Surface and Corrosion Chemistry of PLUTONIUM

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Elemental plutonium, the form in which most of the weapons-grade material exists, is a reactive metal. When exposed to air, moisture, and common elements such as oxygen and hydrogen, the metal surface readily corrodes and forms a powder of small plutonium-containing particles. Being easily airborne and inhaled, these particles pose a much greater risk of dispersal during an accident than the original metal. The present emphasis on enhancing nuclear security through safe maintenance of the nuclear stockpile and safe recovery, handling, and storage of surplus plutonium makes it more imperative than ever that the corrosion of plutonium be understood in all its manifestations.

Metallic plutonium was first prepared at Los Alamos in 1944, during the Manhattan Project. After samples became available, scientists studied the properties of the metal, including its reactions with air, moisture, oxygen, and hydrogen. Extensive investigation of plutonium continued into the early 1970s, and the results of those studies are documented in the handbooks of chemistry, physics, and metallurgy.

In dry air—less than 0.5 part per million (ppm) of water—at room temperature, plutonium behaves much like other active metals, forming a protective layer of dioxide (PuO₂) on the surface. The PuO₂ layer limits corrosion of unalloyed plutonium to the almost imperceptible rate of 20 picometers per hour. The corrosion chemistry of plutonium is therefore assumed to be relatively simple and well understood, but that assumption is far from true. Unexplained phenomena, such as plutonium pyrophoricity (spontaneous ignition in air) and moisture-accelerated corrosion, have been observed in the earliest plutonium studies (Cleveland 1979, Wick 1980, Katz et al. 1986). Recent Los Alamos studies confirm and add to these examples of unexpectedly rapid corrosion. Moreover, our analysis suggests that J. T. Weber, a Los Alamos pioneer in plutonium corrosion, was prescient when he wrote, “Most investigators are inclined to concentrate their attention on PuO₂, which can be well characterized, and to ignore other compounds that may contribute to the overall corrosion behavior” (Wick 1980). Indeed, we believe that plutonium oxides other than PuO₂ and compounds other than oxides play a potent role in plutonium corrosion. These compounds appear to be catalysts, causing anomalous corrosion reactions to proceed at enormous rates.

“Runaway” reactions can occur under fairly routine conditions. In one case, a failed storage package containing a plutonium casting was examined in a glove box, which had a nitrogen-rich atmosphere (less than 3 percent oxygen) normally used for handling plutonium metal. The package remained there for 3 hours after disassembly and initial inspection. When the workers returned to continue their evaluation, they found that the inner container was hot to the touch and its diameter had increased by 50 percent in the region surrounding the casting (Haschke and Martz 1998b). Our subsequent investigation showed that a corrosion reaction involving both oxygen and nitrogen initiated spontaneously at room temperature and advanced into the plutonium metal at a rate of more than 1 centimeter per hour (cm/h), or a factor of 10¹⁰ faster than the corrosion rate in dry air. The reaction generated excessive temperatures and started under conditions that are considered safe for the routine handling of plutonium.

Further studies suggest that the course of corrosion depends very heavily on the chemical condition of the plutonium surface. A surface layer of sesquioxide (Pu₂O₃), which forms in the absence of oxygen, promotes corrosion of the metal by hydrogen. Conversely, a surface layer of hydride (PuHₓ, where 1.9 < x < 3) increases the plutonium oxidation rate in oxygen by a factor of 10¹³ to a value near 3 meters per hour (m/h). Finally, a surface layer of the previously unknown higher oxide PuO₂⁺ₓ, which forms on the PuO₂ layer in the presence of moisture, apparently enhances the bulk corrosion of plutonium metal in moist air. The fact that PuO₂ reacts with water demonstrates that it is thermodynamically unstable in air and environmental media. Evidently, its reaction with water is responsible for both the implosion and the pressurization of sealed containers of plutonium oxide during their extended storage.
In this article, we present our observations and analysis of the chemistry and kinetic behavior of important corrosion reactions involving oxygen, nitrogen, and hydrogen. We observe that PuHₓ, Pu₂O₃, and PuO₂ₓₓ on the metal surface catalyze anomalous corrosion reactions and that the Pu(III) and Pu(VI) oxidation states are important in addition to the predominant Pu(IV) state. We also outline the kinetics that relates specific surface compounds and conditions to anomalous reactions. Our analysis provides explanations for many puzzling phenomena, but it also leaves a multitude of unanswered questions. We conclude that the present understanding of plutonium chemistry is inadequate and that the new evidence presents an immediate technical challenge to the scientific community.

**Plutonium Oxides and Atmospheric Oxidation**

Plutonium corrosion and oxidation are often treated as equivalent topics because plutonium oxides are the only products normally observed during atmospheric corrosion. The metal does not react appreciably with elemental nitrogen even at elevated temperatures, although plutonium mononitride (PuN) is a stable compound. Therefore, we begin with a review of oxide chemistry and oxidation kinetics.

**Diffusion-Controlled Oxidation in Dry Air.** Like aluminum and other reactive metals, plutonium is passivated, or rendered unreactive, upon exposure to air because a coherent (continuous, uncracked) oxide layer rapidly forms over the entire surface. Although oxidation continues despite the protective oxide coating, its rate in dry air at room temperature is extremely low. Evidence from kinetic data demonstrates that the oxidation rate is limited by the rate at which oxygen can diffuse through the oxide surface to the oxide-metal interface.

Figures 1(a)–1(c) show the oxidation process broken down into a sequence of steps. A freshly burned surface of plutonium is exposed to oxygen gas. Oxygen molecules (O₂) adsorb on the oxide surface at a concentration determined by temperature and the partial pressure of oxygen in the gas phase. The adsorbed molecules then dissociate on the oxide surface to form atomic oxygen, a species that either recombines or associates with electrons to form oxide ions (O²⁻). Both adsorption and dissociation depend strongly on the electronic properties of the oxide layer and its ability to transport electrons from the metal to the gas-solid interface. After entering the oxide lattice, O²⁻ diffuses through the oxide layer and ultimately reacts with plutonium to produce oxide, electrons, and heat at the oxide-metal interface. The slowest step in this sequence is called rate limiting because the overall reaction can proceed no faster.

Kinetic data on the thickness τ of the adherent oxide layer as a function of time t demonstrate that classic oxygen diffusion is the rate-limiting step in the Pu + O₂ reaction. On a freshly burned plutonium surface, the thickness τ reflects the extent of the reaction. This thickness exhibits parabolic growth, increasing linearly with the square root of t at a fixed temperature T. The rate of reaction, which is the time derivative of the thickness, must therefore be inversely proportional to the thickness, exactly the behavior expected if the rate of reaction is controlled by the rate at which oxygen diffuses through the oxide layer. As the oxide layer gets thicker, the time to diffuse through the layer becomes longer, and thus the oxidation rate decreases with time. Figure 1(d) shows the decreasing slope of the curve describing thickness versus time during parabolic growth.

Parabolic growth tapers off to a steady-state regime when the low-density oxide—11.45 grams per cubic centimeter (g/cm³)—on high-density plutonium (19.86 g/cm³) begins to induce stresses that lead to the localized spallation of oxide particles from the surface. The thickness of the oxide layer then varies from point to point with regions of thin oxide in recently spalled areas, regions of thick oxide in unsprayed areas, and regions of intermediate oxide thickness in between, as illustrated in Figure 1(c). During this stage, the average thickness of the oxide layer and the isothermal oxidation rate reach constant values as diffusion-controlled oxidation continues. The corrosion rate becomes constant because the continuous spallation of oxide particles and the reoxidation of the surface maintain a steady-state diffusion barrier of constant average thickness. As shown in Haschke et al. (1996), the corrosion rate of unalloyed plutonium in dry air at 25°C is approximately 0.5 nanogram of plutonium per centimeter squared a minute (ng Pu/cm² min), and the steady-state oxide thickness is 4 to 5 micrometers (Martz et al. 1994).

As expected, the diffusion rate through the oxide layer increases strongly with increasing temperature and produces a corresponding increase in the oxidation rate of plutonium. During both the parabolic and constant-rate stages of oxidation, the reaction rate R obeys the classical Arrhenius relationship

\[ R = \exp(-E_p/R^*T) \]

where \( E_p \) is the activation energy for the reaction and \( R^* \) is the gas constant. Activation energies for the parabolic and constant-rate stages are typically derived from the slopes of experimental curves for \( \ln R \).
Surface and Corrosion Chemistry of Plutonium

(a) Freshly Burnished Surface

(b) Parabolic Growth

(c) Steady-State Corrosion

(d) Oxide Thickness versus Time

Figure 1. Standard Chemical Picture of Plutonium Oxidation in Dry Air

(a) A freshly burnished plutonium surface is exposed to molecular oxygen that readily adsorbs onto the metal surface. (b) The oxygen dissociates into atomic oxygen, and combines with plutonium to form a layer of oxide. Oxidation continues, but now the oxygen on the surface must diffuse through the oxide layer before it can react with plutonium and produce more oxide at the oxide/metal interface. The thickness of the oxide layer increases parabolically with time because its growth is limited by the rate of oxygen diffusion through the oxide layer. (c) At a certain oxide thickness (typically, 4–5 µm), at room temperature, surface stresses cause oxide particles to spall from the surface. The oxide layer reaches a steady-state thickness as further oxidation is counterbalanced by spallation. If the reaction is to occur as outlined in steps (a)–(c), the electrons must be transported from the metal to the oxide surface so that O\(^2^−\) ions should form. (d) The plot of oxide layer thickness (τ) versus time (t) shows the two distinct oxidation stages. During parabolic growth, the reaction extent and τ grow as the square root of t, and the oxidation rate (slope of the curve) continually decreases, indicating that diffusion through the oxide layer is the rate-limiting step in the oxidation process. Later on, the extent of the reaction grows linearly with time, and the corrosion rate becomes constant as continuous spallation of oxide particles and reoxidation of the surface maintain a steady-state diffusion barrier of constant average thickness. Diffusion remains the rate-limiting step.
Figure 2. Arrhenius Curves for Oxidation of Unalloyed and Alloyed Plutonium in Dry Air and Water Vapor

Data on the steady-state oxidation rates of alloyed and unalloyed plutonium are summarized for a wide temperature range. Each labeled curve is an Arrhenius plot, showing the natural logarithm (ln) of the reaction rate $R$ versus $1/T$ for a metal or alloy in a specific atmosphere or under a specific condition. The slope of each curve is proportional to the activation energy for the corrosion reaction. Curve 1 plots the well-known oxidation rate of unalloyed plutonium in dry air or dry $O_2$ at a pressure of 0.21 bar. Curve 2a shows the increase in the oxidation rate when unalloyed metal is exposed to water vapor at equilibrium pressures up to 0.21 bar (160 torr), a concentration equal to the partial pressure of oxygen in air. Curves 2b and 2c show the moisture-enhanced oxidation rate at a water vapor pressure of 0.21 bar (160 torr) in the temperature ranges of 61°C–110°C and 110°C–200°C, respectively. Curves $1'$ and $2'$ give the oxidation rates for the $\delta$-phase gallium-stabilized alloy in dry air and moist air (water vapor pressure ≤ 0.21 bar), respectively. Curve 3 indicates behavior in the transition region between the convergence of rates at 400°C and the onset of the autothermic reaction at 500°C. Curve 4 defines the temperature-independent reaction rate of ignited metal or alloy under static conditions. The rate is fixed by diffusion through an $O_2$-depleted boundary layer of $N_2$ at the gas-solid interface. Curve 5 shows the temperature-dependent oxidation rate of ignited droplets of metal or alloy during free fall in air.
versus $1/T$ (see Figure 2). Curve 1 shows the oxidation rate of unalloyed plutonium in dry air during the constant-rate stage of the reaction, and its slope yields an $E_a$ of 17.4 kilocalories per mole (kcal/mol).

**Variations from the Standard Diffusion Picture.** To get a comprehensive picture of the kinetics of plutonium oxidation, we have reviewed published data and prepared a single graph (Figure 2) of Arrhenius curves for corrosion of unalloyed metal and delta-phase gallium alloy in dry air and water vapor during the constant-rate stage of oxidation. The data are normalized to an oxidant pressure of 0.21 bar, the partial pressure of $O_2$ in air at 1 atmosphere.

Curve 1, the oxidation rate of unalloyed plutonium in dry air, spans a temperature range in which the metal exists in four allotropic forms: the alpha, beta, gamma, and delta phases. The variations among curves 1, 2a–2c, 1', and 2' reflect the complex effects of temperature, humidity, and alloying on the oxidation rate. The convergence of these curves near 400°C marks the onset of a region (curve 3) in which the oxidation rate depends only on temperature and oxidant pressure.

The research we describe below shows that all the curves below 500°C in Figure 2 are consistent with control by classic oxygen diffusion through an oxide layer, even those enhanced by the presence of moisture.²

For completeness, we also include Arrhenius data for the very high temperature range—from the 500°C ignition point of plutonium to the boiling point of the liquid metal at 3230°C. (Haschke and Martz 1998a, Martz and Haschke 1998). Activation energies in this region are difficult to measure because the reaction temperature is determined by the rate of reaction and the accumulation of heat by the reacting metal, not by experimental control. The oxidation rate—0.14 gram (g) Pu/cm² min—for ignited metal droplets in static air (curve 4) is independent of temperature and fixed by the rate of $O_2$ transport across a boundary layer of oxygen-depleted nitrogen formed at the gas-solid interface of the burning particle. An example of this behavior is illustrated in Figure 3, in which plutonium is shown spontaneously burning in air under static conditions. The oxidation rate ($E_a = 9.6$ kcal/mol) for ignited metal droplets in air during free fall (curve 5) is temperature dependent because there is no longer a static nitrogen-rich boundary layer limiting the diffusion of oxygen. Self-heating by the oxidation reaction drives droplet temperatures above the boiling point of plutonium and causes an “explosion” in which unreacted metal vaporizes and burns in a flash.

Although the standard picture of diffusion-controlled oxidation given in Figure 1 seems to apply below 500°C, this model does not account either for the spontaneous ignition of metal chips and powder in air at 150°C–200°C or for the large effect of moisture on the oxidation rate (compare curves 2a and 2b with curve 1 in Figure 2). Involvement of compounds other than the dioxide and transport processes other than diffusion must be considered in addressing pyrophoricity, moisture-enhanced oxidation, and other anomalous kinetic behaviors described in subsequent sections of this article.

**Pu₂O₃ and the Oxide Layer.** The presence of oxide is an unavoidable feature of plutonium metal surfaces. Even if one tries to create a perfectly clean surface through repeated cycles of heating and bombardment by an energetic ion beam in ultrahigh vacuum, several atomic percent of oxygen remains on the surface, as measured by x-ray photoelectron spectroscopy (XPS). Such measurements also show that exposure of a cleaned surface to $O_2$ at low pressure (about 1 nanobar) immediately produces a surface layer of Pu₂O₃ (Larson and Haschke 1981). Over time, this sesquioxide oxidizes to

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² This enhancement apparently results from an unanticipated increase in the oxygen gradient rather than from a proposed mechanistic change involving diffusion of hydroxide (OH⁻) ions, but that story is the most surprising, and we save it for last.
PuO$_2$ even at these low oxygen pressures. If the surface is exposed to air at atmospheric pressure, a coherent layer of PuO$_2$ quickly forms over the entire metal surface. Thus, a dioxide layer is present on all plutonium metal surfaces after exposure to air unless extreme measures are exercised.

A surprising feature of the oxide layer at room temperature is the apparent absence of Pu$_2$O$_3$. Studies of the plutonium-oxygen system have shown that both Pu$_2$O$_3$ and PuO$_2$ are stable in the solid phase at room temperature. Moreover, above 700°C, the two oxides coexist as a solid solution in which the average composition varies between these oxides as the oxygen-to-plutonium ratio in the system increases from 1.5 to 2. (The mechanism for forming a solid compound with a continuously varying composition is described in the box “Fluorite and Fluorite-Related Structures in Plutonium Corrosion” on page 260.)

Thermodynamic data presented in the box “Thermodynamics, Kinetics, Catalysis, and the Equilibrium State in the Plutonium-Oxygen System” (page 261) verify that Pu$_2$O$_3$ is a stable oxide in the presence of plutonium metal. Thus, a thin layer of Pu$_2$O$_3$ must be present at the oxide-metal interface of the oxide layer. Because this layer is undetectable at room temperature, we deduce that Pu$_2$O$_3$ is readily oxidized to PuO$_2$ by oxygen and that the observed composition of the oxide layer is determined by the rapid kinetics of PuO$_2$ formation, not by thermodynamics. This conclusion is supported by our observations that a change in conditions, such as an increase in temperature or a decrease in the availability of oxygen, leads to an increase in the fraction of Pu$_2$O$_3$. Indeed, our recent results (Haschke et al. 1998) strongly suggest both the presence and participation of Pu$_2$O$_3$ in the corrosion chemistry of plutonium, possibilities that have not been examined by previous investigators.

The Pyrophoricity of Plutonium.

The first example of Pu$_2$O$_3$ participation is our quantitative model for the pyrophoricity of plutonium chips and powder at 150°C–200°C (Martz et al. 1994). This model grew from understanding the role of Pu$_2$O$_3$ in the kinetics of plutonium oxidation.

Figure 4 shows the revised view of the PuO$_2$ diffusion barrier in the presence of oxygen. By separating a region with excess oxygen from one with excess plutonium, this barrier creates a nonequilibrium condition on the metal surface, in which Pu$_2$O$_3$ is simultaneously consumed and formed by competing reactions. The dioxide is reduced to Pu$_2$O$_3$ by plutonium at the oxide-metal interface.

\[
3\text{PuO}_2(s) + \text{Pu}(s) \rightarrow 2\text{Pu}_2\text{O}_3(s) \quad (1)
\]

Concurrently, Pu$_2$O$_3$ is consumed by reaction with the oxygen that diffuses through the oxide layer:

\[
\text{Pu}_2\text{O}_3(s) + (1/2)\text{O}_2(g) \rightarrow 2\text{PuO}_2(s) \quad (2)
\]

The ultimate equilibrium state of a reaction depends on the molar ratio of oxygen to plutonium in the product. During corrosion in air, excess O$_2$ is present, and equilibrium is reached when all the metal is converted to dioxide. In contrast, when placed in a vacuum, an inert atmosphere, or a sealed container, plutonium consumes any residual O$_2$ and continues to react with the PuO$_2$ surface until only Pu$_2$O$_3$ is present. In the absence of excess oxygen, equilibrium is reached when the PuO$_2$ layer is completely reduced to Pu$_2$O$_3$ by the reaction described by

\[
\text{Pu} + \text{O}_2 \rightarrow \text{Pu}_2\text{O}_3 + \text{heat}
\]
Equation (1). Indeed, data from x-ray diffraction (XRD) and XPS measurements show that, although the surface layer of PuO₂ on a plutonium sample is not detectably altered when the sample is placed in vacuum at room temperature, the dioxide layer is immediately transformed to Pu₂O₃ when that same sample is heated to 150°C in vacuum. Evidently, the rate of Pu₂O₃ formation according to Equation (1) increases sharply with temperature as shown in Figure 5(a).

Data from XRD and XPS measurements also demonstrate that the oxide layer formed in air at 350°C is predominantly Pu₂O₃ beneath a thin layer of PuO₂ and a higher oxide. (The nature of this higher oxide is discussed later.) In other words, the ratio of Pu₂O₃ to PuO₂ at 350°C is essentially the reverse of that observed at room temperature. This observation and the rapid rate of Pu₂O₃ formation at 150°C suggest that the temperature dependence for oxide reduction, Equation (1), is stronger than for the subsequent oxidation of Pu₂O₃, Equation (2). Therefore, we conclude that the fraction of Pu₂O₃ in the oxide layer during oxidation in dry air increases significantly at elevated temperatures as illustrated in Figure 5(b).

This dynamic picture of the oxide layer suggests that plutonium pyrophoricity at 150°C–200°C results from an increased fraction of Pu₂O₃ in the oxide layer at those temperatures.

With this picture in mind, we consider details of plutonium ignition studies performed earlier by numerous investigators, who determined ignition points by heating a metal specimen at a constant rate in flowing air while measuring the metal’s temperature. The ignition temperature was marked by the onset of a self-sustained reaction, indicated by a sharp and sustained increase in specimen temperature above the programmed value. Temperature curves for relatively massive (greater than 0.2 millimeter in thickness) pieces of metal with low specific surface areas—less than 5 centimeters squared per gram (cm²/g)—show small thermal spikes at 150°C–200°C, but those samples do not spontaneously

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**Figure 5. The Oxide Layer on Plutonium Metal under Varying Conditions**

These highly idealized sketches indicate how changes in atmosphere, temperature, and time alter the surface chemistry of the oxide layer. The corrosion rate is strongly dependent on the metal temperature and varies significantly with the isotopic composition, quantity, geometry, and storage configuration of the metal. Serving as reference, the steady-state oxide layer on plutonium in dry air at room temperature (25°C) is shown at the top. (a) Over time, isolating PuO₂-coated metal from oxygen in a vacuum or an inert environment turns the surface oxide into Pu₂O₃ by the autoreduction reaction described by Equation (1). At 25°C, the transformation is slow.

The time required for the complete reduction of PuO₂ depends on the initial thickness of the PuO₂ layer and is highly uncertain because the reaction kinetics is not quantified. At temperatures above 150°C, rapid autoreduction transforms a several-micrometer-thick PuO₂ layer to Pu₂O₃ within minutes. (b) Exposure of the steady-state oxide layer to air results in continued oxidation of the metal. At 25°C in dry air, the layer is essentially unaltered over time. Kinetic data indicate that a one-year exposure to dry air (<0.5 ppm H₂O at 1 bar of total pressure) at room temperature increases the oxide thickness by about 0.1 μm. At a metal temperature of 50°C in moist air (50% relative humidity), the corrosion rate increases by a factor of approximately 10⁴, and the corrosion front advances into unalloyed metal at a rate of 2 mm per year. At 150°C–200°C in dry air, the rate of the autoreduction reaction increases relative to that of the oxidation reaction, and the steady-state condition in the oxide shifts toward Pu₂O₃, a phase that is simultaneously formed by Equation (1) and consumed by Equation (2).
Fluorite and Fluorite-Related Structures in Plutonium Corrosion

Plutonium-oxygen compounds and many other products of plutonium corrosion form in the classic calcium fluoride (CaF$_2$) structure or one of its variations (see figure). As discussed below and in the main text, these particular structures seem to facilitate the remarkable catalytic properties of plutonium compounds.

Within the fluorite-type structure, plutonium cations (the red circles in the figure to the right) form a face-centered-cubic (fcc) lattice; that is, they occupy sites at the corners and centers of all the faces of a cube, where $a_0$ is the length of the cubic cell in angstroms. Atoms at corner sites are shared by eight cubes, and those in faces are shared by two cubes. Thus, four atoms reside within a single unit cell.

Anions such as O$^{2-}$ and H$^{-}$ occupy interstitial sites within the fcc cation lattice. The figure shows two types of interstitial sites: eight tetrahedral sites (gray circles), each surrounded by four cations, and four octahedral sites (white squares), each surrounded by six cations. The CaF$_2$ structure has an fcc lattice of cations with even-integer charge ($2n^+$); all tetrahedral sites are occupied by anions with charge $n^-$.

The fluorite (CaF$_2$) Structure

Cationic positions occupied by plutonium are in an fcc configuration indicated by red circles. Tetrahedral and octahedral sites in the lattice are shown by gray circles and white squares, respectively. The cubic-lattice parameter is defined by $a_0$.

Electrons are apparently removed from the conduction band and bound as H$^+$ on octahedral sites as the hydride composition increases.

Another example of a fluorite-related structure is provided by plutonium monoxide monohydride (PuOH), which is formed by the reaction of plutonium metal with liquid water at 25°C. Charge balance in this ternary compound of Pu(III) is achieved because O$^{2-}$ and H$^+$ occupy tetrahedral sites equally.

These fluorite and fluorite-related structures provide a stable fcc matrix of invariant, immobile plutonium cations, in which anions are surprisingly mobile. Anions move in and out of the stationary metal as chemical reactions occur and the plutonium oxidation state changes. The NaCl type of plutonium mononitride (PuN) forms from PuH$_2$ of the CaF$_2$ type because H$^+$ is displaced from tetrahedral sites and N$^{3-}$ is allowed to occupy all octahedral sites. The rapid rates of ionic transport and exchange are remarkably similar to those of superionic conduction, a phenomenon frequently observed in fluorite materials. Superionic conductors are solids characterized by a very rigid host lattice, light mobile ions, and high direct-current ionic conductivities. Transport in anion-deficient fluorite materials may also be enhanced because both tetrahedral and octahedral sites are vacant and able to participate in the concerted movement of anions. The impact of such factors on the surface and corrosion chemistry of plutonium has not yet been investigated.
Plutonium corrosion chemistry involves the complex interplay of thermodynamic, kinetic, and catalytic factors. Although often misunderstood and incorrectly applied, these concepts are the foundation of all chemistry and chemical processing. It is worthwhile to review those concepts and see how their careful application to the plutonium-oxygen system under varying conditions leads to some surprising results.

Thermodynamic properties quantify the fundamental driving force of chemical reactions. In particular, the drive toward thermodynamic equilibrium is due to the energy differences between atoms in different chemical configurations (or states). Energy differences between initial (reactant) and final (product) states are defined by the free energy change ($\Delta G^\circ$) for the reaction. If a reaction releases energy ($\Delta G^\circ$ is negative), it is thermodynamically favorable and can occur spontaneously. Conversely, if $\Delta G^\circ$ is positive, the reaction is not spontaneous and cannot occur unless sufficient energy is added to the system.

The reaction that occurs in a given system is not necessarily the energetically most favorable one, even though the latter would lead to the equilibrium state of that system. For example, when several reactions with negative free energies are possible, kinetics determines the reaction rate and pathway. In certain systems, energetically favorable reactions never occur at room temperature because their rates are immeasurably slow. Although such systems are not thermodynamically stable, they are kinetically stable, or “metastable.” The reaction observed initially is always the one with the highest rate. If that reaction did not have the most negative $\Delta G^\circ$, it would lead to different metastable states and to the possibility of subsequent reactions that progressively move the system toward equilibrium. A system remains in kinetic control until it reaches the equilibrium state.

Catalysts are substances that change reaction rates but not equilibrium states. By altering a reaction pathway, a catalyst increases the rate of the observed reaction or changes the chemistry of the system by enhancing the rate of a competing reaction. Typical catalysts for the reaction of gases are solids with active surfaces that promote the adsorption and dissociation of reactants and their recombination as products.

Applying these principles to the plutonium-oxygen system requires some care. Table I shows that the free energy for oxidizing 1 mole of plutonium into plutonium dioxide ($\text{PuO}_2$) is more negative than that for forming a half mole of plutonium sesquioxide ($\text{Pu}_2\text{O}_3$). However, contrary to a widely accepted view, coexistence of a $\text{PuO}_2$ layer and plutonium metal is not always the equilibrium configuration. If oxygen is present in excess, the most stable configuration is, indeed, achieved by reaction of the available metal with the maximum amount of oxygen. The relevant measure of stability is the energy released per mole of plutonium reactant, and $\text{PuO}_2$ is the equilibrium oxide. Conversely, if plutonium is present in excess, the most stable configuration is achieved by reacting the available oxygen with the maximum amount of plutonium. The relevant measure is the energy released per mole of atomic oxygen ($\text{O}_2$), and $\text{Pu}_2\text{O}_3$ is the equilibrium oxide on plutonium at room temperature. Thus, the predominance of dioxide on the metal surface at room temperature is clearly controlled by kinetic factors, not by product stability.

Figure 5 in the main text illustrates that exposure of plutonium to the abundant supply of oxygen in the atmosphere creates an inherently nonequilibrium condition. The oxide layer on the metal surface is a diffusion barrier separating an oxygen-rich region and a metal-rich region. Thus, pseudo-equilibrium conditions exist on opposite sides of that barrier. The presence of excess $\text{O}_2$ in the gas phase produces $\text{PuO}_2$ near the gas-oxide interface; the reaction of excess plutonium with the dioxide at the oxide-metal interface yields $\text{Pu}_2\text{O}_3$. At room temperature, the rapid formation of $\text{PuO}_2$ compresses the metal-rich region to the point that $\text{Pu}_2\text{O}_3$ is not observed, but increasing temperature promotes the formation of $\text{Pu}_2\text{O}_3^+$, which becomes dominant above 150°C–200°C.

Although thermodynamic properties define possible reactions and equilibrium states of chemical systems, they fail to predict the strong metastable behavior of the plutonium-oxygen system. A system cannot reach equilibrium if slow kinetics prevents the most stable product from forming. Such is the case for the higher oxide, $\text{PuO}_{2+x}$. After early workers had failed to prepare oxides with compositions greater than $\text{PuO}_2$, they concluded that the dioxide is the equilibrium oxide in air. Their use of strong oxidants such as ozone ($\text{O}_3$) and nitrogen dioxide ($\text{NO}_2$) increased the free energy for reaction, making the thermodynamics unfavorable, but slow kinetics prevented further oxidation. Although water is energetically a less favorable oxidant than $\text{O}_2$, our research has shown that it both participates in and catalyzes the formation of $\text{PuO}_{2+x}$ by altering the chemical pathway.

The need to characterize fundamental behavior in a plutonium-oxygen system and in other chemical systems is evident. We are trying not only to understand how water promotes oxidation, but also to define the phase equilibria and thermodynamic properties of $\text{PuO}_{2+x}$.

**Table I. Free-Energy Data for the Formation of Plutonium Oxides**

<table>
<thead>
<tr>
<th>Reaction</th>
<th>$\Delta G^\circ$ (kcal/mol Pu)</th>
<th>$\Delta G^\circ$ (kcal/mol $\text{O}_2$)</th>
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<tbody>
<tr>
<td>$\text{Pu(s)} + \text{O}_2(g) \rightarrow \text{PuO}_2(s)$</td>
<td>$-239$</td>
<td>$-239$</td>
</tr>
<tr>
<td>$\text{Pu(s)} + 3/4 \text{O}_2(g) \rightarrow 1/2 \text{Pu}_2\text{O}_3(s)$</td>
<td>$-189$</td>
<td>$-252$</td>
</tr>
</tbody>
</table>
ignite until they reach 500°C (±25°C) during continued heating. In contrast, small pieces—turnings, chips, and powder—with maximum dimensions less than 0.07 millimeter and specific areas greater than 15 cm²/g are pyrophoric when heated to 150°C–200°C in air. The thermal spikes observed at 150°C–200°C decrease in size with increasing specimen thickness, suggesting that pyrophoricity is driven by a surface reaction involving oxides. A surface reaction would produce a fixed amount of heat per unit of surface area, resulting in small thermal spikes for massive pieces of metal and large thermal excursions for material forms with high ratios of specific surface area to volume, exactly the pattern observed.

Our model of pyrophoricity applies the idea of surface reaction to our kinetic picture of the oxide layer. According to the model, heating the metal to 150°C–200°C in dry air transforms a large fraction of the steady-state dioxide layer (4–5 micrometers thick) to Pu₂O₃, as indicated by the lower sketch in Figure 5(b). Rapid oxidation of the Pu₂O₃ layer back to PuO₂ produces 54 kilocalories of heat per mole of dioxide, and a constant oxide thickness fixes the amount of heat produced per unit of surface area. The size of the resulting thermal spike is determined by the rate of reoxidation, the metal’s ratio of surface area to volume, and the heat capacities of the oxides and metal.

We have used this model to calculate the thermal excursion of specimens with different dimensions and with geometries having the smallest (spheres) and largest (sheets) ratios of surface area to volume. We assume that the entire oxide layer is Pu₂O₃, reoxidation of Pu₂O₃ is rapid, and all heat from the reaction is instantaneously deposited in the reacting particle. If the excursion for a given metal dimension and geometry exceeds 350°C, we expect the metal temperature to reach or exceed 500°C and predict self-sustained reaction. Results show that pyrophoric behavior is not expected for metal spheres with diameters greater than 0.25 millimeter but is predicted for metal sheets with thicknesses less than 0.09 millimeter. Excellent agreement with experimental observation suggests the validity of our model.

We also propose that a likely mechanism for initiating pyrophoric reaction is exposure of the Pu₂O₃ layer to air by spallation of an oxide particle. A protective PuO₂ layer is not reestablished after ignition because rapid diffusion of oxygen and rapid reduction of PuO₂ to Pu₂O₃ are favored at the elevated temperatures reached during a self-sustained reaction. We also note that diffusion cannot account for the rapid transport of oxygen needed to initiate a self-sustained reaction at 150°C and conclude that Pu₂O₃ promotes transport of oxygen to the oxide-metal interface.

**Conjecture on Oxygen Transport in Plutonium Oxide.** Of the many kinetic factors introduced in this overview of corrosion in air, the mobility of the oxygen in the product layer is especially important and intriguing. One must wonder why diffusion of oxygen in PuO₂ is slow and rate limiting, whereas transport of oxygen in Pu₂O₃ is apparently very rapid. This behavior and other key properties of solid reaction products are strongly influenced by their crystal structures. All corrosion products formed by reaction with oxygen, hydrogen, and water have structures whose plutonium atoms are arranged in face-centered-cubic (fcc) configurations (see the box “Fluorite and Fluorite-Related Structures in Plutonium Corrosion” on page 260). And the chemical and kinetic behaviors described in this article suggest that the fcc plutonium lattice remains stationary whereas anions such as O²⁻ and H⁺ move in and out of the interstitial sites in that lattice with apparent ease. The facile transport of anions in Pu₂O₃ is neither quantified nor understood but is consistent with enhanced anionic mobility in structures with vacant lattice sites.

**Reaction of Oxide-Coated Plutonium with Hydrogen**

We now turn to the reaction of plutonium with hydrogen (hydriding), in which surface compounds are, once again, a determining factor. Plutonium hydride (PuHₓ) is an fcc phase that forms a continuous solid solution for 1.9 < x < 3.0 (Haschke 1991, Ward and Haschke 1994).

\[
\text{Pu(s)} + (x/2)\text{H}_2(g) \rightarrow \text{PuH}_x(s) . \quad (3)
\]

The observed value of x depends on hydrogen pressure and temperature. The hydride is readily oxidized by air, and it decomposes back to its component elements when heated in dynamic (continuously pumped) vacuum.

Hydriding occurs only after the ubiquitous dioxide layer on the metal is penetrated. Unlike oxidation, which proceeds evenly over the entire dioxide surface, the reaction of hydrogen initiates at a limited number of nucleation sites, and a single nucleation site typically appears only after a lengthy, but unpredictable, induction period. Once formed, these sites are the most reactive areas of the surface. They grow like bacterial colonies; that is, the hydriding rate is proportional to the active area covered by the hydride, and it increases exponentially over time to a maximum value as sites grow and ultimately cover the surface. At that point, the rate is enormous and constant because the surface is fully active. For temperatures between −55°C and 350°C and a molecular hydrogen (H₂) pressure of 1 bar, the reaction at a fully active surface consumes plutonium at a constant rate of 6–7 g/cm² min and advances into the metal or alloy at about 20 cm/h.

Curve b in Figure 6 shows that hydriding is 10¹¹ times faster than oxidation of unalloyed metal in dry O₂ at 25°C. Moreover, the hydriding rate is independent of temperature and proportional to the square root of hydrogen pressure, indicating that the rate-controlling step involves dissociation of H₂ at the gas-hydride interface rather than...
than diffusion of hydrogen through the surface layer of the hydride product. On the basis of this extremely rapid rate, we conclude that PuHₓ catalyzes the dissociation of adsorbed H₂ and promotes the transport of atomic hydrogen to the hydride-metal interface.

A dramatically different kinetic behavior occurs if Pu₂O₃ replaces PuO₂ on the surface layer. Instead of forming and growing isolated nucleation sites, hydriding of Pu₂O₃-coated metal proceeds over the entire surface at once. In our studies, we heated Pu₂O₂-coated metal to 150°C–200°C in vacuum to transform the surface to Pu₂O₃—see bottom sketch in Figure 5(a)—then cooled the sample to 25°C, and finally exposed it to H₂ at 1 bar of pressure. We found that the initial (zero time) hydriding rate equals that attained after the dioxide-coated metal is fully activated by the growth and coalescence of PuHₓ sites. These results show that Pu₂O₃ is equivalent to PuHₓ as a catalyst for the dissociation of H₂ and as a medium for hydrogen transport to the product-metal interface. Hydriding may nucleate at specific sites on the surface of the dioxide-coated plutonium metal because Pu₂O₃ is exposed at those locations by the autoreduction of the dioxide layer in thin areas or by the spallation of an oxide particle.

In summary, two independent surface processes catalyze the corrosion rate of oxide-coated plutonium metal by hydrogen. The surface self-activates by the nucleation and growth of PuHₓ sites on the dioxide-coated metal or, before it is exposed to H₂, by the autoreduction of PuO₂ to Pu₂O₃. In both cases, the catalytic enhancements of the dissociation and transport of H₂ increase the initial corrosion rate at 25°C by a factor of 10⁵.

Suppose now that the plutonium has been coated with hydride before being exposed to O₂. We find that PuHₓ and Pu₂O₃ are simultaneously involved in promoting an extreme pyrophoricity of both pure hydride particles and hydride-coated metal. We also relate the pyrophoricity of PuHₓ to its variable stoichiometry, which allows the hydride to absorb additional hydrogen until x approaches 3. The stoichiometry of PuHₓ varies, depending on conditions. Values of x near 2 are favored for surface layers formed on the metal at high temperatures and low H₂ pressures; higher values of x are found on isolated hydride particles at low temperatures and high pressures.

Earlier studies (Haschke 1991) show that particles of PuH₂ react spontaneously and rapidly upon controlled exposure to O₂ at 25°C, and the reaction produces an oxide layer on the
The value of $x$ in the residual $\text{PuH}_x$.

Our recent work (Haschke et al. 1998) shows that self-heating from the reaction raises the temperature of the particle so that the oxide product is $\text{Pu}_2\text{O}_3$ instead of $\text{PuO}_2$. The net reaction is given by

$$x\text{PuH}_2(s) + 3(x - 2)/4\text{O}_2(g) \rightarrow [(x - 2)/2]\text{Pu}_2\text{O}_3(s) + 2\text{PuH}_x(s) \quad (4)$$

Pyrophoricity and rapid oxidation of the hydride are promoted by the absence of gaseous reaction products that would restrict access of $\text{O}_2$ to the surface. As the value of $x$ within a hydride particle approaches 3, the particle raptures while the equilibrium $\text{H}_2$ pressure rises. The particle’s extent of oxidation and ratio of surface area to volume determine the point of rupture.

In recent studies (Haschke et al. 1998), we also focused on hydride-coated metal and alloys and found that they react violently when rapidly exposed to oxygen. We prepared specimens with $\text{PuH}_2$ surface layers (100 micrometers thick) by first heating dioxide-coated plutonium (1–2 millimeters thick) above 150°C in vacuum to form $\text{Pu}_2\text{O}_3$ on the surface and then exposing that lower oxide layer to the requisite quantity of $\text{H}_2$. After exposure to excess $\text{O}_2$, the metal in the hydride-coated specimens was completely consumed in less than 1 second, and the resulting gas-phase temperatures exceeded 1000°C. Hydrogen was not released as $\text{H}_2$ even though excess $\text{O}_2$ was present. Instead, the hydrogen appeared as a small $\text{PuH}_3$ core inside an expanded product shell of $\text{Pu}_2\text{O}_3$ that retained the shape of the starting specimen. The reaction ended so quickly, and the thermal effect was so large that the rate could not be accurately determined by pressure-time data. Based on a reaction time of 1 second, the rate is about 80 g $\text{Pu}/\text{cm}^2 \cdot \text{min}$, corresponding to a corrosion rate of 3 m/h.

We attribute the extraordinarily rapid oxidation of hydride-coated plutonium to catalysis by $\text{PuH}_x$. For both the pure hydride and hydride-coated metal, the reaction of $\text{O}_2$ with the hydride initiates spontaneously, produces oxide, and increases the stoichiometry of the hydride by driving the product hydrogen into the solid phase as described by Equation (4). Excess hydrogen in the hydride is continuously produced at the oxide-hydride interface and consumed at the hydride-metal interface. The reaction generates a large amount of heat (approximately 200 kcal/mol of plutonium). At elevated temperatures, the net reaction is accurately described by Equation (5).

$$2\text{Pu}(s) + 3/2 \text{O}_2(g) \rightarrow \text{Pu}_2\text{O}_3(s) \quad (5)$$

This hydride-catalyzed oxidation of plutonium continues until all the metal is consumed and the composition of the residual hydride reaches $\text{PuH}_3$. Oxidation of that hydride is expected but not observed because the hydride is encased in the thick shell of the silver gray oxide.

The rate of oxidation is definitely much higher than anticipated for a gas-solid reaction. Plutonium hydride is an effective catalyst for the oxidation of plutonium because it forms at a rapid rate via the $\text{Pu} + \text{H}_2$ reaction and also oxidizes rapidly via the $\text{PuH}_x + \text{O}_2$ reaction. Thus, the relatively slow $\text{Pu} + \text{O}_2$ reaction is promoted by participation of the hydride as a reactive intermediate. As detailed in Figure 7, five major processes are...
involved in a hydride-catalyzed oxidation: reaction of O₂ at the gas-solid interface, transport of oxygen across the layer of product oxide, oxidation of the pyrophoric hydride at the oxide-hydride interface, transport of product hydrogen across the hydride layer, and reformation of the hydride at the hydride-metal interface. The slowest of these processes apparently determines the corrosion rate, but it has not been identified.

Other compounds also promote rapid corrosion. For example, diplutonium monoxide monocarbide (Pu₂OC), a surface phase incorrectly identified as plutonium monoxide (PuO) in early literature, forms at elevated temperatures and promotes hydriding. We have recently shown (Allen and Haschke 1998) that plutonium monoxide monohydrate (PuOH) formed by the rapid corrosion of plutonium in salt water also catalyzes the Pu + O₂ reaction.

The unusual transport properties of Pu₂O₃ and PuHₓ discussed in earlier examples are implied here as well. Based on the observed rate of catalyzed oxidation, the hydride advances into the metal at the rate expected for the reaction of plutonium with H₂ at a pressure of 150 bar. If transport across the hydride layer (approximately 100 micrometers thick) is rate limiting, the coefficient for the diffusion of hydrogen in PuHₓ is temperature independent and on the order of 10⁻⁴ centimeter squared per second (cm²/s). The corresponding value for the transport of oxygen in Pu₂O₃ must be of equal magnitude or larger. By comparison, coefficients determined for the self-diffusion of oxygen in Pu₂O₃ at 250°C–500°C are in the range of 10⁻¹⁵ to 10⁻¹² cm²/s (Stakebake 1988). The very rapid temperature-independent transport of both oxygen in Pu₂O₃ and hydrogen in PuHₓ suggests that these compounds may be superionic conductors—solids with ionic mobilities and conductivities typically observed in molten salts.

We have shown that plutonium hydride on the metal activates the surface for the catalyzed corrosion with plutonium. Often the hydride forms from unanticipated sources of hydrogen, such as α-particle radiolysis of organic materials or chemical reactions of water. Regardless of the hydrogen source, the potential consequences are enormous. A comparison of curves c and 1 in Figure 6 shows that hydride-catalyzed corrosion of δ-phase plutonium alloy by O₂ at 25°C is more than 10¹³ faster than oxidation of plutonium in dry oxygen or air.

### Reaction of Hydride-Coated Plutonium with Air

When exposed to air as opposed to oxygen, plutonium hydride reacts with both oxygen and nitrogen. The PuHₓ + N₂ reaction is slow at room temperature but occurs readily at temperatures above 200°C–250°C. According to Equation (4), the reaction of O₂ with pyrophoric hydride produces sufficient heat to promote rapid formation of PuN.

\[
\text{PuH}_x(s) + [(x - 2)/2]\text{N}_2(g) \rightarrow (x - 2)\text{PuN(s)} + 2\text{PuH}_x(s) \quad (6)
\]

Again, the hydrogen (produced by the oxidation and nitriding reactions) remains in the solid and increases the stoichiometry of residual hydride. Thus, the pyrophoricity of PuHₓ is promoted because neither product H₂ nor residual N₂ accumulates and blocks the flow of oxygen to the gas-solid interface.

Plutonium metal coated with hydride also corrodes rapidly when exposed to air, and as in the previous example, the hydride coating catalyzes the reaction. The net reactions for the corrosion of the metal by oxygen and nitrogen in air are defined by Equations (5) and (7), respectively.

\[
\text{Pu(s)} + (1/2)\text{N}_2(g) \rightarrow \text{PuN(s)} \quad (7)
\]

Both reactions occur simultaneously at the gas-solid interface and indiscriminately consume nitrogen and oxygen at the molar ratio (3.71 N₂:1 O₂) of the elements in air. Hydrogen produced by these reactions is transported across the PuHₓ layer and reacts at the hydride-metal interface. Each mole of air consumes 1.86 moles of plutonium and generates approximately 170 kilocalories of heat. The corrosion rate is independent of temperature and alloying, is proportional to the square of the air pressure, and equals 0.7 ± 0.1 g Pu/cm² min in air at 1 bar of pressure. If air gains unrestricted access to surfaces of hydride-coated plutonium, a metal specimen with a thickness of 1 millimeter is completely corroded in about 1.5 minutes. We must further characterize the gray corrosion product to determine if it is an oxide-nitride mixture or a single-phase oxide nitride, but we know it reacts slowly in air at room temperature to form PuO₂.

The presence of hydride on the plutonium surface not only alters the kinetics of corrosion in air, but also changes the chemistry of corrosion. Elemental plutonium and molecular nitrogen (N₂) do not react directly to any significant extent even during their prolonged heating at temperatures above 1000°C. By contrast, nitrogen is the primary reactant during the hydride-catalyzed corrosion of plutonium in air. As a result, a nitrogen boundary layer does not form at the gas-solid interface as it does during the rapid oxidation of ignited plutonium. Thus, the observed rate for hydride-catalyzed corrosion in air is a factor of 5 faster than the rate for the self-sustained oxidation of ignited plutonium in air.

Compared with the rate for hydride-catalyzed corrosion in O₂, the rate in air is 100 times slower. This finding suggests that one or more steps involving reaction or transport of nitrogen limit the rate for air. The data in Figure 6 show that the hydride-catalyzed corrosion rate is 10⁸ times faster than the oxidation rate of unalloyed metal in saturated water-vapor or moisture-saturated air at room temperature. Occurrence of this hydride-catalyzed reaction in a storage container at Los Alamos is described in the box “Catalyzed Corrosion of Plutonium: Hazards and Applications.”
Catalyzed corrosion of plutonium:
Hazards and Applications

John M. Haschke and Joseph C. Martz

Catalyzed corrosion reactions of plutonium metal are not laboratory curiosities produced by careful manipulation of chemical reactants. Several incidents involving the corrosion of plutonium metal, failure of storage containers, and localized release of plutonium-containing particles into the work environment were attributed to catalyzed corrosion reactions (Haschke and Martz 1998b) and led to redesigning the storage package. In addition, catalyzed corrosion reactions have become the basis of efficient methods for converting plutonium metal from classified weapons configurations into simple ingots.

In 1993, a worker at the Los Alamos Plutonium Facility became contaminated with plutonium-containing particles while handling a standard storage package containing a 2.5-kilogram plutonium casting. The packaging configuration was similar to that used worldwide. It consisted of an inner cylindrical steel container to hold the casting, two layers of sealed “bag-out” plastic to contain radioactive contamination, and an outer slip-lid can. The incident occurred after the package had been stored for 11 years. As the outer can was flexed during handling, a puff of air carrying plutonium-containing particles escaped through a break in the taped seal of the can and contaminated the worker. The package was placed in a reduced-oxygen (less than 3 percent O₂) glove box, and the inner cylinder was left there for 3 hours after having been removed from the package. The accompanying photographs were taken during the disassembly. As the plastic-wrapped inner container was lifted out, its end ruptured because of pressure from the continuing rapid formation of plutonium oxides and other low-density corrosion products. A typical plutonium casting would form about 10 grams of PuO₂ per year from normal oxidation in air. Instead, the observed extent of the reaction was many orders of magnitude greater. Also, the inner vessel, which still contained unreacted metal, had become hot, and its diameter had increased by about 50 percent. The vessel was transferred to an argon-filled glove box, where the reaction ceased at once.

Evaluation of this incident shows that hydride catalysis caused by a complex set of physicochemical processes led to rapid corrosion of

(a) (b) (c) (d)
the metal casting during storage and to very rapid corrosion during the 3-hour waiting period in the reduced-oxygen glove box. The weld at the ruptured end of the inner vessel had evidently been defective from the start, allowing gases to be continually pumped in and out of that vessel, as atmospheric pressure changes compressed and expanded the sealed plastic bagging. Plutonium oxide particles were entrained by thermal currents in the vessel, transported through openings in the weld, and deposited on the plastic over several years. During that period, the bagging isolated the metal casting from any external oxygen source, and the normal surface PuO₂ was apparently autoreduced to Pu₂O₃. Over time, the plastic bagging became discolored near the weld, as well as embrittled from the radiolytic decomposition induced by α-particles emitted from the deposited plutonium oxide particles. The molecular hydrogen produced during the radiolysis of the plastic entered the inner vessel and formed hydride on the Pu₂O₃-activated surface of the casting. During these processes, the storage package did not change its mass, and therefore the problem was not detected. When the embrittled plastic failed during storage, air reached the inner vessel, and a hydride-catalyzed reaction of the oxygen and nitrogen with the hydride-coated plutonium ensued at a throttled rate determined by access of air to the reaction zone. The resulting expansion of the solid led to the complete failure of the defective weld. The corrosion rate became rapid during exposure to the glove box atmosphere and was estimated from the dramatic increase in the vessel’s volume during the 3-hour period, the bulk density of the corrosion product, and the approximate surface area of the metal. The result (0.3 g Pu/cm² min) is in excellent agreement with the value measured during studies of hydride-catalyzed corrosion (0.7 ± 0.1 g Pu/cm² min).

The experience and knowledge we gained from evaluating this incident and from studying catalyzed corrosion reactions find broad application in reducing the nuclear danger. The safe storage of plutonium requires using two metal containers with certified seals and ensuring that organic materials are excluded from the package. These two safety criteria are incorporated in the plutonium storage standards adopted by the U.S. Department of Energy and the International Atomic Energy Agency.

Knowledge of plutonium corrosion kinetics and particle-size distributions for a whole range of conditions enables realistic hazard assessments of plutonium dispersal during accidents. Catalyzed hydriding is also the key reaction in the hydride-dehydride recycle process for destroying weapon configurations and recovering plutonium metal—both done in a single step. Hydride-dehydride recycling generates no mixed or liquid waste of any kind. The recovered plutonium is a storage-ready ingot, and the hydrogen gas, which is both created and recycled in a uranium hydride storage system, is never released and does not present an explosion hazard. This process was recognized by the R&D Magazine as one of the best 100 technical innovations of the year and is the foundation technology of the Automated Retirement and Integrated Extraction System (ARIES) for separating plutonium from weapon components. The hydride-catalyzed oxidation of plutonium holds potential for development as a parallel single-step process in the rapid recovery of plutonium as oxide rather than metal.

Failed Storage Container for Plutonium Metal
Each photo shows a step in the disassembly of the container for a 2.5-kg casting of plutonium metal. (a) The outer container is placed in a glove box; (b) when the lid is removed, the plastic bagging shows; (c) when the plastic bagging and inner cylinder are lifted out of the container, discolored, embrittled plastic decomposed by radiolysis is revealed; (d) a rupture in the inner container is visible; (e) the corrosion product pours out of the ruptured end; and (f) an increase in the inner vessel’s diameter near the ruptured end shows the extent of hydride-catalyzed corrosion during a 3-hour period.
**Moisture-Enhanced Oxidation and the Role of Higher Oxides**

The moisture-enhanced corrosion of plutonium in air was first observed during the Manhattan Project, but the mechanism of this process remained a mystery until very recently (Haschke et al. 1996, Haschke and Martz 1998b). We shall first outline that mystery.

At room temperature, the corrosion rate of plutonium by water vapor at its equilibrium vapor pressure is more than 100 times faster than the corrosion rate in dry air. At 100°C, the two rates differ by almost $10^5$. Early work showed that exposure of plutonium to both molecular oxygen and water vapor results in the formation of PuO$_2$.

$$\text{Pu(s)} + \text{O}_2(g) \rightarrow \text{PuO}_2(s) \quad (8)$$

$$\text{Pu(s)} + 2\text{H}_2\text{O}(g) \rightarrow \text{PuO}_2 + 2\text{H}_2(g) \quad (9)$$

Moreover, corrosion rates for moisture-saturated air and moisture-saturated oxygen are equal to those for equilibrium water vapor, suggesting that the oxidation of plutonium metal in the presence of moisture is described by Equation (9). The situation is perplexing because O$_2$, not H$_2$O, disappears from the gas phase during corrosion, and H$_2$ is not observed as a gaseous product if oxygen is present. Similarly, uranium metal oxidizes faster in moist oxygen than in dry oxygen, but again O$_2$ disappears, and the concentration of H$_2$O remains constant, and H$_2$ is not released (Baker et al. 1966). However, after O$_2$ is depleted from the gas phase, the H$_2$O concentration decreases at a rate matched by the formation of H$_2$.

Early attempts to explain the moisture-enhanced oxidation of plutonium metal involved changes in the mechanism of that reaction: The usual O$_2$ diffusion was eliminated as the rate-limiting step of the oxidation process. Most proposed models assumed that the increased rate was due to the relatively fast diffusion of a hydroxide ion (OH$^-$) through the oxide layer. Those models suggested a possible explanation for the enhanced rate but did not account for the appearance of O$_2$ at the rapid rate characteristic of the Pu + H$_2$O reaction nor for the formation of oxide as the only corrosion product in moist air. If OH$^-$ were the diffusing species, an equimolar mixture of PuO$_2$ and PuH$_2$ would form at the product-metal interface.

We solved the mystery by a number of discoveries. Studying the plutonium-water reaction, we observed a previously unknown product and realized it was an oxide higher than PuO$_2$ (Stakebake et al. 1993). In separate studies of PuO$_2$ (Allen and Haschke 1999, Morales et al. 1999), we defined the chemistry of the PuO$_x$ + H$_2$O reaction, in which the higher oxide PuO$_{2+x}$ is formed. We also identified the role of this higher oxide in catalyzing the formation of H$_2$O from H$_2$ and O$_2$. We then realized that PuO$_{2+x}$ would always be found on the surface of plutonium metal in the presence of moisture, increasing the oxygen gradient across the steady-state oxide layer. It then became apparent that O$_2^-$ is indeed the diffusing species and that the rate of plutonium oxidation increases because a higher oxygen gradient increases the flux of O$_2^-$ transport through the layer.

**Plutonium Dioxide and Water Vapor**

We gleaned the key elements of this solution while trying to define safe conditions for the extended storage of surplus plutonium oxide in sealed containers (Haschke and Allen 1995). Our initial concern was the possible radiolysis of water adsorbed on the dioxide surface. Alpha radiation produced in plutonium decay might dissociate water into hydrogen and oxygen and thereby generate unacceptably high gas pressures within the containers. In one test for radiolysis, we exposed PuO$_2$ to saturated water vapor (0.032 bar) at 25°C. In another test, we exposed the oxide to an O$_2$ + H$_2$ mixture in a 1:2 molar ratio at 25°C and 0.2 bar of total pressure.

We measured kinetic data and identified products by chemical analysis. These measurements demonstrated that the dioxide surface is an active template for equilibration of the oxygen-hydrogen system. When the dioxide was exposed to the O$_2$ + H$_2$ mixture, both H$_2$ and O$_2$ were consumed, and H$_2$O was formed as the reaction product. The reaction pathway apparently involves adsorption and dissociation of the diatomic gases on the oxide surface—Equations (10) and (11)—followed by association of the atomic species as water—Equation (12).

$$\text{H}_2(g) = \text{H}_2(ads) = 2\text{H}(ads) \quad (10)$$

$$1/2 \text{O}_2(g) = 1/2 \text{O}_2(ads) = \text{O}(ads) \quad (11)$$

$$2\text{H}(ads) + \text{O}(ads) \rightarrow \text{H}_2\text{O}(ads) = \text{H}_2\text{O}(g) \quad (12)$$

$$\text{H}_2(g) + 1/2 \text{O}_2(g) \rightarrow \text{H}_2\text{O}(g) \quad (13)$$

We recently measured the rate of the net reaction defined by Equation (13). At 25°C, the production of water vapor in the presence of the oxide surface is approximately 6 micromoles per meter squared of oxide surface per day (Allen and Haschke 1999).

Molecular hydrogen appears when the dioxide is exposed to water vapor, but O$_2$ is not observed during these tests. Evidence for the radiolysis of water is absent. If, in fact, water is dissociated by alpha radiation, the O$_2$ and H$_2$ products are transient and immediately recombine on the catalytic oxide surface.

Our detection of hydrogen, but not oxygen, during exposure of the dioxide to water vapor is consistent with results of the earlier study (Stakebake et al. 1993), showing that an oxide with compositions substantially greater than PuO$_2$ forms at the gas-oxide interface during the reaction of plutonium with water vapor. The existence of the higher oxide was surprising and controversial because extensive studies by earlier workers...
PuO$_2$ does not react with oxygen, ozone, or nitrogen dioxide. The earlier workers concluded that PuO$_2$ is the highest oxide composition for plutonium. In contrast, our studies with the dioxide and water vapor at 25°C–350°C and a water vapor pressure of 0.025 ± 0.007 bar (20 ± 5 torr) demonstrate that water reacts with the dioxide to form a higher oxide (PuO$_{2+x}$) and hydrogen (Haschke and Allen 1999, Morales et al. 1999).

PuO$_2$(s) + x H$_2$O(l, g) → PuO$_{2+x}$(s) + x H$_2$(g)   .  \(14\)

We readily determined the reaction rate and oxide composition by quantifying the production of hydrogen over time.

On the basis of this work, we conclude that the PuO$_2$ + H$_2$O reaction in Equation (14) is a normal chemical process and is not promoted by radiolysis. The release of hydrogen from the oxide surface implies that the plutonium oxidation state in the product oxide is higher than Pu(IV). It also implies that mass increases observed by earlier workers during studies under humid conditions cannot be attributed solely to adsorption of water on the oxide surface. We measured a constant reaction rate of 6 nmol H$_2$/m$^2$ of oxide per day at room temperature and found that the rate of hydrogen production increases systematically with temperature. An Arrhenius analysis of the data yields an activation energy of 9.4 kcal/mol. Compositions in excess of PuO$_{2.25}$ are formed, but the maximum value of $x$ is unknown. In the box “PuO$_{2+x}$: The Stable Oxide of Plutonium in Air and the Environment,” we describe additional properties of PuO$_{2+x}$.

**Plutonium Dioxide and Moist Air.**

The studies of PuO$_2$ in moist air unequivocally demonstrate that, relative to PuO$_{2+x}$, PuO$_2$ is thermodynamically unstable in moist air over a substantial range of temperatures. Therefore, PuO$_2$ must be unstable in an atmosphere of oxygen, ozone, or nitrogen dioxide. The failure of early workers to obtain PuO$_{2+x}$ by exposing the dioxide to strong oxidants is undoubtedly a consequence of the very slow kinetics for the oxidation reaction. In contrast, according to Equation (14), the reaction of PuO$_2$ with H$_2$O occurs at a significant rate and is relatively easy to observe. Although we did not detect the formation of the higher oxide during exposure of the dioxide to dry oxygen, we did observe it when we added moisture to the system. Moreover, the behavior of the system was parallel to that encountered during the moisture-enhanced corrosion of plutonium metal: O$_2$ was consumed at the rate observed for the PuO$_2$ + H$_2$O reaction, but H$_2$ was not produced until oxygen was depleted.

Defining the steps in the net reaction between PuO$_2$ and moist air is essential for understanding the moisture-enhanced corrosion of plutonium metal. The observed interactions of PuO$_2$ with water and oxygen-hydrogen mixtures in air suggest that the reaction of PuO$_2$ with moist air proceeds via a catalytic cycle at the gas-solid interface described below and shown in Figure 8.

PuO$_2$(s) + x H$_2$O(ads) → PuO$_{2+x}$(s) + 2xH(ads)    .   \(15\)

2xH(ads) + xO(ads) → xH$_2$O(ads)    .  \(16\)

PuO$_2$(s) + (x/2)O$_2$(g) → PuO$_{2+x}$(s)   .  \(18\)

The reaction of adsorbed water with PuO$_2$ to form PuO$_{2+x}$ described by Equation (15) is faster than the

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Figure 8. Moisture-Enhanced Oxidation of Plutonium Dioxide

When PuO$_2$ is exposed to moist air, a cyclical reaction begins, in which a higher oxide of plutonium is formed. The cycle initiates by adsorption of water on the dioxide and its subsequent reaction to form a higher oxide and adsorbed hydrogen. The cycle continues as oxygen, adsorbed and dissociated on the oxide surface, is consumed either by reacting with the adsorbed hydrogen to reform water or by reacting with the oxide to increase its oxygen content. The amount of water remains constant until all the oxygen is consumed. The reaction of water with the oxide continues in the absence of oxygen until all the water is consumed. Hydrogen formed in this reaction is released as gas.
PuO$_{2+x}$: The Stable Oxide of Plutonium in Air and the Environment

Thermochemical predictions and results of extensive experiments between the 1940s and 1970s led researchers to conclude that plutonium oxides with compositions greater than PuO$_2$ are unstable and cannot be prepared. Recent results discussed in the main article show, however, that a hyperstoichiometric oxide (PuO$_{2+x}$) is the thermodynamically stable plutonium phase in air at temperatures between 25°C and 350°C. The PuO$_2$ + H$_2$O reaction produces the higher oxide and hydrogen, and the standard enthalpy and free energy for the formation of PuO$_{2+x}$ are more negative than –270 and –253 kilocalories per mole (kcal/mol), respectively. Corresponding values for the formation of PuO$_2$(s) are –252.4 and –238.5 kcal/mol. Although favored thermodynamically, the reactions of O$_2$ and other strong oxidants with PuO$_2$ are not observed because they occur extremely slowly. In contrast, the reaction of the dioxide with liquid or gaseous water is kinetically favorable at low temperatures and is easily monitored by measurement of the extent to which H$_2$ forms. Pressure-volume-temperature methods are used for those measurements.

Data from the x-ray diffraction of PuO$_{2+x}$ products obtained from kinetic studies reveal a fluorite-related face-centered-cubic (fcc) structure with lattice parameters near those of PuO$_2$. The graph plots the variation of the lattice parameter $a_0$ versus oxide composition. Below the dioxide stoichiometry (O/Pu = 2), $a_0$ decreases steeply with increasing oxygen content. Above the dioxide stoichiometry, the lattice parameter increases very slowly. The open triangles indicate reference data from literature sources. The open squares show data from the high-purity weapons-grade oxide used in our study. Filled and open circles show lattice parameters for products obtained during microbalance and pressure-volume-temperature (PVT) experiments with water vapor at 0.025 bar of pressure and temperatures in the range of 25°C to 350°C. The data point for the highest reported composition (O/Pu = 2.265) is from a literature source (Haschke 1992) describing PVT experiments of plutonium corrosion in aqueous salt solutions at 25°C. Molecular hydrogen was generated beyond the PuO$_2$ stoichiometry, but the experiment had been terminated before the maximum oxide composition was reached. An appropriate mass loss was observed when the PuO$_{2.265}$ product was heated at 500°C in dynamic vacuum until its mass became constant. The lattice parameter of the fired product (5.395 Å) corresponds to that of PuO$_2$.

The formation of PuO$_{2+x}$ may have far-reaching consequences. In addition to promoting the corrosion rate of plutonium in moist air, the reaction of H$_2$O with the oxide generates hydrogen pressures capable of rupturing sealed containers during extended storage of the oxide. For this reason, plutonium oxide must be fired before it is packaged in sealed containers for extended storage. This process removes water and makes the oxide less soluble. Firing eliminates the higher oxide and thus Pu(VI). The presence of U(VI) increases the solubility of the higher uranium oxide UO$_{2+x}$, and the absence of Pu(VI) is expected to reduce the dissolution rate.

The PuO$_{2+x}$ phase forms in conditions present in natural systems, implying that the higher oxide is the stable phase in the environment. Failure of x-ray diffraction measurements to identify a higher oxide is consistent with the insensitivity of $a_0$ to oxide composition. Recent x-ray absorption fine-structure, or XAFS, results indicate that only PuO$_2$ is present in the environment. Additional work is needed to resolve this discrepancy.
PuO$_2$ + O$_3$ reaction and determines the rate at which the higher oxide is formed. Atomic hydrogen produced by this reaction is not released as H$_2$ but remains on the oxide surface and recombines with O$_2$ to form water. According to Equation (16), adsorption and dissociation of O$_3$ on the dioxide surface, which we first identified in the oxygen-hydrogen-dioxide system, appear here as integral steps of the oxide-catalyzed reaction of H$_2$ with O$_2$ to form water. A water-catalyzed cycle is entered as H$_2$O from Equation (17) reacts with PuO$_{2+x}$ from Equation (15) to progressively increase the x of the higher oxide. Equation (18) gives the net reaction for the process.

The catalytic cycle defined by Equations (15)–(17) accounts for the unusual behavior encountered during the reaction of PuO$_2$ with moist air or moist oxygen. In water vapor, the rate at which PuO$_{2+x}$ forms is determined by the rate of Equation (14). In moist O$_2$, the higher oxide forms at the same rate but by following Equation (15). However, the hydrogen atoms formed by the reaction of H$_2$O molecules immediately recombine with oxygen atoms from O$_2$. Oxygen is consumed according to Equation (18) at the same rate as in the PuO$_2$ + H$_2$O reaction, but H$_2$ is not observed. The net reaction involves two catalytic processes acting in concert: first, H$_2$O acts as a reactive intermediate (or catalyst) in promoting the oxidation of PuO$_2$ by O$_2$, and second, the oxide surface acts as a catalyst in promoting the regeneration of H$_2$O.

We can now explain the unanticipated implosion of storage vessels containing plutonium oxide. Although the reaction of the oxide with residual water shown in Equation (14) is expected to pressurize containers with H$_2$, negative pressures are observed in cans containing unfired oxide that was originally packaged in air. As demonstrated in laboratory tests, the pressure decreases because of the moisture-catalyzed reaction of the oxide with oxygen according to Equations (15) to (17). However, reduced pressure is a transient condition. After the O$_2$ in the storage atmosphere has been depleted, the pressure rises as the reaction of the residual H$_2$O produces H$_2$.

### Plutonium Metal and Moist Air.
Likewise, the catalytic cycle in Figure 8 can explain the moisture-enhanced corrosion of plutonium metal because that cycle creates PuO$_{2+x}$ on the surface of the oxide-coated metal. The result is a gradient in the oxygen concentration across the oxide layer—from a higher concentration at the gas-oxide interface to a lower concentration at the oxide-metal interface. Earlier in this article, we described how the transport of oxygen from the surface to the oxide-metal interface determines the oxidation rate of plutonium in dry air. That rate of transport depends on three factors: the thickness of the coherent oxide layer, the temperature-dependent coefficient of oxygen diffusion in the oxide, and the gradient in oxygen concentration across the oxide layer. In the steady-state situation, when the temperature is fixed and the thickness of the oxide layer is constant because of continual spallation from the surface, the increase in the oxygen gradient produced by the formation of the higher oxide should increase the rate of oxygen transport to the oxide-metal interface and thereby the oxidation rate. In contrast, the higher oxide is not readily formed by dry air, and the lower rate of oxygen transport leads to a lower corrosion rate.

Thus, we find that the chemistry of plutonium corrosion in moist air or moist oxygen is consistent with a catalytic cycle similar to that proposed for the reaction of the dioxide with moist oxygen. The process is driven by the kinetically favored reaction of the metal with water defined by Equation (19).

\[
\text{Pu(s) + H}_2\text{O(ads) \rightarrow PuO}_2\text{(s) + 2H(ads) .} \tag{19}
\]

\[
(1/2)\text{O}_2\text{(g) = O(ads) .} \tag{20}
\]

\[
2\text{H(ads) + O(ads) \rightarrow H}_2\text{O(ads) .} \tag{21}
\]

\[
\text{Pu(s) + (1/2)O}_2\text{(g) \rightarrow PuO}_2\text{(s) .} \tag{22}
\]

Instead of forming H$_2$ by Equation (9), as shown for water vapor, the product hydrogen from the Pu + H$_2$O reaction combines with oxygen to regenerate water on the oxide surface. As described by the net reaction in Equation (22), O$_2$ and plutonium are consumed at the rapid rate characteristic of the Pu + H$_2$O reaction. Retention of hydrogen at the surface implies that O$_2^-$ is the diffusing species in the oxide layer and that the rate is controlled by oxygen diffusion through the oxide layer, as it is in the reaction of plutonium with dry air.

In summary, the moisture-enhanced corrosion of plutonium in air apparently proceeds via a complex catalytic cycle in which plutonium oxide with a high stoichiometry is formed. Enhanced corrosion of plutonium would also occur if both H$_2$ and O$_2$ were present because the oxide surface would catalyze the formation of H$_2$O. Note that hydrogen often forms gradually through the thermal or radiolytic decomposition of organic materials. As shown in Figure 2, the effects of moisture on plutonium corrosion are evident at temperatures of up to 400°C.

### Conclusions and Future Directions

Almost 60 years after it was first made in the laboratory, plutonium is still yielding surprises. We have observed unanticipated reactions and new compounds in various binary and ternary combinations of plutonium, oxygen, and hydrogen. We have also found that the reaction kinetics of those systems is extremely important in determining the reaction products. Our understanding of the chemistry of those systems seems surprisingly inadequate.

Although not reported in this article, we have also observed that the salt-
catalyzed corrosion of plutonium metal by liquid water forms a rich variety of products including oxide hydrides, Pu$_2$O$_3$, mixed-valence oxides of Pu(III) and Pu(IV), and PuO$_2$. The higher oxide PuO$_{2+x}$ forms as well, but it was identified only recently. Like the catalytic compounds Pu$_2$O$_3$, PuH$_x$, and PuO$_{2+x}$ discussed in this article, the products obtained in aqueous media apparently have fluorite-related structures. We wonder about the importance of such materials in determining the solid-state chemistry of plutonium. More work is needed to define corrosion processes and products. We must investigate structures, thermodynamics, conductivities, transport processes, and other fundamental properties of plutonium compounds to understand how they influence the corrosion chemistry of plutonium.

We are intrigued by the consistent appearance of catalytic activity during plutonium corrosion. Although we have demonstrated the effects of surface chemistry and catalytic materials on bulk corrosion kinetics for several systems, studies are far from exhaustive. We must investigate catalytic behavior at the atomic level to augment our understanding and interpretation of kinetic results and other macroscopic observations. The following are some important research efforts in this regard: characterization of adsorption processes and adsorbates, definition of surface reactions at the atomic level, and determination of relationships between a solid's catalytic activity and electronic structure.

Of particular relevance are our continuing efforts to explore moisture-enhanced plutonium oxidation and to define the phase equilibria and thermodynamic properties of PuO$_{2+x}$, the higher plutonium oxide. By measuring the dissolution rates and solubilities of PuO$_{2+x}$ in aqueous media, we hope to understand the role of PuO$_{2+x}$ in plutonium transport. Formation of PuO$_{2+x}$ and leaching of the hexavalent ion are consistent with the appearance of Pu(V) and Pu(VI) in aqueous media.

Another area in which the new findings on plutonium corrosion are indispensable is estimating dispersal hazards during accidents involving nuclear weapons or surplus nuclear material. Metallic plutonium is essentially nondispersible; it therefore presents very little risk of being released into the environment during an accident. In contrast, fine plutonium-containing particles (less than 10 micrometers in diameter) produced by corrosion are at much greater risk of being dispersed. When the solid expands or when gas forms during corrosion, pressures are generated that are known to rupture storage containers and release radioactive materials (Haschke and Martz 1998b). Moreover, mixtures containing hydrogen can form and explode, thus posing additional hazards. To a large extent, this potential hazard depends on the corrosion rate, size distribution of product particles, and time of corrosion, a period that may be as short as hours for accidents or as long as decades for storage.

We conclude that surface compounds are extremely important in determining the course and kinetics of plutonium corrosion and that both the chemical history and external conditions determine the chemical nature of the metal surface. We have been able to identify and characterize surface compounds and determine their effects on the reaction kinetics of plutonium metal and plutonium compounds. But the implications of our recent findings on plutonium technology have yet to be assessed. Certainly, our results are relevant to plutonium recovery, reprocessing, storage, accident assessment, and environmental migration. Continued investigation is therefore a must.

Further Reading


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**John M. Haschke** received his undergraduate degree from Texas Tech University, did postgraduate work as a Fulbright Fellow at Johannes Gutenberg University in Mainz, Germany, and obtained his Ph.D. degree from Michigan State University. His thesis was on the equilibrium and thermodynamic properties of lanthanide compounds at high temperatures. After working as a postdoctoral fellow at Arizona State University, Haschke joined the faculty in the Department of Chemistry at the University of Michigan. Haschke began his career in actinide chemistry as a research scientist for Rockwell International at the Rocky Flats Plant and developed his skills as a visiting scientist at the Institute for Transuranium Elements at Karlsruhe, Germany, before becoming a staff member at Los Alamos National Laboratory. His research interests have focused on fundamental aspects of plutonium corrosion chemistry and on their application to enhancing plutonium process operations, ensuring safe storage of surplus nuclear material, assessing hazards, and maintaining the nuclear stockpile. Haschke has published more than 150 technical papers and has authored several review chapters on lanthanide and actinide chemistry. He is currently working as an independent consultant on plutonium chemistry.