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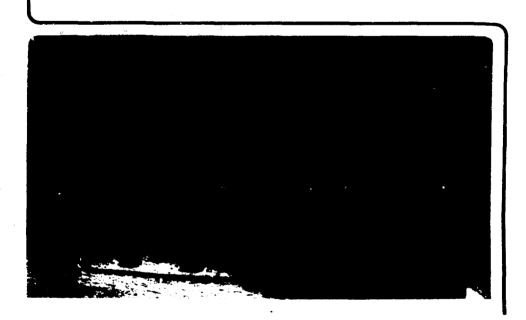
Materials & Molecular Research Division

OXYGEN DIFFUSION IN HYPOSTOICHIOMETRIC URANIUM DIOXIDE



Kee Chul Kim (Ph.D. thesis)

December 1980





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Kee Chul Kim Ph.D. Thesis

December 1980

Materials and Molecular Research Division Department of Nuclear Engineering Lawrence Berkeley Laboratory University of California Berkeley, CA 94720

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ABSTRACT

The tracer oxygen diffusivity in $U0_{2-x}$ has been measured along the lower two phase boundary. The diffusion couple consisted of two matched hypostoichiometric uranium dioxide wafers, one enriched with 13 and the other normal. These two were pressed together with a bond of liquid uranium in between. After a diffusion anneal the $^{18}\mathrm{O}$ concentration profile was determined by ion microprobe mass analysis, from which diffusion coefficients were obtained. The results showed much nigher diffusion coefficients than those of stoichiometric UO2. This directly proved that the major defect species in $\mathrm{UO}_{2-\mathbf{x}}$ is the arrion vacancy. Activation energy of anion vacancy migration was measured to be 11.7 ± 3.0 kcal/mole. A diffusion model established for \mathbb{P}_{2} and $\mathbb{P}_{2\pm \sqrt{2}}$ showed that in stoichiometric 00_2 both interstitials and vacancies contribute significantly to oxygen diffusion and neither can be neglected; at 1400°C their contributions are about equal. This model was extended to nearly stoichiometric $\mathtt{UO}_{2\pm\mathbf{x}}$ to predict oxygen diffusion coefficients in these stoichiometry ranges. Also deduced from the model were the Frenkel defect energy and entropy of 85.6 ± 9.2 kcal/mole and 13.2 ± 7.3 e.u., respectively. Using these values, the

contribution of Frenkel disorder to the excess enthalpy of $\rm UO_2$ was evaluated. Calculation showed that Frenkel disorder accounts for 87 percent of the excess enthalpy at $3000^{\circ} \rm K$. A simple two band model for electronic excitation, with a band gap of 2.0 ev and effective electron mass of 7.6 m_e, accounted for the remainder of the excess enthalpy.



INTRODUCTION

1.1 Statement of the Topic

fransport phenomena in uranium dioxide are of importance in understanding the behavior of fuel elements during reactor operation. EMFBR fuel pins are designed to be operated at night temperatures and much steeper temperature gradients than conventional thermal reactor fuels. Across the 3 mm radius of a fuel pin, the temperature varies from about 2500°C at the center to about 700°C at the cladding. These extreme conditions cause a variety of phenomena, including grain growth, fission product migration, oxygen rediscribution, and actinide redistribution.

Oxygen diffusion is of special interest because of many different contributions of the oxygen-unanium ratio and the oxygen potential to the fundamental properties of the oxide. For example, the thermal conductivity of the oxide changes with 0/3 ratio so that the local 0/3 ratios affect the temperature profile within the fuel element. This temperature profile is directly related to fuel restructuring, pore tips tion, etc., and all these properties and phenomena are observabled.

The oxygen potential also plays an important role in many aspects of the ox defuel, from fuel fabrication to oxygen redistribution and tuel-claiming reaction. The oxygen potential of the fuel determines in large part whether or not the fuel can compute the metallic cladding. In order to minimize this reaction and accomposate the increase of the oxygen obtential by fission processes, the initial composition of the increase of the oxygen obtential by fission processes, the initial composition of the

There have been many studies, both theoretical and experimental, of oxyden diffusion in stoichiometric $\rm UO_2$ and hyperstoichiometric $\rm UO_{2+x}$ [1-11]. However, similar measurements in hypostoichiometric $\rm JO_{2+x}$ have never been attempted mainly because of $\rm UO_{2-x}$ is a defect structure stable only at high temperatures (see Fig. 1) so that its oxygen diffusion coefficient is most likely large enough to render conventional methods unworkable. For example, in the gas-solid isotopic exchange method, the gas phase mass transfer step or the surface isotopic exchange step may be rate-controlling. Also, since the equilibrium oxygen potential of hypostoichiometric uranium dioxide is excremely low, it would be very difficult, if not impossible, to control the gas stream to maintain stoichiometry during the diffusion anneal. Even if this could be achieved, it is doubtful that this small exygen potential could be successfully isotopically monitored for diffusion measurements.

Fogether with thermodynamic information, transport data contribute meetly to the understanding or the defect structure of uranium makes. Nevertheless, considerable uncertainty remains as to the transportal aspects of the defect properties and transport mechanisms in the locate. One of the difficulties lies in the fact that there are no transport data in JO_{2-x} , which is another reason why measurement to oxygen diffusion in JO_{2-x} is urgently needed.

... Review of Previous Work

the of the earliest experiments reported was by Auskern et al. [2]. In the parametering work, the oxygen self-diffusion coefficient in ${\rm UO}_2$

^{.,} was measured using the isotopic exchange reaction between

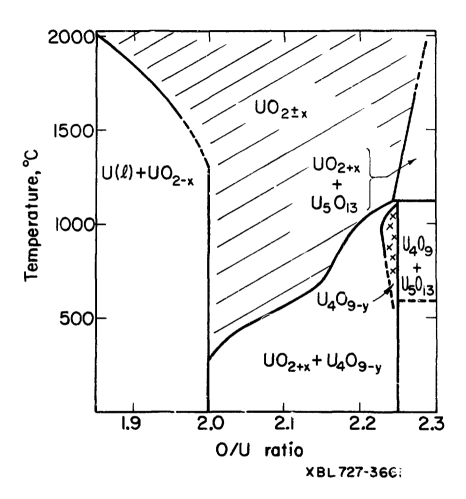


Figure 1. Oxygen-uranium phase-equilibrium bystem [18].

reported that for essentially stoichiometric UO_2 , the oxygen diffusion coefficient in the temperature range $550\text{--}780^\circ\mathrm{C}$ could be represented by $\mathrm{D}=1.2\times10^2$ exp(-65300/RT). The activation energy was verified in later studies [1,7]. However, the pre-exponential factor was abnormally high compared to later studies, both experimental [1] and theoretical [9]. Systematic error due to the large variation in the average particle diameter is a possible reason. Also reported was the activation energy of 29.7 kcal/mole for hyperstoichiometric UO_{2+x} . From these two activation energies Auskern et al. [2] calculated the Frenkel defect formation energy of 70 kcal/mole, assuming that the interstitialcy mechanism of oxygen diffusion applies to both UO_2 and UO_{2+x} .

In an unpublished work, Roberts et al. [7] made a more systematic attempt to obtain diffusion coefficients as a function of stoichiometry. Using single crystal UO₂, they measured activation energies of 23 kcal/mole, 30 kcal/mole and 69 kcal/mole for stoichiometries of 2.03, 2.01, and 2.001, respectively in the temperature range of 1200-1600°C. These values are in fair agreement with those of Auskern and Belle [2].

The two studies used the indirect gas-solid isotopic exchange method, which possesses some inherent problems [1,4]. By performing a direct diffusion couple experiment Marin et al. [1] attempted to overcome the pitfalls of the gas exchange method. They employed massive samples of normal UO_2 which were coated with a thin layer of UO_2 powder

highly enriched in 18 O. Then the couples were annealed, sectioned, and analyzed for 18 O penetration. The results were fitted by D=0.26exp(-59300/RT). The activation energy compares reasonably well with that found by Auskern et al. [2].

Using the same method, Contamin et al. [3] investigated the diffusion of oxygen in hyperstoichiometric UO_{2+x} . In the temperature range of $400-900^{\circ}$ C an activation energy of 21 kcal/mole, independent of stoichiometry, was obtained. They systematically investigated the dependence of diffusion coefficient on stoichiometry as well as temperature, and they observed a sharp increase in diffusion coefficient with x of UO_{2+x} in the vicinity of x=0. An activation energy of 21 kcal/mole was obtained for x \geq 0.006. Also an attempt was made to correlate the D with x at various temperatures.

Similar work was reported by Murch et al. [6] for U_{2+x} . For $U_{2.08}$ they reported D=6.25 x 10^{-4} exp(-23100/RT) over the temperature range of 560-800°C. The pre-exponential factor and the activation energy were in good agreement with those of Contamin et al. [3].

Over the years it was thought that the anion vacancy migration energy might not be so high that the vacancy mechanism to be neglected as a contributor in stoichiometric and near-stoichiometric $\rm UO_2$. This theory is supported largely by the experimental results on anion diffusion in other isostructural anion-deficient oxides such as $\rm CeO_{2-x}$ [12,13] and $\rm PuO_{2-x}$ [14,15]. In these studies the anion migration energy was measured in substantially hypostoichiometric specimen so that the diffusion process was controlled by the vacancy mechanism; also the activation energy would not contain the defect formation

energy. All of the results showed that the vacancy migration energies were within the range of 4–12 kcal/mole, which was compared with $21-30~\rm kcal/mole$ for interstitial migration energy in $\rm UO_{2+x}$ mentioned earlier. In addition, Catlow et al. [16] substantiated these values with a theoretically determined value of 5.8 kcal/mole for vacancy migration in $\rm UO_{2-x}$. From this standpoint, Murch et al. [9] attempted a theoretical calculation of diffusion coefficient of oxygen in $\rm UO_{2\pm x}$, assuming that diffusion in $\rm UO_2$ and $\rm UO_{2\pm x}$ is controlled not only by interstitials but also by vacancies at the same time. Their predictions are reasonably close to the experimental values at high temperature stoichiometric $\rm UO_2$ and to the data of Contamin et al. [3] for $\rm UO_{2+x}$. Similar attempts were made by Breitung [8] using a simpler model.

THEORY

2.1 Methods of Diffusion Experiment

It was indicated in the previous chapter that conventional methods are difficult to apply in hypostoichiometric UO_{2-x} . In this chapter these methods will be reviewed and the theoretical basis of the present experiment will be introduced.

2.1.1. Gas-Solid Isotopic Exchange Method

As applied in the works of Auskern et al. [2] and Roberts et al. [7], the diffusion anneal is followed by a mass spectrometric analysis of the $^{18}0/^{16}0$ ratio of the gas phase, one of the two constituents (gas or solid) being tagged with $^{18}0$. For this indirect method to be workable in uranium dioxide, several experimental conditions must be satisfied:

- Gas-solid surface exchange step should not be the rate controlling step.
- (2) Gas phase mass transfer rould not be the rate controlling step.
- (3) The partial pressure of oxygen in the gas should be matched to the equilibrium oxygen potential of the solid urania in order to maintain the original stoichiometry throughout the experiment

Employing this method in nypostoichiometric UO_{2-x} might violate the first and second conditions because the UO_{2-x} phase exists only at high temperatures and therefore the diffusion in solid phase is probably extremely fast. In regard to the third condition, since the exide should be in the form of powder in order to provide large surface

area, the stoichiometry of the samples would be extremely vulnerable to the surrounding gas, especially at the high temperatures.

The most convenient way of achieving a desired oxygen potentials is by a mixture of ${\rm CO_2-CO}$ or ${\rm H_2O-H_2}$. For example, the equilibrium oxygen potential of ${\rm UO_{1.98}}$ at $1700^{\circ}{\rm C}$ is -170 kcal/mole, which is equivalent to ${\rm CO_2/CO_0} \simeq 4 \times 10^{-13}$ and ${\rm H_2O/H_2} \simeq 10^{-6}$ at this temperature. Ine ${\rm CO_2/CO}$ ratio is so low that it would be extremely difficult to achieve. The ${\rm H_2O/H_2}$ ratio could be controlled in this range fairly well, but no easily. However, since the ${\rm H_2O}$ is the only oxygen-carrying species, it would be difficult to detect minute changes in the $^{18}{\rm O}$ fraction in such a small amount.

from these standpoints, this method does not seem to be appropriate for $\mathrm{UO}_{2-\mathbf{v}}$.

2.1.2 Diffusion Couple (layer-solid) Method

In this method two UO $_2$ pellets, one enriched with 18 O and the other normal, are placed in contact. 18 O and 16 O interdiffuse when the couple is neated. Unlike the gas-solid isotopic exchange method, this is a direct—surement of diffusion and there are none of the innerent systematic errors or difficulties in using gases. However, accomplishing a truly good contact between two solids is difficult. Contamin et al. [3] overcame this difficulty by depositing a trin layer of 18 O $_2$ (10-20 um) by decanting a suspension of 18 O $_2$ in ethylaiconol. Pairs of samples with enriched layers in contact were annealed in vacuum for several hours under pressure at low temperature

in order to promote good contact. They were then diffusion annealed in hydrogen and argon, and then sectioned to determine 18 0 profile using spark mass spectrometry and ion mass analysis.

Murch et al. [6] used a technique that differed only in that a thicker deposit (~150 $\mu m)$ of $^{18}0-enriched~UO_{\rm J+v}$ was used.

There are numerous ways to investigate the concentration profile other than by spark mass spectrometry and ion mass analysis, which are described elsewhere $\{11\}$.

2.2 Experimental Technique of the Present Study (Solid-Solid Method)

In order to eliminate the experimental difficulties and uncertainties of a layer-bulk diffusion couple, a bulk-bulk couple was utilized instead. The diffusion couple consisted of two $\rm UO_{2-x}$ wafers, one of which was enriched with $\rm ^{18}O$. As mentioned previously, this technique has the difficulty in achieving good contact between two wafers. Calculation showed that the vapor pressure of $\rm UO_2$ is too low for sufficient oxygen transport from one side to another through a vacuum gap.

To avoid this interfacial resistance, the wafers were bonded together by liquid uranium. Liquid uranium is believed to have a sufficiently high solubility [17-20] so that the liquid metal bond should transport oxygen from one wafer to the other quite efficiently. This technique is equivalent to reduction of the heat transfer resistance in the fuel-cladding gap of carbide fuel pins by sodium bonding.

However, since the diffusion coefficient in liquid uranium is different from that in solid urania, the problem had to be analyzed with this effect included.

In order not to perturb the stoichiometry of the wafers by the presence of liquid uranium, the experiments were performed only in $U(\cdot)+U_{2-x}$ two-phase region. In this system, the stoichiometries to be studied were fixed automatically by the temperatures and therefore the diffusion measurements would be only on 0/U ratios along the lower phase boundary (Fig. 1).

?.3 Ineoretical Analysis

In order to measure the diffusion coefficient, the ¹⁸0 profile has to be fitted to an analytic solution of the diffusion equation, which should include the effect of the liquid uranium layer at the interface. This can be obtained by solving the diffusion equation with appropriate boundary conditions. The diffusion equation can be written in dimensionless form:

$$\frac{\partial \phi}{\partial x} = \frac{\partial^2 \phi}{\partial n^2} \tag{1}$$

where
$$\phi = \frac{y - y_0}{y_1 - y_0}$$
 , $y = \frac{18}{0}$ isotopic ratio = $\frac{18}{18} \frac{1}{0} + \frac{16}{0}$

 $v_{\rm p}$ = initial isotopic ratio of $^{18}{\rm O}$ in $^{13}{\rm O-enriched}$ wafer

 r_0 = initial isotopic ratio of $^{18}0$ in normal wafer.

 $r = 0 t/r^2$, D is diffusion coefficient, t is time, R is the thickness of one wafer, $\eta = \frac{z}{R}$, z is the distance measured from the surface of the enriched wafer.

The geometry of the diffusion couple is depicted in Fig. 2. The initial condition can be written as:

I.C.
$$\phi(n,0) = 1$$
 , $0 < n < 1$ (2) $\phi(n,0) = 0$, $1 < n < 2$

Since both ends are insulated;

B.C.1
$$\frac{\partial \phi}{\partial n} = 0 \text{ at } n = 0$$

$$\frac{\partial \phi}{\partial n} = 0 \text{ at } n = 2$$
(3)

Throughout the calculation it is assumed that the liquid uranium thickness δ is very small, i.e., $\delta << 2$ and that the wafers are infinite slabs.

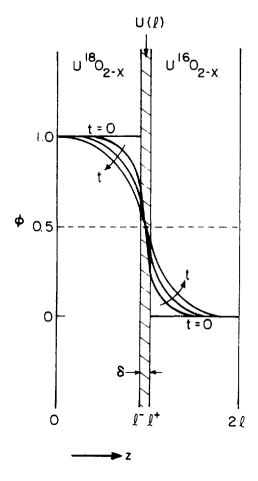
At the interface, the $^{18}0$ flux is continuous:

$$J = -0 \left[\frac{3C_{13}}{3Z} \right]_{2} = -0 \left[\frac{3C_{18}}{3Z} \right]_{2} + \tag{4}$$

where $\hat{\mathbf{S}}_{13}$ is the concentration of $^{13}\mathbf{0}$ in $\mathbf{U0}_{2-\mathbf{x}}$.

$$\mathcal{I}_{13} = \mathcal{I}_{0} y = \mathcal{I}_{0} [y_{0}^{+} (y_{1}^{-} y_{0}^{-}) \phi]$$
 (5)

where $\mathbf{S}_{\mathbf{0}}$ is the total oxygen concentration in $\mathbf{UO}_{2-\mathbf{x}}.$



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Figure 2. Geometry of the diffusion couple bonded by liquid uranium.

Thus,

$$J = -\frac{\partial C_{O}(y_{1} - y_{0})}{2} \left[\frac{\partial \phi}{\partial y}\right]_{1}^{1} = \frac{DC_{O}(y_{1} - y_{0})}{2} \left[\frac{\partial \phi}{\partial y}\right]_{1}^{1}$$
 (6)

The ¹³J flux in the liquid uranium can be written as:

$$x = y_0^2 - \frac{y_{13}^2 y_{13}^2 - y_{13}^2 y_{13}^2}{5}$$

 θ_{9}^{d} is exygen diffusion coefficient in liquid uranium, θ_{13}^{d} is the 13) concentration in the liquid uranium, and 3 is the thickness of the liquid uranium. Assuming isotopic equilibrium at the interface of $3\theta_{2-x}$ and liquid uranium,

$$z_{13}^{u} = z_{0}^{u} y = z_{0}^{u} y_{0}^{+} (y_{1}^{-} y_{0}^{-}) \phi]$$
 (3)

where \mathcal{C}_0^u is the solubility of oxygen in liquid uranium. Thus, from Eq. (7)

$$z = \frac{3}{3} (A^{T} - A^{2}) (A^{T} - A^{2})$$
 (4)

From the symmetric nature of the problem, at the center of the liquid unanium layer, $\phi = 1/2$ at all times, and Eq. (9) can be neplaced by:

$$J = \frac{9_0^{d}c_0^{d}}{(5/2)} (y_1 - y_0)(\phi_{1-} - \frac{1}{2}) = \frac{9_0^{d}c_0^{d}}{(5/2)} (y_1 - y_0)(\frac{1}{2} - \phi_{1-})$$
 (19)

Thus, from Eqs. (6) and (10), we have:

3.2.2
$$\left[\frac{3\phi}{3n}\right]_{1^{+}} = -B(\phi_{1^{-}} - \frac{1}{2})$$

$$-\frac{3\phi}{3n}\Big]_{1^{+}} = -B(\frac{1}{2} - \phi_{1^{+}})$$
(11)

where
$$B = \frac{D_0^u}{D} \frac{f}{(5/2)} \frac{G_0^u}{G_0^u}$$
 (12)

The parameter 3 represents the overall conductance of the liquid unanium layer for oxygen. The higher the value 8, the less resistance is affected by the liquid unanium layer to oxygen transport across the interface. This parameter depends mainly upon the solubility of oxygen in liquid unanium, which is not very well established experimentally and shows a large disagreement among different investigators [17-20]. However, even use of the most pessimistic data yields 3 values that permit modest diffusion rates. The only factor in 3 that can be controlled as 5. Therefore, it is important to minimize the thickness of the liquid unanium layer.

From Eqs. (1), (2), (3), and (12), ϕ can be solved to yield [21]:

$$\phi = \frac{1}{2} + \sum_{n=1}^{\infty} e^{-\partial a_n^2 t} = \frac{\left\{ (i x_n)^2 + \beta^2 \right\} \cos(a_n z) \sin(a_n z)}{\left\{ (i x_n)^2 + \beta^2 + \delta \right\}} = \frac{\sin(a_n z)}{a_n z}$$
(13)

 x_n 's are positive roots of $(x_i)\tan(\alpha x)-B=0$. The best values of D and B are sought to fit the data points from the diffusion experiments to Eq. (13).

3. EXPERIMENTAL - SAMPLE PREPARATION

As was indicated previously, two reduced $\rm UO_{2-x}$ wafers are needed: the enriched in $^{13}\rm O$ and the other normal. The overall procedure for sample preparation can be described as follows:

- (1) Preparation of $\theta^{13} \theta_2$ Resultion of θ_2 and $\theta_3^{13} \theta_2$
- $^{\circ}$ Determination of stoichiometry of the reduced sample.

tech step is described in detail below.

1. Preparation of Ulbo

i...i inesny

 16) atoms in 90_2 were replaced by 18 0 using 18 0-enriched water (95.1 percent 13 0-enriched water was obtained from Mound Laboratory), at high temperature:

$$J^{16} J_2 + H_2^{18} J_2 + H_2^{16} 0 \tag{14}$$

In thing so, it was necessary to maintain the original stoicniometry, $\text{This was achieved by mixing H}_2 \text{ and H}_2^{13}0 \text{ in the ratio that}$ which is the equation of the equation of the stoichiometric dO $_2$. This was reformulated in the following way:

$$r = \frac{3}{3 + \frac{5}{2} \cdot \frac{3}{2} \cdot \frac{3}{2} \cdot \frac{1}{2}} = \exp(1 - \delta G^0 / RT) .$$
 (16)

P's are partial pressures and ΔG^{Ω} is standard free energy of reaction (16).

Thus,

$$P_{0_{2}}^{1/2} = \frac{P_{H_{2}0}}{P_{H_{2}}} \exp(\Delta G^{0}/RT)$$
 (17)

ine oxygen potential is defined as:

$$\frac{\overline{\Delta a_0}}{2} = RT \ln P_{0_2} \tag{13}$$

From Eqs. (17) and (18),

$$\overline{\Delta G}_{02} = 2RT \ln \frac{P_{43}U}{P_{H_2}} + 2\Delta G^{0}$$
 (19)

 ΔG^{0} , in cal/mole, is very well known [22]:

$$aa^{0} = -57250 = 2.21 \text{ T} + 1.943 \text{ TinT}$$

since the oxygen potential of 30_2 is also well known [23-13], the matth $\rm H_2O/H_2$ can be determined from Eq. (14).

Except potential of $30_{2\pm\chi}$ is shown graphically in the 3. As an persent, the oxygen potential changes very snarply in the vicinity in storahiometric 30_{2} , e.g., from -14° to -70 kcal/mole at 13070. The to this effect, vintually any oxygen potential within this range would be satisfactory for maintaining storahiometric 30_{2} . Due to this effect, vintually any oxygen potential within this range would be satisfactor, for maintaining storahiometric 30_{2} .

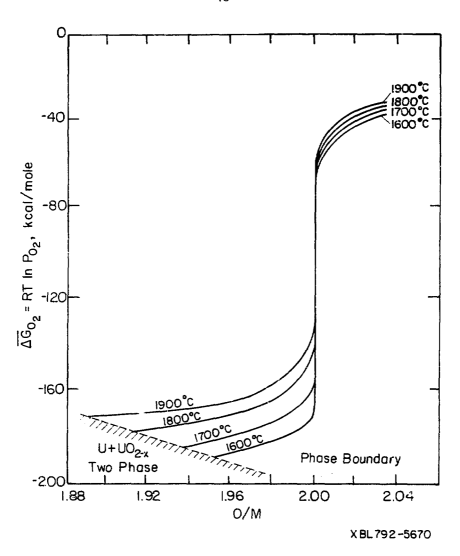


Figure 3. Variation of oxygen potential with temperature and O/U ratio [54]. The two-phase boundary is based on Ref. [38].

To promote a fast reaction, nigh temperature and high ratio of $H_2^{18}0/H_2$ were preferred, yet the temperature had to be low enough to prevent a significant evaporation. To satisfy these requirements, the conditions of $H_2^{13}0/H_2=10^{-2}$ and T = 1500°C were chosen, which would yield $\overline{\Delta G}_{0_2}=-103$ kcal/mole.

The desired $\mathrm{H}_2^{18}\mathrm{O/H}_2$ ratio could be obtained by saturating H_2 with $\mathrm{H}_2^{13}\mathrm{O}$. The saturation was achieved by flowing H_2 gas at 1 atm through $\mathrm{H}_2^{18}\mathrm{O}$ the temperature of which was controlled so that it would yield the predetermined $\mathrm{H}_2^{18}\mathrm{O/H}_2$ ratio. Although the H_2 flowed through the $\mathrm{H}_2^{13}\mathrm{O}$ in the form of tiny bubbles, the exit gas might nave been slightly undersaturated. However, due to the wide range of equilibrium oxygen potential of stoichiometric UO_2 , this slight uncertainty was acceptable.

3.1.2 Apparatus

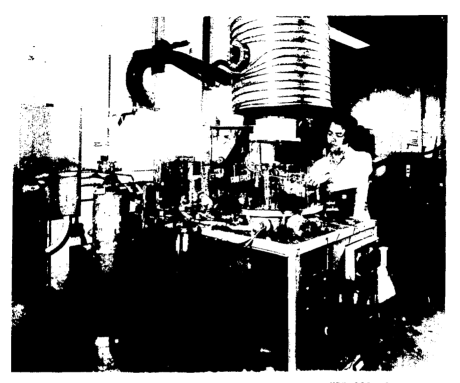
Figure 4 depicts the overall system which was used for all aspects of sample preparation.

The gas lines were made from 1/4 in. 0.D. stainless steel tube. To minimize contamination, high purity gas valves were used exclusively and only stainless steel Swagelok-type fittings were used for connections. The use of 0-ring type fittings and valves was avoided. This design was particularly important for the UO_2 reduction step, because even a trace of H_2O in H_2 would inhibit or limit the capability of reduction (see Fig. 3).

To prevent a back-diffusion of air into the system the gas was verted through diffusion pump oil which separated the system from the atmosphere.

Figure 4. Line diagram of the system.

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Figure 5. Photograph of overall apparatus.

3.1.2.1 Furnace

Shown in Fig. 6 is a detailed view of the furnace. The $\rm U0_2$ specimen was placed inside a molybdenum crucible, which was 1 in. 0.0. and 6 in. long, and electron beam welded to a 3.5 in. wide molybdenum flange.

A tungsten mesh element, 3 in. diameter and 6 in. long, was used to heat the furace. The temperature was controlled by the voltage applied to the heating element which was surrounded by a series of tungsten radiation shields to minimize the heat loss and to protect the outer shell of the furance, which was cooled by water.

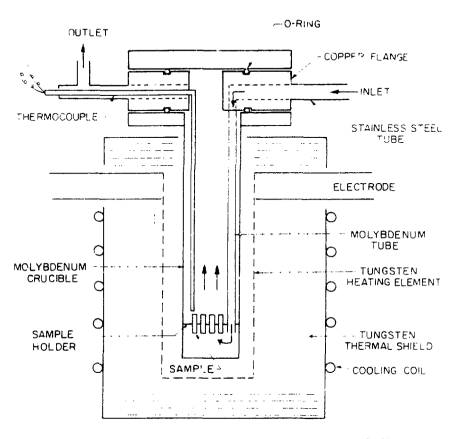
The gas was fed into the bottom of the crucible by a 1/8 in. molybdenum tube and flowed upward from then on. A rhenium rig was designed to hold several 100 wafers in an upright position.

The entire furnace was contained in a belljar which was under a vacuum for operation. A pressure of 10^{-6} torr could be obtained using a 6 in. diffusion pump. During furnace operation $5-9 \times 10^{-6}$ torr could be maintained.

The temperature was measured by W3%Re-W25%Re thermocouple which was adjacent to the specimen inside the molybdenum crucible.

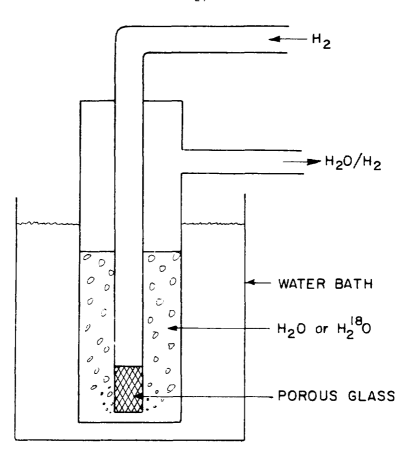
3.1.2.2 H_2^{18} 0 - H_2 Controlling System

The $\rm H_2$ was saturated with $\rm H_2^{18}0$ in a Pyrex tube where tiny pubbles were generated in a $\rm H_2^{18}0$ by flowing $\rm H_2$ through a porous glass frit (see Fig. 7). The cube was immersed in a water bath which was temperature controlled by Neslab PBC-2 bath cooler within $\pm 0.5^{\circ}\rm C$.



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Finure 6. Detailed view of the furnace.



XBL 8010-6099

Figure 7. H₂0 - H₂ controller.

Since only a small portion of $\mathrm{H}_2^{18}0$ in the mixture reacted with UO_2 , it was necessary to collect and recycle this valuable water from the outlet. First, the outlet from the furnace went through a cold trap the temperature of which was maintained slightly above 0°C. The temperature was chosen to prevent blocking the tube by ice formation. After this, the gas went through a liquid nitrogen cooled trap which collected the remaining $\mathrm{H}_2^{18}0$. Of course this recycled water was slightly less enriched than the original $\mathrm{H}_2^{18}0$.

3.1.3 UO2 Wafer Preparation

 ${
m UO}_2$ pellets of 1.17 cm diameter, 1.5 cm nigh were provided by the General Electric Co., Vallecitos.

Wafers of approximately 1.4-1.1 mm thickness were cut from the pellets and polished using silicon carbide abrasives and diamond paste. Polishing promoted a good contact in the diffusion couple. It was equally important that the thickness of a wafer should be uniform. Otherwise the contact of the two wafers would not be parallel. Thicknesses of the two wafers were matched within 3.02 mm for each set, one of which was later enriched with 180.

Single crystal UO_2 pellets of 1 in. diameter were obtained from the Georgia Institute of Technology. A single crystal portion, about 1.2 cm in the center, was cut out after slicing them into 1.1 mm thick wafers. The same procedure employed for polycrystalline specimens was used for grinding and polishing.

lement	bed	Element	PPM	Element	РРМ
Мg	<1	Сu	<0.4	Sn	< l
Sí	10	Fе	13	V	<5
€d	<0.3	Pb	<0.3	Zn	<1.5
В	0.4	Mn	<0.3	aZ	<0.5
Αl	<5	Мо	<5	Ве	< 0.5
Ja	1	Ni	<2	Вi	<0.2
0r	<2	Аg	<0.1	ې	<3.7
Jo	<2.7	Na.	<8	Τi	< 0.3

Table 1. Impurities in UO, pellets

3.1.4 Procedure for Enrichment of UO2 in Oxygen-18

- Measure the weight of the sample: Before placing in the furnace, the weight was measured using a Mettler microbalance to compare with the weight after the exchange reaction.
- (2) Pump out the entire system: Before each run the entire system was pumped out while heating up the samples to 300°C in order to degas them.
- (3) Cool down to room temperature and fill the system with helium.
- Start flowing helium at a rate of 10 cc/sec; with valves
 No. 1 and No. 2 in Fig. 4 closed and valves No. 3 and No. 4
 open.
- (5) Turn on the furnace and heat up to 800°C; To prevent the samples from cracking by the thermal stress, the furnace was neated up slowly.
- (6) Snut off helium flow.
- (7) Start flowing hydrogen.

- (3) Set the hydrogen flow rate at 5 cc/sec.
- (9) Heat up to 1500°C.
- (10) Flow $H_2^{18}0-H_2$ for 38-48 nours; with valves No. 1 and No. 2 open and No. 3 and No. 4 closed. The temperature of $H_2^{18}0$ was maintained at 7.5°C which would yield $H_2^{18}0/H_2 = 10^{-2}$
- (11) Stop flowing H_2^{13} O/ H_2 and anneal for 5 nours; Samples were annealed in order to achieve a uniform 13 O concentration in UO_2 . Since they were to be reduced at higher temperature (~2000°C), where the diffusion would be very fast, even if there had been a slight honuniformity at this stage it was considered to be immaterial.
- (12) Cool down to 300°C.
- (13) Start flowing nelium.
- (14) Cool down to room temperature.
- (lo) Take out the sample and measure the final weight.

Exactly the same procedure, except for using normal water instead of $\pm\frac{13}{2}$ 0, was applied to the counterpart of the matched sample in order to impose an identical history. Since substantial grain growth was observed after 48 nours of reaction, this process was considered to be an essuitial step, especially for the polycrystalline samples. As was expected, no significant weight changes were observed when normal water was used. Therefore, the following conclusions were drawn:

- There had been no storoniometry change due to this procedure.
- (2) There had been no significant vaporization.
- (3) All the weight change observed when using $A_2^{13}5-A_2$ should be attributed to the substitution of 16 by 18).

1.1.5 Results of the Isatopic Englamment Process

The degree of enrichment can be calculated from the weight unange and the fractional weight increase for 100 percent substitution of $\frac{10}{100}$) by $\frac{10}{100}$ for 1 gram $\frac{10}{100}$, sample $\frac{10}{100}$, $\frac{10}{100}$.

enc inhert =
$$\frac{(M_1 - M_1)^7 M_1}{0.001481}$$
,

where w. I initial weight

W_F = final weight.

Shown in the following Table 2 are results of two different vatines. The thinner the wafer, the higher the enrichment, which indicates that the diffusion of oxygen ${\rm JO}_2$ was one or the rate conscious steps. This result means that there had been a nonuniformity in ${\rm JO}_2$ concentration in the sample. However, at ${\rm JOOOO}_2$ the oxygen outfusion suefficient is ${\rm JOC}_2$ is approximately or ${\rm JOOO}_2$ in ${\rm JOOO}_2$ and with this diffusion coefficient, 5 hours of annealing should have relieved most of the nonuniformity and the subsequent 4 hours reduction at high temperature (2000°C) would virtually eliminate the homomorphismity. This expectation was proved later when the diffusion couple was analyzed after the diffusion experiment, and showed a flat

 13 C profile near the edge where the diffusion had not yet penetrated. This will be discussed in detail later.

Table 2. Typical weight changes after oxygen exchange reaction, and the halculated enrighment.

Batth No. 1				
^r การkness, แก	w ₁ , gn	we, gun	enrichment, ≟	
7.465 7.440	// # * / 1.182345	1.120030	73.5 71.3	
7.940 J.914	112347 19334U	1.134875	73	
1.339	. 15±3)()	1.030175	`ɔ.ĺ	
4.364	. +4 * 11	1. (45)	7 7	
1.333	4,444147	J. 9003 TH	30.3	
4.3);	. yn * 45	. 477795	41.	
	Batun	Yo. ?		
244	;o#3‡z	:'96J'	44.5	
1.22) 168	1.422315 1.354 9 20	1.432321 1.17425.	45.1 15.1	
1.155	351141	1.370571	47.3	

1.2 High Temperature Bas Phase Mass Transfer Study of Jour

The axygen deficient, hypostoloniumethic $\mathcal{D}_{3-\mathbf{x}}$ can be prepared by hydrogen reduction at () at high temperature:

$$(32) * (44) * (3) = (44)$$

The course of the reduction process was followed by a thermobalance. At high temperatures, nowever, the sample loses weight not only by reduction but by vaporization. Therefore, in order to study the reduction/evaporation process and to understand the capability of the system, it was necessary to conduct a systematic mass transfer study.

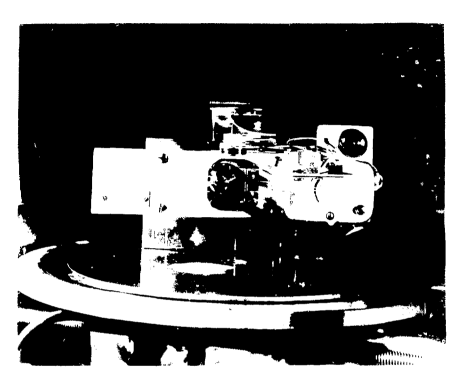
3.2.1 Apparatus

3.2.1.1 Inermobalance

The furnace was basically the same as the one used in the oxygen exchange reaction except that it was equipped with a Cann RG Electro-balance. The balance consisted of a control unit and a weighing assembly. A Heath dual per chart recorder adopted for 1 my signal was used for the readout of the balance. It had a capacity of 2.5 gm, readability of 0.3 percent of 1 my recorder (equivalent to ultimate readability of 0.1 ug).

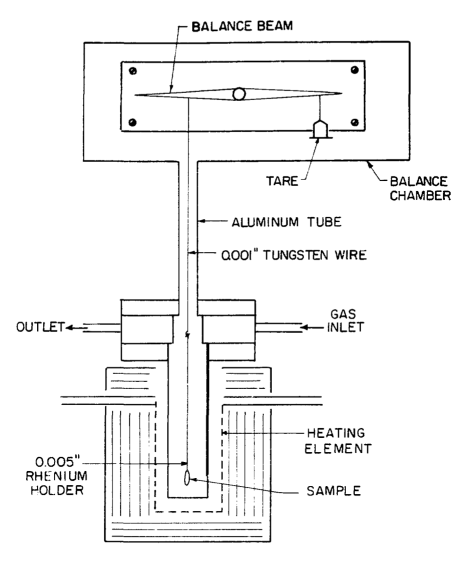
The weighing unit was noused in a leak-tight aluminum chamber which was haintained at a constant temperature by cooling water. An aluminum hangdown tube which was connected to a 3-1/2 in. Flange was used to suspend the sample below the balance into the furnace (see Figs. 9 and 10).

charmen, the hangdown tube needed to be harrow and long. Yet, at the same time the tube should be wide enough to keep the suspension wire from no bing against the wall which would cause enormous noise in the reacout. A 0.4 in. 1.3., 1/2 in. 0.3., 5 in. long aluminum tube was used. It was essential to have an absolutely straight wire to prevent hupbing against the tube wall, yet very light in weight. Therefore,



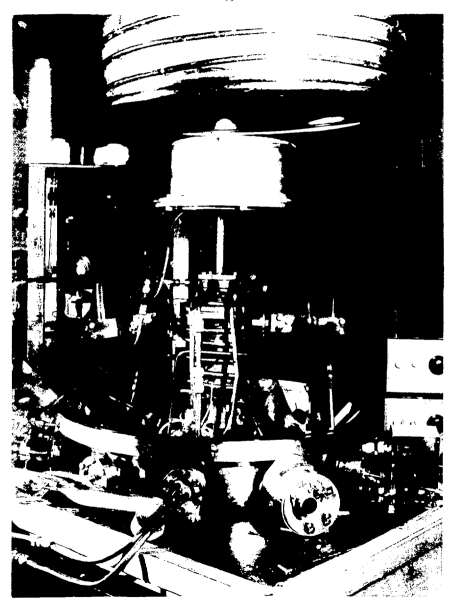
 $\lambda 98 = 0.1129 \, v .$ For the unit of Table D1. No feel labels.

•



XBL 8010-6100

Figure 9. The thermobalance setup.



VIB 100 1, 1, 1

them. 1). Photograph of the furnace with themsetalance

from the balance beam to the end of the hangdown tube, a 0.001 in. thick tungsten wire was used which was thin and flexible enough to be straightened by the weight of the sample. Then a 0.005 in. thick rhenium wire was connected to it to hold the sample.

Since the continuous gas flow provided substantial noise to the balance it was necessary to use an extra stage noise filter. A second stage filter essentially eliminated noise from the output.

3.2.1.2 Gas Purifier

To reduce $U0_2$ the hydrogen had to be as free of H_20 as possible. For example at 1900°C , 10~ppm of $H_2^{18}0$ in H_2 would limit the thermodynamic capability of reduction to 0/U=1.975.

To remove $\rm H_2^{\,0}$ and other contaminants from $\rm H_2^{\,}$, the gas was passed through a liquid nitrogen cooled trap consisting of activated charcoal and a molecular seive.

3.2.2 Procedures for Reduction/Evaporation Tests

 ${\rm UO}_2$ specimens were essentially same in weight and shape as the ones used in the previous experiments. For the purpose of comparison, iron samples of similar geometry were tested, of which equilibrium vapor pressures are well known [29]. ${\rm UO}_2$ samples were tested in argon and hydrogen, and for iron, helium and hydrogen were used. The balance was calibrated for each run.

3.2.2.1 UO, in Ar

- (1) Pump out the entire system; The sample was degassed at 300°C for 2 hrs at the same time. During the degassing step the typical weight loss was approximately 0.1 mg for 1 gm sample.
- (2) Fill the entire system with Ar and set the flow rate.
- (3) Heat up to the predetermined temperature; measure the rate of weight loss for different flow rates and temperatures.
- (4) Cool down.

3.2.2.2 UO2 in H2

- (1) Pump out the entire system.
- (2) Fill the system with Ar and start flowing.
- (3) Heat up to the predetermined temperature; measure the rate of weight loss.
- (4) Maintain the temperature until the rate reaches a steady state.
- (5) Shut off Ar and switch to H₂; The temperature was changed due to the difference in the properties of these two gases. The voltage had to be lowered to maintain the temperature.
- (6) Flow ${\rm H_2}$ until the weight loss rate reaches a steady state.
- (7) Cool down to 1000°€.
- (8) Flush the system with Ar; It was necessary to cool down the reduced ${\rm UO}_{2-x}$ in an atmosphere free of hydrogen because the precipitated uranium would react with ${\rm H_2}$ to form hydride, which would result in a total destruction of urania.
- (9) Cool down.

3.2.2.3 Iron Samples

Iron samples were tested in the streams of He and $\rm H_2$. The same procedure was taken as $\rm UO_2$ except that lower temperatures were employed.

3.2.3 Results and Discussion

3.2.3.1 UO, Vaporization in Argon

Table 3 shows the weight loss rates of ${\rm UO}_2$ in argon streams at three different temperatures and different flow rates.

Table 3. UO2 Vaporization in Ar

Γ°Κ	Flow Rate cc(stp)/sec	v cm/sec	W mg/min	κ _m	κm
1340	20	13.8	<i>≃</i> 0	0	
2173	3	3.9	0.009 ± 0.001	24.4	4.2
2173	10	11.1	0.110 ± 0.001	27.1	4.7
2173	20	22.2	0.014 ± 0.002	39.0	5.6
2273	20	23.3	0.048 ± 0.005	38.0	7.0

w = weignt loss rate

$$Sh = \kappa_m 1/0$$

l = diameter of sample

T = temperature

 $k_m^2 = \text{experimental mass transfer coefficient, defined by } J=k_m^2 P_{eq}^2/RT$

 $[\]kappa_{\rm m}^{'}$ = mass transfer coefficient calculated from theory

D = diffusivity of $UO_2(g)$ in Ar

 P_{eq} = equilibrium vapor pressure of 90_2

d = mass flux

Re = v1/v

v = velocity

v = kinematic viscosity

Sc = v/0

From the analogy between heat and mass transfer laminar boundary layer theory for a flat plate, the mass transfer coefficient $k_{\rm m}$ is predicted by:

$$Sn = 0.664 \ Sc^{1/3}Re^{1/2}$$
 (23)

As can be seen in Fig. 11, mass transfer of UO_2 vapor in argon was as much as six times faster than predicted. Using least squares fitting, the data at 2173°K are represented by: Sh = 3.91 Sc $^{1/3}$ Re (0.47 ± 0.01) . The possible errors may be: (i) underestimation of diffusivity of $UO_2(g)$ in argon. (The transport properties estimated from theory [30] are given in Table 4.) (ii) flat plate boundary layer theory was not very accurate for a thick disk hung in a gas stream. (The effect of decreasing flow path length from the center to the periphery was small.) Nevertheless, the data still seem to follow the theoretical dependence on Reynolds number, as shown in Fig. 11.

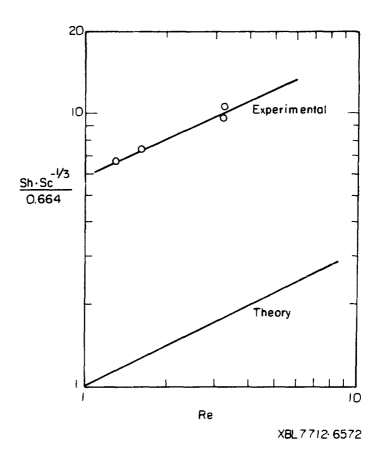


Figure 11. Comparison of UO, vaporization in argon with flat plate boundary layer theory for Reynolds number variations.

т°к	D _{UO2} -Ar, cm ² /sec	μ _{Ar} , ^{Poise}	D _{UO2} -H ₂ ,cm ² /sec	H _{H2} ,Poise
2073	2.31	7.7 x 10 ⁻⁴	12.53	3.2 × 10 ⁻⁴
2173	2.48	7.8 x 10 ⁻⁴	13.59	3.3 × 10 ⁻⁴
2273	2.71	7.9 x 10 ⁻⁴	14.54	3.4 × 10 ⁻⁴

Table 4. Diffusivity and Viscosity of Ar and H2

The effect of temperature on UO_2 vaporization can be estimated from the data in Table 3 at 2173°K and 2273°K. From Eq. (23) and $J_{vap} = \kappa_m P_{eq}/RT \text{ and } \dot{w} = JA \text{ where } A = \text{surface area of sample,}$

$$\frac{\dot{W}}{A} = \kappa_{m} \frac{eq}{RT}$$
 (24)

or
$$\frac{\dot{W}T}{DSC} \frac{1}{3} \frac{1}{8e^{1/2}} = \frac{P}{eq}$$
 (25)

In Fig. 12, the temperature dependence of $\dot{W}T/DSc^{1/3}Re^{1/2}$ is compared with equilibrium vapor pressure curve. Least squares fitting yields a slope corresponding to a heat of vaproization $\Delta H_{vap} = 123.3 \pm 4.5 \text{ kcal/mole}$ which is in fair agreement with $\Delta H_{vap} = 143.1 \text{ kcal/mole}$ obtained from the equilibrium vapor pressure curve [31].

3.2.3.2 UO, Reduction in Hydrogen

Table 5 snows the results in ${\rm H_2}$ streams.

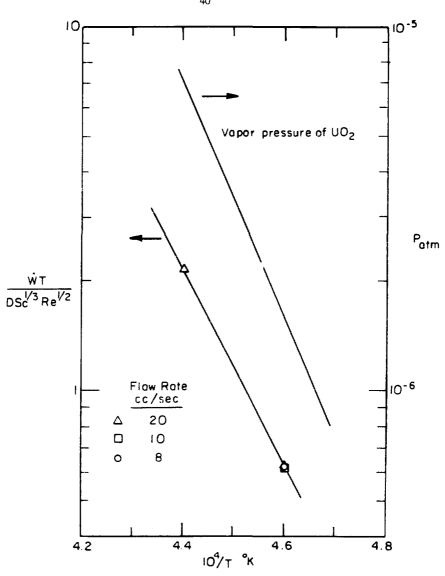


Figure 12. Temperature dependence of the UO_2 vaporization rate.

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Table 5. UO2 reduction in hydrogen.

т°к	Flow Rate			Expected W
	cc(STP)/sec	₩ in H ₂ , mg/min	W in Ar, mg/min	in H ₂ , mg/min
2173	10	0.011 ± 0.0005	0.0 0 ± 0.001	0.021
2173	20	0.050 ± 0.0025	0.014 ± 0.002	J.029

Fig. 13 is a typical weight loss curve from thermopalance output in which one can see the snarp change of the slope after the argon stream was replaced by hydrogen.

Also snown in Table 5 are the weight loss rate in Ar for comparison. Using Eq. (23), the ratio of $\kappa_{\rm m}$ in H₂ to $\kappa_{\rm m}$ in Ar is estimated to be ~2.1 at the same temperature and flow rate, and the weight loss rates due to pure evaporation are listed as "Expected W in H_2 " in the table. However, the data at a flow rate of 10 cc/sec show a slower rate than expected, especially considering the contribution of reduction. Inis is believed to be experimental error. Another possibility is that the rate 0.011 mg/min was measured before steady state was reached. For a flow rate of 20 cc/sec, the weight loss rate 0.050 - 0.029 = 0.021 mg/min could be attributed to the reduction of JO_2 by H_2 . At this rate, JO_2 could have been reduced to JO_1 977 in an hour, assuming that the vapor pressure was independent of stoichiometry within this range. This rate of reduction was significant and may have peen mass transfer-limited (rather than limited by a surface chemical reaction or solid state diffusion of oxygen in the $y_{0,x}$

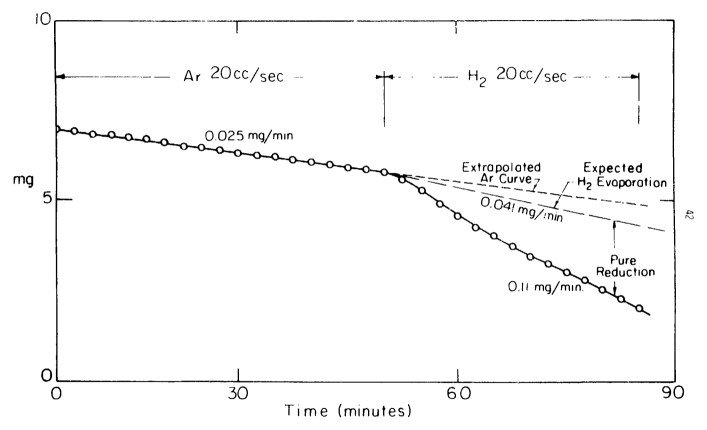


Figure 13. Typical weight loss data for 100_2 in flowing argon and hydrogen. XBL7712-6574

3.2.3.3 Iron Vaporization

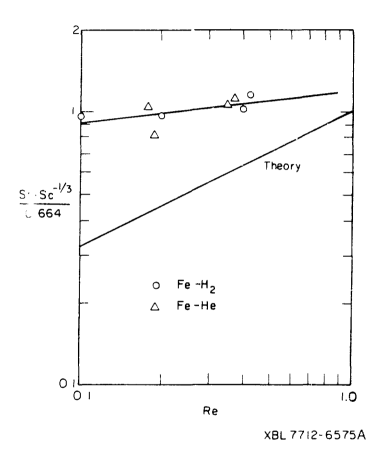
Iron evaporation results are snown in Table 7. Fig. 14 shows that mass transfer was 2-3 times higher than expected from the theory. Using least squares fitting, the experimental data can be represented by: $Sn = 0.3 \cdot 5c^{1/3} Re^{(0.115\pm0.3)}$. Diffusivities of Fe in H₂ and He calculated by theory [30] are tabulated below:

Table 6. Diffusivity and Viscosity of He and Hy

ťķ	J _{Fg-m2} ,cm ² /sec	O _{Fe-He} ,om ² /sec	J _H ,Poise	₄ _{He} ,Poise
1693°	14.48	13.76	2.7 x 10 ⁻⁴	5.3 x 10 ⁻⁴
1773	15.75	14.88	2.9 x 10 ⁻⁴	5.5 x 10 ⁻⁴

The dependence of the Sherwood number on the Reynolds number does not agree with the value of 0.5 predicted by laminar flat plate boundary layer theory. This discrepancy may be due to the sensitivity of the hydrodynamics to slight misalignment of the hanging specimen from the vertical axis. Note also that the magnitude of the discrepancy between theory and experiment is a factor of \approx 2 instead of \approx 6 for JO₂.

The heat of vaporization $\Delta H_{\rm vap} = 91.2 \pm 32.1 \, {\rm kcal/mole}$ was obtained from Fig. 1b, which is in good agreement with $\Delta H_{\rm vap} = 90.1 \, {\rm kcal/mole}$ mole obtained from the equilibrium vapor pressure curve [29].



From 14. Comparison of iron vaporization in hydrogen and in herost.

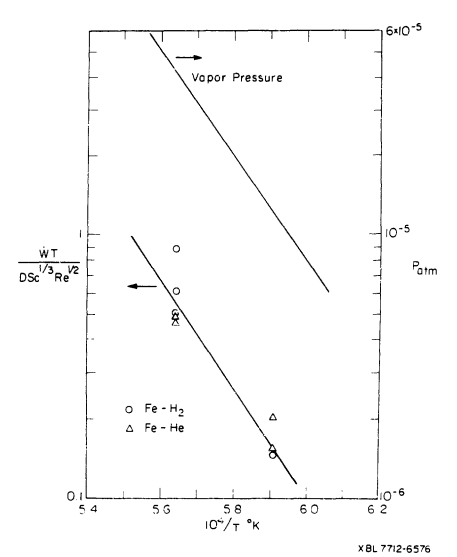


Figure 15. Temperature dependence of iron vaporization.

Table /. Fe Vaporization in 12 and He

	(Conditions			rłą	l	н	<u>!</u>	
т•к	Peq,aim	Flow cc/sec	v cin/ sec	₩ mg/min	k _m cm/sec	k in cin/sec	₩. mg/min		K'n :/sec
1486	~	20	13.96	≃ 0	-				
1693	1.21×10^{-5}	10	7.95		. - .		0.012 ± 0.003	22.9	9.1
1693	1.21 × 10-5	50	15.90	0.013 ± 0.004	24.8	13.63	0.013 * 0.002	24.8	12.9
1773	4.12 x 10 ⁻⁵ 4.12 x 10 ⁻⁵	5	4.16	0.038 ± 0.003	22.2	3.64			
1773 1773	4.12 x 10-5	10 20	8.33 16.65	0.038 ± 0.007 0.041 ± 0.005	22.2 23.6	7.30 14.59	0.030 ± 0.005 0.039 ± 0.002	17.5 22.8	9.3 13.6

3.3 Reduction of UO2

Uranium oxide at elevated temperature exists as a single phase over a broad range of stoichiometry (see Fig. 1). In the oxygen deficient region, this extends down to 0/U ratio of about 1.46 at a monotectic temperature of 2425°C.

As was mentioned earlier, hypostoichiometric urania can be prepared by reduction of UO_2 in hydrogen. In this technique the purity of hydrogen is essential. The ultimate stoichiometry is limited by the oxygen potential of the stream (see Fig. 3).

3.3. Apparatus

The thermobalance was not needed in the UO₂ reduction step.

From the previous experience a standard procedure was established.

Since hydrogen free of oxygen and water is essential to the capability of reduction, in addition to the activated charcoal filled liquid nitrogen trap, H_2 was passed through an oxygen getter consisting of copper turnings at 650°C. This served to remove any oxygen in the hydrogen stream by oxidation of copper. Copper turnings were contained in a 1-1/4 in. 0.D., 10 in. long stainless steel tube which was heated from outside. On some occasions fused titanium lumps were used as the getter instead of copper.

3.3.2 Procedure for Reducing UO2

One set of matched ${\rm U0}_2$ wafers (one ${\rm U}^{16}{\rm O}_2$ and the other ${\rm U}^{18}{\rm O}_2$) that had gone through the oxygen exchange steps was placed in the furnace together. The rhenium rig used in the oxygen exchange step was used here. The two wafers were separated as far as possible in order to prevent (or minimize) premature isotope exchange via vapor phase

transport. Also present was another identical UO₂ wafer, which was to serve the purpose of stoichiometry determination after the reduction. This was necessary because a nondestructive method of stoichiometry determination was not available.

- (1) Pump out the entire system; samples were simultaneously degassed at 300° C. This time all tubing was baxed as were the molecular seive and the activated characoal-filled liquid nit ogen moisture trap. This step was to remove any $\rm H_2O$ inside the tubing. It was most essential when the system had been previously exposed to $\rm H_2O-H_2$ mixture.
- (2) Fill the system with helium; oxygen-free helium was used.

 Flush the system for 30 mins.
- (3) Heat up the copper getter to 650°C.
- (4) Start heating up the furnace to 1000°C ; Again, slow heatup was essential to prevent cracking of the UO_2 . Approximately 30 mins. were requied.
- (5) Shut off helium.
- (6) Flow hydrogen at 20 cc/sec; Hydrogen was passed through the liquid nitrogen trap and the copper getter.
- (7) Raise the temperature to $1900 \sim 2000^{\circ}\text{C}$; Normally 2-4 hrs. were required to substantially reduce UO_2 .
- (3) Stop flowing hydrogen and anneal the samples for 2 hrs.; this was to eliminate any oxygen concentration gradient in the samples.
- (9) Cool down the temperature to 1000°C.

- (10) Pump out the entire system; Before cooling down to room temperature the system had to be absolutely free of hydrogen to avoid hydriding the precipitated uranium which would lead to a complete destruction of the samples.
- (11) Fill the system with helium and flow.
- (12) Cool down to room temperature.

3.3.3 Results

Shown in Table 8 are several results of reduction. The degree of reduction was controlled by the reaction time and temperature.

Table 8. Typical Results of Reduction in H2 Flow.

			Final O/U		
T°C	H ₂ cc/sec	time, hrs	poly. crystal	single crystal	
1850	20	1	1.972	_	
1850	10	1	1.970	_	
1850	20	2	1.954	1.970	
1950	20	2	1.951	_	
1950	20	4	1.955	_	
1950	20	4	1.954	_	
2000	20	4	1.950	1.953	
2000	20	4	1.955	1.950	
2020	20	4	1.948	_	
2040	20	4	1.945	1.944	
2040	20	4	1.951	· · · · <u>-</u>	

The stoichiometries were measured by a thermogravimetric method which will be described in the next chapter.

Apparently the stoichiometry approached 1.95 rather rapidly and reached a plateau. It looks, nowever, as if higher temperatures may lead to lower stoichiometries, which is consistent with the fact that at higher temperature the equilibrium oxygen potential is higher (see Fig. 3).

However, temperatures above 2050°C were not imposed because of a severe distortion of the snape of the sample due to the extremely high vaporization rates. The reduced samples were not to be ground or polished again because of the possibilities of contamination and change in stoichiometry. Therefore, a severe distortion in the snape of the samples could not be tolerated.

Both single crystal and polycrstalline samples showed similar stoichiometries after the reduction in similar conditions. At this stage this was interpreted as (1) exygen diffusion in ${\tt U0}_2$ was not rate controlling or (2) reaction had reached equilibrium.

At the conditions of $T=2000^{\circ}\text{C}$ and H_2 flow of 20 cc/sec for 4 nrs, approximate stoichiometries of 1.95 could be consistently obtained.

Shown in Fig. 16 is a photomicrograph of a reduced sample. The uranium metal precipitates are immediately visible (bright areas); many of them associated with voids. Using EDAX (Energy Dispersive X-ray Analysis), the bright areas were confirmed to be uranium metal. Figure 17 shows one of the uranium precipitates under SEM.

The reduced wafers that were to be used in diffusion experiment already had gone through oxygen exchange steps described previously. This process, which took ~40 hrs., combined with 4 hrs. reduction at high temperature, yielded very large grains, approximately 200 microns. Figures 18 and 19 show the morphologies of $\rm U^{160}_{2-x}$ and $\rm U^{180}_{2-x}$ specimens, respectively. As expected they show approximately same grain size.

3.4 Stoichiometry Determination

3.4.1 Survey of Methods

There are a number of different methods for measuring the stoichiometry of urania. Methods commonly used are:

- 1. X-ray diffraction [32]: The lattice parameters are correlated with the O/U ratio. By measuring this parameter the stoichiometry is obtained.
- 2. Solid State Electrolytic Cell [33]: High temperature galvanic cells using Ni-NiO mixture and heavy metal oxides $(TnO_2, UO_2, etc.)$ are utilized. The equilibrium oxygen potential can be measured by the emf generated between the two electrodes.
- 3. Gas Equilibrium Method [34]: Using appropriate gas mixture of ${\rm CO-CO_2}$ or ${\rm H_2O-H_2}$ the specimen is brought to stoichiometric ${\rm UO_2}$ and the total accumulated change in the ratio of the ${\rm CO/CO_2}$ or ${\rm H_2O/H_2}$ during this process is recorded to calculate the original stoichiometry.
- 4. <u>Thermogravimetric Method</u> [35]: By measuring the weight change of the specimens when they are brought to a known, standard stoichiometry (00_2 or 0_30_8), the original stoichiometries can be obtained.

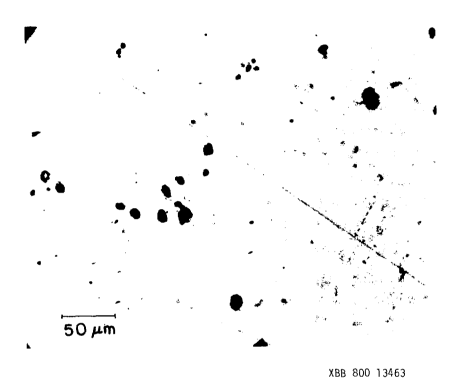
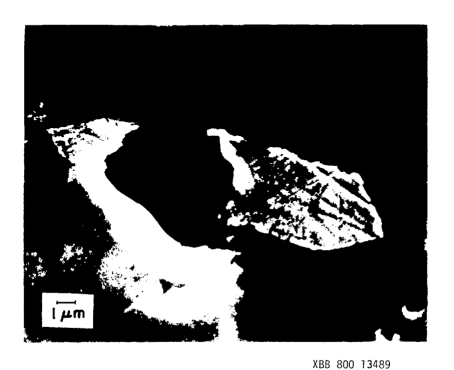


Figure 16. Photomicrograph of the reduced urania. Bright spots are the uranium metal precipitates.



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Figure 17. Photomicrograph of the uranium precipitates under SEM.

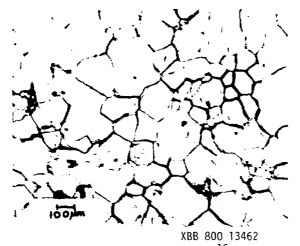


Figure 18. Photomicrograph of the $v^{16}o_{2-x}$ surface.

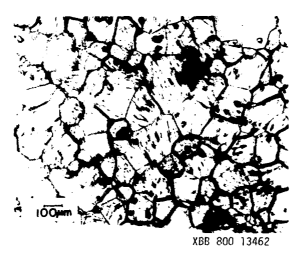


Figure 19. Photomicrograph of the $u^{18} o_{2-x}$ surface.

Of all these methods, the thermogravimetric method is the most convenient and reliable for routine experiments. Also it was readily available since the system was equipped with thermopalance.

Since there was a substantial amount of vaporization at high temperature, the degree of reduction could be measured directly by the total weight change during the reduction step. This problem was overcome simply by having another piece of ${\tt UO}_2$ present during the process and subsequently oxidizing it back to ${\tt UO}_2$ at low temperature while measuring the weight gain, from which the stoichiometry can be calculated by:

$$\frac{0}{U} = 2.0 - \frac{270}{16} \frac{W_f - W_i}{W_f}$$
 (25)

where \mathbf{W}_{i} and \mathbf{W}_{f} are weight of the sample before and after the reaction, respectively.

3.4.2 Procedure for Stoicniometry Measurement

The atmospheric conditions for this oxidation were almost the same as the ones used in the oxygen exchange step described previously. Here, normal water was used and in order to minimize the chance of vaporization, lower temperatures were employed ($\sim1300^{\circ}$ C rather than 1500° C). At these temperatures the amount of evaporation was negligible, yet reaction was fast enough. As in the oxygen exchange procedure, the water was kept at 7.5° C, which would yield an oxygen potential of -105 < cal/mole at 1300° C.

Sometimes the specimen surface spalled due to a sudden violent reaction between uranium precipitation and H₂O, which caused a large weight change. Therefore, the specimen had to be contained in a rhenium basket so that the small particles were not lost from the weight measurement. This basket in turn was suspended from the thermopalance to follow the weight change continuously throughout the experiments. Although spallation did not happen very often, the precautions were taken each time.

Before and after the reaction, the specimen was weighed butside the system using the Mettler microbalance to compare with the output of the thermobalance. Most of the time those two readings were in good agreement. However, the calibration of the thermobalance seemed easy to disturb when a sudden large force was exerted, for example a sudden change in flow rate. Thus, whenever there were significant discrepanties between those two readings, the Mettler's reading overruled that of the thermobalance. The main function of the thermobalance was to provide the indication that the reaction was completed by showing a steady weight. The following is the detailed procedure:

- 11 Measure weight of the specimen before loading.
- 2. Pumb ti it invstem and degas the specimen.
- (3) Fill the sy lium.
- 1' Flow neliu | cc/sec.
- (5) Heat up to 1000°C; The reduced samples should not be exposed to hydrogen at low temperature.
- .6) Shut off helium.

- (7) Flow hydrogen 10 cc/sec.
- (8) Heat up to 1300°C.
- (9) Start flowing H₂0-H₂ 5 cc/sec.
- (10) Observe the weight reaches a steady state.
- (11) Flow hydrogen 10 cc/sec
- (12) Cool down to 1000°C.
- (13) Snut off nydrogen.
- (14) Flow helium 10 cc/sec.
- (15) Cool down to room temperature.
- (16) Shut off nelium.
- (17) Take out the specimen and measure the final weight.

Since the balance was extremely sensitive, the valve operations and flow rate changes gave considerable perturbations. Also for different conditions (temperature, gas species, flow rate), the readout was different even though the weight of the sample remained unchanged. Changes occurred because (1) for different gas species and/or flow rates the buoyancy and skin friction forces are different and (2) for different temperatures the brightness in the balance chamber changes due to the rapiation and this contributes to a change in the readout because the balance has a photoelectric tube the current of which is directly proportional to the readout. Therefore, it was essential to compare the readout at identical conditions (i.e., at the same temperature, same gas species, and same flow rate), to obtain a true weight change of the specimen. It was also essential to take the reading after the system reached a steady state following each perturbation.

To satisfy these requirements the weight increase was calculated by comparing the weights at steps (9) and (10), (11) and (8), (12) and (17), (14) and (15), and (15) and (4). These points are depicted in Fig. 20. In most cases these values were in agreement to within ± 0.005 mg. An average of these values represented the thermobalance measurement.

In order to make sure that the chenium wire and basket holding the specimen did not react with $\rm H_2O-H_2$ during the process, a dummy experiment was performed without any sample in the basket. In the basket was a piece of rhenium foil of approximately the same weight as the $\rm JO_{2-x}$ specimen. The test consisted of exactly the same steps as the actual stoichiometry determination, except that the temperature was raised from 1300°C to 1500°C gradually in the $\rm H_2O-H_2$ stream. No weight change was observed through 1500°C. This proved that the conditions for the stoichiometry determination were inert to rhenium and all the weight change observed could be attributed to the reaction of $\rm JO_{2-x}$ specimen with $\rm H_2O$.

Shown in Fig. 20 is a typical output of the thermobalance during the experiment. Each step of the procedure was marked by a numbered arrow. Step 3 shows a sudden huge perturbation when the pre-evacuated system is filled with helium. Similar, but smaller, spikes are seen on many other occasions whenever there were valve operations or changes in the flow rate. Also observed is a difference of ~0.4 mg in readout before and after filling the system with helium, although the weight of the specimen should have remained unchanged. This change was due to the buoyancy force exerted to the whole balance system by helium gas.

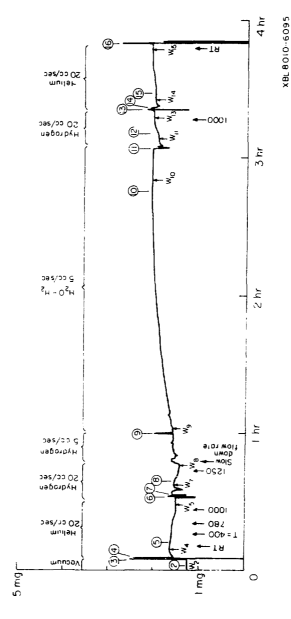


Figure 20. Typical thermobalance output during the stoichiometry determinations.

Also, as the temperature increased from room temperature to 1000° C the readout decreased by ~ 0.15 mg due to the radiation into the balance chamber.

Before feeding $\rm H_2O-H_2$, it was verified that the weight remained constant under steady $\rm H_2$ flow. This test was necessary to make sure that (i) evaporation of the sample was negligible and (ii) all the weight increase in the $\rm H_2O-H_2$ atmosphere could be attributed to the reaction of $\rm H_2O$ and $\rm UO_{2-x}$.

In the example of Fig. 20 it took 1.5 hrs to complete the reaction. The time required for the reaction varied from 1.5 hrs to 3 hrs in most cases. It is very likely that this range was due to the different geometries of the samples.

Table 9 shows the weight increase measured at different conditions.

As mentioned earlier, for each measurement the conditions of comparison were identical.

Table 9. Weight increase measurements at different condtions.

₩j₩k	gas	Condition: flow cc/sec	T°C	∆w mg
W10-W0	Н20-Н2	5	1250	0.475
₩10-₩g ₩11-₩8	H2 τ	10	1250	0.475
W12-W7	H2	10	1000	0.48
W14-W5	Hē	10	1000	0.48
W15-W4	He	10	room temp.	0.475
WF-Wi	air	0	room temp.	0.485

Weights $W_i = 0.178370$ gm and $W_f = 0.178855$ gm were measured by the Mettler microbalance for this example. The measured weight

increases in Table 9 are in good agreement with the measurement by Mettler microbalance. Using a 0.485 mg increase, the initial stoichiometry was calculated from Eq. (26) to be 1.954.

4. DIFFUSION EXPERIMENT

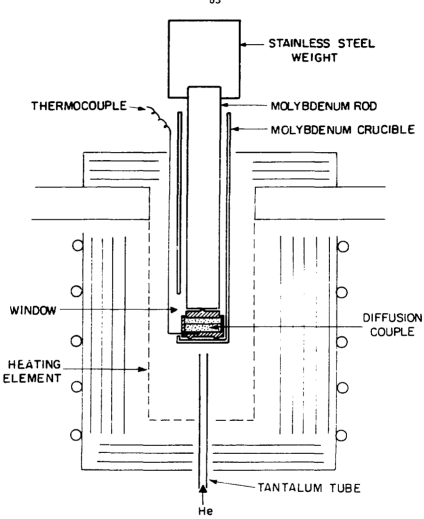
4.1 Apparatus and Procedure

Following the preceding steps, two identical nypostoicniometric $\rm UO_{2-x}$ wafers (one with ^{18}O and the other normal) were prepared for each diffusion experiment. These two matched wafers were put together with a uranium foil 0.003 in. thick in between. Shown in Fig. 21 is the experimental setup. Experiments were performed in a glass belljar filled with high purity helium. Throughout the experiment the nelium flowed continuously from underneath the sample to keep the surrounding atmosphere as clean as possible.

The furnace was heated by a tungsten mesh heating element of 3 in. diameter and 6 in. high. The diffusion couple was positioned in the center of the heating element to establish as uniform a temperature as possible in the diffusion couple. Since the samples were approximately 1 mm thick, which is very small compared to the size of the heater, it was assumed that the temperature was uniform throughout the diffusion couple.

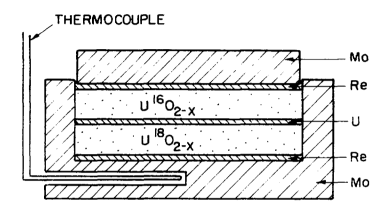
The diffusion couple was enclosed by a molybdenum crucible of 0.5 in. I.D. (see Fig. 22). It was necessary to make the crucible wide enough to accommodate the thermal expansion of urania.

In order to promote good contact between the two wafers and to minimize the thickness of the liquid uranium layer, the diffusion couple was put under compression. This was achieved by a weight on top of the molybdenum crucible.



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Figure 21. Diffusion experiment setup.



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Figure 22. Diffusion couple arrangement.

Temperature was measured by a W3%Re-W25%Re thermocouple the not junction of which was placed in the center of the molypdenum cruciple (see fig. 22). The distance between the junction and the end of 90_{2-x} wafer was less than 1/16 in. The thermocouple was connected to a digital indicator from which the temperatures were read directly. In order for the thermocouple to respond quickly to the temperature changes, unsheatned pare wire thermocouple was used.

Prepared samples were stored in an inert gas or vacuum until they were used in the diffusion experiment.

The following steps were taken for each run:

- (1) Sump out the entire system.
- (2) Heat up to 400° C to degass the system for one nour.
- (3) Fill the system with high purity oxygen-free helium.
- (4) Flow melium at 20 cc/sec.
- (5) Slowly neat up to 1100° C; Slow heatup was essential to prevent the samples from cracking. Since the uranium remained solid up to the meiting point of 1132° C, it acted as a parrier to the premature diffusion.
- (6) Rapidly heat up to the desired temperature; Once the uranium melted, diffusion through it would occur. Increfore, it was necessary to reach the temperature as quickly as possible. It usually took 40 sec.-2 mins., which was short compared to the total annealing time.
- (7) Maintain the temperature for a predetermined time period; For the first experiment, the diffusion coefficient had to be guessed in order to determine the appropriate annealing time.

from the result of this first experiment, diffusion coefficients at other temperatures could be estimated and the annealing time which would yield an appropriately developed diffusion profile was determined. The criteria were that the time should be long enough for diffusion to penetrate at least half of the thickness, yet no more than 2/3 of the thickness. By having a less than fully penetrated diffusion profile, the original $^{18}0$ concentration on both sides of the diffusion couple could be checked. This is discussed in greater detail in the next section.

(3) Turn off the power supply to cool the couple as quickly as possible; Since sintering of the two wafers could occur in some contact areas of the sample, diffusion could take place even below the melting point of uranium. Time required to reach 1100°C was measured, which was also short compared to the overall operation time.

After each run, the couple was taken out and was cut in half using a low speed diamond saw. Each half was mounted in a copper filled conductive thermosetting epoxy and polished to 6 micron grade using champed paste.

4.2 Sample Analysis and Results

 13 O/(13 O+ 16 O) profiles were determined by Hanford Engineering Development Laboratory and Argonne National Laboratory using ion microprope mass analyzer (IMMA). The basic operating principles of IMMA are described elsewhere [36,37]. Basically, an ion beam is accelerated to an adjustable focal spot on the sample surface and the sputtering ions

are analyzed by mass spectrometer. This technique was used by Marin et al. [1] and Contamin et al. [3] for urania and by Valencourt et al. [37] for porous $UO_{2+\nu}$.

For the present analysis an $^{28}N_2^+$ primary beam was used in 15-20 kV accelerating potential. The area analyzed was about 5x6 microns for each spot. Each area was sputter cleaned for 30 seconds before collecting data to eliminate any surface effects.

Since $\rm H_2O$ also has mass 13, it is indistinguishable from $^{13}\rm O$ in the mass spectrometer. Therefore, water had to be avoided in grinding and polishing the samples and they had to be degassed in a high vacuum (5x10⁻⁹ torr) for 2-3 days before each analysis. Also samples were stored in vacuum after experiments and shipped in a small leak-tight stainless steel containers filled with slightly pressurized helium.

Experiments were conducted at eight different temperatures in the range 1257 - 1597°C, each of them corresponded to stoichiometries in the range of 1.993-1.955 following the lower phase boundary. An empirical equation developed by Fryxell et al. [38] was employed to determine the stoichiometries of the oxide in the two phase region:

$$lnx = 3.073 - 12675/T^{\circ} K \tag{27}$$

Snown in Table 10 are $(0/J)_0$, the stoichiometries of the samples used in each experiment, annealing temperatures, and (0/J), the stoichiometries of the oxide phase in the two phase region corresponding to each temperature.

Table 10. Experimental Temperatures, stoichiometries, initial $^{\frac{1}{18}}$ 0 compositions of the two wafers, and diffusion times.

RUN	r*c	(0/U) ₀	(0/11)	130 (/) enriched wafer		original ¹³ 0 enrichment ()	t _i mins.	t ₂ mins.	t _o mins.	-
1 2 3 4 5 7 8 9 1J* 11* 12*	1190 1257 1330 1400 1400 1530 1565 1597 1330 1400 1490 1565	1.970 1.973 1.954 1.955 1.975 1.948 1.948 1.945 1.950 1.956 1.944	1.993 1.990 1.945 1.980 1.970 1.965 1.960 1.955 1.985 1.980 1.970	70.9 70.6 70.0 33.6 54.1 45.7 47.4 44.3 45.2 61.0 58.1 58.7 65.1	1.2 2.2 0.3 4.9 2.1 1.4 2.1 1.5 4.9 5.0 2.1 2.7	71.5 73.5 72.3 40.1 61.5 40.0 51.2 48.4 47.9 69.8 65.2 65.2	1.0 1.3 0.8 2.2 3.0 2.5 2.1 2.5 3.0 1.6 2.2 3.0 2.5	1.0 1.4 0.8 1.8 1.5 2.2 2.0 2.5 2.3 1.5 1.8 3.0 2.5	340 180 20 45 30 50 35 20 20 45 45 28 15	68

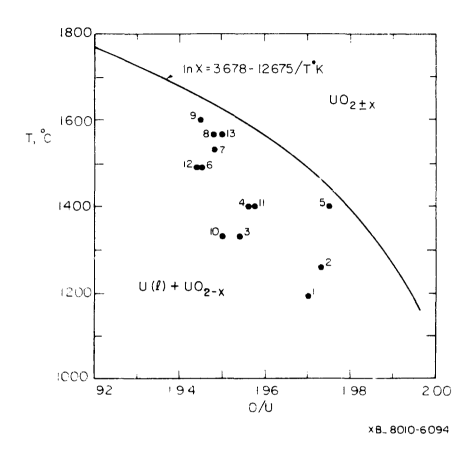
^{*}Single Crystals

In Fig. 23 the experimental points are depicted in the phase diagram. In order to make sure that the experimental points were in the two phase region, annealing temperatures were kept at least 50° C below the two phase boundary.

Calculation snowed that under these conditions the effect of liquid phase on diffusion in the oxide is insignificant.

Snown in Figs. 24-37 are normalized profiles of 13 Or 13 O+ 15 Or analyzed by IMMA. For normalization of the naw data it was imperative to have correct values of original isotopic concentrations on both sides of the diffusion couple. Although the isotopic enrichment had been measured from weight increase after the oxygen exchange step, it was essential to confirm these measurements because any error in this value was directly bassed along to the diffusion coefficient calculations. Since the wafers had gone through a high temperature reduction step, it was suspected that the original enrichments have have been altered by vapor phase isotope transport.

This could be verified by companing the original isotopic enhichment with the IMMA data hear the end of $\frac{100}{120}$, wafer where diffusion had not vet benetrated. From the unnormalized IMMA data, it was polyrous that in every experiment at least one third of the wafer had not been affected by diffusion. In some of the samples, No. 3, 4, 10, 11, 10; more than half of the wafer was unchanged. Average values of these unchanged, flat profile regions are tabulated in Table 10. Also shown in the table are $\frac{18}{100}$ original enhichment values calculated from weight inchease.



fraure 33. Experimental points in phase diagram.

IMMA data near the ends of $U^{16}O_{2}$ wafers showed substantially higher values of 18 O concentration (1-5 percent) than natural abundance (0.2 percent), which is easily detectable by IMMA (see Table 10). Yet, the profiles were also flat at least up to one third of the thickness, i.e., the maximum depth of diffusion penet ation was two thirds of the thickness, i.e., the overall ¹³0 profiles were symmetric in every experiment. This implies that the ¹³0 phoentrations were already higher than natural abandance in $\mathbb{S}^{16}\mathrm{O}_{-3}$ wafers even before the diffusion anneal. The immediate explantion is that there had been oxygen exchange between the two wafers through the vapor phase during the 4 hours of reduction at high tempera are. This interpretation was strengthened by the fact that the sum of average values of both sides of the diffusion couple were close to the original enrichment (see Table 10). Based on this interpretation, the two 18 0 fraction on the sides of the diffusion couple aw a from the interface obtained by IMMA were used for normalization or the data (see the definition of **a** in Eq. (1)).

Experiments 10 through 13 are on single Hystals.

In most experiments two different lines perpendicular to the interface were proped in order to average out six difference in the ¹³0 profiles. They were marked as triangles and squares in the Figs. 24-37. First one was usually near the center and the other was near half way to the edge from the center. Each position of the lines was carefully chosen under the microscope in such a way that the line of probe would not choss any voids or chacks which would cause IMMA to pick up values from different planes. As can be seen, the profiles of the two thaverses were very close to each other in most experiments.

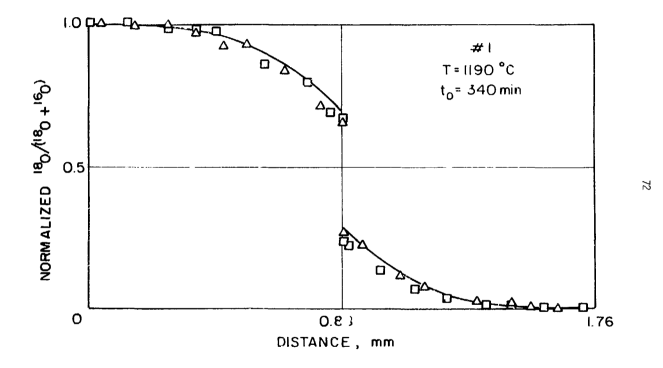
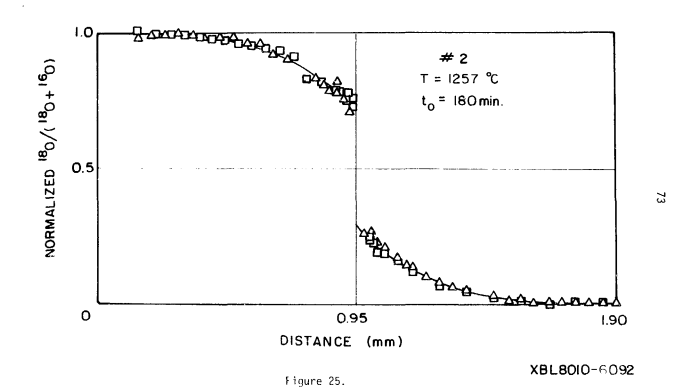


Figure 24. Normalized profile of ¹⁸0 concentration. Squares and triangles are for traverses near the center and near half way to the edge from the center, respectively.

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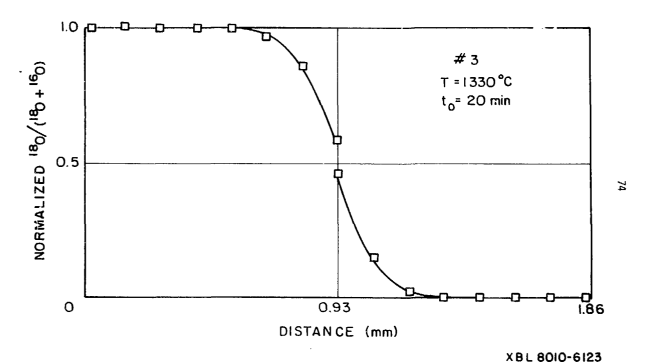


Figure 26.



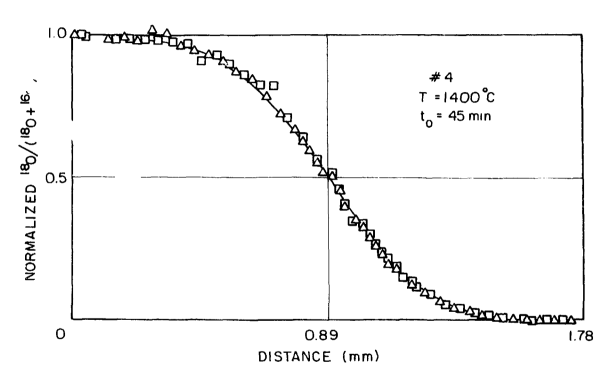
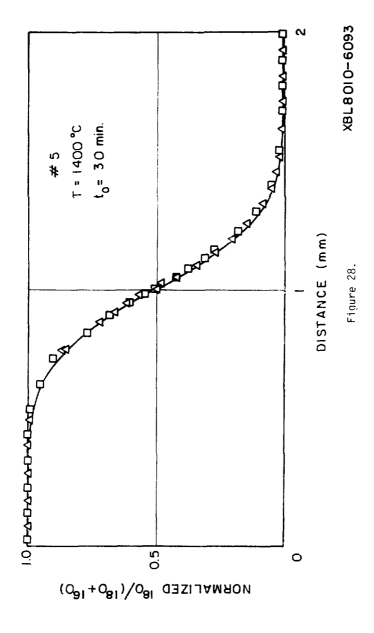
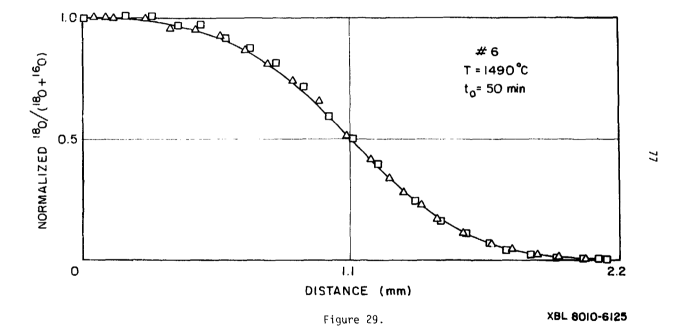


Figure 27.

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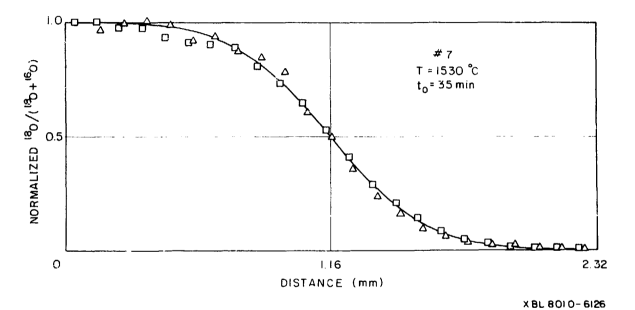


Figure 30.



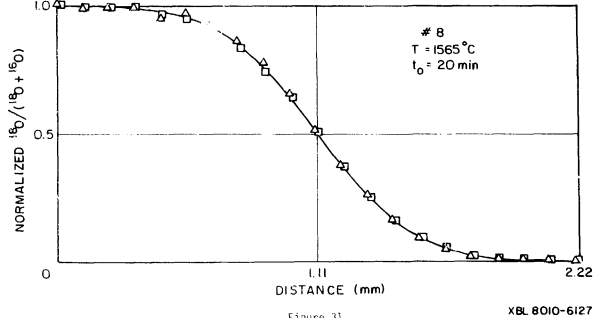
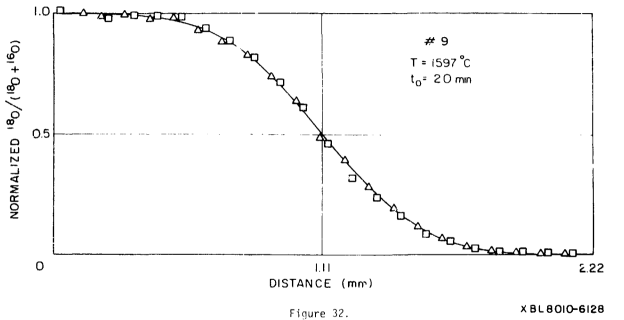
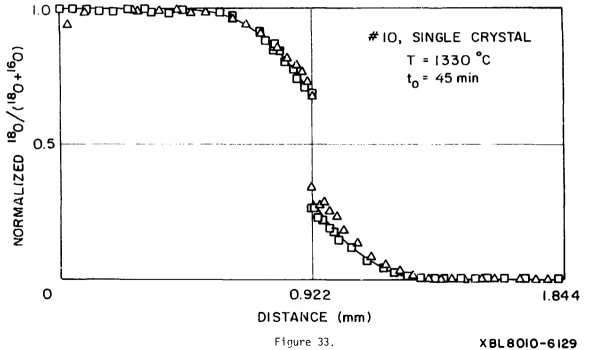


Figure 31.

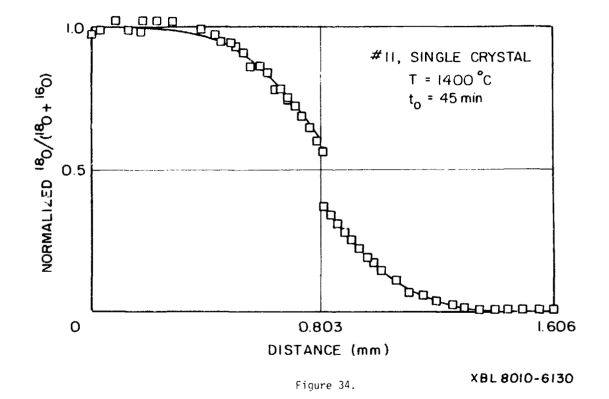




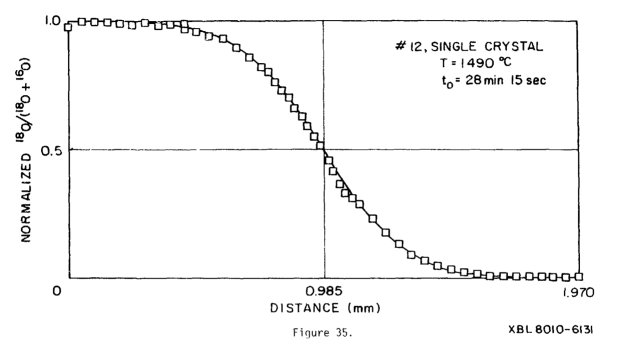




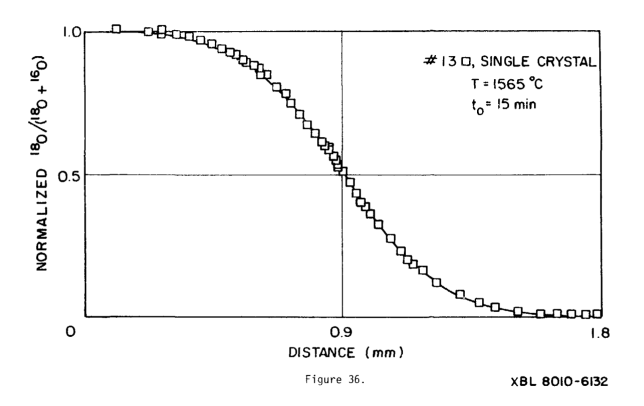




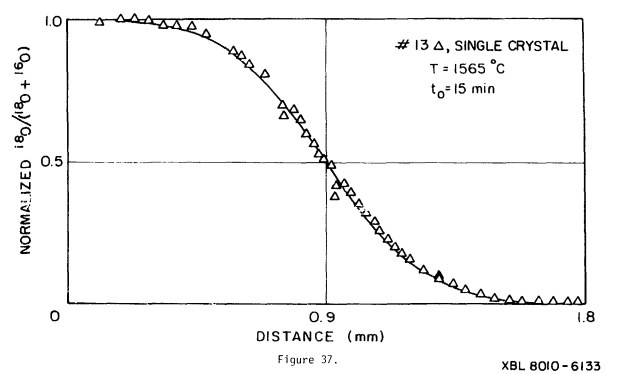












There were basically three different types of bonds observed between the two wafers.

- (1) Continuous thick uranium layer (5~30 $_{\rm um}$) across the entire sample: As expected, samples of this type of bond yielded discrepancies in 18 0 concentration across the interface. Samples No. 1, 2, 3, 10, 11 belong to this type. Shown in Fig. 38 are the photomicrographs of the interface of samples No. 2 and 11. The uranium layer is distinctly visible as a white hand.
- (2) Sintered interface: Naturally this type showed no ¹⁸0 discrepancies. Although two wafers were sintered together, the original interface was easily identifiable decause a number of uranium particles remained along the interface (see Fig. 39). This type was observed in samples such as No. 8 and 12 which were annealed at high temperatures.
- (3) Combination of (1) and (2): This type of specimen snowed some areas which were sintered and some which had a uranium layer (see Fig. 40). It is likely that this structure occurred because the wafers were not truly parallel and therefore the contact was not uniform. However, in every case of this type, the uranium layer was extremely thin (2-3 microns). For this type of samples one of the probe lines traversed the sintered area and the other crossed the uranium layer for the purpose of comparisor. Those two profiles were always in good agreement and showed no

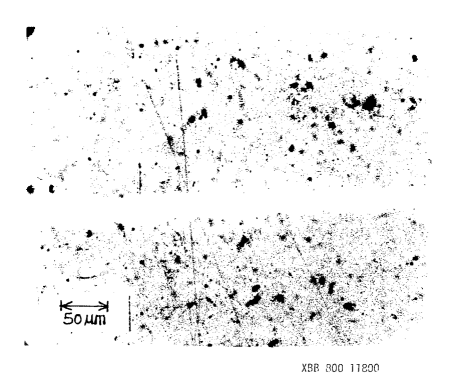
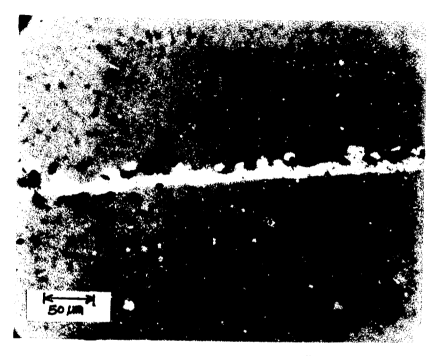
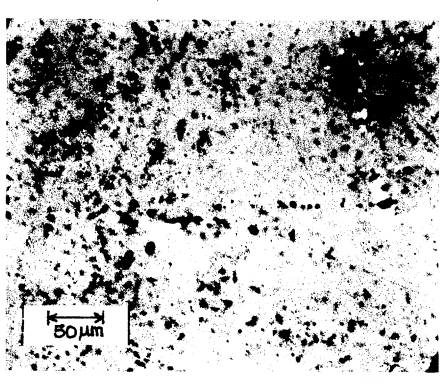


Figure 38A. Thick cranium interface (sample #11).



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Figure 38B. Thick uranium interface (sample #2).



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Figure 39. Sintered interface (sample #8).

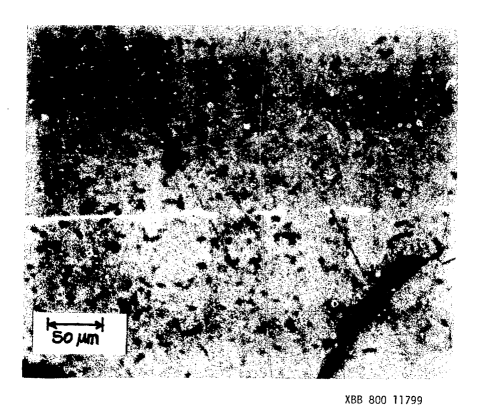


Figure 40. Half-sintered interface (sample #5).

significant discrepancies at the interface, most likely because the uranium layer was extremely thin. Samples No. 4, 5, 6, 7, 9, 13 belong to this type.

In every sample, whenever the probe line crossed the uranium layer, the $^{18}\mathrm{O}$ concentrations of the points immediately adjacent to both sides of the uranium layer were measured in order to detect any discontinuity in $^{18}\mathrm{O}$ concentration across the uranium layer.

The contact between liquid uranium and urania was excellent (see Fig. 38a), which was essential to minimize the interfacial resistance. No gap or voids were observed even when the surface was not smooth (see Fig. 38b).

The normalized data were fitted to Eq. (13) using a computer code MINUIT to find the best values of D and B. However, since time was required to reach the operating temperature and to cool down after the experiment, diffusion during these stages had to be taken into account.

Corrections were significant in higher temperature experiments where the annealing time was relatively short and diffusion was fast.

As was indicated, the temperature could be raised slowly up to the melting point of uranium without causing any premature diffusion. From 1100°C (near the melting point of uranium), the temperature was raised rather rapidly up to the annealing temperature, which took at most 3 mins. Since this is a temperature range where urania is highly plastic, this operation could be carried out rapidly without cracking the samples.

The times required to heat the couple from 1100° C to the annealing temperature, t_1 , are shown in Table 10. Also shown are the cooling

times to 1100° C (t₂). Usually 2-3 mins, were taken for cooling from 1100° to 500° C so that even for the sintered diffusion couples it was assumed that below 1100° C diffusion annealing was negligible.

The nominal annealing times, t_0 , are also tabulated.

To take these thermal transients into account, however, it is apparent from the nature of the problem that only an approximation is possible. Here, an effective $\mathbf{t}_{\mathrm{eff}}$ was sought which would accommodate the diffusion anneal during these transient periods. Since the degree of diffusion anneal was a function of the product Dt, effort were concentrated on the value Dt itself to deduce the approximate value of $\mathbf{t}_{\mathrm{eff}}$.

The teff was defined as:

$$\int_{0}^{t} t \, D \, dt = D_{T} \cdot t_{eff}$$
 (28)

 D_{\uparrow} is the diffusion coefficient at the annealing temperature and t_{\uparrow} = t_{1} + t_{2} + t_{0} , the total time for the experiment.

To calculate $t_{\mbox{eff}}$, D(t) needs to be known. Therefore, $t_{\mbox{eff}}$ and 3 should be obtained by an iterative process.

It was assumed that the temperature T' changed linearly with time:

$$T' = 1373 + (T-1373)(t/t_1), \quad 0 \le t \le t_1$$

$$T' = T - (T-1373)(\frac{t-t_1-t_0}{t_2}), \quad t_1+t_0 \le t < t_1+t_2+t_0$$
(29)

and T is the annealing temperature in °K.

As temperature changes, the corresponding stoichiometry of the two phase boundary also changes. In near-stoichiometric material, the following equation can be assumed for oxygen self-diffusion by the vacancy mechanism [49].

$$D = D_0^{\mathsf{v}} \Theta_{\mathsf{v}} (1 - \Theta_{\mathsf{v}}) \exp(-\Delta H_{\mathsf{v}} / \mathsf{RT})$$
 (30)

 θ_v is vacancy concentration (x/2 in UO_{2-x}), ΔH_v is the activation energy of vacancy migration, and D_0^V is a constant. The equation will be discussed in greater detail in the next chapter.

The iteration was started by fitting Eq. (13) to the normalized data using $t=t_0$, the nominal annealing time. Fitting yielded diffusion coefficients for each experiment. This first diffusion coefficient obtained is shown in Table 11 as $D_{(1)}$. The $D_0^{\rm V}$ and $\Delta H_{\rm V}$ of Eq. (30) were calculated by plotting $\ln[D_{(1)}/\theta_{\rm V}(1-\theta_{\rm V})]$ vs 1/T using linear fitting.

from Eq. (27) and (30),

$$D_{(1)} = 0.5D_{0}^{V} \exp(3.678-12675/T^{T})$$

$$\left\{1-0.5\exp(3.673-12675/T^{T})\right\} \exp(-\Delta H_{V}/R^{T})$$
(31)

Combining Eqs. (31) and (29), $D_{(1)}(t)$ was obtained and from Eq. (28) $t_{\rm eff}$ was determined. The integration was done graphically. The first $t_{\rm eff}$ values are shown in Table 11 as $t_{\rm eff}$. Using $t_{\rm eff}$, new diffusion coefficients were obtained ($D_{(2)}$ in the table) and second iteration started. After the third iteration it was apparent

Table II. Effective times and diffusion coefficients during the iteration and the final θ values and effective times.

RUN	r°c	t _o mins	$D_{(1)} \times 10^{7} \text{ cm}^{2} \text{ sec}$	t'eff	D ₍₂₎ ×10 cm ² /sec	^t eff	$0 \times 10^7 \text{ cm}^2/\text{sec}$
1	1190	340	0.25	341	0.249	341	0.249
2	1257	180	0.47	181	0.467	181	0.466
3	1330	20	0.86	20.8	0.828	20.8	0.823
4	1400	45	1.45	46.8	1.398	46.6	1.400
5	1400	30	1.43	31.8	1.352	31.6	1.357
Ó	1490	50	2.43	52.1	2.237	51.8	2.346
7	1530	35	.65	36.5	2.536	36.4	2.551
8	1565	20	3.58	21.8	3.278	21.6	3.279
9	1597	20	4.48	21.9	4.087	21.7	4.129
10	1330	45	0.85	46.4	0.820	46.4	0.820
11	1400	45	1.11	46.8	1.072	46.6	1.072
12	1490	28	1.93	30.4	1.791	30.3	1.799
13	1565	15	4.19	ló.8	3.741	16.6	3.786

that values had converged adequately to the $t_{\mbox{eff}}$ and D figures shown in the table.

Values of D were used to deduce D_{0}^{V} and ΔH_{V} of Eq. (30). In Figs. 41 and 42, $\ln[D/\Theta_{V}(1-\Theta_{V})]$ vs 1/T is plotted for the polycrystalline samples and single crystal samples, respectively. Linear least squares fitting yielded $D_{0}^{V}=4.4 \times 10^{-4}$ and $\Delta H_{V}=11.7 \pm 3.0 \times 10^{-4}$ mole for polycrystalline samples;

$$0 = 4.4 \times 10^{-4} \, e_{y}(1 - e_{y}) \, \exp(-11700/RT)$$
 (32)

Similarly, for single crystals $\theta_0^{V} = 5.9 \times 10^{-4}$ and $\Delta H_{V} = 13.0 \pm 10.2$ kerroller where obtained:

$$D = 5.9 \times 10^{-4} e_{v}(1-e_{v}) \exp(-13000/RT)$$
 (33)

It should be noted that two separate experiments (No. 4 and No. 5 in Table 11) were conducted at 1400°C for different annealing time; one for 30 mins, and the other for 45 mins. This was to check the reproducibility. As can be seen in Table 11, the values of 0 obtained are very close to each other.

In order to clearly demonstrate the difference between single crystal and polycrystalline samples, one of the 1400°C polycrystalline experiments (No. 4) and single crystal experiment No. 11 were conducted simultaneously. This was achieved by simply stacking the single crystal diffusion couple on top of the polycrystalline couple in a

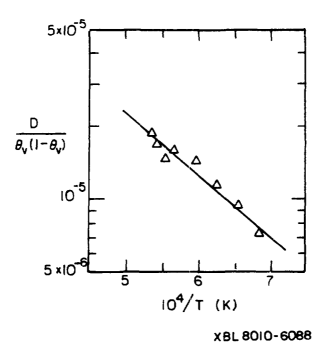


Figure 41. Plot of polycrystalline data to determine D_0^{ν} and $\Delta H_{\nu}.$

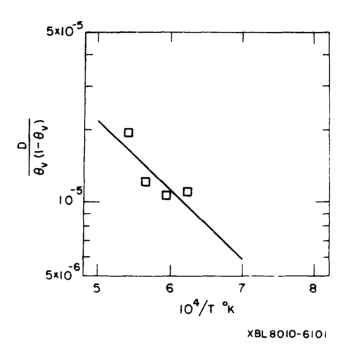
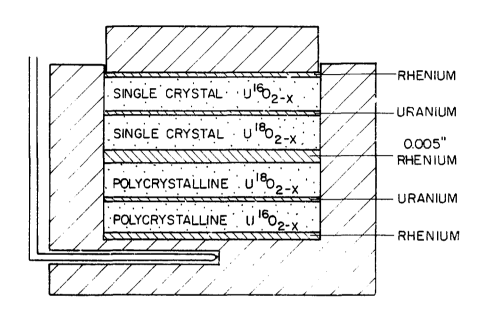


Figure 42. Plot of single crystal data to determine $D_{\rm p}^{\rm V}$ and ${\rm CH}_{\rm V}$.

crucible deeper than ordinary ones (see Fig. 43). A 0.005 in. thick rhenium foil was used to separate the two couples. By this arrangemen⁺, identical temperature history and all other conditions could be imposed to the two couples. The results show clearly that diffusion is slower in the single crystal (see No. 4 and No. 11 in Table 11). The difference is not very large because of the unusually large grain size ($\sim 200 \, \mu m$) of the polycrystalline samples.

Since the single crystal correlation has fewer data points and is not as good as the polycrystalline correlation, only Eq. (32) will be used in subsequent analysis.



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Figure 43. Double diffusion couple arrangement.

5. DISCUSSION

5.1 Comparison with Other Materials of Fluorite Structure

Due to the similarity in their crystal structure it has been widely believed that UO_{2-x} has the same type of crystal defect (anion vacancy) and diffusion mechanism (vacancy) as CeO_{2-x} and PuO_{2-x} . However, this supposition has never been confirmed experimentally. On the contrary, thermodynamic studies favor the excess uranium model [39].

Since the stoichiometry and temperature were changed simultaneously in the present experiment, it is difficult to separate the stoichiometry contribution to the enhanced diffusion at higher temperature. However, comparing the present data for \mathtt{UO}_{2-x} with existing data of stoichiometric \mathtt{UO}_2 [1] at the same temperature, it is readily seen that the present data are almost two orders of magnitude higher than those of stoichiometric \mathtt{UO}_2 . This effect can only be attributed to non-stoichiometry, thereby demonstrating the existence of defects in the anion sublattice, i.e., oxygen vacancies in \mathtt{UO}_{2-x} . Based on this fact the diffusion coefficient is analyzed in terms of vacancy contribution and migration energy.

It is well known from the random walk theory that for a vacancy mechanism the diffusion coefficient can be expressed as [49]:

$$D_{v} = D_{o}^{V} \Theta_{v} (1 - \Theta_{v}) \exp(-\Delta H_{v}/RT)$$
 (35)

where D_0^V is the pre-exponential factor, ΔH_V is the vacancy migration energy, Θ_V is vacancy concentration, and $(1-\Theta_V)$ is diffusion path probability.

In substantially hypostoichiometric $U0_{2-x}$, the concentration of oxygen interstitials is negligible compared to vacancies, and therefore interstitial contribution to diffusion is also negligible. In addition, thermally generated vacancies are negligible. The analysis of the diffusion data in the previous chapter to yield Eqs. (32) and (33) was based on these two assumptions.

Experimental results on ${\rm CeO}_{2-{\rm x}}$ and ${\rm PuO}_{2-{\rm x}}$ are compared with the present work at different stoichiometries in Table 12. For the present work the result of polycrystalline samples were used. It is seen that both the activation energies and pre-exponential factors are in excellent agreement with those of ${\rm CeO}_{2-{\rm x}}$ and ${\rm PuO}_{2-{\rm x}}$.

The current model is based on random walk theory with no interactions among defects, which is valid only for small nonstoichiometries. Therefore, the disagreement in Table 12 between ${\rm CeO}_{1.80}$ and the remaining values is attributed to the large deviation in the stoichiometry of the former and therefore should be explained in terms of defect clustering or microdomains of ordered regions [12].

5.2 Oxygen Diffusion in Near-Stoichiometric UO_{2±x}

5.2.1 Introduction

As indicated earlier, there have been several studies [1,2,4,7] of uxygen diffusion in stoichiometric UO_2 and they are in reasonably good agreement. The diffusion mechanism, however, is not very well established primarily due to the lack of experimental data on vacancy migration.

Table 12. Comparison of oxygen diffusion in UO_{2-x} with CeO_{2-x} and $\text{PuO}_{2-x}.$

X	CeO _{2-x} [12,13]	Pu0 _{2-x} [15]	uo _{2-x}
0.005 0.01 0.03 0.05 0.08 0.2	l.51x10 ⁻⁵ exp(-11900/RT) 6.16x10 ⁻⁶ exp(-3600/RT)	0.2x10 ⁻⁵ exp(-10900/RT) 0.5x10 ⁻⁵ exp(-11100/RT) 1.3x10 ⁻⁵ exp(-11300/RT) 1.6x10 ⁻⁵ exp(-10800/RT)	0.11x10 ⁻⁵ exp(-11700/RT) 0.22x10 ⁻⁵ exp(-11700/RT) 0.65x10 ⁻⁵ exp(-11700/RT) 1.07x10 ⁻⁵ exp(-11700/RT) 1.68x10 ⁻⁵ exp(-11700/RT)

In their early study Auskern et al. [2] simply assumed the applicability of the interstitialcy mechanism to stoichiometric $\mathrm{U0}_2$ as well as to hyperstoichiometric $\mathrm{U0}_{2+x}$ and calculated the energy to form Frenkel defects to be 70 kcal/mole, which was later supported by the heat capacity measurement of Szwarc [40]. More recently, Breitung [8] and Murch et al. [9] attempted to include both interstitial and vacancy contributions in the diffusion model. However, due to different estimates of the vacancy migration energy, their results were quite different, supporting interstitialcy and vacancy mechanisms, respectively. By theoretical calculation, Catlow et al. [16] obtained 5.8 kcal/mole for the energy of vacancy migration. Also calculated was a Frenkel energy of 11.5 kcal/mole. Assuming a vacancy mechanism in staichiometric $\mathrm{U0}_2$ instead of an interstitialcy mechanism, these two theoretical values yielded a diffusion activation energy of 65.6 kcal/mole, which was in good agreement with the experimental result.

As can be seen, the disagreements arise from lack of reliable data on vacancy migration energy or pled with uncertainty in Frenkel energy.

Since experimental results in $\rm UO_{2-x}$ are now available from the present work, it should be useful to look into this problem in detail.

5.2.2 Diffusion Model

Throughout the work it is assumed that the dominant defect in ${\tt UO}_{2\pm x}$ is the anion Frenke! defect; Schottky defects are neglected. The defect model is based on the following relations:

(1) Defect formation:

$$0_0 + V_1 + V_0 + 0_1$$
 (36)

$$K_{F} = \frac{\theta_{V}}{1 - \theta_{V}} \frac{\theta_{i}}{1 - \theta_{i}} = \exp(\Delta S_{F}/R) \exp(-\Delta H_{F}/RT)$$
 (37)

where $0_n = \text{oxygen ion in regular lattice site}$

 V_i = unoccupied interstitial site

O; = oxygen in interstitial site

V = vacancy in regular lattice site

$$\theta_{v} = \frac{\text{number of anion vacancies}}{\text{number of anion lattice sites}} = \frac{N_{v}}{N_{o}} = \frac{N_{v}}{2N_{U}}$$
 (38)

$$\theta_i = \frac{\text{number of interstitials}}{\text{number of interstitial sites}} = \frac{N_i}{\alpha N_{ii}}$$
 (39)

 $N_v = number of anion vacancies/cc$

 N_{ij} = number of cation sites/cc

 N_0 = number of anion sites/cc

 ΔS_{\sharp} = entropy of Frenkel defect formation

 $\Delta H_{p} = enti$ by of Frenkel defect formation.

The number of available anion interstitial sites of $u0_2$ and $u0_{2-x}$ is equal to number of uranium atoms. In $u0_{2+x}$, however, as the number of interstitials increases, occupiable interstitial sites gradually become selective. α is the parameter that accommodates this effect in $u0_{2+x}$, for which Contamin et al. [3] developed the semi-empirical expression;

$$\alpha = \frac{2}{(2+x)(1+10^3x^3)^{1/2}} \tag{40}$$

For $U0_2$ and $U0_{2-x}$, $\alpha=1$.

(2) Electroneutrality:

$$X = \alpha \theta_1 - 2\theta_{v} \tag{41}$$

Solving Eqs. (37) and (41) for θ_i and θ_v ,

$$e_{i} = \frac{-B + (B^{2} - 4AC)^{1/2}}{2A}$$
 (42)

and

$$\mathbf{e}_{\mathbf{v}} = \frac{1}{2} \left(\alpha \mathbf{e}_{\mathbf{j}} - \mathbf{x} \right) \tag{43}$$

where

$$A = \frac{\alpha}{2} (1 - K_F)$$

$$B = K_F (1 - \frac{\alpha}{2} + \frac{x}{2}) - \frac{x}{2}$$

$$C = -K_F (1 + \frac{x}{2})$$
(44)

Assuming that in $\rm UO_{2\pm x}$ (i) oxygen diffusion can proceed by both vacancy and interstitial migration simultaneously, and (ii) the movements of these two species are independent of each other, the total oxygen diffusion coefficient can be expressed as the sum of the two terms:

$$D = D_{(v)} + D_{(i)}$$
 (45)

In substantially hypostoichiometric UO_{2-x} , $D_{(i)} \approx 0$. Thus, from Eq. (35)

$$D \simeq D_{(v)} = D_0^{v} \Theta_v (1 - \Theta_v) \exp(-\Delta H_v / RT)$$
 (46)

Similarly, in substantially hyperstoichiometric U_{2+x} , $D_{(v)} = 0$, and

$$D = J_{(i)} = D_0^i e_i(1-e_i) \exp(-\Delta H_i/RT)$$
 (47)

From the present experiment, $D_0^V = 4.4 \times 10^{-4}$ and $\Delta H_V = 11.7$ kcal/mole were obtained. D_0^i and ΔH_i were obtained by re-analyzing the existing oxygen diffusion data of $U0_{2+x}$. Only the data of Contamin et al. [3] and Murch [6] were used because their experimental methods were considered to be most accurate.

In substantially hyperstoichiometric UO $_{2+x}$, the thermally generated interstitials are negligible. Thus, from Eq. (43), $\theta_1 \simeq \frac{x}{\alpha}$. In Fig. 44, $\ln[D_{(i)}/\theta_i(1-\theta_i)]$ is plotted vs. 1/T. Least squares fitting yielded $D_0^i = 4.7x10^{-3}$ and $\Delta H_i = 21.8 \pm 13.0$ kcal/mole. Thus,

$$D_{(i)} = 4.7 \times 10^{-3} e_i (1 - e_i) \exp(-21800/RT)$$
 (48)

Combining Eqs. (32), (48), and (45),

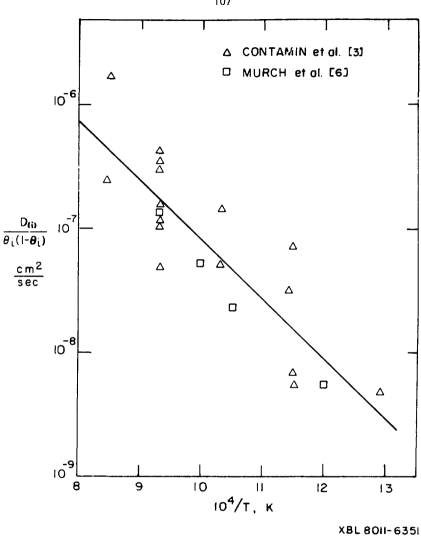


Figure 44. Re-analysis of the diffusion data of UO_{2+x} .

$$0 = 4.4 \times 10^{-4} e_{v} (1 - e_{v}) \exp(-11700/RT) + 4.7 \times 10^{-3} e_{i} (1 - e_{i}) \exp(-21800/RT)$$
(49)

Using the computed code MINUIT, data for stoichiometric UO_2 were fitted to Eq. (49) combined with Eqs. (42) and (43) with $\alpha=1$ and $\alpha=0$ to obtain the Frenkel energy and entropy. Only the data of Marin et al. [1] were used; Data of Auskern et al. [2] and Roberts et al. [7] were excluded because of their unusually large pre-exponential factor which probably resulted from other rate controlling processes affecting the gas-solid isotopic exchange method [1,6]. Also, the stoichiometries of their samples were not as close to 2.0 as those of Marin et al. [1].

Figure 45 shows the data points of Marin et al. [1] and the fitted curve (solid line) for $\Delta H_F=85.6\pm9.2$ kcal/mole and $\Delta S_F=18.2\pm7.2$ e.u. These may be compared with Szwarc's calculation of $\Delta H_F=71.3\pm2.2$ kcal/mole and $\Delta S_F=14.8\pm0.84$ e.u. [40].

Also shown in Fig. 45 are the contribution of interstitials and vacancies, $D_{(i)}$ and $D_{(v)}$. At very low temperatures, vacancies are the primary species that contributes to the total diffusion in UO_2 . In the temperature range of $800-1800^{\circ}\mathrm{C}$, however, neither of the species is completely predominant; and at $1400^{\circ}\mathrm{C}$ the contributions of the two species are approximately equal. Above this temperature $D_{(i)} > D_{(v)}$, and below this temperature $D_{(i)} < D_{(v)}$. The fractional contributions of the two species are depicted in the upper portion of Fig. 45.

Since one of the two terms in Eq. (49) becomes negligible in a substantially nonstoichiometric region, this equation provides a

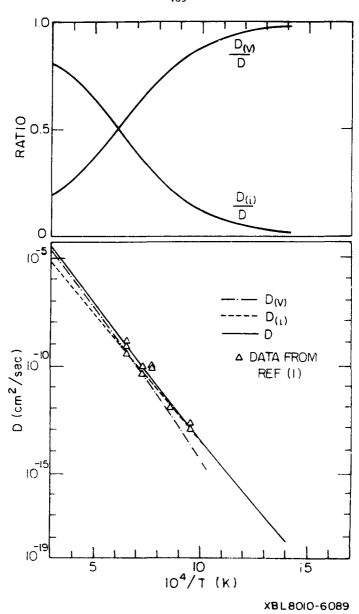


Figure 45. Fitted curve to the diffusion data of stoichiometric UO₂ using optimum values of Frenkel energy and entropy. Absolute and fractional contributions of interstatials and vacancies are also plotted.

unified diffusion model in $\mathrm{UO}_{2\pm x}$ for all stoichiometries. However, it should be emphasized that the current model is valid only when the point defects are independent of each other.

Using Eq. (49), diffusion coefficients in near-stoichiometric ${\rm JO}_{2-x}$ and ${\rm JO}_{2+x}$ were calculated and the results are snown in Fig. 46 and Fig. 47, respectively.

Compared to Fig. 46, Murch et al. [9] predict 2-3 orders of magnitude higher values of D in $\rm MO_{2-x}$, due to the low $\rm \Delta H_v$ and high pre-exponential factor they employed.

5.3 Excess Enthalpy and Frenkel Energy of UO2

It is well known that the specific heat and the enthalpy of $\rm UO_2$ display unusually rapid increase from $1500^\circ K$ to $3100^\circ K$ [41-44], which cannot be explained by lattice vibrations. Szwarc [40] attributed all of this excess enthalpy to Frenkel disorder to deduce the Frenkel energy and entropy. Supporting this interpretation was the oxygen diffusion data in stoichiometric $\rm UO_2$ with the assumption of an interstitialcy mechanism [2]. However, in light of the present work, the diffusion model should be altered, thereby undermining Szwarc's interpretation of the excess enthalpy.

Recently, a series of attempts were made to re-interpret the excess enthalpy in terms of electronic excitation [46-48]. Although quantitative results could not be obtained because the electronic structure of ${\rm JO}_2$ is not well established, it was demonstrated that the electronic contribution to the enthalpy could be significant.

In the previous section, the Frenkel energy of $\Delta H_F = 85.6$ kcal/mole and entropy of $\Delta S_F = 18.2$ e.u. were obtained by fitting

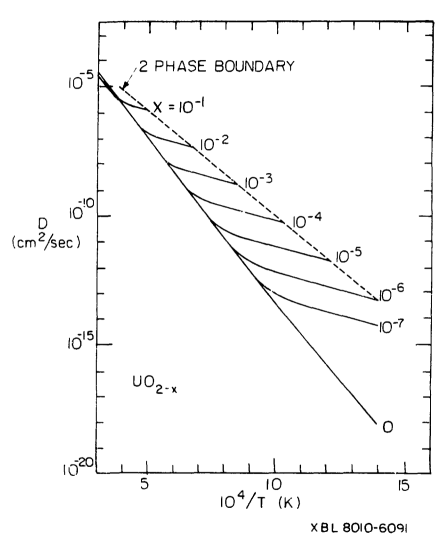
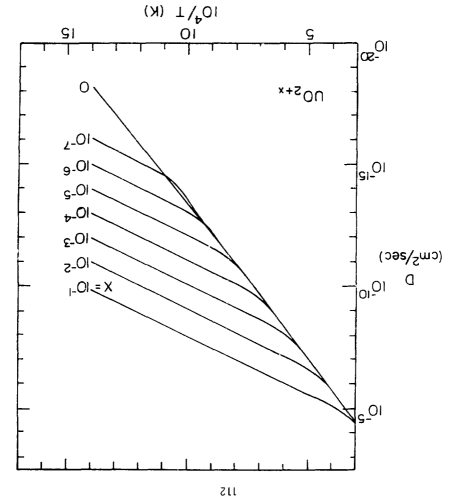


Figure 46. Predicted oxygen diffusion coefficients in $\frac{\text{UO}_{2-x}}{}$



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Figure 47. Predicted oxygen diffusion coefficient in $\text{UO}_{\text{Z+x}}.$

the oxygen diffusion data of stoichiometric UO_2 to the present model. This Frenkel energy is higher than Szwarc's calculation of 71.3 kcal/mole [40]. This means that the ropulation of Frenkel defects is less than predicted by Szwarc's model and therefore their contribution to the excess enthalpy is also smaller.

Her the rest of the excess enthalpy is the contribution. Therefore, the observed enthalpy, $\Delta H_{\rm p}$ of 80_2 can be written as:

$$\Delta H = \Delta H_{ph} + \Delta H_{Fr} + \Delta H_{el}$$
 (50)

where ΔH_{pn} , ΔH_{Fr} , ΔH_{el} are enthalpy due to the lattice vibration, Frenkel disorder, and electronic exciation, respectively.

Collecting data from various studies, Kerrisk et al. [44] developed equations for enthalpy and heat capacity of UO₂ that represent the data. Although they also attributed all of the excess enthalpy to Frenkel disorder, the phonon terms in their equations still represent the low temperature data very well:

$$\Delta H_{DN} = K_1 \Theta \left\{ \left[\exp(\Theta/T) - 1 \right]^{-1} - \left[\exp(\Theta/298) - 1 \right]^{-1} \right\} + K_2 (T^2 - 298^2)$$
 (51)

where
$$K_1 = 19.1450 \text{ cal/mole-}^2 K$$

$$K_2 = 7.84733 \times 10^{-4} \text{ cal/mole-}^2 K$$

$$\Theta = 535.285^2 K .$$

The Frenkel disorder term can be derived as [40]:

$$\Delta H_{Fr} = \sqrt{2} \quad \Delta H_{F} \exp(\frac{\Delta S_{F}}{2R}) \exp(-\frac{\Delta H_{F}}{2RT})$$
 (52)

where $\Delta H_F = 85.6 \text{ kcal/mole}$

$$\Delta S_{F} = 18.2 \text{ e.u.}$$

MacInnes [46] calculated the electronic contribution by introducing two models using two-band structure; valence band and conduction band, with a band gap of $\tilde{\epsilon}_g$. Using standard technique of the semiconductor theory, the energy absorbed by the band structure per unit volume at a given temperature was obtained as:

$$E = \frac{1}{2\pi^2} \left(\frac{2m^*}{h^2} \right)^{3/2} \left(\frac{N_d}{n_0} \right)^{1/2} \bar{L} \left(\frac{3}{2} \right) (kT)^{5/2} +$$

$$E_g \left(\frac{1}{2} \right) ! (kT)^{3/2} \right] \exp(-E_g/2kT)$$
(53)

 N_d = number of electrons per unit volume in the valence band $n_0 = 2(2\pi m \star \kappa T/n^2)^{3/2}$

h = Planck's constant

 $h = h/2\pi$

m* = electronic effective mass

 $m_e = electron mass.$

Following Catlow [45], MacInnes assumed that the valence band was comprised of 4f orbitals, and accordingly assigned 14 electrons to it. However, as Thorn et al. [47] indicated, to use 5f orbitals as the valence pand may be more plausible than 4f. Thorn et al. [47] found

that two electrons could be assigned to the 5f orbitals. Using 5f orbitals for valence band in Eq. (53), $\Delta H_{el}(cal/mole)$ can be written as:

$$\Delta H_{el} = \left(\frac{m^*}{m_e}\right)^{3/4} \left[1.789 \times 10^{-3} T^{7/4} + 6.003 \times 10^{-4} \Delta E \cdot T^{3/4}\right] \exp(-\Delta E/2RT)$$
 (54)

ΔE is the electronic activation energy in cal/mole.

The reported enthalpy data [42,43] were fitted to Eq. (50) using Eqs. (51), (52), and (54) to obtain the optimum values of (m^*/m_e) and ΔE . In Fig. 48, the fitted ΔH curve and the data points are shown. Also shown are ΔH_{ph} , and the excess enthalpy ΔH_{ex} (= ΔH - ΔH_{ph}). Fitting yielded (m^*/m_e) = 7.6±0.1 and ΔE = 45.0±0.9 kcal/mole, respectively. The ΔE value obtained is equivalent to a band gap of 2.0 ev, which is in good agreement with the reported values: 1.8~2.3 ev obtained from electrical conductivity data Letween 1400°K and 3000°K [52] and 2.1 ev obtained from reflectivity data [53].

Electronic contribution ΔH_{el} and Frenkel contribution ΔH_{Fr} are calculated seperately and shown in Fig. 49. Also snown on the same temperature scale are the fractional contributions of the two effects; $\Delta H_{el}/\Delta H_{ex}$ and $\Delta H_{Fr}/\Delta H_{ex}$. As temperature increases the fraction of electronic excitation decreases; at 3000°K it accounts for approximately 13 percent of the total excess enthalpy.



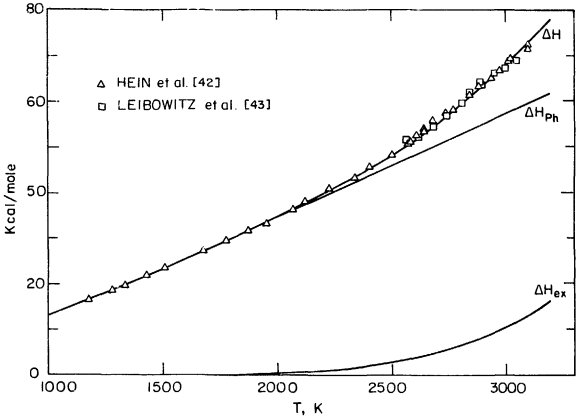


Figure 48. Fitted curve to the enthalpy data of UO, between 1000⁰K and 3000⁰K. Lattice vibrational contribution and excess enthalpy are also plotted.

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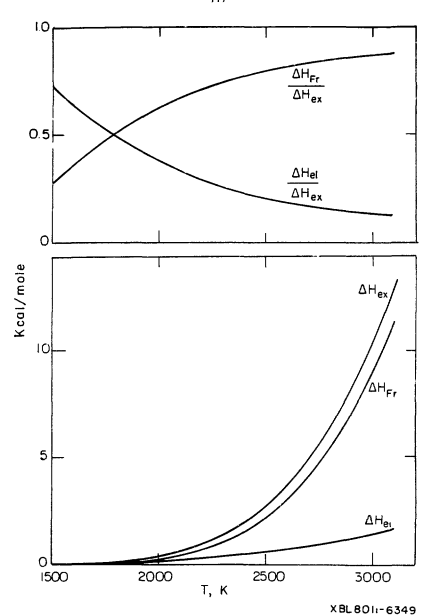


Figure 49. Absolute and fractional contributions of Frenkel disorder and electronic excitation to the excess enthalpy.

CONCLUSIONS

- 1. Much faster oxygen diffusion in UO_{2-x} than in UO_2 was observed, which proves that the primary defect in UO_{2-x} is the anion vacancy.
- 2. The measured migration activation energy of the anion vacancy, $11.7 \, \text{kcal/mole}$, is lower than the migration energy of interstitials. However, it is not as low as predicted by theoretical calculation [16]. The activation energy and the pre-exponential factor of the oxygen diffusion coefficient are in good agreement with those of other materials of fluorite structure, e.g., CeO_{2-x} , PuO_{2-x} . This confirms the similar oxygen migration mechanisms in these matertials.
- 3. In stoichiometric UO_2 and near-stoichiometric $UO_{2\pm x}$, both vacancies and interstitials contribute significantly to oxygen diffusion. At 1400° C, contributions of the two species are approximately equal in stoichiometric UO_2 . When T > 1400° C, the interstitial contribution is higher; when T < 1400° C, the vacancy contribution is higher.
- 4. The Frenkel energy and entropy deduced from measured diffusivities in 90_{2-x} , 90_{2} and 90_{2+x} are $$\Delta H_F = 85.6$ kcal/mole and $$\Delta S_F = 18.2$ e.u. These values yield lower anion Frenkel defect concentration than predicted by Szwarc's model [40]. This deviation is consistent with the theory that not all of the excess enthalpy of 90_{2} can be attributed solely to the Frenkel disorder. Use of the 90_{2} can be attributed solely to the Frenkel disorder. That at 900_{2} k electronic excitation can account for 13 percent of the excess enthalpy.

5. McInnes' [46] two-band model for electronic excitation is modified to quantify the electronic enthalpy. Data fitting yields a band gap of 2.0 ev, which is in good agreement with the reported values [52,53], and effective mass of conduction band electrons of 7.6 m $_{\rm e}$.

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I wish to dedicate this to my parents.

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