THE DIFFUSION OF KRYPTON-85
FROM URANIUM DIOXIDE POWDER

A. B. Auskern

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The diffusion of Kr$^{85}$ in two UO$_2$ powders from different sources has been studied by performing a series of post-irradiation anneals on the powders. The emanation data were analyzed in a manner that considered the effect of sintering as well as the effect of a distribution of particle sizes within the sample. Measurements were made in the temperature range of 900 to 1500°C. The time at a temperature was between 8 and 24 hours. The diffusion coefficients for Kr$^{85}$ in the two powders can be represented by the equations: 

\[ D = 2.65 \times 10^{-4} \exp(-65,500/RT) \]  

for UO$_2$ prepared from crushed UO$_2$ pellets and, for a chemically prepared UO$_2$ powder, 

\[ D = 4.9 \times 10^{-4} \exp(-73,800/RT) \]

THE DIFFUSION OF KRYPTON-85 FROM URANIUM DIOXIDE POWDER

A. B. Auskern

I. INTRODUCTION

The present investigation forms part of a larger program concerned with the release of fission gases from uranium dioxide. In this study the release of krypton-85 from powdered UO$_2$ was considered. Experiments were performed to determine the kinetics and temperature dependence of the release process.

Extensive discussions of the work done in this area are presented in reviews by Lustman (Ref 1) and by Eichenberg, et al. (Ref 2). Evidence indicates that the release of fission gases from UO$_2$ is one of diffusion through the lattice. The measurements of Booth and Rymer (Ref 3) on powders prepared from fused UO$_2$ give some support for this mechanism. They showed that the release of xenon-133, in agreement with an appropriate solution to the diffusion equation and within the limitations of the experiment, varied directly as the reciprocal of the particle radius, and directly as the square root of the annealing time. For this powder, the diffusion coefficient variation for xenon-133 from 1000 to 1500°C is represented by the equation \[ D = 1.5 \times 10^{-8} \exp(-46,000/RT) \]. Booth and Rymer also reported that there was no significant difference between xenon and krypton diffusion rates.

Post-irradiation diffusion anneals on dense UO$_2$ sintered pellets have been performed (Ref 4) in the temperature range of 1000 to 1575°C. Diffusion coefficients varied from about 3.5 \times 10^{-20} at 1000°C to 1 \times 10^{-16} cm$^2$/sec at 1575°C. The activation energy calculated from the results is 58.5 kcal/mole. For this work the absolute value for the diffusion coefficient is in some doubt because of the difficulty in assigning an average spherical "radius" to a sintered pellet.

A series of experiments has been performed to measure the solubility and diffusion of helium in UO$_2$ powder (Refs 2 and 4). The helium diffusion data fit the spherical diffusion model very well. Diffusion coefficients ranging from 1 \times 10^{-17} cm$^2$/sec at 700°C to 2 \times 10^{-15} cm$^2$/sec at 1000°C were calculated. The activation energy is 46 kcal/mole, the same as that found by Booth and Rymer (Ref 3) for xenon. Diffusion of the helium is, however, at least an order of magnitude faster than the xenon diffusion.
Recent work by Lindner and Matzke (Refs 5 and 6) on xenon diffusion from uranium oxide powder has been published. They report in their initial study, with UO\(_{2} \cdot 0\) and UO\(_{2} . 67\) powders in the temperature range of 500 to 1100°C for relatively short periods of time, an activation energy of about 34 kcal for UO\(_{2}\) and somewhat less for UO\(_{2} . 67\). Diffusion in UO\(_{2} . 67\), however, is about two orders of magnitude faster than in UO\(_{2}\). In a second study which concerned the effect of O/U ratio on the diffusion of xenon, the activation energy and diffusion coefficients found with UO\(_{2} . 00\) were very similar to those of Booth and Rymer (Ref 3). About an order of magnitude increase in D occurred when the oxygen content was raised from UO\(_{2} . 00\) to UO\(_{2} . 02\), and from UO\(_{2} . 02\) to UO\(_{2} . 12\).

The Booth and Rymer work is most similar to the present investigation. Some data from Booth and Rymer will be discussed in detail for their general applicability. For the calculation of diffusion coefficients in the fused UO\(_{2}\) powder, a particle radius of 28 microns was used. This is the average radius given by the screen fraction used, which was -200 to +325 mesh. However, the assumption that an average screen radius represents the true particle radius can be misleading. It has been shown that the spherical radius when calculated from surface area and density measurements is smaller than that given by sieve sizes. For example, for a powder prepared from fused UO\(_{2}\) (liquid density, 10.95 g/cc) and screened between 80 and 120 mesh screens, the surface area and density measurements resulted in an average spherical radius of 15 microns. However, the average radius for the screen fraction is 7.5 microns. Since such a difference may be present in the powders studied by Booth and Rymer, then their diffusion coefficients can be too large.

The Booth and Rymer report contains much information on the amount of gas released from sintered compacts. For a given compact type, the fraction of xenon released varies inversely as the compact density. However, for compacts of the same density, but from different sources, there are large differences in the amount of gas released. What is surprising about these data is the number of samples which gave release rates much greater than that for the fused powder. For the fused powder, for example, 2.1% xenon diffused out at 1400°C in three hours. For some low density pellets (85% theoretical), the fraction that diffused out under identical experimental conditions was much higher—ranging from 15 to 35%. If, as it is assumed, the rate controlling step is diffusion through the lattice and the diffusion coefficient of xenon through the UO\(_{2}\) lattice is independent of the powder source or treatment, then, in order for the gas release to be greater in the compact, the diffusion path length in the compact must be smaller than that in the fused powder. This further implies that the surface areas of these compacts are greater than that of the fused -200 to +325 mesh UO\(_{2}\) powder. Of course, the rate controlling step for diffusion from sintered compacts containing some open porosity may not be pure lattice diffusion. If this is the case, then the greater release rate for some compacts may be reasonable if the data are analyzed with some other appropriate model. Unfortunately, Booth and Rymer report no surface area measurements which might help clear up some of the apparent discrepancies between the powder and compact data.

The absolute magnitude of diffusion coefficients is very sensitive to the particle radius used. For powders like UO\(_{2}\) which can be porous, highly irregular in shape, and which usually consist of groups of fine crystallites, the use of microscopic measurements or sieve screen estimates for particle radii is unrealistic because much of the available surface may not be directly visible. Flugge and Zimens (Ref 7), in their paper on the Hahn emanation method, consider the surface as measured by gas adsorption to be the best estimate of an emanating surface.

With fine powders undergoing anneals at high temperatures, the possibility that sintering can occur and reduce the powder surface area must be considered. The occurrence of sintering during

* A. B. Auskern, unpublished data, Bettis Laboratory.
an anneal makes the effective spherical radius of the powder larger and results in the emanation rate being reduced. Either of two things is essential then to make emanation data meaningful: the powder must be prepared in such a way that it will undergo no surface area change during an anneal or, if this is difficult, account must be taken of the surface area change in order to correctly evaluate the diffusion coefficients.

Neglecting sintering results in calculated diffusion coefficients that are too small. Also, if sintering occurs over a range of temperatures, the constantly changing particle radius indicates that the activation energy for the process would be higher than that found if the effect of sintering were neglected.

A further consideration is that the sample does not consist of a single size particle, but varies over a range of sizes. During an anneal the fine particles become depleted more rapidly than the coarse, and the emanation kinetics deviate from the ideal case where a single uniform particle size is assumed.

In this study two powders from sources other than fused UO$_2$ were investigated. The analysis of the emanation rate data included the effect of sintering and depletion of fine particles.

II. EXPERIMENTAL

A. Powders Investigated

Two uranium dioxide powders were used in this study. The powder designated NUR was prepared by the conversion of uranyl nitrate hexahydrate to UO$_3$, and then the UO$_3$ to the dioxide by heating in hydrogen at 800°C. The UO$_2$ formed was annealed in hydrogen at 1650°C for 21 hours, and then ball milled for 16 hours in a rubber lined mill using uranium balls. The surface area of this powder, as determined by gas adsorption, is 1.3 m$^2$/g and the liquid density, 10.71 g/cc. Chemical analysis showed the material to be UO$_2$ and the impurity level, estimated by spectrographic analysis, to be 0.02%.

The second powder studied was prepared by crushing a number of sintered UO$_2$ pellets (92% theoretical density) and collecting the -200 to +325 mesh screen fraction. The surface area of this powder is 0.1 m$^2$/g and the density is 10.67 g/cc. The material was analyzed as UO$_2$. Spectrographic analysis gave an impurity content of about 0.05%, with the principal impurity being iron.

The average spherical radii of the powders were calculated from surface area and density measurements. The calculation for the spherical radius involves the assumptions that the particles are perfect spheres and are of uniform size. The relationship

$$a = \frac{3}{s \, d}$$

is used, where "a" is the average particle radius in cm, s is the surface area (cm$^2$/g), and d is the density (g/cc). For the NUR powder, $a = 0.22$ microns; for the crushed UO$_2$, $a = 2.8$ microns.

B. Irradiation of Samples

Irradiations of the samples were carried out at the Brookhaven National Laboratory reactor. Reactor hole N-5 was used for all irradiations. This is an air cooled facility (ambient temperature about 50°C) designed to permit sample insertion and withdrawal while the reactor is in full operation.

The uranium dioxide samples, about three grams in weight, were contained in quartz vials under a helium atmosphere. The samples were irradiated to an nvt of about 4 x 10$^{16}$ thermal neutrons/cm$^2$. This resulted in an approximate burnup of 0.004% of U$^{235}$ atoms present, and produced about 3 x 10$^{12}$ Kr$^{85}$ atoms in the three gram sample.

The amount of Kr$^{85}$ actually produced in the uranium dioxide during irradiation was determined by total dissolution of a sample in concentrated nitric acid and collection of the released fission gases.
In all cases the quantity of Kr\textsuperscript{85} measured was less than the amount calculated from the conditions of the irradiation. For the NUR powder the amount of Kr\textsuperscript{85} determined was about 65\% of that calculated; for the crushed UO\textsubscript{2}, 52\% of the amount calculated was found. These differences may be the result of errors in neutron flux estimates and the recoil of fission fragments from the UO\textsubscript{2} powder during irradiation.

A diagram of the dissolving and gas collection system is shown in Fig. 1. In operation, a flow of helium gas is established through the complete system, including the activated charcoal traps. The crucible containing the UO\textsubscript{2} sample is upset with a magnetic stirring rod and the rapid UO\textsubscript{2}-HNO\textsubscript{3} reaction begins. The acid is frequently agitated and then boiled gently for about 15 to 20 minutes to facilitate complete removal of the gases. The ferrous sulfate solution in the train is used to remove nitric oxide and nitrogen dioxide from the gas stream. Finally, the charcoal traps are isolated, evacuated, and sealed off directly above the charcoal. In no case was any Kr\textsuperscript{85} detected in the second charcoal trap.

C. Diffusion Anneals

A drawing of the annealing system is shown in Fig. 2. Emanation data over the complete temperature range are obtained without changing the sample. With the system evacuated, the sample is suspended above the hot zone and the furnace brought to the lowest annealing temperature. The system is then isolated from the vacuum station and the sample is lowered into the hot zone. After heating for the desired length of time, the sample is raised from the hot zone and the gases evolved during the anneal are collected on activated charcoal cooled with liquid nitrogen. Usually, for anneals up to 1100°C, there is enough degassing of the furnace and UO\textsubscript{2} powder to raise the system pressure to around 500 microns. By following the decrease in the pressure during gas collection, the point at which essentially all the gas is collected is easily determined. At this point the sampling bulb is sealed off. The sample is then lowered for the next anneal at the same temperature.

With the sample raised out of the hot zone, the furnace temperature is raised to that required for the next anneal. Anneals at 100°C increments were made up to 1400°C for the NUR powder and up to 1500°C for the crushed UO\textsubscript{2} powder. The total time at a temperature was between 8 and 24 hours; three or four gas samples were collected during this time.

The furnace was controlled with a Bristol-Pyrovane temperature controller to within ±5°C, and temperatures were read with a Pt-Pt, 10 Rh thermocouple located outside the combustion tube at the sample level.

Separate anneals were made on unirradiated samples of the two powders to estimate the amount of sintering that could occur during an emanation anneal. The anneals were performed in hydrogen at 1400°C for 22 hours. The results of surface area measurements showed a decrease, by a factor
of 5, for both powders. Thus, the spherical radii of the powders can be estimated to have increased by roughly the same factor (density changes have been neglected). The sintered radius of the crushed UO$_2$ powder is then about 14 microns, and that of the NUR powder, about 1 micron.

D. Analytical Procedure

The bulbs containing the collected gases were analyzed for Kr$^{85}$ content by gamma ray spectroscopy methods. A spectrum profile in the vicinity of 0.52 Mev was obtained with a 100-channel analyzer employing a thallium-activated KI well-type crystal. A peak of this energy is characteristic of the decay scheme involving Kr$^{85}$. The sample activity at this energy may be related, with consideration for the counter efficiency (approximately 9%) and the branching ratio ($6.5 \times 10^{-3}$), to the number of Kr$^{85}$ atoms present. The experiments were completed within a few months after the sample was irradiated. Since krypton-85 has a 10.3 year half-life, no corrections for decay were made.

The counting for the total dissolution experiments was handled in the same manner, except that the "U" tubes containing the fission gases were counted using a flat 3-in. diam crystal.

III. RESULTS

The data for the complete heat treatment of a powder can be plotted as the cumulative fractional release vs the cumulative time. Discontinuities in the curve result when the temperature is raised. These plots for the two uranium dioxide powders are shown in Fig. 3.

For the calculation of diffusion coefficients, the model assumed is that of diffusion from a group of uniform spheres at some initial Kr$^{85}$ concentration, with the surface maintained at zero concentration. A solution to the diffusion equation (Ref 4) for this case is

$$F = 1 - \frac{6}{\pi^2} \sum_{n=1}^{\infty} \frac{1}{n^2} e^{-n^2 \pi^2 \frac{D}{a^2} t}.$$
where

\[ F = \frac{6}{v \sqrt{\pi}} + \frac{3}{\nu^2}, \]

where \( \nu^2 = \frac{(\alpha^2)}{D \tau} \), may be used.

In calculating diffusion coefficients at different temperatures, account must be taken of the experimental condition that the zero time for the anneal is not known for all temperatures after the first. Each isothermal portion of the cumulative fractional release vs cumulative time curve represents only the measured segment of a hypothetical continuous anneal which started at some undefined zero time. Thus, the equations just listed are directly applicable only to the first anneal, since this is the only anneal to which the boundary conditions of the diffusion equation solution directly apply. However, diffusion coefficients at temperatures above the first are readily determined.

From a master plot of \( \frac{D \tau}{\alpha^2} \) vs \( F \), values of \( \frac{D \tau}{\alpha^2} \) for the fractional release obtained during an anneal are read. These values are plotted against time. Since the zero time is not known, only the time differences between samples at a particular temperature can be plotted, with the initial point (equivalent to the last point at the previous lower temperature) set at an arbitrary position on the time scale. The relationship between \( \frac{D \tau}{\alpha^2} \) and \( t \) is linear, and the slope of the line through the experimental points is \( \frac{D}{\alpha^2} \). From a knowledge of \( \alpha \), \( D \) is calculated directly. Extrapolating the curve to \( \frac{D \tau}{\alpha^2} = 0 \) fixes a theoretical zero time. The significance of the theoretical zero time is that it is the amount of time at temperature the sample would take to reach the concentration distribution defined by the first experimental point, assuming that the concentration in the sample was uniform at \( t = 0 \).

The effects of a changing particle radius, as well as depletion of \( \text{Kr}^{85} \) in the fine particles, must be evaluated. The occurrence of either or both conditions changes the shape of the emanation curve.

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*J. M. Markowitz, personal communication, Bettis Laboratory.*
Consider first that sintering is occurring. During the anneal the spherical radius increases from its initial value to some final value at the end of the anneal. Thus, the rate of emanation of the gas does not follow a specific diffusion curve. At the start of the anneal, emanation follows the curve calculated with the initial small radius. However, as sintering progresses and the particles become larger, the emanation rate slows down and leaves the initial rate curve, passing through a family of rate curves (each corresponding to a larger particle size). For long enough times, the emanation rate curve approaches the curve given by the final particle size. Thus, a true diffusion curve does not result. The emanation data lie between the curves defined by the initial and final particle sizes.

When plotting the data as $Dt/a^2$ vs $t$ to calculate diffusion coefficients, a similar type of curve is observed. Had no sintering taken place, the $Dt/a^2$ vs $t$ curve would be a straight line of slope $D/a^2$. With the occurrence of sintering, the experimental curve moves from the initial slope, crosses a family of lines, each corresponding to a larger "a" value (and smaller slope), and for long enough times, parallels the line, the slope of which is given by the final sintered radius. Thus, the experimental $Dt/a^2$ vs $t$ curve is a curved line lying between the lines defined by the initial and final particle radius. The calculation of the diffusion coefficient must then be made from the initial slope of the $Dt/a^2$ vs $t$ curve since, for this region at least, a reasonable value for the particle radius is known.

An analogous set of curves results when emanation from a range of particle sizes is considered. Initially, the emanation rate is rapid because of diffusion from the fine particles. As the amount of gas in these particles becomes depleted, the rate slows down and becomes controlled by emanation from the larger particles. Thus, the resulting $Dt/a^2$ vs $t$ plot is also not linear. It is a curved line similar to that resulting from sintering. The initial slope is steep, corresponding to emanation from the finer fractions. As these particles become depleted, the slope becomes smaller and smaller and approaches finally the slope given by the limiting emanating particles.

The situation is complex when sintering and depletion are occurring together. Detailed knowledge of the rate of sintering is required to estimate the combined effects, as well as information on the initial size distribution and the manner in which the distribution changes as a function of time and temperature.

The shape of an experimental emanation curve is usually modified by both sintering and particle depletion. As in the case of sintering, the best estimate for the diffusion coefficient is given by the initial slope of the $Dt/a^2$ vs $t$ curve and the initial particle radius. If the particle size distribution of the sample contains many very fine particles and these control the initial rate of emanation, then the diffusion coefficient calculated using the average particle radius will be too large. It would be difficult to correct this without knowing the particle size distribution.

Since the emanation data in the experiments performed here were obtained from one sample for each powder for the whole temperature range, the value of the particle radius must be constantly revised at each temperature to account for particle sintering and depletion.

Both the diffusion coefficient and the new number for the particle radius are determined from the experimental $Dt/a^2$ vs $t$ curve. Starting with the lowest annealing temperature, the first diffusion coefficient is calculated from the initial slope and the known initial particle radius. With this diffusion coefficient and the slope observed at the end of the anneal, a value for "a" can be calculated. This "a" value is not necessarily the average particle radius at the end of the anneal; it is a fictitious radius which refers only to the condition of the material that is emanating at the end of the anneal and includes the effects of sintering as well as depletion of the fine particles. This radius is then used for the calculation of the diffusion coefficient for the next highest temperature anneal. The calculation is repeated again and another "a" value is determined from the slope at the end of the higher temperature anneal. The data from the anneals of the two uranium dioxide powders have been treated
TABLE I

DIFFUSION COEFFICIENTS OF Kr$^{85}$ IN UO$_2$

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Crushed UO$_2$</th>
<th>NUR UO$_2$</th>
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<tbody>
<tr>
<td>900</td>
<td>6.7 x 10^{-17}</td>
<td>3.9 x 10^{-18}</td>
</tr>
<tr>
<td>1000</td>
<td>1.3 x 10^{-15}</td>
<td>5.3 x 10^{-17}</td>
</tr>
<tr>
<td>1100</td>
<td>8.3 x 10^{-15}</td>
<td>7.6 x 10^{-17}</td>
</tr>
<tr>
<td>1200</td>
<td>6.6 x 10^{-14}</td>
<td>5.3 x 10^{-15}</td>
</tr>
<tr>
<td>1300</td>
<td>2.2 x 10^{-13}</td>
<td>1.8 x 10^{-14}</td>
</tr>
<tr>
<td>1400</td>
<td>5.9 x 10^{-13}</td>
<td>1.0 x 10^{-13}</td>
</tr>
<tr>
<td>1500</td>
<td>2.2 x 10^{-12}</td>
<td>--</td>
</tr>
</tbody>
</table>

in this manner. Diffusion coefficients have been calculated at the various temperatures and are listed in Table I.

At temperatures above 1200°C for the crushed UO$_2$, the Dt/a$^2$ vs t curves were linear for the duration of the anneals. For these higher temperatures no further adjustment of "a" was required. For the NUR powder a continuous adjustment of "a" was necessary, although at the higher temperatures the amount of change became less and less.

The variation of log D with 1/T is shown in Fig. 4. The variation is quite linear over the whole temperature range investigated, with no breaks noted in the curve. The diffusion coefficients for the crushed UO$_2$ powder are about an order of magnitude higher than those for the NUR powder. The slopes of the curves correspond to an activation energy of 65.5 kcal/mole for the crushed powder, and 73.8 kcal/mole for the NUR powder. The variation in D with temperature can be represented by the equations $D = 2.65 \times 10^{-4} \exp(-65.500/RT)$ for crushed UO$_2$, and $D = 4.9 \times 10^{-4} \exp(-73.800/RT)$ for the NUR powder.

IV. DISCUSSION

A comparison should be made between the "a" values of the powders after the sintering anneals and the final "a" values that were calculated from the analysis. These are given in the following table.

<table>
<thead>
<tr>
<th>Powder</th>
<th>Initial Radius</th>
<th>After Sintering</th>
<th>Final Emanation Radius</th>
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<tr>
<td>NUR</td>
<td>0.22 microns</td>
<td>1 micron</td>
<td>7 microns*</td>
</tr>
<tr>
<td>Crushed</td>
<td>2.8</td>
<td>14</td>
<td>43**</td>
</tr>
</tbody>
</table>

*This radius is still changing at 1400°C.
**No further change after 1200°C.

These numbers should not be the same. The radius calculated from the data is based on an emanation curve which is weighted (because of depletion of the fine particles) toward the coarser particle fraction. The radius after the sintering anneals is determined by a gas adsorption method and includes all the particles, fines as well as coarse. Thus, this radius should be smaller than the emanation radius and this is observed. For the crushed UO$_2$, the sintered radius is 14 microns and
KRYPTON-85 DIFFUSION IN UO₂ POWDER

Fig. 4 Temperature Dependence of the Diffusion Coefficient

The calculated emanation radius is 7 microns. The experimental results are therefore consistent with a diffusion model which includes both sintering and a distribution of particle sizes within the sample. However, the data do not permit the emanation curves to be analyzed into their sintering and depletion components. No data are available on the rate of sintering of the powders at each temperature studied and, as mentioned previously, the initial and final distribution of particle sizes is not known.

Ideally, diffusion experiments of this type should be performed on well sintered powders of uniform particle size. However, if data are desired at low temperatures (900°C) on low burnup materials, a small particle size powder must be used. For example, the anneal at 900°C of the crushed UO₂ powder resulted in 1% gas release after about 5 hours. The particle radius of this material was 2.8 microns. If the material had been sintered before annealing to 14 microns (the radius found after the supplementary 1400°C anneal), the amount of gas released would have been about 0.04%. Because of the short neutron exposure the samples received, a quantity of Kr₈⁵ corresponding to 0.04% of the total could not have been readily detected.

The results found here for Kr₈⁵ diffusion are among the highest that have been reported for rare gases in UO₂. The diffusion coefficients are significantly higher than those reported for UO₂ compacts (Ref 4), and the ones given by Booth and Rymer (Ref 3), and by Lindner and Matzke (Refs 5 and 6) for xenon in UO₂. At 900 and 1000°C, helium diffusion coefficients are greater than those found for Kr₈⁵, but the activation energy is lower (Ref 4).

The results of thoron diffusion experiments in UO₂, using the Hahn emanation technique, give estimated diffusion coefficients and activation energies in the same range as those found here. In the work of Anderson, et al. (Ref 8), an activation energy of 81 kcal/mole was found above 1150°C. The process responsible for this activation energy is considered to be bulk diffusion through the lattice. A diffusion coefficient estimated from the emanation data is about 10⁻¹⁴ cm²/sec at 1200°C. Recent work with the Hahn emanation method on UO₂ compacts gives activation energies between 60 and 85 kcal/mole above 1100°C. * Estimated diffusion coefficients are around 10⁻¹⁴ cm²/sec at 1400°C.

In the temperature range investigated, diffusion rates for the crushed UO₂ are about an order of magnitude greater than those of the NUR powder. Since no provisions were made for the maintenance of a reducing atmosphere during the anneals, the possibility exists that some oxidation can take place and result in enhanced diffusion coefficients for the crushed UO₂ powder. Lindner and Matzke (Ref 6) have shown that very slight amounts of excess oxygen significantly increase the xenon diffusion.

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*J. C. Clayton, personal communication, Bettis Laboratory.
rate. A similar effect was also observed for oxygen diffusion in \( \text{UO}_2 \), especially for nearly stoichiometric compositions.

Further work is needed before the mechanism of fission gas release can be precisely described. For example, the effect of burnup has not been evaluated. In these experiments the powders received a short reactor exposure compared to the exposure for some pellet work. Also, in the work performed here and in the referenced works, the annealing times were relatively short, i.e., less than 24 hours. Bostrom** has demonstrated that, for dense \( \text{UO}_2 \) plates, the emanation rate for xenon for the first 10 to 30 hours usually corresponds to a diffusion coefficient greater than that observed for the final 30 to about 120 hours. This may be associated with a "burst" release that has been observed on initially heating sintered \( \text{UO}_2 \) materials. For the emanation curves in the experiments reported here, a model of pure diffusion was assumed, complicated by particle sintering and depletion. However, the possibility that "bursts" occurred cannot be excluded, especially from the data for the low temperature anneals.

The question of a low-temperature, low activation energy release process for fission gas in \( \text{UO}_2 \) has not been resolved. It was not observed in this study nor in the work of Booth and Rymer (Ref 3) or Lindner and Matzke (Ref 6). However, work with the Hahn emanation method and with gas release from sintered pellets indicates that a different diffusion mechanism may be operative at lower temperatures.

REFERENCES

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