\(^\text{10}\) The path of an individual particle is an unpredictable zigzag. The length of an individual step in this zigzag is determined in a solid by its jump length, b, i.e., the distance the particle is able to move when it acquires enough energy to make one jump.

The frequency of jumping, \(f\), is the vibrational frequency multiplied by the probability that the vibrating atom has the activation energy to make the jump;

\[
f = \frac{1}{3} \omega \exp \left( \frac{G_D}{RT} \right)
\]

(4)

where \(\omega\) is the vibrational frequency, \(1/3\) allows roughly for the fact that only about \(1/3\) of the randomly directed vibrations are in the required direction, and \(G_D\) is the free energy of activation.\(^\text{11}\) From this we can obtain an equation for the diffusion coefficient, \(D\),

\[
D = \frac{1}{2} b^2 f
\]

(5)
B. Diffusion Mechanisms

How the substance diffuses through the material is just as important as the diffusion coefficient formula itself. There are quite a variety of mechanisms by which an atom can move from one position to another in a crystalline structure.

**Vacancy Mechanism:** In all other than perfect crystals, some of the lattice sites are unoccupied. These unoccupied sites are called vacancies. If one of the atoms on an adjacent site jumps into the vacancy, the atom is said to have diffused by a vacancy mechanism, see Figure 1(b).

**Interstitial Mechanism:** When small atoms dissolve in metallic lattices as impurities so as to occupy interstitial positions between solvent atoms and an atom passes from one interstitial site to one of its nearest neighbor interstitial sites without permanently displacing any of the matrix atoms an interstitial mechanism is said to have taken place. The interstitial mechanism is thought to operate in alloys for those solute atoms which normally occupy interstitial positions. It will be dominant in any nonmetallic solid in which the diffusing interstitial doesn't distort the lattice too much, see Figure 1 (a).

**Interchange or Exchange Mechanism:** This is the direct exchange of two nearest neighbor atoms.

**Ring Mechanism:** A more general form of the exchange mechanism consisting of a number (three or more) of
DIFFUSION MECHANISMS

Figure 1
atoms forming a closed ring. Atomic diffusion then takes place by rotation of the ring.

**Interstitialcy Mechanism:** An interstitial atom moves from an interstitial site to an adjacent normal lattice site displacing the atom that had occupied that site into a new interstitial site, see Figure 1 (c).

**Crowdion Mechanism:** A line imperfection consisting of \( n \) nearest neighbor atoms compressed into a space normally occupied by \( (n-1) \) atoms. Diffusion then takes place by movement along the line of atoms, see Figure 1 (d).

**Dislocation Mechanism:** Dislocations can provide easier paths for diffusion than a perfect lattice. The diffusion of dislocated atoms can be produced in two ways: 1) the diffusion of the dislocated atoms over the interstices and 2) displacement of the dislocated atom from the interstice into a normal site of the lattice, and the atom situated in the lattice goes over into the interstice.

C. **Theoretical Methods of Diffusion Coefficient Calculations**

While the mechanisms of diffusion may be summarized from purely qualitative consideration, the problems involved in quantitatively calculating a diffusion coefficient for a specific solute-solvent system are quite complex. A convenient first step towards solving these problems is the consideration of the rate at which the solvent atoms themselves move around in the solvent lattice, or the self-diffusion coefficient of the solvent material. After a solution
has been found to the problem of self-diffusion, the effects
of a solute impurity are treated as a superposition problem.
The relationship between impurity diffusion coefficient and
the self-diffusion coefficient is discussed in section C-2.

C-1 Self-Diffusion

Introduction: In a pure metal, diffusion operates
by the vacancy mechanism and the self-diffusion coefficient
is determined by the frequency with which an atom will jump
into a vacant neighboring site, $\omega$, and by the probability
that a given neighboring site is vacant, $P_v$. These experi-
ments are usually performed by plating a very thin film of
radioactive tracer on a pure metal, annealing, sectioning,
and using the thin film solution to determine $D_s$.\textsuperscript{13} This
$D_s$ is called the self-diffusion coefficient of the solute
in the given solvent, and the experiments are done ideally
at "infinite dilution." For the infinitely dilute alloy, the
problem is then to estimate whether and by how much $\omega$ and
$P_v$ for a solute atom differs from $\omega$ and $P_v$ for a solvent
atom.

C-1.1 Mathematical Model: A general approach to the
derivation of the diffusion coefficient is to consider the
problem as a whole sequence of jumps that result in paths of
atoms migrating through the lattice rather than merely
jumps between two planes. This concept is referred to as
the "random walk" approach.
Consider that successive atom jumps are vectors \( \vec{r}_1, \vec{r}_2, \text{etc.} \). If \( r \) is the total number of completed jumps which an atom makes per unit time, then after a time \( t \), a total of \( n = rt \) completed jumps will have occurred, and the atom will have moved an average distance, \( \overline{R(t)} \)

\[
\overline{R(t)} = \Sigma \vec{r}_i \tag{6}
\]

from its initial position. It may be shown that the diffusion coefficient, \( D \), is obtained by

\[
D = \frac{\overline{R(t)^2}}{6t} \tag{7}
\]

where

\[
\overline{R(t)^2} = |\overline{R(t)} \cdot \overline{R(t)}|.
\]

Now

\[
\overline{R(t)^2} = \Sigma (\vec{r}_i)^2 = \vec{r}_i^2 + 2\Sigma \Sigma \vec{r}_i \cdot \vec{r}_{i+j} \tag{8}
\]

For the case of the fcc, bcc, and hcp crystal systems, all the jump vectors will be of equal magnitude, \( r \). Then equation (8) may be written as

\[
\overline{R(t)^2} = nr^2 + 2r^2 \Sigma (n-j) \cos \theta_j \tag{9}
\]

where \( \cos \theta_j \) is the average value of the cosine of the angle between the \( i \)th and the \( (i+j) \)th atom jump.

If the direction of each jump is independent of all earlier jumps, then \( \cos \theta_j = 0 \) and substituting equation (9) into equation (7) we obtain

\[
D = \left( \frac{1}{6} \right) nr^2. \tag{10}
\]

The factor \( \left( \frac{1}{6} \right) \) come from the fact that only 1/3 of the randomly directed vibrations are in the required direction, and then substituted into equation (5).
As an example, consider an fcc lattice. For this case \( r = 12\omega N \), where 12 is the number of nearest neighbors, \( \omega \) is the jump frequency, and \( N_v \) is the fraction of vacant sites. With a lattice constant \( a_0 \), for vacancy diffusion

\[
D = a_0^2 N_v \omega ,
\]

while for interstitial diffusion

\[
D = \delta a_0^2 \omega
\]

with \( \delta \) being a geometric constant.\(^{14}\)

The problem of determining \( D \) has now been reduced to finding a way of calculating \( N_v \) and \( \omega \).

C-1.2 Correlation Effects: The assumption of uncorrelated motion made in the derivation of equation (9) is valid for interstitial diffusion in dilute alloys. However, it is not valid in the case of vacancy or interstitialcy diffusion. Consider vacancy diffusion, after an exchange between an impurity atom and a vacancy all of its neighbors are not identical; one of them consists of a vacancy, it is most probable that the impurity atom will jump back to the position occupied by the vacancy. In other words, the mean square displacement for the impurity atom after \( n \) jumps, \( R_i^2 \), will be less than that for a vacancy which took the same number of jumps, \( R_v^2 = nr^2 \). The ratio of these two quantities defines the correlation factor, \( f \), as

\[
f = \lim_{n \to \infty} \frac{R_i^2}{R_v^2} \quad .
\]
The correlation factor has been calculated by Compaan and Haven. They demonstrated that the correlation factor could be written as

\[ f = \frac{1 + \cos \theta_i}{1 - \cos \theta_i} \]  \hspace{1cm} (12)

where \( \cos \theta_i \) refers to the average of the cosine of the angle between consecutive jumps. For vacancy diffusion in a fcc lattice, where the terms have been previously defined,

\[ D = a_0^2 f \omega N_v . \]

C-1.3 Calculation of \( N_v \): According to classical thermodynamics, a given system tends to assume a configuration such that the free energy of the system is a minimum. This free energy \( G \) is related to the enthalpy, \( H \), temperature in degrees Kelvin, \( T \), and entropy, \( S \), of the system by

\[ G = H - TS \]  \hspace{1cm} (13)

The entropy of any system may be divided into two parts, the thermal entropy \( S_t \) and the configurational entropy \( S_c \). The thermal entropy \( S_t \) is determined by the number of different ways \( W_t \) in which the thermal energy of the crystal may be distributed over the possible vibrational modes of the individual atoms. \( W_t \) is related to \( S_t \) by the Boltzmann relation, where \( k \) is Boltzmann's constant,

\[ S_t = k \ln W_t . \]  \hspace{1cm} (14)

The configurational entropy of a crystal \( S_c \) is determined by the number of different ways \( W_c \) in which the atoms of the
crystal may be arranged over the available number of lattice sites. For the case of vacancies in a crystal, let $N_a$ be the number of atoms in the crystal and $n_v$ be the number of vacancies, and let all of the lattice sites be equivalent. It can be then shown that this case\textsuperscript{16}

$$W_C = \frac{(N_a + n_v)!}{(N_a! \cdot n_v!)}$$ \hspace{1cm} (15)

Again the relation between $S_C$ and $W_C$ is given by the Boltzmann relation

$$S_C = k \ln W_C = k \ln \left( \frac{(N_a + n_v)!}{(N_a! \cdot n_v!)} \right)$$ \hspace{1cm} (16)

Now assume the energy to create a vacancy in a crystal to be $h_v$. The enthalpy of a single crystal containing $n_v$ vacancies is increased by a factor $n_v h_v$ over that of a perfect crystal. For reasons given below, the thermal entropy will increase by a factor $n_v A_s$ when $n_v$ vacancies are added to the crystal. Thus, from these considerations and from equations (13) and (16), the free energy of an imperfect crystal as a function of $n_v$ and $T$ becomes,

$$G(n,T) = G_{\text{perfect}}(T) + n_v h_v - n_v T A_s - k \ln \left( \frac{(N_a + n_v)!}{(N_a! \cdot n_v!)} \right)$$ \hspace{1cm} (17)

In order to find the equilibrium value of $n_v$, we note that at equilibrium $\frac{\partial G}{\partial n_v} = 0$. Using this fact, and the approximation that $\ln x! = x \ln x$ for $x < 1$, we have from (17)\textsuperscript{17}

\footnote{Note: Lower case letters indicate a property of a single crystal, while a capital letter indicates a property of an entire crystalline structure.}

18
\[
\frac{n_v}{n_v + N_a} = \frac{n_v}{N_a} = \exp \frac{\Delta S_t}{k} \exp \frac{-H_v}{RT} .
\]  

(18)

Here, \((n_v/N_a)_{eq}\) is the equilibrium atomic fraction of vacancies in the crystal. In terms of the equilibrium mole fraction of vacancies \(N_v\), equation (18) may be written

\[
N_v = \exp \frac{\Delta S_t}{R} \exp \frac{-H_v}{RT} .
\]  

(19)

In equation (19), \(\Delta S_t\) and \(H_v\) refer to the thermal entropy and enthalpy changes of the entire crystal due to the addition of \(N_v\) vacancies.

It should be noted that the foregoing derivation assumed that \(s_t\) and \(h_v\) are independent of \(n_v/N_a\), which would be true if \(n_v/N_a\) or \(N_v\) were very small. Experimentally, \(N_v\) has been found to be about \(10^{-4}\) in pure metals, so this assumption is a reasonable one.

The enthalpy of a crystal should change with the addition of a vacancy, and this energy change equals approximately the energy necessary to remove an atom from the interior of the crystal and place it on the crystal surface. The reason for the change in \(S_t\), however, is more subtle. One way to estimate this change is to assume an Einstein model for the solid consisting of \(N\) atoms equally distributed over \(3N\) harmonic oscillators. When \(v\) is the Einstein frequency of the atoms in a perfect crystal and \(h v \ll k\), where \(h\) is Planck's constant, and \(k\) is Boltzmann's constant, then \(S_t\), the thermal entropy can be shown to be

19
\[ S_t = 3N_a k \left( |1 + \ln \left( \frac{kT}{\hbar \nu} \right)| \right) \]

for that perfect crystal. In a crystal containing \( n_v \)
the atoms around a vacancy will have a vibrational frequency \( \nu' \) less than \( \nu \) because the atomic restoring forces are reduced. Again using the Einstein model, each atom neighboring a vacancy is assumed equivalent to 3 harmonic oscillators of frequency \( \nu' \). For \( N_a \) atoms and \( n_v \) vacancies in a crystal, we have

\[ 3n_v x \text{ oscillators of frequency } \nu \]
\[ (3N_a - 3n_v x) \text{ oscillators of frequency } \nu' \]

where \( x \) is the number of nearest neighbor atoms around each vacancy. Applying this to equation (20), we have that the thermal entropy of an imperfect crystal is:

\[ S_t = 3n_v x k \left| 1 + \ln \left( \frac{kT}{\hbar \nu} \right) \right| + (3N_a - 3n_v x) k \left| 1 + \ln \left( \frac{kT}{\hbar \nu'} \right) \right| \]

(21)

The change in thermal entropy due to the addition of \( n_v \) vacancies to a perfect crystal can be estimated by subtracting (20) from (21) and dividing by \( n_v \) to obtain

\[ S_t = 3x k \ln \left( \frac{\nu}{\nu'} \right) \]

(22)

Thus the change in thermal entropy is due primarily to the change in vibrational frequency of the atoms around the vacancies.
Calculation of $\omega$: While the calculation of $N_v$ is a straightforward one, accurate determination of the atomic jump frequency for a tracer atom is quite difficult. In fact, the accuracy of the present calculations of $\omega$ is so poor that such calculations provide no real verification of experimentally determined diffusion coefficients. However, the following relatively simple theoretical treatment of $\omega$ yields a numerical value of this parameter within at least an order of magnitude, and it further provides some insight into the temperature dependence of this quantity.

First, assume a simplified sequence of atomic motions required for a tracer atom to jump from one lattice site to another, as shown in sequence Figures 2a, 2b, and 2c. Before such a sequence can occur, the tracer atom must attempt a jump in the direction of the vacant site, with the two restraining atoms simultaneously moving far enough apart to let the tracer atom slide between them. The atomic configuration shown in Figure 2b is called an activated complex, and consists of a region containing a diffusing atom between two equilibrium sites. The number of atoms diffusing per second in one mole of solvent material equals the mole fraction of activated complexes $N_m$, multiplied by the average velocity of the diffusion atoms moving between the restraining atoms $v$, divided by the width of the barrier created by these restraining atoms $\delta$. 

21
Free Energy Variation of Lattice During Sequence of Atomic Motions Shown in a, b, and c

MECHANISM FOR AN ATOMIC JUMP INTO A VACANCY

Figure 2
The average jump frequency per atom, \( \omega \), becomes

\[
\omega = \frac{N_m \tilde{v}}{\delta}
\]  

(23)

To find an approximate value of \( N_m \), it is assumed that diffusing atoms enter, remain at, and leave the activated complex slowly so that the remainder of the lattice has time to continually relax to an equilibrium state. Using this assumption, the work done during the resulting reversible, isothermal process of moving an impurity atom from one site to another is just equal to the change in the Gibbs free energy for the region, \( G_m \). From these assumptions, at any temperature, there will always be an equilibrium number of activated complexes. From the definition of \( G_m \), in equation (13)

\[
G_m = H_m - T S_m
\]

(24)

Since the lattice was assumed to be in thermal equilibrium, \( G_m \) will have all of the properties of \( G_v \) of equation (17). By a process identical to the derivation of equation (19) from equation (17), \(^{22}\) we can obtain the equilibrium mole fraction of activated complexes at any instant:

\[
N_m = \exp \left| \frac{-H_m + T \Delta S_m}{RT} \right| = \exp \left| \frac{-\Delta G_m}{RT} \right| 
\]

(25)

In equation (23), a dimensional analysis will reveal that \( \tilde{v}/\delta \) is a frequency. This frequency is conveniently taken to be the mean vibrational frequency of an atom about its
equilibrium site, $v$, and represents the frequency with which an atom attempts to change sites. From this and from equations (23) and (25) we find

$$\omega = v \exp \left(\frac{-H_m + T\Delta S_m}{RT}\right)$$

(26)

It has been determined experimentally that the activated complexes do not exist long enough for much of the surrounding lattice to reach equilibrium. Thus, it would seem that the basis for the derivation of equation (26) is open to serious question. However, it does yield an order of magnitude agreement with experiment.

C-2 Relationship Between Self-Diffusion and Dilute Impurity Diffusion

The problems involved with impurity diffusion are more complex than those of self-diffusion, since the atoms of both solvent and solute migrate simultaneously. The migration of solute atoms is influenced by that of solvent atoms, and the presence of impurity atoms in the lattice, changes the energy conditions, and thus influences the rate of self-diffusion.

C-2.1 Correlation Effects: In dilute alloys, the vacancies are partially bound to the solute atoms, so that there is in addition to the correlation between the jumps of the solute atoms a correlation between the successive jumps of the vacancies. Thus, it is no longer possible for $P_v$ to
equal $N_v$, although these two quantities are still closely related. Also, the correlation coefficient becomes much less than unity. In this discussion of impurity diffusion, it is assumed that the solute concentration is so small that the diffusion coefficient of the solute in the solvent lattice, or the impurity diffusion coefficient, is independent of the solute concentration. In analyzing the behavior of a solute atom-vacancy pair in a solvent lattice under these conditions, the frequency with which the solute atom and vacancy exchange positions may be taken as $\omega_1$, the frequency with which the vacancy exchanges places with a nearest neighbor atom to the solute atom as $\omega_2$, and the frequency with which the vacancy exchange places with a solvent atom which is not a nearest neighbor to the solute atom as $\omega_3$. Thus $\omega_3$ is proportional to the frequency at which the solute atom-vacancy pair dissociates. Now if the vacancy is tightly bound to the solute atom, $\omega_3$ will be nearly zero, and $\omega_1$ will be much greater than $\omega_2$. In other words, if a vacancy exchanges places with a solute atom with a given jump, the probability that it will reverse that exchange with its next jump is nearly unity. If this is so, the vacancy movement is nearly completely correlated, and the solute atom will move through the lattice only as fast as the vacancy exchanges with the solvent atoms. Thus, the impurity diffusion coefficient for small solute concentrations, $D_I(0)$, will not be given by
\[ D_{l}(0) \sim a_{0}^{2} \omega_{1} N_{V} \]

but by
\[ D_{l}(0) \sim a_{0}^{2} \omega_{2} P_{V} = 2f a_{0}^{2} \omega_{1} P_{V} \ll a_{0}^{2} \omega_{1} P_{V} \] (27)

The relationship between \( P_{V} \) and \( N_{V} \) is discussed on page 35.

In the above equations \( f \ll 1 \), and the correlation effect becomes quite important. Note, the relation between the correlation coefficient, \( f \), and the quantities \( \omega_{1} \), \( \omega_{2} \), and \( \omega_{3} \) is a function of the lattice geometry. As an example, we can consider a face centered cubic lattice, where the expression for \( f \) is given by\(^{24}\)

\[ f = \left| \frac{\omega_{2} + 7 \omega_{3}/2}{\omega_{1} + \omega_{2} + 7 \omega_{3}/2} \right| \] (28)

Here the return of the dissociated vacancy is ignored. The expression for \( D_{l}(0) \) thus becomes

\[ D_{l}(0) = \left| \frac{a_{0}^{2} \omega_{1} (\omega_{2} + 7 \omega_{3}/2)}{\omega_{1} + \omega_{2} + 7 \omega_{3}/2} \right| P_{V} \] (29)

Present day calculations of \( \omega_{2} \) and \( \omega_{3} \) yield ambiguous results, so the best practical approach is to estimate the value of \( D_{l}(0) \) for several limiting cases.\(^{25}\) These are:

(1) the rate of dissociation of solute atom vacancy pairs is much less than the exchange rate between the vacancy and the nearest neighbor to the solute atom, or \( \omega_{2} > \omega_{3} \). Then we have from equation (29)

\[ D_{l}(0) \sim a_{0}^{2} \left| \frac{\omega_{2} \omega_{1} P_{V}}{\omega_{1} + \omega_{2}} \right| \] (30)
(2) the vacancy-solute atom exchange rate is much more rapid than the two vacancy-solvent atom exchange rates, or the vacancy-solute atom pair is tightly bound. Then $\omega_1 \gg (\omega_2 + 7\omega_3/2)$ in (28), and we have

$$D_I(0) \sim a_o^2(\omega_2 + 7\omega_3/2)P_V$$

(31)

(3) the solute atom jumps slowly, or $\omega_1 \ll (\omega_2 + 7\omega_3/2)$ in equation (29). Thus

$$D_I(0) \sim a_o^2\omega_1 P_V$$

(32)

(4) the "solute" is a solvent tracer, so that $\omega_1 = \omega_2 = \omega_3 = \omega$. Then, since here $P_V = N_V$, and $D_s(0) = D_I(0)$

$$D_s(0) = 2fa_o^2N_V\omega.$$ (33)

It must be emphasized that the above results are for a face centered cubic lattice. Other lattice geometries will give results of essentially the same form, but with different numerical constants.

C-2.2 Valence Effects: Due to the electronic effect of an impurity atom with a valence different than that of the surrounding host lattice a vacancy is often attracted to the solute atom. Consider the case of a divalent impurity in a monovalent lattice. According to the free electron theory of metals the free electron density in the immediate environment of the impurity will be greater than that throughout the lattice as a whole, but it will not be great.
enough to completely neutralize the excess positive charge of the solute ion. This ion is said to be "screened," and it is surrounded by an electrostatic potential of the form

$$V(r) = \frac{(Ze/r)}{exp (-qV)}.$$  \hspace{1cm} (34)

Where $r$ is the radial distance from the impurity ion, $Z$ is the number of excess electrons per impurity ion and $q$ is the screening parameter, which gives a measure of the neutralization of the excess charge of the impurity ion.

Let a solvent atom be removed from a site adjacent to the solute ion, and make the assumption that the electron distribution throughout the lattice will not be appreciably distributed by this removal. If this is the case, the vacancy resulting from this removal will have an effective charge of $-e$. Its energy on the site next to the impurity ion then will be reduced by

$$eV(r) = E(r) = (Ze^2/r) \ exp (-qr).$$ \hspace{1cm} (35)

as compared to a site surrounded by normal solvent atoms.

Since the energy of the vacancy is lower next to the impurity ion, the concentration of vacancies on any nearest neighbor site will be increased to

$$P_v = N_v \ exp (E(r)/kT).$$ \hspace{1cm} (36)

This effect will also decrease $H_v$ on a site next to a solute atom by $E(r)$ as compared to $H_v$ on a site surrounded by solvent atoms. This decrease in potential energy also makes
H_v for the solute less than H_v for the solvent. Finally, there is a force e(dV/dr) tending to draw the vacancy and the impurity ion together. If the impurity ion and the vacancy begin to move together, this force tends to pull them together until H_v is decreased by roughly r_o e(dV/dr)_{r=r_o}.  

C-2.3 **Size Effects:** The effects on the impurity diffusion coefficient of the size of the solute atom relative to that of the solvent atom have been considered by Swalin and Overhauser. 

Swalin assumes that the solute atoms are compressible spheres and the solvent lattice is an elastic continuum. For example, if the solute is larger than the solvent atom, H_v will be increased by the additional strain required at the saddle point. This change in H_v is a balance between the strain on the diffusing solute atom and the lattice continuum. Although it would seem that a large solute atom would elastically strain the lattice and thereby attract vacancies and lower H_v, Swalin concluded that the actual change in H_v is negligible to a first order approximation. 

Overhauser considered the effects on the impurity diffusion coefficient of the presence of interstitials in the solvent lattice. Since these interstitials dilate the lattice locally, the average interatomic distance between the matrix atoms will be increased. This lattice expansion will reduce the work required to squeeze
an atom through the saddle point, so $H_v$ should be decreased by the addition of an interstitial solute whose atoms are larger than those of the solvent. Likewise, the addition of an interstitial solute which contracts the solvent lattice should increase $H_v$.

Although the impurity diffusion concept has been studied in detail for many elements, no single correlation between diffusion parameters $D_o$, the frequency factor, and $Q$, the activation energy and the properties such as valence and size has been found to fit more than one or two elements.

C-2.4 Solute Concentration Effects: When the solute concentration becomes greater than some threshold value, (which varies for different systems), the solute atoms begin to interact and the diffusion coefficient for the solute as well as the self-diffusion coefficient for the solvent material becomes a function of the solute concentration. In order to obtain a qualitative concept of the nature of these functions, we consider the previously presented equations for the self-diffusion of the pure solvent, (33), and the dilute impurity diffusion coefficient for the solute, in a face centered cubic solvent lattice, (28), i.e.,

$$D_s(0) = 2f\omega_o^2\omega_o N_v$$

and

$$D_I(0) = 2a_o^2 \frac{\omega_1}{\omega_1 + \omega_2 + \frac{7\omega_3}{2}} P_v .$$
Now if $P_v = N_v$ and $\omega_1 \approx \omega_2 - \omega_3$, then no matter how large $\omega_1$ is, $D_s(0)$ will about equal $D_I(0)$. However, if $D_I(0)$ is greater that $D_s(0)$ in very dilute alloys, it follows that the impurity atoms attract vacancies so that $P_v > N_v$. Then the solvent jump frequencies $\omega_2$ and $\omega_1$ are increased near a solute atom. But if $P_v > N_v$, or $\omega_2 > \omega_0$ near a solute atom, then $D_s$ is also greater near a solute atom. As the solute concentration increases, so does the number of affected solvent atoms. Thus, if $D_I(0) < D_s(0)$, an addition of solute will increase $D_s$. Similarly, if $D_I(0) > D_s(0)$, an addition of solute will decrease $D_s$.

To provide an estimate of the magnitude of the change in the self-diffusion coefficient, $D_s$, with solute concentration, the following empirical equations have been derived for solute concentration $\leq 0.01$:

$$D_s = \left( 1 - A N_I \right) D_s(0) + A B N_I D_I$$

$$D_I = D_I(0)$$

(37)

(38)

Here, $N_I$ is the solute concentration, $A$ is the number of nearest neighbor sites around each atom, and $\beta$ is an arbitrary constant $< 1.0$ which must be experimentally determined. Thus, $D_I$ does not vary much with solute concentration for solute concentrations less than 0.01. For solute concentrations greater than 0.01, the relation between the impurity diffusion coefficient and solute concentration is given by

$$D_I = D_I(0) \left( 1 + \mu N_I \right)$$

(39)
Here, $\mu$ is a constant $< 1.0$. The values of the constants $\beta$ and $\mu$ vary from system to system. Specific theoretical calculations of their values are quite difficult and are not usually attempted.

D. Summary of Equations for Diffusion Coefficient

The following comprise a summary of the most useful previously derived diffusion relationships which are most useful in comparing experimentally observable parameters with theoretical models for the diffusion process.

For vacancy self-diffusion in a pure solvent with correlated tracer jumps, the self-diffusion coefficient, $D_s$, is given by

$$D_s(0) = 2f a_o^2 N_v \omega$$  \hspace{1cm} (40)

where $f$ is the correlation coefficient, $N_v$ is the equilibrium vacancy concentration, $\omega$ is the tracer jump frequency, and $a_o$ is the lattice constant. For dilute impurity vacancy diffusion, the diffusion coefficient of the solute in the solvent lattice is

$$D_1(0) = 2f a_o^2 \omega_1 P_v$$  \hspace{1cm} (41)

Here $\omega_1$ is the frequency with which a solute atom changes places with a nearest neighbor vacancy, and $P_v$ is the probability that a nearest neighbor site to the impurity atom is vacant. Because of attractive forces between the vacancies and solute atoms have already been discussed, $P_v$
in equation (27) will not equal \( N_V \). An estimate of the value of \( P_v \) in terms of \( N_V \) for a given solvent-solute system may be obtained from equations (33), (34), and (35). The coefficient of correlation \( f \) is a function of lattice geometry. For self-diffusion, \( f \) lies between 0.50 and 0.79;\(^\text{15}\) for a face-centered cubic solvent structure, \( f \) for impurity vacancy diffusion is given by

\[
f = \frac{\omega_2 + 7\omega_3/2}{\omega_1 + \omega_2 + 7\omega_3}
\]

(42)

Here \( \omega_2 \) is the frequency with which a vacancy exchanges position with a nearest neighbor atom to a solute atom, and \( \omega_3 \) is the frequency with which a vacancy exchanges position with a solvent atom, which is not a nearest neighbor to a solute atom.

From equilibrium thermodynamics, the equilibrium vacancy concentration in a crystal was found to be

\[
N_V = \exp \frac{\Delta S_t}{R} \exp \frac{-H_V}{RT} \quad \text{.}
\]

(43)

In a similar manner, an equilibrium approximation allowed us to arrive at an expression for

\[
\omega = \nu \exp \frac{-H_m + T\Delta S_m}{RT} \quad \text{.}
\]

(44)

Combining equations (19), (26), and (40) allows us to write for the vacancy self-diffusion coefficient in a pure or very dilutely contaminated solvent
Likewise, combining equations (19), (26), (27), and (35), we find for dilute impurity vacancy diffusion

\[
D_L(0) = 2fa_0^2 v \exp \left( \frac{\Delta S_f + \Delta S_m}{R} \right) \exp \left( \frac{H_f - H_m}{RT} \right).
\]  

(46)

Here \( E(r) \) may be estimated from equation (34), and \( f \) for a fcc lattice is given by equation (28). Generally, \( f \) will be given in terms of the various jump frequencies described above. Of these frequencies, only \( \omega_1 \) can be calculated with reasonable certainty; because of uncertainties in the values of \( \omega_2 \) and \( \omega_3 \), the usual practice is to consider the behavior of \( D_L(0) \) in the limiting cases given in equations (30), (31), and (32).

Equations (37), (38), and (39) give the variation in the solvent self-diffusion coefficient \( D_S(0) \), and vacancy impurity diffusion coefficient, \( D_I(0) \), with solute concentration. Finally, for interstitial diffusion, the diffusion coefficient, \( D \), is given by

\[
D = a_0^2 v \exp \left( \frac{\Delta S_m}{R} \right) \exp \left( \frac{H_m}{RT} \right).
\]  

(47)

Thus, interstitial diffusion can be considered a much simpler process than vacancy diffusion. For the present investigation it is probable that both vacancy and interstitial diffusion mechanisms take place.
Equations (45), (46), and (47) are useful in calculating values of $D$ from theoretical models of the diffusion process. To check the validity of a given model, a comparison is made of the calculated values of the factors comprising $D$ with corresponding experimentally determined values. Several workers have devised experimental methods for determining the individual terms of equations (45), (46), and (47), such as $\Delta S_f$, $\Delta S_m$, $H_m$, etc.. However, the most common procedure is first to express these equations in the form

$$D = D_0 \exp \left( \frac{-Q}{RT} \right) \quad (48)$$

Then, after determining the diffusion coefficients of the system under consideration at different temperatures, $Q$ may be found from the slope of the plot of the natural logarithm of $D$ versus (temperature,$^0K$)$^{-1}$, while the intercept of this plot with the ordinate axis yields $D_0$. The quantity $Q$ is called the activation energy, and is the sum of the enthalpy terms in each of the equations for $D$, (45), (46), and (47). This quantity is thus associated with the energy necessary for the solute ion to make a single jump in the solvent lattice. The symbol, $D_0$, called the frequency factor, contains the correlation coefficient, jump attempt frequency, lattice parameter, and the entropy changes associated with an atomic jump. Hence, $D_0$ gives a measure of the disorder induced in the solvent lattice by an atomic jump.
The "random walk" is not the only approach for the explanation of the phenomena of diffusion. Some of the more interesting theories are presented by: Feit discusses a dynamical theory, Huntington and Seitz with a discussion from the modern theory of metals, Lazarus discusses the effect of "screened" electrons on diffusion, Johnson presents a "hole" theory for diffusion, while Zener presents a model for ring diffusion. All of these studies present arguments which are valid for some very limited cases and were therefore not used in this work. For further reading on diffusion, the studies by LeClaire, Wert and Zener, Turnbull and Hoffman, and Zener are suggested.
This research depended upon mass spectrometric instrumentation of high sensitivity. It was also conditioned by the type and form of samples which were amenable to diffusion analysis by mass spectrometric means.

The samples of zircaloy-2, tantalum, and 304 stainless steel ribbon were obtained from H. Cross Company of Weehawken, New Jersey, with a detailed analysis of the composition of each sample material. These appear in Tables I, II, and III.

The mass spectrometer used in this research was a 90° double focusing instrument with a resolution of approximately 1/1000, designed and built by F. A. White and J. C. Sheffield.40

A. Description of the Filament

A single filament thermal ionization source, Figure 3, was used in all experiments. Each of the three sample materials (304 stainless steel, zircaloy-2, and tantalum) were used as the filament material. The ribbons of the zircaloy-2 and tantalum were 0.030" (0.076 cm) x 0.001" (0.0025 cm), while the 304 stainless steel ribbon was 0.030" (0.076 cm) x 0.0005" (0.0013 cm).

Hermetically sealed, insulated feedthrus, on which the filament ribbons were spot-welded, were obtained from Electrical Industries, Murray Hill, New Jersey. The
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Table II

COMPOSITION OF TANTALUM

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<td>&lt; 0.3</td>
</tr>
<tr>
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<td>0.5</td>
<td>Mn</td>
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<td>Ag</td>
<td>&lt; 0.4</td>
<td>Co</td>
<td>&lt; 0.3</td>
</tr>
<tr>
<td>Si</td>
<td>0.7</td>
<td>Fe</td>
<td>10.0</td>
<td>Cd</td>
<td>&lt; 0.3</td>
<td>Al</td>
<td></td>
</tr>
<tr>
<td>P</td>
<td>0.3</td>
<td>Ni</td>
<td>1.0</td>
<td>In</td>
<td>&lt; 0.2</td>
<td>others</td>
<td>&lt; 0.1</td>
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METHOD OF PREPARATION: Electron Beam Zone Refined

STANDARD SINGLE CRYSTALS

<table>
<thead>
<tr>
<th>DIAMETER</th>
<th>$/IN. (1&quot; Minimum Order)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1/4&quot;</td>
<td>$100.00</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>$125.00</td>
</tr>
<tr>
<td>1/4&quot;</td>
<td>$150.00</td>
</tr>
<tr>
<td>1/4&quot; (.450&quot;)</td>
<td>$225.00</td>
</tr>
</tbody>
</table>

Random Orientation. Oriented crystals, $75. additional orientation charge per crystal, independent of length. (100), (110), and (111) available.

SPECIAL APPLICATION CRYSTALS

Cut from 1/8" nominal diameter (elliptical cross-section) to approximately 1/16" thickness, one face polished strain-free to expose specified crystallographic plane:

Within 1° $650./slice; additional slices of same orientation $200. each.
Within 4° $325./slice; additional slices of same orientation $150. each.

POLYCRYSTALLINE RODS

<table>
<thead>
<tr>
<th>DIAMETER</th>
<th>IN./LB.</th>
<th>GMS/IN.</th>
<th>&lt;100 GMS</th>
<th>100-300 GMS</th>
<th>1-3 LBS</th>
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<tr>
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<td>$3.00</td>
<td>$1.70</td>
<td>$550.</td>
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<tr>
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<tr>
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<td>350.</td>
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>1 lb. on request.
Minimum Order: $50.00
TABLE III

COMPOSITION OF ZIRCALOY-2

ZIRCALOY-2 INGOT
HEAT NO. 387266
COMPOSITION IN PERCENT

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<tbody>
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<td>Zr</td>
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IMPURITIES IN PPM

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<td>&lt;25</td>
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</tr>
<tr>
<td>V</td>
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<td>&lt;25</td>
</tr>
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</table>

INGOT HARDNESS, BHN

<p>| | |</p>
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<tr>
<td>Average</td>
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40
SINGLE FILAMENT THERMAL IONIZATION SOURCE

FIGURE 3
feedthrus are designated as EI type A-40W-SS Mod H.

B. The Mass Spectrometer

The mass spectrometer separates chemical elements according to their mass to charge ratio. The mass spectrometer shown in Figure 4 was used exclusively in this study. A schematic of this two-stage, double focusing instrument is shown in Figure 5. The major components of the instrument include:

(a) Ion source
(b) Electrostatic analyzer
(c) Magnetic analyzer
(d) Beam defining slit system
(e) Ion detector and counting system
(f) Vacuum system

Thermal Ionization. For mass spectrometric analysis, some method is needed to vaporize and ionize the sample into a beam of positively charged ions before entering the electrostatic and magnetic filters. For the analysis of metals, thermal ionization is an effective means of producing gaseous ions especially if the metal has a low ionization potential compared to the filament's work function. As the sample vaporizes, some atoms or molecules will leave the surface of the filament with the loss of one or more of their electrons. The hot filament surface has a higher affinity for electrons than the metal itself. This is due in part, to the intrinsic nature of the metallic crystal along with the low binding energy of the outer shell electrons of most metallic atoms.
TWO-STAGE MASS SPECTROMETER

FIGURE 4
Two Stage, Double Focusing Mass Spectrometer

Figure 5
The probability of an atom evaporating from a surface in the ionic state rather than as a neutral atom is expressed by the relation,

\[
\frac{n^+}{n^0} \propto \exp \left( \frac{e(\phi - \text{IP})}{kT} \right) \propto \exp \left( \frac{11606(\phi - \text{IP})}{T} \right)
\]  

(49)

where \(n^+\) is the number of positive ions, \(n^0\) is the number of neutral specie, \(e\) is the electron charge, IP is the ionization potential of the sample in electron volts, \(\phi\) is the surface work function of the filament material in electron volts, \(k\) is Boltzmann's constant, and \(T\) is the surface temperature in \(^{0}\text{K}\).

It can be seen that for elements where \(\text{IP} > \phi\) the relative number of ions to neutrals produced will increase with temperature.

Ions resulting from thermal ionization on the filament surface are accelerated to high energy and defined by collimating slits in the ion source. These high energy ions pass through the electrostatic analyzer where they are energy filtered. Ions with a discrete kinetic energy (accelerating voltage) including a small energy spread pass through the center defining aperture. The defining slit determines the energy spread accepted. This filtered beam of ions then enters the magnetic analyzer where the beam is momentum filtered. Those ions of appropriate energy and proper momentum (essentially mass) come to a focus at the final slit and strike an electron multiplier.
Current pulses result from ion-electron conversion and subsequent secondary electron amplification in the electron multiplier tube. Output pulses are further amplified and counted by an amplifier/discriminator/scalar system. Thus, ions of the same mass are individually counted and the exceedingly small ion currents can be quantitatively measured.

By properly changing the accelerating potential of the ion source, ions of other mass numbers are focused on the electron multiplier. By comparing counting rates for each mass position the isotopic ratio can be quantitatively determined.
Part IV
EXPERIMENTAL METHOD

A. General Experimental Techniques

From the literature, the most common method for obtaining diffusion coefficients is that of serial sectioning. However, a mass spectrometric approach was chosen for this work. The basic concept of the mass spectrometric technique for determining diffusion coefficients is that a correlation exists between the solute atoms moving through a solvent lattice and the fraction of these atoms which are evaporated, as ions, from the crystal surface at a given temperature.

In this work, a thin ribbon filament of solvent material is used with its own impurity or solute concentration. No doping of the solvent was required and thus there was no alteration of the crystal lattice. Since the mass spectrometer source chamber pressure is well below the vapor pressure of the solute material in the temperature range considered, the solute atoms diffuse to the filament surface, and evaporate immediately. This results in a zero solute concentration at the filament surface after the filament temperature is raised to the desired level. Further, as the ribbon thickness is much less than its length or width, the diffusion problem may be considered to be one dimensional.

Application of Fick's Second Law for Diffusion in a Thin Plate: Since the solute concentrations
under consideration are very small, the diffusion process occurring in the ribbon may be described by

\[
J(x,t) = -D \frac{\partial C(x,t)}{\partial x} \quad (50)
\]

and

\[
\frac{\partial C(x,t)}{\partial t} = -D \frac{\partial^2 C(x,t)}{\partial x^2} \quad (51)
\]

where \(J(x,t)\) is the solute flux, \(C(x,t)\) is the solute concentration, and \(D\) is the diffusion coefficient. For the experimental conditions previously described, the solution to Fick's second law for impurity evaporation from a thin plate yields the solute concentration, as a function of position in the filament and evaporation time, \(t\).

Solving equation (51) by the method of separation of variables, the solute concentration may be found to be

\[
C(x,t) = ( A \sin \lambda x + B \cos \lambda x ) \exp(-\lambda^2Dt) \quad (52)
\]

where \(\lambda^2 = -(1/dT)(dT/dt)\) or \(-(1/X)(d^2X/dx^2)\), \(D\) is the diffusion coefficient, \(t\) is time, and \(A\) and \(B\) are constants of integration.

For solute depletion from a thin solvent plate with an initial solute concentration \(f(x)\) across its thickness, the boundary and initial conditions become

\[
C = 0 \quad \text{at} \quad t \geq 0, \quad \text{for} \quad x = h \quad \text{and} \quad x = 0
\]

\[
C = f(x) \quad \text{at} \quad t = 0, \quad \text{for} \quad h > x > 0
\]

With these conditions, equation (52) becomes

\[
C(x,t) = \frac{2}{h} \sum \sin \frac{n \pi x}{h} \exp \left\{ -\frac{D\pi^2 n^2 t}{h^2} \right\} \int f(x') \sin \frac{n \pi x'}{h} \, dx' \quad (53)
\]
Where \( h \) is the filament thickness. Thus from Fick's first law, equation (50), the solute atom flux at the filament surface (\( x=0 \)) becomes

\[
J(0,t) = \frac{2}{h} \sum \exp \left( -\frac{D\pi^2 n^2 t}{h^2} \right) f(x') \sin \frac{n\pi x}{h} dx' \tag{54}
\]

or more conveniently, where \( n \) is an integer,

\[
J(0,t) = A \sum \exp \left( -\frac{D\pi^2 n^2 t}{h^2} \right) \tag{55}
\]

where \( A \) is a constant. For \( h^2 < 16D t \) the error in using only the first term is less than 1\%.\(^{41}\)

**Positive Ion Emission from a Hot Metal Filament:**

The combination of high temperature and solvent work function causes a constant fraction of the evaporating solute atoms to ionize at the filament surface at a given temperature. According to the Langmuir-Saha equation, the ratio of the number of emitted positive ions \((n^+)\) to emitted neutrals \((n^0)\) is

\[
n^+/n^0 = B \exp \left( \frac{(\phi - IP)}{kT} \right) \tag{56}
\]

where \( B \) is a proportionality constant, \( \phi \) is the filament work function, \( IP \) is the ionization potential for the solute, \( k \) is Boltzmann's constant, and \( T \) is the temperature of the filament in degrees Kelvin.\(^{42}\)

The mass spectrometer accelerates, mass analyzes, and detects these emitted solute ions while excluding all other ionic species from its detector. By combining equations (54) and (55), and assuming that the atomic transport time is greater than \( h^2/16D t \), the time variation of the
detected ion current is

\[ J(0,t) = A \exp \left( -\frac{\pi^2 D t}{h^2} \right) \]  

(57)

B. Experimental Procedures

The procedures used in this work are detailed below to allow reproduction of the experiments. A general procedure followed at all times was the wearing of lintless gloves when handling anything going into the vacuum environment. The procedures described are:

1. Cleaning
2. Spotwelding
3. Mass spectrometry
4. Calibration of mass spectrometer
5. Filament temperature measurement
6. Data acquisition
7. Treatment of data
8. Error analysis

1. Cleaning:
   
   **Filament holders**: Clean metal surfaces with 4M HNO₃, and rinse with de-ionized water. Dry under the infrared lamp.
   
   **Filament legs**: Place in an alcohol bath immersed in the ultrasonic cleaner. Rinse with de-ionized water and dry under an infrared lamp.

2. Spotwelding:

   By visual inspection, make sure electrodes are clean
   
   Set to 18-25 watt-seconds
   
   Place one end of filament next to the supporting legs and then spotweld
   
   Do the same for the other end of the filament and leg

50
3. **Mass Spectrometer:**

**Opening Source Region:**

- Close main gate valve, isolation valve, and the roughing valve
- Open the foreline valve
- Connect up nitrogen gas, turn on, and let it into the source chamber
- Let chamber come to atmosphere, put clips on O-ring
- Before dismantling anything, make sure power is OFF
- Screw filament assembly onto the top plate of the ion source.

**Closing Source Section and Vacuum Pump Down:**

- Make sure no lint is on O-ring or glass, and then put glass back on
- Turn nitrogen gas off and close inlet valve
- Open roughing valve, let it run for a few minutes
- Close roughing and open foreline valve
- Turn on first ionization gauge
- Make sure liquid nitrogen is in cold trap
- Open up the diffusion pump valve, do this SLOWLY and watch pressure gauge
- Turn on automatic ionization gauge

**Opening System and Operation:**

- Open isolation valve to system, after chamber pressure has reached $5 \times 10^{-7}$ torr
- Set electron multiplier tube to -4500 volts
- Close gate valve on multiplier tube's ion pump
- Turn on polarity switch to minus, on the electrostatic lens power supply
- Turn on main power to mass bridge (ion source voltage divider)
Turn voltage control on filament heater, adjust current control for the temperature setting

Turn up voltage to desired setting, CAREFULLY

Begin data acquisition

System Shut Down:

Turn all voltage settings OFF
Close isolation valve
Open ion pump to electron multiplier tube
Turn multiplier tube voltage to -500 volts
Close diffusion pump gate valve
Turn OFF all power

Note: Liquid nitrogen will evaporate away

4. Calibration of the Mass Spectrometer:

It is necessary to calibrate the mass spectrometer before undertaking any experiments measuring diffusion coefficients. Using equation

$$B^2R^2 = (144)^2(mV_a)$$

where $B$ is the magnetic field in gauss, $R$ is the radius of curvature (in cm.) of the magnetic analyzer, $m$ is the mass of the isotope of interest in amu., and $V_a$ is the accelerating potential in volts, the appropriate magnetic field-accelerating potential values for a given mass can be determined.

Since the alkali metal atoms were to be examined, calibrations were made using filaments which had solutions of either rubidium or cesium evaporated on them. Rubidium has two naturally occurring isotopes, mass 85 and mass 87. Detection of one mass and calculation of the magnetic field allowed the determination of the accelerating potential of
the other isotope with the same magnetic field. In practice, an accurate calibration requires many repetitive measurements. Cesium-133 is the only stable isotope of cesium, and provides an accurate determination of the magnetic field.

Ion implanted, as well as micro-pipetted filaments of Cs-133 were also used for calibration purposes. The identical method was followed.

5. Filament Temperature Measurement:

Using a precision optical pyrometer, filament temperatures were determined by visually matching the color of the filament with the color of a calibrated, resistively heated wire within the pyrometer. It is accurate to within ±3°C.

6. Data Acquisition Procedure:

The mass spectrometer data was obtained as follows:

1. The sample filament was placed in the ion source and the source chamber pressure was evacuated to a pressure of about 10⁻⁷ torr.

2. The source voltage was set at the accelerating potential of the desired diffusing isotope.

3. The filament was raised in temperature until ions were detected at the given accelerating potential. The filament temperature was then determined.

4. The ion emission rate as a function of time was monitored until a slowly decreasing emission rate was observed. The latter indicated that the maximum emission rate had occurred and the exponential decay portion predicted by diffusion theory, had been reached.

5. At this point, the ion current was recorded for one second every four seconds utilizing a frequency-period-counter which has an accuracy of ±1 count.
6. The procedure was repeated for successively higher temperatures.

7. Treatment of Data:

A graph of the natural logarithm of the impurity ion current (represented by counts per second) from a particular solvent lattice versus time, for each temperature was plotted.

The slope of the "linear best fit" line through the data was calculated.

The diffusion coefficient was then calculated by

\[ \text{slope of the line} \cdot \left( \frac{\text{thickness of filament}}{\pi} \right)^2 = -D \]

A graph was made of the ln D versus \(10^4/T{\,^{\circ}\!\!\!{\text{C}}}\).

Utilizing a PDP 8/I computer, the linear best fit line to the data points was made. The y-intercept of this line is the frequency factor, \(D_0\), and was calculated by the computer. The correlation coefficient of the data to the line was also calculated.

The activation energy, Q, was determined by multiplying the slope of the line in the preceding step by the Rydberg gas constant in units of kcal/mole/\(^{\circ}\!\!\!{\text{C}}}.

8. Error Analysis:

Programs were written to provide the linear best fit line from a set of data points. The program also gave the degree of fitness in the form of a correlation coefficient. All error values were determined by calculating the product-moment coefficient of correlation and using a 95% confidence level. A final program was written to obtain the standard deviation for the activation energies. These programs appear in Appendix 1 and 2.
Part V
RESULTS AND DISCUSSION

Following the procedure previously indicated, graphs of the ln (alkali metal ion count rate) versus time were obtained. Figure 6 is an example of how the ion count rate varies with time. These superposed curves illustrate the detection of singly charged, mass resolved ions at several temperatures. The counting rates are relative, but most experimental measurements were in the region of \(10^{-14}\) to \(10^{-17}\) amperes, corresponding to counting rates of \(10^5\) to \(10^2\) ions per second.

Tables IV, V, and VI show the experimental results for the diffusion of the alkali metals in 304 stainless steel, tantalum, and zircaloy-2. All five naturally occurring alkali metals were found in tantalum and in 304 stainless steel; however, lithium was not found in sufficient quantities to determine its diffusion parameters in the zircaloy-2 sample.

Activation energy plots for these diffusion systems are shown in Figures 7 thru 20. The frequency factor, \(D_o\), and the activation energy, \(Q\), were calculated from these plots and are presented in Tables VII, VIII, and IX.

The data derived from the activation energy plots can be applied to the metallurgy of the system. For example, a sharp discontinuity can be attributed to an allotropic phase change in the host material. Since pure zirconium has
DIFFUSION OF Rb IN TYPE 304 STAINLESS STEEL

RUN NO.  TEMP °C
1     903
2     968
3     1052
4     1118

TIME (5 SECOND INTERVALS)

DIFFUSION OF Rb IN TYPE 304 STAINLESS STEEL

FIGURE 6
TABLE IV
DIFFUSION COEFFICIENTS OF THE ALKALI METALS IN TANTALUM

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<thead>
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<th>Li-7</th>
<th>Na-23</th>
<th>Rb-85</th>
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<tr>
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<td></td>
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<tr>
<td>$T(\circ K)$</td>
<td>$D(cm^2/sec)$</td>
<td>$T(\circ K)$</td>
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<tr>
<td>1033</td>
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<tr>
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<td>1.32 $\times 10^{-8}$</td>
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TABLE IV CONCLUDED

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<th>$T (^\circ K)$</th>
<th>$D (\text{cm}^2/\text{sec})$</th>
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<th>$D (\text{cm}^2/\text{sec})$</th>
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<td>D(cm²/sec)</td>
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<td>D(cm²/sec)</td>
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**TABLE V**

DIFFUSION COEFFICIENTS OF THE ALKALI METALS IN ZIRCALOY-2
<table>
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<tr>
<th>$T(\text{K})$</th>
<th>$D(\text{cm}^2/\text{sec})$</th>
<th>$T(\text{K})$</th>
<th>$D(\text{cm}^2/\text{sec})$</th>
<th>$T(\text{K})$</th>
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**TABLE VI**

**DIFFUSION COEFFICIENTS OF THE ALKALI METALS IN 304 STAINLESS STEEL**

**Li-7**

**K-39**

**Na-23**

**Rb-85**
**TABLE VI CONCLUDED**

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Table VII

$D_0$ and $Q$ for IMPURITY DIFFUSION in 304 STAINLESS STEEL

<table>
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<tr>
<th>Impurity</th>
<th>$D_0$</th>
<th>$Q$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li-7</td>
<td>$(1.86 \pm 2.01) \times 10^0$</td>
<td>$52.96 \pm 0.54$</td>
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<td>Na-23</td>
<td>$(6.11 \pm 0.56) \times 10^{-7}$</td>
<td>$14.22 \pm 0.19$</td>
</tr>
<tr>
<td>K-39</td>
<td>$(9.53 \pm 0.67) \times 10^{-7}$</td>
<td>$14.09 \pm 0.15$</td>
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<tr>
<td>Rb-85</td>
<td>$(3.19 \pm 0.58) \times 10^{-1}$</td>
<td>$41.92 \pm 0.38$</td>
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<tr>
<td>Cs-133</td>
<td>$(6.75 \pm 0.70) \times 10^{-4}$</td>
<td>$25.04 \pm 0.22$</td>
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928 $< T(°K) < 1228$

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<th>Impurity</th>
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<th>$Q$</th>
</tr>
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<tbody>
<tr>
<td>Li-7</td>
<td>$(1.73 \pm 0.31) \times 10^5$</td>
<td>$84.42 \pm 0.56$</td>
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<td>Na-23</td>
<td>$(1.74 \pm 0.59) \times 10^4$</td>
<td>$74.08 \pm 0.93$</td>
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<td>K-39</td>
<td>$(1.72 \pm 0.10) \times 10^3$</td>
<td>$75.22 \pm 0.18$</td>
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<tr>
<td>Rb-85</td>
<td>$(5.56 \pm 1.75) \times 10^6$</td>
<td>$98.16 \pm 1.01$</td>
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<tr>
<td>Cs-133</td>
<td>$(5.03 \pm 0.49) \times 10^{-5}$</td>
<td>$24.50 \pm 0.26$</td>
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1228 $< T(°K) < 1789$

$D_0$ is in units of cm$^2$/sec

$Q$ is in units of kcal/mole
Table VIII

D₀ and Q for IMPURITY DIFFUSION in TANTALUM

990 < T(K) < 1230

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<td>Li-7</td>
<td>((5.15 \pm 0.91) \times 10^{-5})</td>
<td>20.29 ± 0.41</td>
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<tr>
<td>Na-23</td>
<td>((2.07 \pm 0.27) \times 10^{-6})</td>
<td>11.90 ± 0.37</td>
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<tr>
<td>K-39</td>
<td>((2.71 \pm 0.50) \times 10^{-3})</td>
<td>32.19 ± 0.43</td>
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<tr>
<td>Rb-85</td>
<td>((1.22 \pm 0.37) \times 10^{-1})</td>
<td>36.46 ± 0.72</td>
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<tr>
<td>Cs-133</td>
<td>((8.53 \pm 0.61) \times 10^{-6})</td>
<td>15.07 ± 0.16</td>
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</table>

1230 < T(K) < 1790

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</tr>
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<td>((3.38 \pm 0.35) \times 10^{-5})</td>
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<td>((1.13 \pm 0.17) \times 10^{-6})</td>
<td>15.46 ± 0.47</td>
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<td>((1.06 \pm 0.07) \times 10^{-5})</td>
<td>22.25 ± 0.20</td>
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<td>((2.33 \pm 0.24) \times 10^{-7})</td>
<td>14.78 ± 0.31</td>
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<tr>
<td>Cs-133</td>
<td>((1.18 \pm 0.12) \times 10^{-6})</td>
<td>13.48 ± 0.26</td>
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</table>

D₀ is in units of cm²/sec
Q is in units of kcal/mole
### Table IX

**\(D_0\) and \(Q\) for IMPURITY DIFFUSION in ZIRCALOY-2**

#### \(900 < T(\text{K}) < 1045\)

<table>
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<th>Impurity</th>
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<th>(Q) ((\text{kcal/mole}))</th>
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<tbody>
<tr>
<td>Na-23</td>
<td>((9.76 \pm 2.99) \times 10^{-3})</td>
<td>(26.44 \pm 0.72)</td>
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<td>((2.12 \pm 0.31) \times 10^{-2})</td>
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<tr>
<td>Rb-85</td>
<td>((1.11 \pm 0.26) \times 10^1)</td>
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<tr>
<td>Cs-133</td>
<td>((8.91 \pm 2.96) \times 10^2)</td>
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#### \(1045 < T(\text{K}) < 1345\)

<table>
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<th>(Q) ((\text{kcal/mole}))</th>
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<tbody>
<tr>
<td>Na-23</td>
<td>((1.92 \pm 0.25) \times 10^{-1})</td>
<td>(40.77 \pm 0.37)</td>
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<td>K-39</td>
<td>((3.08 \pm 0.41) \times 10^0)</td>
<td>(46.10 \pm 0.38)</td>
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<td>Rb-85</td>
<td>((3.53 \pm 0.54) \times 10^{-4})</td>
<td>(25.79 \pm 0.44)</td>
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<td>((7.74 \pm 1.01) \times 10^{-2})</td>
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#### \(1345 < T(\text{K}) < 1520\)

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<td>K-39</td>
<td>((5.35 \pm 0.82) \times 10^9)</td>
<td>(111.06 \pm 0.58)</td>
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<td>Rb-85</td>
<td>((1.11 \pm 0.22) \times 10^{-2})</td>
<td>(37.59 \pm 0.75)</td>
</tr>
<tr>
<td>Cs-133</td>
<td>((1.21 \pm 0.30) \times 10^{-2})</td>
<td>(36.74 \pm 0.94)</td>
</tr>
</tbody>
</table>

\(D_0\) is in units of \(\text{cm}^2/\text{sec}\)

\(Q\) is in units of \(\text{kcal/mole}\)
ACTIVATION ENERGY DIAGRAM OF NA-23 IN ZIRCALOY II

Figure 7
ACTIVATION ENERGY DIAGRAM OF K-39 IN ZIRCALOY-II

Figure 8
ACTIVATION ENERGY DIAGRAM OF Ra-85 IN ZIRCALOY-II

Figure 9
ACTIVATION ENERGY DIAGRAM OF Cs-133 IN ZIRCALOY-II

Figure 10
Figure 11

Activation Energy Diagram of Li-7 in 304 Stainless Steel
ACTIVATION ENERGY DIAGRAM OF Na-23 IN 304 STAINLESS STEEL

Figure 12
Figure 13

ACTIVATION ENERGY DIAGRAM OF K-39 IN 304 STAINLESS STEEL
ACTIVATION ENERGY DIAGRAM OF Re-85 IN 304 STAINLESS STEEL
ACTIVATION ENERGY DIAGRAM OF Cs-133 IN 304 STAINLESS STEEL

Figure 15
ACTIVATION ENERGY DIAGRAM OF Li-7 IN TANTALUM

Figure 16
ACTIVATION ENERGY DIAGRAM OF Na-23 IN TANTALUM

Figure 17
ACTIVATION ENERGY DIAGRAM OF Rb-85 IN TANTALUM

Figure 19
ACTIVATION ENERGY DIAGRAM OF Cs-133 IN TANTALUM

Figure 20
both a hexagonal close packed, h.c.p., and a body centered cubic, b.c.c., geometry depending on the temperature, zircaloy-2, which has a composition of ~98.5% zirconium, will probably have a h.c.p. and b.c.c. structure. However, instead of having a single discontinuity, there is a large transition region which possesses its own diffusion characteristics. Pande, Naik, and Agarwala\textsuperscript{43} have also observed this three region activation energy diagram in their study of the diffusion of chromium and iron in zircaloy-2. Their transition region held limits of 780 and 980°C, while this work extends the transition region from 772 to 1072°C. This large transition region can be explained by a statistical average of the diffusivity of both phases present at a particular temperature.

When examining the stainless steels, the results in this work did not completely agree with literature values. A reason for the discrepancy may involve the fact that the composition of stainless steel has a wide range in the percent abundance of its constituent elements. For example, Delong\textsuperscript{44} has found that 304 stainless steel changes from a face centered cubic, f.c.c., to a body centered cubic, b.c.c., structure at 910°C, while the results in this study indicate a phase change at 955°C.

While carbon,\textsuperscript{45} deuterium and hydrogen,\textsuperscript{46} molybdenum,\textsuperscript{47} and sulphur\textsuperscript{48} have been studied as they diffused through tantalum the diffusion of the alkali metals have not been studied in tantalum. Tantalum has a body centered cubic
structure, thus there will be no phase transformation. However, from the data obtained in this experiment it appears that there has been a phenomenological occurrence at approximately 1000°C. There have been many possible explanations suggested, but the one that presents the greatest problem in discounting it is that of cold working. For tantalum to be processed into the thin ribbons used in this study, tantalum would undergo a great deal of cold working. It is possible to remove these defects by annealing, and it so happens that the annealing temperature for tantalum is 1000°C. The resultant reduction in lattice disorder would tend to decrease the diffusion coefficient. However, it would increase when the temperature was again increased. This is exactly what occurs in figures 16 - 20.

These results provide an insight into the basic diffusion mechanism in addition to supplying D, D₀, and Q. For example, the relationship between the diffusion coefficient, D, and the ionic radius at a constant temperature can be compared, by considering the alkali metal series; Li, Na, K, Rh, and Cs, in the same host material. Plots of these five impurity ions as observed in this study are shown in Figure 21 for tantalum, Figure 22 for zircaloy-2, and Figure 23 for 304 stainless steel. Although there is no phenomenological relationship, the diffusion coefficient versus mass is somewhat analogous to sputtering yield data in which very large differences in secondary ion yield vary with atomic mass and ionic radius.

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Each set of figures must be considered separately, as they represent completely different crystalline structures and alloy compositions.

**Tantalum:** Referring to Figure 21 A and B, there is a similarity in the pattern of the constant temperature lines. Independent of the ambient temperature of the material, the diffusion coefficient seems to follow the same pattern of maxima and minima with increasing ionic radius. Similar diffusion patterns can be expected if there were no change in either/or both the diffusion mechanism and the geometric structure of the crystal. There has been no reported phase change in tantalum, which always possesses body centered cubic geometry. Note that as both the ionic radius of the impurity, \( r_{\text{imp}} \) (Å), and \( T(\text{O}K) \) increase, there appears to be an upper limit on the order of \( 10^{-8} \text{ cm}^2/\text{sec} \) for \( D \). At the same time there is a "collimating" effect on \( D \) as \( r_{\text{imp}} \) goes from approximately 1Å to approximately 1.33Å. In an attempt to determine the reason for the behavior at these two points, an interstitial mechanism for diffusion was assumed and the largest sized impurity which would fit through the confining lattice ions for a b.c.c. system was calculated. For a b.c.c. system the following formula was used

\[
r_{\text{impurity ion}} < \frac{a}{\sqrt{2}} - r_{\text{lattice ion}}
\]

(59)

where \( a \) is the edge size. For tantalum, it was found that \( r_{\text{imp}} \) should be less than 1.66 Å. This is extremely close to
Figure 21

D (cm²/sec) vs $r_{\text{imp}}$ (Å) at constant T (°C) in tantalum.
the 1.67 Å point on the graph for the Cs⁺ ion. This implies that the lattice structure of the host material plays a role in limiting the size of the impurity ion's diffusion. To go beyond this size would require additional energy. The region of $1 < r_{\text{imp}}(\text{Å}) < 1.33$ shows what might be considered inhibited diffusion. This could be caused by the simultaneous diffusion of all the other impurities in this dynamic system while being analyzed for just one particular impurity.

**Zircaloy-2:** A very interesting pattern of diffusion can be observed in Figures 22 A, B, and C. Using the 900°K line in Fig. 22 A as a reference, in a h.c.p. structure, as the ionic radius of the impurity increases, the diffusion coefficient decreases. However, as the temperature is increased it is evident that the lattice vibrations cause D to increase and the larger impurity ions to diffuse at a greater rate. This increase in acceptable ionic radius appears to have an upper limit of about 1.47 Å which corresponds closely to the radius of the Rb⁺ ion.

Figure 22 B represents the transition region where the crystal structure changes from h.c.p. to b.c.c.. This graph is not simply understood as the crystal structure changes with temperature. It does appear that the disorder induced in the crystal by the phase change increases the probability of diffusion.

Figure 22 C shows the zircaloy-2 system in its b.c.c. state. Note that D(1250°K) is less than D(1040°K) even though the temperature has been increased into the higher temperature
phase region. A possible mechanism to qualitatively explain this sudden drop in diffusion rate is that as the transition region is approached from a lower temperature, lattice defects begin to occur before the transition point is actually reached. These lattice defects are greatly multiplied in number in the phase transition region due to the increased disorder in the lattice structure. This increases the impurity diffusion rate in the phase transition region. However, once the phase transition is passed, the crystal lattice became more ordered, and many of the lattice defects retreat to defect "sinks," such as grain boundaries. This results in an initial decrease in the diffusion rate. As the defect sinks will release these defects into the crystal lattice as the temperature increases, the diffusion rate will again begin to increase.

Figure 22 C shows an interesting occurrence as both T(°K) and r_{imp} (Å) increase. In the region r_{imp} > 1.47 Å, D is not only relatively invariant as r_{imp} increases, but also relatively invariant as the temperature increases. It seems that as T increases D is constrained to be of the order of magnitude of 10^{-8} cm^{2}/sec. A partial explanation is proposed: if it is assumed that the solute will diffuse by an interstitial mechanism, a relationship for the largest impurity ion accepted before configuration problems are encountered is again equation (59). Using r_{lattice ion} of zirconium, it is determined that r_{imp} < 1.47 Å. From r_{imp} = 1.47 Å in Figure 22C, the diffusion coefficient changes very little.
D(CM^2/SEC) vs $r_{\text{imp}}$ (Å) at constant T(°K) in ZIRCALOY-II

Figure 22
with increasing \( r_{\text{imp}} \). Even increasing the temperature does not cause much of an increase in \( D \) after \( r_{\text{imp}} \) reaches 1.47 Å.

The changing behavior of \( D \) in the range of 1.33 < \( r_{\text{imp}} \) (Å) < 1.47 could possibly be explained by the fact that the zircaloy-2 has a certain percentage (~1.5) of tin, which has an ionic radius of 0.93 Å. This means that for a given diffusion jump through four restraining lattice ions, if one is a tin ion, impurity ions of larger size will be able to diffuse through easier than if all four lattice ions were zirconium. This would represent only a minor perturbation as the amount of tin is limited. At higher temperatures, this represents a negligible effect as \( r_{\text{imp}} \) gets closer to 1.47 Å the size plays an increasingly inhibiting effect.

304 Stainless Steel: Stainless steel 304 presents a special challenge for determining diffusion parameters using a mass spectrometer. No longer is just a pure metal being considered, as in the case with tantalum, or a very dilute alloy as with zircaloy-2, but an alloy which has major constituents of iron, nickel and chromium. The activation energy plots have shown the phase change representing a transformation from f.c.c. to b.c.c. along with the corresponding activation energies. On the other hand, Figures 23 A and B represent a more complex picture in regard to \( D \) as both, \( r_{\text{imp}} \) and \( T \) are varied.

Figure 23 A represents the f.c.c. structure of the stainless steel. As each succeeding temperature line is examined, a pattern in the region of approximately 0.70 Å
and 1.5 Å takes place with an apparent change of slope and vortex, respectively. This can be explained by the fact that as the temperature is increased the lattice vibrations are increased so that it is easier for the impurity of a larger size to diffuse. There is a limit as to the size that will be accommodated. This is primarily controlled by the crystallographic structure of the host lattice. Stainless steel is affected in that there may be unit cells of different host ions, representing the major constituents of stainless steel, and grain boundaries may differ for specific samples. When the effect of Ni in the f.c.c. structure is considered \( r_{imp} \) should be less than 1.07 Å. Configurational limitations do not appear to be significant until \( r_{imp} \) is approximately 1.30 Å and that the region from 1.0 Å to 1.3 Å is a relatively temperature and radius independent region. When \( r_{imp} \) becomes greater than 1.33 Å, the larger size ion cannot diffuse as readily until more energy is supplied to increase the lattice vibrations.

Figure 23 B represents the b.c.c. structure of 304 stainless steel where there are again inflection points which indicate an area of physical constraints. The point \( \sim 1 \) Å is caused by the influence of Ni, while the area around 1.3 Å is caused by the influence of Cr (\( r_{imp} \) should be less than 1.23 Å) and Fe (\( r_{imp} \) should be less than 1.34 Å). After \( r_{imp} \) becomes greater than 1.33 Å, D drops unless more
Figure 23

D(CM²/SEC) vs $r_{imp}$(Å) at constant T(°K) in 304 STAINLESS STEEL

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energy is given to increase the lattice vibrations. There appears to be a significant size effect at approximately 1.67 Å. The increase in D at lower temperatures is probably due to the fact that there are unit cells consisting of different sized lattice ions, so that larger ions than normal could diffuse through. However, this is a limited effect as T increases, the increased lattice vibrations no longer tend to increase D and the larger size of the impurity ion becomes a more dominant parameter.

In fact, there are two aspects to note. The first is that along with the other samples, as both r_{imp} and T increase it appears there may be an upper limit for impurity diffusion coefficients in these b.c.c. structures of approximately $10^{-8}$ cm²/sec. The second is that as in the case of zircaloy-2, after the phase change the diffusion coefficient drops initially and then proceeds to rise with temperature.

As there are no reported values for the diffusion of the alkali metals in 304 stainless steel, it was thought that a comparison between Na-23 and C-14 diffusion in 304 and 316 stainless steel would be desirable. This would enable one to determine whether the results obtained via mass spectrometry were reasonable in comparison to the determination of diffusion coefficients by other methods for a different element. Table X illustrates that the Na-23 diffusion is within an order of magnitude of the diffusion of C-14. The Na-23 also seems to diffuse at a slower rate than the C-14 as would be expected since the Na-23 has a larger ionic radius than C-14. The C-14 diffusion coefficients were determined by Perkins and Carlson.
Table X

C-14 & Na-23 Diffusion in 304 Stainless Steel

<table>
<thead>
<tr>
<th>$T(\text{\textdegree} \text{K})$</th>
<th>$D(\text{ref 49})$</th>
<th>$D(\text{ref 50})$</th>
<th>$D(\text{This Study})$</th>
</tr>
</thead>
<tbody>
<tr>
<td>873</td>
<td>$4.30 \cdot 10^{-10}$</td>
<td>$4.27 \cdot 10^{-10}$</td>
<td></td>
</tr>
<tr>
<td>923</td>
<td>$1.81 \cdot 10^{-10}$</td>
<td>$1.66 \cdot 10^{-10}$</td>
<td></td>
</tr>
<tr>
<td>943</td>
<td></td>
<td>$3.02 \cdot 10^{-10}$</td>
<td></td>
</tr>
<tr>
<td>963</td>
<td></td>
<td>$4.57 \cdot 10^{-10}$</td>
<td></td>
</tr>
<tr>
<td>973</td>
<td>$6.54 \cdot 10^{-10}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>993</td>
<td></td>
<td>$8.71 \cdot 10^{-10}$</td>
<td></td>
</tr>
<tr>
<td>1023</td>
<td></td>
<td>$1.91 \cdot 10^{-9}$</td>
<td></td>
</tr>
<tr>
<td>1049</td>
<td>$3.36 \cdot 10^{-9}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1063</td>
<td></td>
<td>$3.98 \cdot 10^{-9}$</td>
<td></td>
</tr>
<tr>
<td>1093</td>
<td></td>
<td>$7.25 \cdot 10^{-9}$</td>
<td></td>
</tr>
<tr>
<td>1123</td>
<td>$9.54 \cdot 10^{-9}$</td>
<td>$1.02 \cdot 10^{-8}$</td>
<td></td>
</tr>
<tr>
<td>1153</td>
<td></td>
<td>$2.19 \cdot 10^{-8}$</td>
<td></td>
</tr>
<tr>
<td>1198</td>
<td></td>
<td>$4.36 \cdot 10^{-8}$</td>
<td></td>
</tr>
<tr>
<td>1209</td>
<td>$2.83 \cdot 10^{-8}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1223</td>
<td></td>
<td>$6.61 \cdot 10^{-8}$</td>
<td></td>
</tr>
<tr>
<td>1269</td>
<td>$9.74 \cdot 10^{-8}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1273</td>
<td></td>
<td>$1.45 \cdot 10^{-7}$</td>
<td></td>
</tr>
<tr>
<td>1333</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1373</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1473</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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and by Agarwala, Naik, Anand, and Paul.\textsuperscript{50}

In the further analysis of the data, two additional relationships merit examination. Based on equation (48)

\[ D = D_0 \exp\left(-\frac{Q}{RT}\right) \]

it would seem that there should be some type of direct relationship between $D_0$ and $Q$. Using the PDP 8/I computer a best fit least squares program was run utilizing data on $D_0$ and $Q$ for h.c.p., f.c.c., and b.c.c. structures and at all temperatures. It was found that the correlation coefficient for the best fit line was 0.98 where 1.0 is a perfect fit. Subsequently, it was determined that the relation between $D_0$ and $Q$ was

\[ D_0 = 3.3 \times 10^{-8} \exp\left((0.36)(Q)\right). \]  \hfill (60)

Whether or not this equation applies only to a dilute solute diffusion of the alkali metals in these particular matrices, or for any other diffusing system requires further investigation.

A second interesting consequence of this work involves the possibility of using this method to further develop concepts on ultra-fast diffusion. Ultra-fast diffusion has been considered when the solute diffusion is much greater than self-diffusion in the host material. An example is the diffusion analysis of zircaloy-2 in this work. The self-diffusion rates of pure $\alpha$-zirconium have been determined by Dyment and Libanati.\textsuperscript{51} For a temperature range of 1036°K to 1130°K, they found $D$ to be on the order of $10^{-12}$ to $10^{-13}$ cm$^2$/sec, while in this study, the diffusion rates of the alkali
metals for the same temperature range were on the order of $10^{-8}$ to $10^{-9}$ cm$^2$/sec. This is approximately four to five orders of magnitude faster. Ultra-fast diffusion has been suggested by Hood and Schultz$^{52}$ to take place by an interstitial mechanism. They studied both Co-60 and Fe-59 diffusion in pure α-zirconium and found that $D$ was of the order of $10^{-7}$ to $10^{-9}$ cm$^2$/sec which corresponds well with this work. That ultra-fast diffusion would take place by an interstitial mechanism coincides with the mechanism for determining the largest diffusing ion size; a concept used to explain figures 22 and 23.

It should be further noted that in Part II, concepts of the jump frequency $\omega$, the mean vibrational frequency, $\nu$, and the change in free energy, $\Delta G$, were all discussed. With the results obtained in this study, it now becomes important to determine some of these quantities.

Zener$^{39}$ has stated that while it would not be impossible for $\Delta G$ to increase with increasing temperature, it would be highly improbable. The reason for this is that a negative temperature coefficient of the elastic moduli exists. Zener also found that the change in entropy, $\Delta S$ can be related to the activation energy, $Q$, by

$$\Delta S = \lambda gQ/T_m$$

(61)

where $\lambda$ is a numerical coefficient essentially equal to
unity for interstitial, $\beta$ is a dimensionless quantity tabulated by Zener for some pure metals, and $T_m$ is the melting temperature for the host material. The value of $\Delta S$ can be used to calculate $\nu$ using

$$D_o = \frac{1}{6} a_o^2 \nu \exp (\Delta S/R)$$

(62)

provided that $D_o$ is known.

Of the three sample materials used in this study, $\beta$ has been determined for tantalum only. It has a value of 0.40. Using values for $D_o$ obtained experimentally, $\nu$ can be determined. Once the values of $\nu$ have been determined, $\omega$ can be calculated using equation (23).

$$\omega = \nu N_m = \nu \exp (-Q/RT) \exp (\Delta S/R)$$

Table XI gives the calculated values for $\omega$ for the five diffusing impurity isotopes in both temperature regimes.
Table XI
Calculated Values of $\nu$ and $\omega$ For Tantalum

$990 < T(\text{°K}) < 1230$

<table>
<thead>
<tr>
<th>Impurity</th>
<th>$\nu$ (sec$^{-1}$)</th>
<th>exp(-Q/RT)</th>
<th>exp($\Delta S/R$)</th>
<th>$\omega$ (sec$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li-7</td>
<td>$2.83 \times 10^{11}$</td>
<td>.997</td>
<td>1.001</td>
<td>$2.82 \times 10^{11}$</td>
</tr>
<tr>
<td>Na-23</td>
<td>$1.14 \times 10^{10}$</td>
<td>.998</td>
<td>1.0007</td>
<td>$1.13 \times 10^{10}$</td>
</tr>
<tr>
<td>K-39</td>
<td>$1.49 \times 10^{13}$</td>
<td>.995</td>
<td>1.002</td>
<td>$1.48 \times 10^{13}$</td>
</tr>
<tr>
<td>Rb-85</td>
<td>$6.71 \times 10^{14}$</td>
<td>.994</td>
<td>1.002</td>
<td>$6.68 \times 10^{14}$</td>
</tr>
<tr>
<td>Cs-133</td>
<td>$4.69 \times 10^{10}$</td>
<td>.998</td>
<td>1.001</td>
<td>$4.68 \times 10^{10}$</td>
</tr>
</tbody>
</table>

$1230 < T(\text{°K}) < 1790$

<table>
<thead>
<tr>
<th>Impurity</th>
<th>$\nu$ (sec$^{-1}$)</th>
<th>exp(-Q/RT)</th>
<th>exp($\Delta S/R$)</th>
<th>$\omega$ (sec$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li-7</td>
<td>$1.86 \times 10^{11}$</td>
<td>.996</td>
<td>1.002</td>
<td>$1.85 \times 10^{11}$</td>
</tr>
<tr>
<td>Na-23</td>
<td>$6.22 \times 10^{9}$</td>
<td>.998</td>
<td>1.001</td>
<td>$6.21 \times 10^{9}$</td>
</tr>
<tr>
<td>K-39</td>
<td>$5.83 \times 10^{10}$</td>
<td>.997</td>
<td>1.001</td>
<td>$5.82 \times 10^{10}$</td>
</tr>
<tr>
<td>Rb-85</td>
<td>$1.28 \times 10^{9}$</td>
<td>.998</td>
<td>1.001</td>
<td>$1.27 \times 10^{9}$</td>
</tr>
<tr>
<td>Cs-133</td>
<td>$6.50 \times 10^{9}$</td>
<td>.998</td>
<td>1.001</td>
<td>$6.49 \times 10^{9}$</td>
</tr>
</tbody>
</table>
Part VI
CONCLUSIONS

It was the purpose of this work to illustrate a new technique in the general field of materials characterization. The transport of impurities, in the p.p.m. to p.p.b. range and less, is becoming increasingly important. Since the mass spectrometer is an instrument which can detect impurity concentrations on this scale, it offers another view of the high temperature diffusion properties in materials which are of engineering usefulness. This is instead of being confined to the world of the single crystal.

As a consequence of this work, not only were the diffusion coefficients of the alkali metals in the sample materials calculated, but known phase transformations were determined in the samples of 304 stainless steel and zircaloy-2. This, of course, lends a measure of credibility to the results obtained.

While the primary intent of this work has been accomplished, it has been shown that what used to be a chemist's tool can be applied in metallurgical research. It is not to be suggested that the mass spectrometer is the ultimate instrument, but rather that it allows one more unique way of looking at material's properties on the atomistic level. It is conceivable that by determining the relationship between the diffusion coefficient and the atomic radius of all the impurities in a material that a better understanding of diffusion processes could be obtained. It is hoped that
further funding will allow this laboratory to pursue this and to see how ion implantation of selected species affect the diffusion process in certain materials - i.e., semiconductors.
APPENDIX 1

Program to Determine Least Squares Fit of Data to Exponential Curve, Correlation Coefficient, and 95% Confidence Limit

C-8K FOCAL #1969

01.10 T !!!"THIS PROGRAM FITS PAIRS OF POINTS TO AN EXPONENTIAL
01.11 T !!!"LEAST SQUARES CURVE FIT - TO THE EQUATION: Y=A*EXP(B*X)
01.12 T !!!" X IS THE RECTANGULAR ABSYSSA
01.13 T !!!" Y IS THE RECTANGULAR ORDNATE (INTERNAL CONVERSION TO LOG Y
01.14 T !!!" A IS THE Y-INTERCEPT
01.15 T !!!" B IS THE SLOPE ON SEMI-LOG PAPER
01.16 T !!!" R IS THE CORRELATION COEFFICIENT
01.17 T !!!" CL IS THE 95% CONFIDENCE LIMIT +/- AT X=(TEST POINT)
01.18 T !!!"PROGRAM OPERATION:
01.19 T !!!" AFTER FIRST PROMPT ENTER THE NUMBER OF ORDERED PAIRS
01.20 T !!!" THEN -ENTER X & Y AS REQUESTED BY THE PROGRAM
01.21 T !!!" -ENTER T(0.125) FOR THE SPECIFIED D.F. WHEN REQUESTED
01.22 T !!!" -ENTER THE NUMBER OF TEST POINTS
01.23 T !!!" -ENTER THE TEST POINTS
01.30 A !!!"ENTER THE NUMBER OF VALUES: "N
01.35 S SX=0;S SY=0;S XT=0;S XY=0;S YT=0
01.40 I (N-1) 9.9,1.6
01.50 F I=1,N;D 2
01.55 G 3.1
01.60 T !!!"TRY AGAIN -- 1 IS NO GOOD
01.65 G 1.3

02.10 A !!!"ENTER X: "X,"ENTER Y: ",Y;S Y=FLOG(Y)
02.15 S Sx=Sx+x;S Sy=Sy+y;S Xt=Xt+x+2;S XY=XY+x+y+2;S YI=YI+y+2
03.10 S XS=XS/N;S YS=YS/N
03.15 T !!!"THE AVERAGE VALUE OF X IS: ",%X$S
03.20 S YY=YI-<SY+2/N)
03.30 S XX=XT-(SX+2/N)
03.40 S IT=XY-<SY*SX/N)
03.50 S SS=<TT+2)/XX
03.55 S MS=(YY-SS)/(N-2)
03.60 C CALCULATION OF THE SLOPE -- B
03.65 S B=TT/XX
03.70 C CALCULATION OF THE Y-INTERCEPT -- A
03.75 S A=FEXP(YY-B*X$)
03.80 C CALCULATION OF THE CORRELATION COEFFICIENT -- R
03.85 S R=TT/FSQRT(XX*YY)
03.90 G 5.10
04.10 C PRINT OUT THE RESULTS
04.20 T % "RESULTS:
04.30 I " THE SLOPE IS: ",B
04.40 I " THE Y-INTERCEPT IS: ",A
04.50 I " THE CORRELATION COEFFICIENT IS: ",R
04.60 R
04.70 T " IF YOU WISH TO STOP -- AT THE NEXT PROMPT TYPE (-1)
04.80 T " OTHERWISE CONTINUE
04.90 G 1.3

05.10 C INPUT FOR THE DETERMINATION OF THE 95% C.L.'S FOR EACH TEST PT
05.15 T %2.0, "ENTER THE VALUE OF T(0.025) FOR ",N-2, " DEGREES
05.20 A " OF FREEDOM: ",DF
05.25 F I=1,M;D 6
05.30 DO 4
05.35 F I=1,M;S CL(I)=DF*FSGT(MS*(1/N+(X(I)-0)*2/XX))
05.36 F I=1,M;S Y(I)=FEXP(B*X(I)))*A
05.37 F I=1,M;D 7
05.38 D 9.04
05.39 D 9.05
05.40 F I=1,M;D 9.1
05.45 G 4.70

06.05 C INPUT OF TEST POINTS -- X(M)
06.10 I %2.0, "ENTER TEST POINT X("I",":"
06.20 A X(I)

07.10 S ZA=FEXP(FLOG(Y(I))+CL(I))
07.20 S ZB=FEXP(FLOG(Y(I))-CL(I))
07.30 S ZC(I)=FABS(ZA-ZB)/2

08.04 T " TEST POINTS AND THEIR RESPECTIVE 95% C.L.'S
08.05 T " M ",X(M) Y(M)
08.10 I ",Z2.0, I", X(I), Y(I), ZC(I)
08.90 T " THE END";Q
APPENDIX 2

Program to Determine D for a Given Temperature
Provided D₀ and Q are Supplied

C-8K FOCAL @ 1969

01.01 C PROGRAM: DIFF
01.02 C
01.10 A !!!"WHAT IS THE SAMPLE MATERIAL? ",A
01.20 A !!!"WHAT IS THE DIFFUSING ISOTOPE? ",A,B
01.30 A !!!"WHAT IS THE TEMPERATURE RANGE? ",A,"TO ",B,"DEGREES KELVIN."
01.35 A !!!"WHAT IS THE VALUE OF D(0)? "D0
01.40 A !!!"WHAT IS THE VALUE OF THE ACTIVATION ENERGY? ",Q
01.60 T !%2

02.10 A !!"T= ",T
02.15 I (T) 2.6,2.5
02.20 S D=D0*EXP(-Q/(2E-3)*T)
02.25 G 3.1
02.30 T ","D= ",D
02.31 G 2.1
02.35 T ","D= ",D
02.40 G 2.1
02.50 G 1.1
02.60 T !!!!;Q

03.10 I (1000-T)2.3,2.3,2.35

*
REFERENCES


14. Walling, op cit., p. 35.


17. Shewmon, op cit., p. 56.

18. Shewmon, op cit., p. 56.
20. Dekker, op cit., p. 62
22. Dekker, op cit., p. 62.
23. Shewmon, op cit., p. 60.
36. LeClaire, A. D.; Acta Metal; 1, 438, 1953.
This report focuses on the high temperature diffusion of trace elements in metals and alloys. Specifically, these measurements have been made by high sensitivity mass spectrometry in which individual atoms were detected, and quantitative data was obtained for zircaloy-II, 304 stainless steel, and tantalum. Additionally, a mass spectrometer was also an analytical tool for determining an allotropic phase change for stainless steel at 955°C, and a phase transition region between 772°C and 1072°C existing for zircaloy-II.

Diffusion rates were measured in thin (0.001" (0.0025 cm) and 0.0005" (0.0013 cm)) ribbons which were designed as high temperature thermal ion sources, with the alkali metals as naturally occurring impurities. In the temperature and pressure regime where diffusion measurements were made, the solute atoms evaporated from the ribbon filaments when the impurities diffused to the surface, with a fraction of these impurity atoms ionized according to the Langmuir-Saha relation.

While these measurements were made on only a few important materials, the techniques developed can be clearly applied to many other alloys important to space vehicles and supersonic transports; and, with appropriate modifications, to the diffusion of impurities in composites.