Analysis Techniques for Tracer Studies of Oxidation

Soumendra Nath Basu
Case Western Reserve University
Cleveland, Ohio

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CHAPTER 1

INTRODUCTION

The double oxidation technique using O-18 as a tracer has been successfully used to identify the transport mechanisms in growing oxide scales. The detailed mechanism of oxidation is of more than academic interest as it determines growth stresses and oxide spallation. The aim of this study was to develop analysis techniques to obtain quantitative diffusion data from the tracer concentration profiles in growing oxide scales. An oxide can grow inward at the metal-oxide interface by inward oxygen diffusion; it can grow outward at the gas oxide interface by outward cation diffusion or it can grow by a combination of both the mechanisms. The problem of determining the fraction of inward and outward growth of the scale from the tracer concentration profile was solved. The process of tracer diffusion in a growing polycrystalline scale was theoretically modelled.
The analysis techniques developed were then applied to a FeCrAl-Zr alloy by using SIMS to obtain the tracer profile.
CHAPTER 2

LITERATURE SURVEY

2.1 Mathematics of diffusion

2.1.1 Diffusion

Atoms in a solid are in constant motion about their lattice sites. Occasionally, a few atoms gain sufficient energy to jump into neighbouring sites. This random movement of atoms is called diffusion. Self diffusion is the phenomenon of migration of atoms in the absence of concentration gradients. Self diffusion can be studied by introducing an isotopic tracer into the system such that the diffusing atoms can be identified. Strictly speaking the diffusion coefficients of isotopes are related by the equation:

\[
\frac{D_2}{D_1} = \left( \frac{M_2}{M_1} \right)^{1/2}
\]  

...(2.1)

Where \( M \) and \( M_1 \) are the masses and \( D \) and \( D_1 \) are the diffusion coefficients of the isotopes. However, generally the isotopic masses are so close to each other that the diffusion coefficients are assumed to be the same. When O-18 is used as a tracer to study oxygen diffusion, the difference
is about 6%.

2.1.2 Fick’s laws

Matter flows in a manner to decrease its concentration gradient or more precisely its chemical potential gradient. The flux density, $J_i$ (kg/m sec), of the species $i$ in 3 dimensions is given by:

$$J_i = -D_i \rho \nabla C_i$$  \hspace{1cm} ...(2.2)

Where $D_i$ is the diffusion coefficient (m$^2$/sec), $\rho$ is the density (kg/m$^3$) and $C_i$ is the mass fractional concentration gradient (m$^{-1}$). This is Fick’s first law. In 1-dimension Equation 2.2 becomes:

$$J_i = -D_i \rho \frac{\partial C_i}{\partial x}$$  \hspace{1cm} ...(2.3)

From Equation 2.3 and using the principle of mass conservation [1] it can be shown that:

$$\frac{\partial C_i}{\partial t} = \frac{\partial}{\partial x} \left(D_i \frac{\partial C_i}{\partial x} \right)$$  \hspace{1cm} ...(2.4)

where $t$ is the time. This is Fick’s second law.

If $D_i$ is independent of position, Equation 2.4 becomes:

$$\frac{\partial C_i}{\partial t} = D_i \frac{\partial^2 C_i}{\partial x^2}$$  \hspace{1cm} ...(2.5)

2.1.3 Diffusion mechanisms
Diffusion generally occurs by a vacancy or an interstitial mechanism. An atom adjacent to a vacancy can exchange sites with the vacancy and thus migrate through the crystal. Some atoms occupy positions in the interstices of the crystal lattice. They can move either by jumping from an interstitial site to another (interstitial mechanism) or by displacing a lattice atom into an interstitial position and taking up the lattice site itself (interstitialcy mechanism).

The predominant point defects in an ionic crystal are [2]:

i) Schottky disorder- Stoichiometric proportion of cations and anion vacancies.

ii) Frenkel disorder- Equal number of cation vacancies and interstitials.

iii) Anti Frenkel disorder- Equal number of anion interstitials and vacancies.

iv) Anti Schottky disorder- Stoichiometric proportion of anion and cation interstitials.
It should be noted that the experimental diffusion coefficient can be represented by an Arrhenius type equation:

\[ D = D_0 \exp \left( -\frac{Q}{RT} \right) \]  \hspace{1cm} \text{(...)2.6)}

where \( D \) is the preexponential term (m$^2$/sec) and \( Q_0 \) is the activation energy (kJ/mole).

2.2 Grain boundary diffusion

2.2.1. Introduction

Enhanced grain boundary diffusion has been observed in several oxides [3-4]. The observed tracer concentration profile in a polycrystalline sample is due to both fast diffusion along the grain boundaries and slower diffusion into the grains from the surface and the grain boundaries. Some of the models proposed to mathematically treat the combined grain boundary and volume diffusion problem are discussed in the next section.

2.2.2 Mathematical treatment
Fisher [5] modelled a grain boundary as a thin slab of thickness \( \omega \) extending into a semi-infinite crystal (Figure 1). The diffusion coefficient \( D_b \) in the grain boundary is much greater than the diffusion coefficient in the crystal. Whipple [6] used the Fisher model and obtained exact solutions of the average tracer concentration profile using Fourier – Laplace transforms. He used a constant source condition, i.e.,

\[ C(x, y, t) = C \quad \text{for } t > 0 \text{ and } y = 0 \]

He obtained that the logarithm of the average tracer concentration (\( \ln \bar{C} \)) varies linearly with \( y \) with a slope that can be related to the grain boundary diffusion coefficient by the equation:

\[ D_b \omega = \left( \frac{\ln \bar{C}}{y^{6/5}} \right) \left( \frac{4D}{t} \right) \left( \frac{\ln \bar{C}}{(n^2 - 1/2) 6/5} \right)^{5/3} \]

\[ \ldots (2.7) \]

where:

\[ \eta = \frac{y}{\sqrt{Dt}} \quad , \quad \beta = \frac{(\Delta - 1) \omega}{\sqrt{D}} \quad , \quad \Delta = \frac{D_b}{D} \]

Levine and MacCallum [7] solved the grain boundary diffusion equation directly for the average concentration in polycrystalline samples.
FIGURE 1 - Fisher's model of a grain boundary.
and found that for $\theta >10$ the last term in Equation 2.7 is independent of $\eta^{1/2}$ and is equal to 0.78. This was later confirmed for Whipple's solution by LeClaire [8].

Suzuoka [9] solved the same problem, but assumed an instantaneous source condition ie.,

$$C(x, y, t) = K \delta(y) \quad x=0, \ t=0$$

where $K$ is the surface density of the planer source. His solution also predicted a linear plot between $\ln C$ and $y$ with:

$$\frac{\partial \ln C}{\partial (\eta^{1/2})} = 0.75$$

Thus to obtain the value of $D \omega$, experimental $b$ values of $\ln C$ can be plotted against $y$. A straight line can be fitted to the data points of deeper penetrations. The slope of this line can be used to determine the value of $D \omega$ using Equation 2.7.

Zaeschmar [10] treated the grain boundary diffusion problem by considering the polycrystal as a 3-dimensional brickwork of cylindrical grains. He introduced another empirical parameter - the filling time of a grain. Oishi and Ichimura [11] treated the problem as the homogeneous filling of a grain pinned to the grain boundary.
at an arbitrary distance from the surface. As this author has used the Oishi-Ichimura geometrical configuration for modelling, it will be described in detail later.

2.3 Previous studies of oxygen diffusion in alumina

2.3.1. Lattice diffusivity

Oishi and Kinsey [12] studied oxygen diffusion in single crystal alumina using gaseous exchange technique. Their plot of \( \log(D) \) versus \( 10^{\frac{10}{\text{RT}}} \) showed a break at 1650 °C which was interpreted as an extrinsic to intrinsic transition. The high temperature data showed that:

\[
D = 1.9 \times 10^{-12} \exp\left(-\frac{636 \text{ kJ/mole}}{RT}\right) \text{ m}^2/\text{sec} \quad (2.8)
\]

while the fit to the low temperature data suggested that:

\[
D = 6.3 \times 10^{-12} \exp\left(-\frac{241 \text{ kJ/mole}}{RT}\right) \text{ m}^2/\text{sec} \quad (2.9)
\]

This was later shown to be an experimental artifact sensitive to the surface preparation of the specimen. When the samples in the low temperature region were preannealed before the diffusion anneal, the measured diffusion coefficients fell on the extrapolation of the high temperature data.
Oishi, Ando and Matsuhiro [13] studied high purity vapour grown crystals using the same method as Oishi and Kingery. Their results were in approximate agreement with Oishi and Kingery’s results.

Reed and Weunsch [14] studied single crystal alumina using ion probe. Their data suggest that:

\[ D = 6.4 \exp\left(\frac{-787 \text{kJ/mole}}{RT}\right) \text{m}^2/\text{sec} \quad (2.10) \]

Reddy and Cooper [15] exchanged single crystal alumina in enriched O-18 gas and measured the tracer concentration profile using proton activation. Their data fit the equation:

\[ D = 2.66 \times 10^{-2} \exp\left(\frac{-615 \text{kJ/mole}}{RT}\right) \text{m}^2/\text{sec} \quad (2.11) \]

Cawley, Halloran and Cooper [16] exchanged single crystal alumina in Reddy’s gas exchange apparatus. The tracer profiles were determined by both proton activation and Secondary Ion Mass Spectrometry (SIMS). Cawley found that the diffusion coefficient of oxygen was insensitive to impurity levels and oxygen partial pressure and could be described by:

\[ D = 1.51 \times 10^{-3} \exp\left(\frac{-527 \text{kJ/mole}}{RT}\right) \text{m}^2/\text{sec} \quad (2.12) \]
Recently Laserlof, Heuer and Mitchell studied the annihilation of dislocation loops by annealing deformed cylinders of Al O . Based on the assumption that the self diffusion of oxygen was rate controlling they calculated the diffusion coefficient of oxygen in undoped alumina as:

\[
D = 6.8 \times 10^{-4} \exp\left(-\frac{587 \text{ kJ/mole}}{RT}\right) \text{m}^2/\text{sec} \quad \ldots (2.13)
\]

It should be noted that the data of Oishi and Ando, Reddy and Cooper, Cawley, Halloran and Cooper, and Laserlof and Heuer, taken by using different techniques, agree well in the temperature range of 1250 to 1800°C. So the lattice diffusion coefficient of oxygen in pure alumina is well established.

2.3.2 Grain boundary diffusivity

Mistler and Coble [17] calculated grain boundary diffusion coefficients and widths from the results of grain growth, creep, sintering and diffusion. Mathematical models of sintering and creep gave \(D_b\) while grain growth models were assumed to give \(D/W_b\). Thus \(D\) and \(W\) were calculated separately. The grain boundary width for alumina was reported to be between 9-12 nm.
while the strain boundary diffusion coefficient was reported as:

\[ D_b = 5.15 \times 10^{-4} \exp\left(-\frac{451 \text{ kJ/mole}}{RT}\right) \text{ m}^2/\text{sec} \quad \text{(2.14)} \]

There is a lot of uncertainty in these results as the effect of impurities on strain growth was neglected. The model assumed an effective diffusion coefficient \( D_{\text{eff}} \) given by:

\[ D_{\text{eff}} = \varepsilon D_b + (1-\varepsilon)D \]

where \( \varepsilon \) is the volume fraction of the strain boundaries. The above expression does not take into consideration the diffusion from the strain boundaries into the grains.

Johnson and Berrin [18] used initial stage sintering studies to obtain:

\[ D_{b0} = 5.2 \times 10^{-4} \exp\left(-\frac{653 \text{ kJ/mole}}{RT}\right) \text{ m}^3/\text{sec} \quad \text{(2.15)} \]

However their model neglected the contribution of surface diffusion to the neck growth.

Lessins and Gordon [19] used creep results of iron doped alumina samples to obtain:

\[ D_{b0} = 4.38 \times 10^{-7} \exp\left(-\frac{439 \text{ kJ/mole}}{RT}\right) \text{ m}^3/\text{sec} \quad \text{(2.16)} \]

However they had to guess the rate controlling diffusion process. At large concentrations of divalent iron and large grain size the creep was
assumed to be limited by oxygen grain boundary diffusion.

Reddy and Cooper [15] used proton activation on undoped polycrystalline alumina to obtain:

$$D_b = 4.4 \times 10^{-3} \exp(-690 \text{KJ/mole} / RT) \text{ m}^2/\text{sec} ...(2.17)$$

The boundary widths assumed were typically 15 to 40 nm. This assumption is too large as the grain boundary width of alumina is typically closer to 10 Å. There is thus a great degree of disagreement between the reported values of grain boundary diffusivity and boundary width.

2.4 Oxidation of superalloys

2.4.1 Superalloys

The need to develop materials for high temperature applications like gas turbine engines led to the development of superalloys [20]. MCrAl alloys (M=Fe,Ni,Co) are promising for use up to 1200°C. The high strength of these alloys is achieved by solid solution and precipitation hardening while good oxidation resistance is obtained by the formation of adherent protective oxide layers. These alloys owe their oxidation resistance...

* Personal communications with N.L. Peterson of Argonne National Laboratories.
resistance to the formation of an alumina scale since it is the most stable oxide and has the lowest free energy of formation. Alumina is a stoichiometric oxide with low diffusion rates and is not attacked by either \( \text{H}_2 \) or \( \text{CO} \). Giddins and Pettit [21] showed that the external alumina scale forms for certain compositions of the alloy only. The amount of aluminum needed to form a continuous scale can be reduced by increasing the chromium content as chromium acts as an oxygen getter.

2.4.2. Transient scales

Though the superalloys owe their oxidation resistance to the formation of a protective alumina scale, it has been found that initially transient oxides are formed. Smialek and Gibala [22] studied the structure of transient scales formed on pure, \( \text{Y} \) doped and \( \text{Zr} \) doped \( \text{NiCrAl} \) alloys. Oxidation of these alloys for 0.1 hour at 1100°C produced oriented \( \gamma - \text{Al}_2 \text{O}_3 \), \( \alpha - (\text{Al},\text{Cr})_2 \text{O}_3 \) or a spinel like \( \text{Ni}((\text{Cr},\text{Al})_2 \text{O}_4 \) before the random \( \alpha - \text{Al}_2 \text{O}_3 \) layer was formed. The scales were densely populated with internal precipitates and there was evidence of plastic flow in the underlying metal due to growth.
stresses. The formation of coherent layers of aluminium depleted phases indicated the selective removal of aluminium at these short times.

Golightly, Wood and Stott[23] studied the oxidation of FeCrAl alloys with various Y additions at 1100°C and 1200°C. They observed the formation of transient oxides rich in iron and chromium in all cases which was followed by the steady state α-alumina scale. For the Y free alloy it was found that the steady state was reached faster if the chromium content or the temperature was increased. The amount of transient oxide formed was also determined by the specimen surface topography since the development of the α-alumina layer was less rapid at the base of the alloy asperities than at a flat alloy oxide interface.

2.4.3 Mechanism Of Oxidation

Inert marker experiments by Hindam and Smeltzer [24] indicated that alumina scales grow inward by oxygen diffusion. Recent O-18 tracer studies by Reddy [25] have also confirmed that the
Primary transport is oxygen inward and the profiles are most consistent with the grain boundary transport model.

Golightly et al. [26] observed that the growth morphology of alumina scales on FeCrAl alloys was different for Y free and Y containing alloys. They suggested that in the Y free alloy, the scale grew by a combined inward oxygen and outward aluminum transport mechanism which caused lateral growth within the scale. This was responsible for the highly stressed oxide on the Y free alloys. The absence of growth stresses and lateral growth in scales with Y was attributed to the blocking of outward aluminum diffusion.

Whittle and Stringer [27] proposed a nucleation model to explain the 'Rare earth effect'. They suggested that the rare earth additions acted as nucleation sites for all the oxide phases, thus producing a finer grained initial oxide. This caused the steady state oxidation to be reached more rapidly and at a lower aluminum or chromium content.
2.4.4 Oxide adherence

As discussed earlier, the introduction of small amounts of rare earth elements to the superalloys caused a dramatic improvement in the scale adherence. Some of the mechanisms put forward to explain this phenomenon are as follows [27,28]:

a) Y (or other rare earth elements) improve the chemical bond at the metal-oxide interface as they have a better affinity for oxygen than the alloying elements.

b) Yttride precipitates or yttria particles can induce mechanical pressings of the oxide scale on the substrate.

c) Y promotes the formation of an inner yttria layer which acts as a diffusion barrier and also decreases the mechanical and thermal stresses between the oxide and the alloy.

d) Soluble Y in the alloy can trap the metal vacancies and consequently avoid vacancy coalescence at the metal oxide interface.
e) Y present in the oxide scale can influence oxide defect mobility and consequently the diffusion rate in the oxide scale, oxide thickness and plasticity.

Delaunay et al. studied the mechanical stresses generated in the oxide scale during the oxidation of superalloys. They concluded that Y improves the oxide adherence by trapping the vacancies and preventing them from precipitating at the metal oxide interface [29]. It may also improve the oxide plasticity by increasing the concentration of oxygen vacancies [30].

2.4.5 Growth kinetics and microstructure of FeCrAl-Zr alloy

Smialek studied the microstructure of the oxide grown on the FeCrAl-Zr alloy used by this author. He saw monoclinic zirconia precipitates at the grain boundaries at the gas surface. There were deep depressions in the large grains at the metal-oxide interface which represented areas where the grain boundary regions had advanced faster than the interior of the grains due to
short circuit diffusion \[31,32,33\]. At \(1100\ C\), the growth rate of the oxide was found to obey the law:

\[
0.46 \\
X = (K t)^P
\]

\[\text{...(2.18)}\]

Where \(X\) is the oxide thickness and \(t\) is the oxidation time. The grain growth in the alumina scale was found to obey the law:

\[
0.2 \\
\text{Grain size} = (K t)^G
\]

\[\text{...(2.19)}\]

There was a variation of grain size and shape across the scale. The grains near the gas surface were spherical and small in size while the grains near the oxide-metal interface were larger in size and columnar in shape.

2.5. Previous O-18 studies

O-18 has been successfully used as a tracer to identify the growth mechanisms of oxide scales. The double oxidation technique consists of successive oxidation of the alloy in natural oxygen and O-18 tracer. The concentration profile of the tracer is determined by some convenient technique like proton activation or SIMS. The tracer can then be used to identify the regions of new growth. The shape of the tracer concentration...
Profile can provide insight into the mechanism of oxide growth.

O-18 has been successfully used as a tracer by many workers. Lees and Calvert [34] used the double oxidation technique on FeCr alloys in 1976. In 1978, Sheasby and Brown [35] used SIMS to obtain O-18 concentration profiles in CoO scales. In 1980, Hartley et al. [36] at Harwell oxidized 18 FeCr alloys in a CO atmosphere. They used proton activation to obtain the tracer concentration profiles. In 1982 Lees et al. studied the mechanism of corrosion of low silicon [37] and high silicon [38] ferritic steels in high pressure CO using O-18 tracer.

Reddy [15] used the double oxidation technique to measure oxygen diffusivities in various oxides. He used the proton activation technique to measure the tracer concentration profiles. Reddy also studied the oxidation mechanism of NiCrAl alloys by the double oxidation technique. Very recently Cawley studied the self-diffusion coefficient of oxygen in alumina using O-18 tracer. He determined the tracer profiles by both proton
activation and SIMS. Cawley also studied the thermal oxidation of silica and silicon carbide using the same technique. He was able to obtain good fits of his data to the theoretical calculations if he assumed different diffusion coefficients for his near surface and deep profiles.

2.6. SIMS

In SIMS [39], the sample is bombarded with high energy primary ions (Ar in this case). These primary ions penetrate into the solid where binary collisions with target atoms occur. The primary atoms come to rest at the so-called penetration depth and are thus implanted in the solid. These primary ions cause the target atoms to be displaced from the surface due to various generation target recoils. These target or secondary ions are passed through a mass analyzer where they are separated by mass to charge ratios. An electron beam is also focussed on the sample to keep it neutral.
The number of sputtered secondary ions of a given type is a complex function of the surface concentration, electronic structure and sputtering conditions. However, as all the variables for isotopes are almost the same, the ratio of counts of isotopic ions gives the true value of isotopic concentration.

The resolution of SIMS is limited by non-uniform sputter etching which may yield sputtered species from a range of depths [40,41]. This problem is particularly severe at the edge of the crater. To avoid this problem the raster sating technique is used where the beam scans over a large area and detects secondary ions from the central portion of the crater only. Another problem faced when sputtering insulators is that of charging due to ion removal. If the charging is stable it can attenuate the secondary ion signal [42]. The electron beam can neutralize the secondary ions, causing further complications.
The advantage of using SIMS over proton activation is its depth resolution which is less than 10A. Also there is no limit on the depth of the sputtered crater except for the machine time. SIMS also provides the data in a very convenient form and can monitor up to 9 elements at a time.

However, the problems of non uniform sputtering, especially in polycrystalline samples can severely limit the resolution of SIMS and can actually make the data worthless. This point will be discussed in detail later. The conversion of counts to concentration with the help of standards can be extremely unreliable as the ion yield depends on, among other things, the sample surface preparation, the incident angles of the ions and the sample microstructure. So it is extremely important to apply stringent tests on the quality of data before any quantitative information is obtained from it.
3.1 Statement of the Problem

O-18 has been successfully used as a tracer to identify the growth mechanisms in oxide scales. The double oxidation technique is used to identify the diffusion processes involved. In this technique the alloy is sequentially oxidized, first in natural oxygen and then in O-18 tracer. The oxide grown during the natural oxygen oxidation is referred to as the old oxide while the oxide grown during the tracer oxidation is referred to as the new oxide. The position of the new oxide growth is determined from the concentration profile of the tracer across the oxide scale. The tracer concentration profile is obtained by SIMS.

It has been shown that the primary transport mechanism operative during the growth of α-alumina scales on superalloys is inward grain boundary diffusion of oxygen [15,32]. Thus the protective oxide scale has been found to be primarily inward.
 growing. However, as the oxygen tracer diffuses down the grain boundaries, it exchanges with O-16 existing in the grains. To get any quantitative information from the tracer concentration profile, it is necessary to model this process of inward tracer grain boundary diffusion with exchange. The advantage of a quantitative model is that the grain boundary and the lattice diffusion coefficients can be obtained from the shape of the tracer concentration profile. The effect of structural parameters like the oxide grain size on the tracer profile can also be studied. It can also be used to determine if the scale is inward or outward growing. In the case where scale growth is a combination of both the mechanisms, a simple mass balance test can be used to separate the two processes. To model the diffusion process, Oishi and Ichimura's geometrical configuration of the grain boundaries and grains have been used.

In the model, the shape of the tracer concentration profile depends on the three parameters:

1) the grain size \( r \), 2) the grain boundary
diffusion coefficient $D$ and 3) the lattice diffusion coefficient $D$. By choosing the appropriate values of the three parameters, a family of curves can be generated which define a range of O-18 profiles which are quantitatively compatible with the inward oxygen diffusion mechanism. Conversely, given an experimental tracer concentration profile, it should be possible to pin down the values of the three parameters. It should be noted that the grain size of the oxide scale can be obtained by SEM studies. A good estimate of the grain boundary diffusion coefficient can be obtained from the growth kinetics of the oxide while values of the lattice diffusion coefficient can be obtained from the literature. Thus there is independent information existing for all the three model parameters. A major assumption made during the modelling of the diffusion process is that the oxide scale grows in a planar fashion.

3.2 Expected Tracer Profiles
Figure 2 shows the schematic O-18 concentration profiles for various transport mechanisms. If the outward aluminum transport is the primary mechanism the scale will be outward growing. The tracer concentration profile will be a step function due to the formation of $\text{Al}_0$ at the gas-oxide interface. If the transport mechanism is inward oxygen grain boundary transport, the scale will be inward growing and the concentration profile will be a step function due to the formation of $\text{Al}_0$ at the metal-oxide interface. This is however an ideal case. When the tracer in the grain boundaries comes in contact with O-16 in the grains of the old scale, exchange will take place and the concentration profile will look like Figure 2c). Finally if the diffusion mechanism is a combination of outward aluminium and inward oxygen transport the tracer profile would look like Figure 2d).

3.3 Mass Balance

Given a tracer concentration profile across the oxide scale a simple mass balance test can be used to determine the growth mechanism. As shown
FIGURE 2 - Schematic tracer concentration profiles for various transport mechanisms.
in Figure 3, a normalized concentration profile can be plotted such that the X-axis is a plot of the fraction of the total scale thickness while the Y-axis is the plot of the concentration of the tracer as a fraction of the maximum tracer concentration in the scale.

We know that every atom of O-18 in the oxide scale got there when the sample was exposed to the tracer. Thus the number of lattice sites occupied by the tracer atoms in the oxide scale must be the same as the number of oxygen lattice sites created in the oxide during the tracer oxidation. This concept can be used to calculate the thickness of the new oxide $Y_{\text{new}}$ as shown in Figure 3b). $Y$ is so chosen, that area A in new Figure 3a) (amount of O-18 in the total oxide) is equal to the area B in Figure 3b) (amount of oxygen in the new oxide). Having established the new oxide thickness, it can be determined if the growth mechanism is primarily inward oxygen transport or a combination of outward aluminum and inward oxygen transport. If the diffusion mechanism is inward transport of oxygen through the grain boundaries with exchange, the scale is inward growing. Then
FIGURE 3 - Determination of transport mechanisms by mass balance.
the number of 0-18 atoms retained in the old oxide must be the same as the number of 0-16 atoms found in the new oxide as they are both involved in the same exchange process. Thus if area C is equal to the area D in Figure 3c) the transport mechanism is inward oxygen diffusion. However, if area C is greater than area D some of the tracer atoms in C did not set there due to exchange but rather due to outward aluminium diffusion leading to the formation of Al\textsubscript{2}O\textsubscript{3} at the gas-oxide interface. In this case, it is possible to determine, at least in principle, what fraction of the scale growth is due to each mechanism. Thus Y can be split up into in and out new Y and Y such that area E=area F in Figure 3d). This allows the growth kinetics to be split up into inward growth and outward growth and both these can be separately analysed. So with profiles of sufficient quality, one can obtain detailed information by simple mass balance analysis. It should be noted that this is model free.

It should be mentioned that the mass balance criterion is valid if the number of tracer atoms in the grains at the oxide-gas interface due to direct surface exchange with the gas is neglected. This
is a valid assumption because the grain boundary diffusivity is much higher than the lattice diffusivity in the oxide and the penetration depth of the tracer atoms due to volume diffusion at the surface is negligible.

Much more information can be obtained from the actual shape of the tracer concentration profile, if the actual equations describing tracer incorporation during scale growth could be directly solved. Even in the simple case of inward oxygen diffusion it is a formidable problem of combined lattice and grain boundary diffusion with a moving boundary under an imposed field. However a relatively simple but still realistic model permits the diffusion equations to be solved.

3.4 Geometrical Configuration

Figure 4 shows an idealised drawing of the old scale with hexagonal grains arranged in a close packed two dimensional network. These grains are formed during the O-16 oxidation and contain natural oxygen atoms. When the sample is exposed to the tracer, O-18 diffuses rapidly down the grain
FIGURE 4 - Schematic of grain boundary diffusion of tracer with exchange.
boundaries to form new oxide at the oxide-metal interface. However, while diffusing down the grain boundaries the tracer comes into contact with the grains which have O-16 in them. This causes a tracer exchange between the grain boundaries and grains. It should be noted that the grains close to the gas surface have more tracer in them due to greater exchange time than the grains at the old-oxide new-oxide interface.

To model this process of grain boundary diffusion of tracer with exchange, the Oishi-Ichimura geometrical configuration has been chosen as shown in Figure 5. The polycrystalline oxide is assumed to consist of equal sized spherical grains and to possess a grain boundary diffusion coefficient that is much greater than the lattice diffusion coefficient. Diffusion of the tracer proceeds from the gas surface preferentially along the grain boundaries and from the grain boundaries radially inward towards the center of the grains. The grain boundaries of thickness are assumed to be perpendicular to the oxide surface. The grains of radius $r$ are assumed to be attached to the grain boundaries at any arbitrary
FIGURE 5 - Geometrical configuration of oxide grain and grain boundary.
depth in the oxide and are assumed to be completely surrounded by the grain boundaries. Basically, the grains act as sinks for the tracer and are attached to the grain boundary diffusion path. The complicated two dimensional combined lattice and the boundary diffusion problem is transformed into a simple one dimensional diffusion problem in the grain boundary with the grains acting as distributed sinks. The strength of the sinks is given by the well known case of diffusion into a sphere.

It should be noted that when the oxygen flux \( J_{\text{TOT}} \) enters the grain boundary at the gas surface, it is all \( O_{18} \). However, as it comes in contact with a grain, there is a flux of tracer, \( J \), into the grain due to exchange. However, as the number of oxygen sites in the grains are fixed, there is an equal and opposite flux \( J \) of \( O_{16} \) atoms out of the grain and into the grain boundary. Thus as the tracer diffuses down the grain boundary, it becomes more and more diluted in \( O_{16} \) due to exchange. The grains act as a sink drawing \( O_{18} \) atoms from the grain boundary and as a source supplying \( O_{16} \) to the grain boundary. The grain
FIGURE 6 - Coordinate system for the model.
boundary oxygen flux at the old oxide-new oxide interface which is used to form new oxide is actually a mixture of tracer and natural oxygen. This suggests that the isotopic content of the new oxide grains preserves the information on the amount of exchange occurring during the transport of the tracer through the old oxide.

Figure 6 shows the coordinate system used in the mathematical treatment of the diffusion process. The list of symbols used are as follows.

18

$J_y$: Flux of tracer down the grain boundary (kg/m$^2$/sec)

18

$J_x$: Flux of tracer out of the grain boundary (kg/m$^2$/sec)

$D_b$: Grain boundary diffusion coefficient (m$^2$/sec)

$D$: Lattice diffusion coefficient (m$^2$/sec)

$R$: Gas constant (Joules/K mole)

$T$: Temperature (K)
\( \mu^0 \): Oxidation chemical potential (Joules/K mole)

\( Y \): Oxide thickness (m)

\( K \): Parabolic rate constant for oxide growth \( \left( \frac{m^2}{s} \right) \)

\( t \): 0-18 oxidation time (sec)

\( \tau \): 0-16 oxidation time (sec)

\( \phi_y \): Tracer concentration in the grain boundary at depth \( y \) (kg/m

\( C \): Tracer concentration in the grain (kg/m

\( C_i \): Initial tracer concentration in the grain (kg/m

\( \bar{C} \): Average tracer concentration at depth \( y \) (kg/m

\( \delta \): Grain boundary width (m)

\( r \): Grain size (m)

\( \bar{C}_s \): Average tracer concentration in oxide grain (kg/m

40
3.5 Mathematical Modelling Of Diffusion

The flux of tracer down the grain boundary $18 (J)$ is due to two gradients, the oxidation chemical potential gradient $\nabla \mu_0 \frac{\Delta \mu_0}{Y}$ which drives oxidation and its own concentration gradient $\frac{\partial \phi}{\partial y}$ due to isotopic mixing.

$$J_{18}^{18} = \frac{DB}{RT} \frac{\Delta \mu_0}{Y} - \frac{2\phi}{D \partial y} \quad ...(3.1)$$

The diffusion coefficient $D$ is different from the self diffusion coefficient of oxygen as jumps in the direction of the gradient are favoured. There is also a difference in correlation coefficients. However this difference in diffusion coefficient is small [43].

The flux of tracer out of the grain boundary and into the grain $(J)$ is proportional to the concentration gradient of the tracer in the grain surface and the lattice diffusion coefficient $D$.

$$J_{18}^{18} = -D \left( \frac{\partial C}{\partial x} \right)_{x=\delta} \quad ...(3.2)$$

The grain boundary is assumed to be thin enough to neglect the variation of $\phi_y$ in the $x$ direction. Fick's second law gives the time dependence of the grain boundary tracer content.

$$\frac{\partial \phi_y}{\partial t} = - \frac{\partial J_{18}^{18}}{\partial y} - \beta \quad ...(3.3)$$
where \( \beta \) is a sink term representing the rate of tracer loss into the grain. \( \beta \) is given by [5]:

\[
\beta = \frac{2}{5} J_x^{18}
\]

Substitution of the Equation 3.2 in Equation 3.4 gives:

\[
\beta = -\frac{2D}{\delta} \left( \frac{\partial C}{\partial x} \right)_{x=\delta/2}
\]

Thus the time dependence of the tracer content in the grain boundary is given by substituting Equation 3.1 and Equation 3.5 in Equation 3.3.

\[
\frac{\partial \phi_x}{\partial t} = -\frac{D_b \Delta \mu^0}{RT \gamma} \frac{\partial \phi_x}{\partial \gamma} + D_b \frac{\partial^2 \phi_x}{\partial \gamma^2} + \frac{2D}{\delta} \frac{\partial C}{\partial x} \left( \frac{\partial C}{\partial x} \right)_{x=\delta/2}
\]

The last term of the Equation 3.6 can be evaluated if the concentration gradient of the tracer at the grain surface is known. The increasing rate of the grain boundary concentration is much larger than the increasing rate of the tracer concentration within the grain for a given time, since the grain boundary diffusion coefficient is much larger than the lattice diffusion coefficient. Thus the concentration variation within the grain can be considered for a constant \( \phi_y \) with a relatively small error. Thus the problem of tracer exchange reduces to the standard problem of diffusion into a sphere having a constant surface concentration \( \phi_y \).
The standard solution to this equation is given in Equation 3.7.

\[
\frac{C - C_i}{\phi_y - C_i} = 1 + \frac{2r}{\pi \ln(r - x)} \sum_{n=1}^{\infty} \frac{(-1)^n}{n} \sin \left( \frac{n \pi (r - x)}{r} \right) \exp \left( -\frac{Dn^2 \pi^2 t}{r^2} \right) \quad (3.7)
\]

Differentiation of Equation 3.7 by \( x \) gives Equation 3.8, an expression for the tracer concentration gradient at the grain surface (if \( \delta \ll r \)).

\[
\frac{\partial C}{\partial x} \bigg|_{x=\delta} = -\left( \frac{\phi_y - C_i}{\phi_y - C} \right)^2 \sum_{n=1}^{\infty} \frac{Dn^2 \pi^2 t}{r^2} \exp \left( -\frac{Dn^2 \pi^2 t}{r^2} \right) \quad (3.8)
\]

Substitution of Equation 3.8 in Equation 3.6 gives Equation 3.9.

\[
\frac{\partial \phi_y}{\partial t} = -\frac{D_b \Delta u^o}{RT} \frac{\partial \phi_y}{\partial y} + \frac{D_b \Delta u^o}{R \bar{\delta} y^2} \frac{\partial^2 \phi_y}{\partial y^2} - \frac{4D}{b_y \bar{\delta}^2} \sum_{n=1}^{\infty} \frac{\phi_y - C_i}{\phi_y - C} \exp \left( -\frac{Dn^2 \pi^2 t}{r^2} \right) \quad (3.9)
\]

This is a differential equation involving only the grain boundary concentration and its variation in the \( y \) direction and time. The scale thickness is time dependent. Thus, if a parabolic growth rate is assumed with \( K \) as the parabolic rate constant,

\[
Y = (K(t+\tau))^{1/2} \quad (3.10)
\]

where \( t \) and \( \tau \) are the 0-18 and 0-16 oxidation times respectively. Equation 3.9 can thus be written as:

\[
\frac{\partial \phi_y}{\partial t} = \frac{D_b \Delta u^o}{b_y \bar{\delta}^2} \frac{\partial \phi_y}{\partial y} - \frac{D_b \Delta u^o}{RT(K(t+\tau))^{1/2}} \frac{\partial \phi_y}{\partial y} - \frac{4D}{b_y \bar{\delta}^2} \sum_{n=1}^{\infty} \frac{\phi_y - C_i}{\phi_y - C} \exp \left( -\frac{Dn^2 \pi^2 t}{r^2} \right) \phi_y + \frac{4D \phi_y}{\bar{\delta}} \sum_{n=1}^{\infty} \frac{\phi_y - C_i}{\phi_y - C} \exp \left( -\frac{Dn^2 \pi^2 t}{r^2} \right) \quad (3.11)
\]

This differential equation can be easily solved.
numerically to give the grain boundary tracer concentration $\phi_y(y, t)$. Note that $\phi_y$ depends upon three transport parameters $D$, $\delta$ and $D_b$, the oxide microstructure through the grain size $r$ and the oxide growth kinetics through the parabolic constant $K$. When Equation 3.11 is divided by the total oxygen concentration in $kg/m^3$, which is the same throughout the oxide scale, the tracer concentration can be converted from $kg/m^3$ to fractional oxygen concentration. Thus $C$, $C_i$ and $\phi_y$ all become unitless and are actually:

$$\frac{kg\ of\ tracer\ per\ cubic\ meter}{kg\ of\ oxygen\ per\ cubic\ meter}$$

As the SIMS determines the tracer concentration profile by the sectioning technique, the average concentration of the tracer at any depth $y$ should be determined. The average tracer concentration in the oxide grain at depth $y$ at a time $t$ is given by [11]

$$\bar{C} = \frac{M_t}{M_\infty} (\phi_y - C_1) + C_1 \quad \ldots (3.12)$$

where $M$ and $M_\infty$ are the total amounts of tracer $t$ diffusing into the grain at times $t$ and $\infty$, respectively, $\frac{M_t}{M_\infty}$ is not a function of the grain
boundary diffusion coefficient $D_b$ but only of the lattice diffusion coefficient $D$ and is given by the standard form [11]:

$$\frac{M_t}{M_\infty} = 1 - \frac{6}{\pi^2} \sum_{n=1}^{\infty} \frac{1}{n^2} \exp\left(-\frac{Dn^2t}{r^2}\right)$$

...(3.13)

Thus $\overline{C}$ can be written as:

$$\overline{C} = 1 - \frac{6}{\pi^2} \sum_{n=1}^{\infty} \frac{1}{n^2} \exp\left(-\frac{Dn^2t}{r^2}\right) \left(\phi_y C_i + C_i\right)$$

...(3.14)

If the contribution of the tracer in the grain boundary is considered, the average tracer concentration at depth $y$ and time $t$ is given by:

$$\overline{C}(y,t) = \frac{4\pi r^3}{3} \overline{C} + 4\pi r^2 \frac{\delta y}{2} \phi_y$$

$$\overline{C}(y,t) = \frac{4\pi r^3}{3} + 4\pi r^2 \frac{\delta y}{2}$$

...(3.15)

Due to the rastering technique $C(y,t)$ is the quantity measured by SIMS.

3.6 Boundary Conditions

Equation 3.11 is solved by a numerical solution using the finite difference method. ($y,t$) is determined by using the first central difference form of the derivatives [45]. If $X$ is the thickness of the old oxide and $Y(t)$ is the total thickness of the oxide at any time $t$, the initial conditions are:

$$\phi_y = C_i, \ y \leq X, \ t=0$$

$$C = C_i, \ 0 < x < r, \ y \leq X, \ t=0$$

...(3.16)
It is assumed that the natural isotopic concentration of the tracer is negligible i.e.,
\[ C_1 = 0 \] ... (3.17)
The boundary condition at the oxide-sas interface is:
\[ \phi_y = C_s \text{, } t>0 \text{, } y=0 \] ... (3.18)
where \( C \) is the tracer concentration in the sas during the 0-18 oxidation. \( C \) is assumed to be constant during the diffusion.

The oxide-metal interface is a moving boundary. As new oxide is being formed at the oxide-metal interface due to the flux of tracer through the grain boundaries, the tracer concentration in the new oxide grains being formed is the same as the grain boundary tracer concentration at the metal-oxide interface. Figure 7 shows the new oxide growth \( \Delta Y \) during the time \( \Delta t \). The boundary conditions at this interface are:
\[ C(0<x<r, Y<y<Y+\Delta Y, t=t+\Delta t) = \phi_y(y=Y, t=t+\Delta t) \]
\[ \phi_y(y=Y+\Delta Y, t=t+\Delta t) = \phi_y(y=Y, t=t+\Delta t) \] ... (3.19)
where limit : \( \Delta Y, \Delta t \to 0 \)

In the following section some of the quantities necessary for the numerical solution of Equation 3.11 will be determined.
FIGURE 7 - Boundary conditions at the oxide-metal moving boundary.
3.7 Determination of oxidation chemical potential gradient

Let the gas oxide interface be denoted as 1 and the oxide metal interface be denoted as 2 as shown in Figure 8. Let $\mu_0^1$ and $\mu_0^2$ be the oxygen chemical potential in the oxide at 1 and 2 respectively. At 1 the chemical potential of oxygen in the gas must be the same as the chemical potential in the oxide \[46] i.e.,

$$\mu_0^1 = \mu_{gas}^1$$

As the sample is exposed to a closed atmosphere of pure oxygen during the tracer oxidation,

$$p_{O_2}^{gas} = 1$$

This gives

$$\mu_0^1 = \mu_{O_2}^1 = RT \ln p_{O_2}^{gas} = 0 \quad \ldots (3.20)$$

i.e., we choose the standard state as 1 atmosphere oxygen gas. At 2 the oxygen chemical potential must be the same at the oxide and the metal i.e.,

$$\mu_0^2 = \mu_{metal}^2$$

At 2 the reaction:

$$2Al(s) + \frac{3}{2} O_2(g) \rightarrow Al_2O_3(s) \quad \ldots (3.21)$$
FIGURE 8 - Oxidation chemical potential gradient across the oxide scale.
takes place which has a free energy change of \( \Delta G^o \). An equilibrium constant \( K \) for this reaction can be defined as:

\[
\Delta G^o = -RT \ln K = -RT \ln \frac{a_{Al_2O_3}}{(a_{Al})^2(a_{O_2})^{3/2}} \quad \ldots (3.22)
\]

\( a_{Al_2O_3} \) can be assumed to be unity. Thus Equation 3.22 becomes:

\[
a_{metal} = \exp(-\frac{2|\Delta G^o|}{3RT})(a_{Al})^{-4/3} \quad \ldots (3.23)
\]

Since \( \mu_{O_2} = RT \ln a_{O_2} \), Equation 3.23 gives:

\[
\mu_{metal} = -\frac{2}{3} |\Delta G^o| - \frac{4}{3} RT \ln (a_{Al}) \quad \ldots (3.24)
\]

In the absence of any thermodynamic activity data, the metal can be assumed to be a single phase ideal solution. Then \( a \) is the atomic percent of \( Al \) aluminum in the alloy \([Al]\). Thus the Equation 3.24 becomes:

\[
\mu_{O_2}^2 = \mu_{metal} = -\frac{2}{3} |\Delta G^o| - \frac{4}{3} RT \ln [Al] \quad \ldots (3.25)
\]

Thus the oxidation chemical potential difference across the oxide scale is given by:

\[
\Delta \mu^o = \mu_{O_2}^1 - \mu_{O_2}^2 = \frac{2}{3} |\Delta G^o| + \frac{4}{3} RT \ln [Al] \quad \ldots (3.26)
\]

Substitution of values of \( [Al] \) in Equation 3.26 gives: \( \Delta \mu^o = 787915 \text{ J/mole at } 1100^\circ \text{C.} \)
3.8 Relationship between $D$ and $K_b$

If the mechanism of oxide growth is primarily inward oxygen diffusion through the grain boundaries, the grain boundary diffusivity and the growth rate kinetics can be related by considering a simple model.

The sample is exposed to natural oxygen and an oxide scale is grown. The oxide is assumed to have cubic grains of dimension $r$ surrounded by the grain boundaries of thickness $\delta$ as shown in Figure 9. The cross section of the sample itself is assumed to be a square with each side being $r$ grains long.

Let the thickness of the oxide at any time $t$ be $Y$ and let the scale grow by an amount $\Delta Y$ in time $dt$. Then the flux of oxygen $J$ (kg/m sec) down the grain boundaries is due to the oxygen potential gradient.

The amount of oxygen transferred in time $\Delta t$ through the grain boundaries, $W$ (kg) is given by:

$$ W_T = J^0 \cdot \text{area of grain boundaries} \cdot \text{time} $$

The shaded area in Figure 9 is the grain boundary area through which the oxygen is transported ($b \ll r$).

$$ W_T^0 = \frac{D_b C}{RT} \frac{\Delta Y^0}{Y} \cdot 2\pi r \delta \cdot \Delta t \quad ...(3.27) $$
FIGURE 9 - Schematic of model geometry used to relate scale growth kinetics to grain boundary diffusion.
This amount of oxygen transported must be the same as the amount of oxygen incorporated in the net growth of oxide, $W_N$ (ks).

$$W_N = \text{volume of new growth in time } \Delta t \cdot \text{concentration of O in scale}$$

$$W_N^O = (nr)^2 \cdot \Delta Y \cdot C$$

where $C$ is the oxygen concentration ($\text{ks/m}^3$) in the scale. As $W = W_N$, Equation 3.27 and Equation 3.28 gives:

$$\frac{D \cdot \Delta C^O}{RT} \cdot \frac{\Delta L}{\Delta t} = 2n^2 r \cdot 6 \cdot \Delta t = n^2 r \cdot 2 \cdot \Delta Y C$$

$$\frac{2\delta D \cdot \Delta C^O}{RT \cdot r \cdot \Delta Y} = \Delta t$$

$$\frac{\Delta Y}{\Delta t}$$

is the growth rate of the oxide. If a parabolic growth rate is assumed with growth rate constant $K_p$

$$Y = (K_p t)^{1/2}$$

$$\frac{dy}{dt} = \frac{1}{2} K_p t^{-1/2}$$

Combining Equations 3.29 and 3.30 gives:

$$\delta D_b = \frac{rK_p}{4} \frac{RT}{\Delta C^O}$$

Substitution of appropriate values in Equation 3.31:

$$r = 0.3 \text{ microns, } T = 1100 \text{ C, } K_p = 1.2 \times 10^7 \text{ m/sec}$$

$$\delta = 10 \text{ A, } \Delta C^O = 78915 \text{ J/mole}$$

sives:
\[ D_b = 1.3 \times 10^{-5} \text{ m}^2/\text{sec} \] \((3.32)\)

This gives a good estimate of the value of \( D_b \) to be used in the numerical solution of the Equation 3.11.

3.9 Tracer concentration profile in the new oxide

The tracer concentration profile in the new oxide is obtained by the principle of mass balance. It is assumed that the amount of exchange of \( O^{16} \) in the oxide grains at the gas surface with the \( O^{18} \) in the tracer due to volume diffusion at the gas-oxide interface is negligible. Then for an inward growing scale once the sample is exposed to pure tracer, the amount of \( O^{16} \) in the entire oxide must remain constant. This means that the amount of \( O^{18} \) retained in the old oxide must be the same as the amount of \( O^{16} \) in the new oxide. Let \( X \) be the thickness of the oxide after an \( O^{16} \) oxidation for time \( T \) and let \( Y \) be the oxide thickness after a further tracer oxidation for a time \( t \). Let an additional tracer oxidation for a time \( dt \) cause a new oxide growth of \( dY \) as shown in Figure 10. The tracer concentration profile till depth \( Y \) at time \( t+dt \) is also shown.
FIGURE 10 - Determination of oxide growth kinetics by mass balance.

\[ \frac{dY}{dY} = (Y, t = t + \Delta t) \]

\[ A = B + E \]

new oxide growth \( dy \):

old oxide

tracer concentration

metal
Area A denotes the amount of tracer in the old oxide and area B denotes the amount of 0-16 in the new oxide grown till time $t$. Let $A-B=E$. Then by mass balance, the amount of 0-16 contained in the new growth $dY$ must be $E$. As mentioned in Section 3.6, the tracer concentration in the new growth is:

$$\phi_y = (y=Y, t=t+\Delta t)$$

which gives the 0-16 concentration in the new growth as:

$$1 - \phi_y(y=Y, t=t+\Delta t)$$

Thus the growth $dY$ during time $dt$ can be calculated as:

$$dY = \frac{(A-B)}{(1-\phi_y(y=Y, t=t+\Delta t))} \quad ...(3.33)$$

3.10 Flow chart of computer solution

Equation 3.11 is solved on the computer using the finite difference technique. The old oxide is divided into 19 equal parts with the oxide-gas interface being called 1 and the metal-oxide interface being called 20, as shown in Figure 11. Each part is considered to be 1 unit. The growth rate of the oxide is determined by mass balance as discussed in Section 3.9. The value of $D$ is chosen such that the growth rate is comparable.
FIGURE 11 - Division of oxide scale for finite difference solution.
with the parabolic growth rate kinetics. The flow chart for the computer program is shown in Figure 12. The computer program is listed in appendix A.

3.11 Computer plots of tracer profiles

Figures 13-15 show the theoretical tracer concentration profiles in the old oxide for various values of lattice diffusivity, grain boundary diffusivity and grain size. A higher value of lattice diffusivity and causes a fast exchange rate to be set up. The grains in the old oxide are filled up quickly. A higher value of the grain boundary diffusivity causes more tracer to be retained in the grains of the old oxide as the tracer diffuses quickly down the grain boundaries increasing the exchange time for the grains. A smaller grain size increases the surface area for exchange and causes more tracer to be retained in the old oxide. The variation of the tracer profiles when each parameter is varied is different. Thus theoretically given an experimental tracer profile, it should be possible to come up with unique values of the three parameters to fit the curve. It should be noted that the values of $D$ and $r$ are
FIGURE 12 - Flow chart for the computer program.
0-16 oxidation time - 18 hours
0-18 oxidation time - 14 hours
at 1100°C
D_2 = 1.3 \times 10^{-16} \text{ m}^2/\text{sec}
grain size = 0.3 \text{ microns}

FIGURE 13 - Effect of lattice diffusion coefficient on the tracer profile in the old scale.
FIGURE 14 - Effect of grain boundary diffusion coefficient on the tracer profile in the old scale.
0-16 oxidation time-18 hours
0-18 oxidation time-14 hours
at 1100°C
$D_b = 1.3 \times 10^{-16} \text{m}^2/\text{sec}$
$D_b = 5.0 \times 10^{-20} \text{m}^2/\text{sec}$

**FIGURE 15** - Effect of grain size on the tracer profile in the old scale.
not independent, but are related to the $K$ value by Equation 3.31. Thus if one of the above parameters is fixed, there is a unique value of the other parameter to satisfy the parabolic growth rate kinetics. Figures 13-15 are plotted to show how sensitive the profiles are to the three parameters.

Figure 16 shows the computer simulation of how the tracer concentration profile develops across the entire oxide scale when the scale grown by a 18 hour O-16 oxidation followed by a 14 hour O-18 oxidation. The values of the three parameters chosen are:

$$D = 1.41 \times 10^{-2} \text{ m/sec}, \quad D = 10^{-20} \text{ m/sec}$$

$$b = 0.3 \text{ microns}$$

The value of the grain boundary width chosen is $0.10 \text{A}$. Here we can see the versatility of the model. The oxide microstructure can be incorporated in the program by feeding in the real grain size variation across the scale. In case the lattice diffusion coefficient varies across the side, that too can be incorporated in the program. The actual value of
0-16 oxidation time = 18 hours
1 unit = X/19

X=19 units

old oxide

new oxide

gas

a) 0-18 oxidation time = 4 hours

19 units

old oxide

new oxide

gas

21 units

b) 0-18 oxidation time = 8 hours, 25 minutes

19 units

old oxide

new oxide

gas

23 units

c) 0-18 oxidation time = 14 hours

19 units

old oxide

new oxide

gas

25.3 units

FIGURE 16 - Development of tracer concentration profile.
the oxide growth kinetics can be experimentally found and the real value of $Y(t)$ can be used in the program. The value of $D$ can be so adjusted that the growth rate predicted by mass balance is the same as the experimental growth rate. So long as the oxide is inward growing, the tracer profile for any growth kinetics and microstructure can be modelled.

It should be noted that the model is based on a planar oxide growth mechanism and any quantitative information derived from the experimental profiles assumes good resolution and high quality of the data itself. In case the oxide metal interface is irregular, it is difficult to identify the interface for the purpose of mass balance. The same problem arises if the sputtering itself is non uniform and information at the metal-oxide interface is being gathered simultaneously from both metal and oxide grains. Non uniform sputtering can severely limit the resolution of the data such that no quantitative information can be obtained from the shape of the profile.
CHAPTER 4

EXPERIMENTAL PROCEDURE

4.1 Sample Preparation

The FeCrAl-Zr alloy used in the experiment was obtained through NASA Lewis Research Center. The alloy used was Fe-21.9Cr-5.8Al-0.22Zr and was obtained as cold rolled strips about 1.5mm thick. Samples of size 6mm x 6mm x 1.5mm were cut out by a Buehler diamond saw. The specimens were then mounted and polished on 240, 320 and 600 grit polishing paper followed by 15, 3 and 1 micron diamond paste polishing. Some of the samples were prepared at NASA and were polished by 600 grit polishing paper followed by 3 micron and 1 micron diamond paste polish.

4.2 Exchange Apparatus

Figure 17 shows a schematic of the apparatus used for the gas exchange process. The tracer is kept in a zeolite storage tank.

* Courtesy J. Smialek.
FIGURE 17 - Schematic of gas exchange apparatus.
The sample is placed in a quartz tube with the polished side up. Oxygen flow into the quartz tube can be controlled by valve 5 and the stopcock. The thermocouple vacuum gauge measures the pressure inside the system when it is evacuated. The pressure gauge measures the pressure in the system when the tracer is introduced. Normally the tracer pressure in the system is maintained slightly above the atmospheric pressure such that no air leaks into the system during the tracer anneal. The sample is annealed by placing the quartz tube in a single zone horizontal Lindberg furnace. The temperature of the furnace is monitored by a Pt-6%/Rh/Pt-30%Rh thermocouple.

4.3 Exchange Procedure

Some of the samples are annealed in vacuum after polishing to remove the polishing damage. The samples are placed in the quartz tube and the system is pumped down to a pressure of 0.03 torr. The samples are then annealed at 1100 °C for 30 minutes. This also helps the sample and the quartz tube to heat up and come into thermal
equilibrium with the furnace. For the 0-16 anneal, valve 2 is closed to cut off the vacuum pump and valve 3 is opened to introduce air into the system. So the old scale is grown in air at 1 atmosphere pressure. The transfer of O-18 into the system for the tracer anneal is done without cooling down the system. At the end of 0-16 anneal, valve 3 is closed and the system is evacuated with the vacuum pump by opening valve 2. Meanwhile the temperature of the zeolite tank is raised by using the heating coil. When the pressure of the system falls to 0.03 torr, valve 2 is closed and the tracer is introduced into the system through valve 1. The O-18 is kept at a slightly higher than atmospheric pressure to ensure no leakage of atmospheric air into the system which can cause dilution of the tracer. At the end of the tracer anneal, the system is evacuated again by cooling the zeolite tank with liquid nitrogen. On cooling it acts as a pump and sucks out all the tracer from the system.

4.4 SIMS
The GATAN SIMS at Pennsylvania State University was used to obtain the data for the tracer concentration profile. The primary beam used was Ar \(^+\) under a beam acceleration voltage of 7KV. The complete instrument settings are listed below:

<table>
<thead>
<tr>
<th>Component</th>
<th>Value 1</th>
<th>Value 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beam Accelerator</td>
<td>7 KV</td>
<td>1.53 mA</td>
</tr>
<tr>
<td>Ion Source Filament</td>
<td></td>
<td>17.2 A</td>
</tr>
<tr>
<td>Anode</td>
<td>123 V</td>
<td>3.65 mA</td>
</tr>
<tr>
<td>Beam Deflector</td>
<td>Initial - 43 V</td>
<td>Final - 204 V</td>
</tr>
<tr>
<td>Lens Control Condensor</td>
<td>6.16 KV</td>
<td></td>
</tr>
<tr>
<td>Velocity filter</td>
<td>Electric Field- 0.81 V</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Magnetic field- 0.59 A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Objective - 4.18 KV</td>
<td></td>
</tr>
<tr>
<td>Energy filter</td>
<td>Outer potential- 40 V</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Inner potential- 113 V</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bias - 39 V</td>
<td></td>
</tr>
</tbody>
</table>

* Operated by Dr. C. Hauser under the guidance of Prof. C. Pantano.
The list of species simultaneously monitored and their mass to charge ratio is given below:

<table>
<thead>
<tr>
<th>Element</th>
<th>Mass/Charge ratio monitored</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>13.9</td>
</tr>
<tr>
<td>16</td>
<td>16.4</td>
</tr>
<tr>
<td>18</td>
<td>18.4</td>
</tr>
<tr>
<td>O</td>
<td>13.9</td>
</tr>
<tr>
<td>Cr</td>
<td>52.9</td>
</tr>
<tr>
<td>Fe</td>
<td>56.9</td>
</tr>
<tr>
<td>Zr</td>
<td>91.7</td>
</tr>
</tbody>
</table>

Due to the insulating properties of alumina, it was necessary to compensate for the positive charge of the primary beam by spraying electrons on the sample surface. These electrons can cause desorption of light elements from the near surface region. This gives rise to a background intensity of the monitored ions and is termed as Electron Stimulated Desorption (ESD). The ESD level is evaluated by shutting off the primary beam and measuring the secondary ions due to the electron beam only. This background level is subtracted out from the actual reading.
A sample holder for SIMS can hold up to 6 samples and has in addition 2 standards. The first standard is Ta O on Ta which is used to aim the primary beam and to check if the rastering is correct. The second standard is ZnS which is used to aim the electron beam such that it is coincident with the primary beam.

The samples are then sputtered one at a time and the output of the machine is fed directly into a computer. The output is in the form of the number of counts of a species versus the sputtering time. The conversion of counts to concentration of tracer \( \bar{C}_{\text{18}}(t_s) \) at a time \( t \) is given by:

\[
\bar{C}_{\text{18}}(t_s) = \frac{h_{\text{18}}(t_s) - \text{0-18esd}}{(h_{\text{18}}(t_s) - \text{0-18ESD}) + (h_{\text{16}}(t_s) - \text{016ESD})} \quad ... (4.1)
\]

where \( h_{\text{18}}(t) \) and \( h_{\text{16}}(t) \) are the counts of 0-18 and 0-16 at a sputtering time \( t \). 0-18ESD and 0-16ESD are the ESD values of 0-18 and 0-16 measured by shutting off the primary beam. The sputtering time is converted to sputtering depth by measuring the depth of the crater by a profilometer and dividing it by the total time under the primary beam giving the sputtering rate.
S such that:

\[ Y = S \cdot t \]  \hspace{1cm} \text{(4.2)}

where \( Y \) is the sputtered depth corresponding to the sputtering time \( t \).
CHAPTER 5

RESULTS AND DISCUSSION

5.1 Reasons for choosing the alloy and technique

The theoretical modelling of the oxidation process of the FeCrAl-Zr alloy was done on the assumption that the oxide growth is planar, i.e., the oxide metal interface is sharp. Smialek studied the microstructure of the oxide formed on the same alloy. He found that at 1100 °C, the oxide formed had closely packed polycrystalline grains and the oxide-metal interface was reasonably planar, at least for long oxidation times. At 1100 °C there were very few pores observed in the oxide. Reddy [15] used the double oxidation technique on a NiCrAl alloy and concluded that the oxide was inward growing with the mechanism being oxygen grain boundary transport. There was no reason to believe that the behaviour of FeCrAl-Zr alloy would be drastically different. So the FeCrAl-Zr alloy seemed like a good alloy to apply the double oxidation technique. Cawley [16] used
the double oxidation technique very successfully on Si and SiC. He used the SIMS to obtain the tracer concentration profile. He was able to experimentally obtain a sharp oxide interface as the oxygen signal in the SIMS fell down rapidly at the interface. He was also successful in determining the diffusion coefficients of oxygen from the observed concentration profile. His success was inspirational in choosing the double oxidation technique and SIMS for these experiments.

5.2 Lacey growth

When the tracer concentration profile in the oxide on a FeCrAl-Zr sample was obtained, it differed from the profiles obtained by Reddy on his NiCrAl-Zr samples. To try to interpret the difference the oxide was looked at under the SEM. The microstructure as shown in Figure 18 varied drastically from that observed by Smialek on the same alloy. The sample had been polished by 3 micron diamond paste, annealed in vacuum at 1100 °C for 30 minutes and oxidised for 40 hours at 1100 °C.

*Courtesy J. Smialek and J. Cawley at NASA Lewis and M. Occhionero at Case Western Reserve University.
FIGURE 18 - Top view of the oxide at the gas-oxide interface. The sample has been polished to a 3 micron diamond paste finish, annealed in vacuum for 30 minutes at 1100°C followed by a 40 hour oxidation at 1100°C.
Instead of a compact polycrystalline scale, a lacy structure of about 2 micron wide ridges was observed at the oxide-gas interface. This shows that the oxide growth is not planar.

To understand this difference in microstructure a systematic study was undertaken. The effect of three variables on the microstructure was studied: the oxidation time, the surface finish of the sample and the time of vacuum anneal. The vacuum anneal is done before the oxidation to remove the polishing damage.

The development of the oxide microstructure with time is shown in Figure 19. The alloy is polished to a 1 micron diamond paste finish and oxidized at 1100°C for times of 3 hours, 10 hours, 20 hours and 36 hours, 25 minutes. The lacy structure at the oxide-gas interface is evident at least for the shorter oxidation times. The ridges form a network of cells and the average cell size seems to decrease with time. The average cell size decreased from about 5 microns at 3 hours to about 3 microns at 10 hours and goes down to about 1.5 microns at 20 hours. At 36 hours and 25 minutes,
19 a) 3 hour oxidation at 1100°C.
19 b) 10 hour oxidation at 1100°C.
19 c) 20 hour oxidation at 1100°C.
19 d) 36 hours, 25 minutes oxidation at 1100°C.

FIGURE 19 - Changes in microstructure of oxide at gas-oxide interface with increasing oxidation time. The samples have been polished to a 1 micron diamond paste finish.
the lacy structure cannot be seen and the microstructure looks like that reported by Smialek. So if only the long time microstructure was examined it would seem to be a planar scale and would encourage the application of the model to derive quantitative information. However, such a procedure would lead to errors as the oxide did not start off looking that way and at least the initial period of oxidation did not lead to a planar oxide growth.

A detailed description of the oxidation mechanism leading to the lacy structure will not be attempted here. However it is important to note that the outward aluminum diffusion must occur to change the microstructure at the gas-oxide interface. So at least for oxidation up to 20 hours, the oxide is not completely inward growing. Hindam and Smeltzer [49], observed lace on Al O scales grown on NiAl. They explained this occurrence by stating that the initial oxide formed was Y-Al O by inward oxygen diffusion. The scale then went through a to tranformation accompanied by a 14% reduction in volume. The α-alumina formed was for most part oriented. However, this volume
reduction gave rise to some faulted oxide regions. The scale was thus a combination of oriented alumina and misoriented oxide which provided the path for combined inward oxygen and outward aluminium diffusion. This gave rise to ridges at the gas-oxide and the oxide-metal interface corresponding to misoriented oxide grains. However, this does not explain why the cell size of the lacy structure should decrease with time as the transformation occurs only once.

Hindam and Smeltzer observed that the structure of lace itself can be a function of the alloy grain orientation and can be vary drastically over two alloy grains in the same sample. Douchak [50] also observed a grain orientation dependency of the lacey structure during his study of NiAl oxidation. Work is presently being done by Smialek at NASA to understand the occurrence of lace. The dependency of lace on the alloy grain orientation raises the question whether the observed decrease in cell size with time is real or an artifact because the four samples were so chosen that the alloy grain orientation led to a decreasing cell size. Though this is a possibility, the odds
Figure 19b) and Figure 20 show the effect of surface finish on the lacy structure. The sample in figure 20 has been polished by a 600 grit polishing paper while the sample in Figure 19b) has been polished by 1 micron diamond paste. Both the samples have been oxidised for 10 hours at 1100 °C. No dramatic difference is observed in the lacy structure which suggests that surface finish may not play a big role in lace formation.

The effect of the vacuum anneal on the lacy structure can be seen by studying Figure 19b) and Figure 21. The sample in Figure 19b) has not been treated to the vacuum anneal while the sample in Figure 21 has been annealed in vacuum for 30 minutes at 1100 °C. The sample with no vacuum anneal shows the lacy structure as a network of cells with the average cell size around 3 microns. However, the structure of the sample with vacuum anneal looks drastically different. The lace takes up a spider like form with a nucleus and the lace spreading in radial directions from it. This shows up in certain areas of the oxide surface. Regions
FIGURE 20 - Top view of oxide at the gas-oxide interface. The sample has been polished by 600 grit polishing paper and annealed for 10 hours at 1100°C.
FIGURE 21 - Top view of oxide at gas-oxide interface. The sample has been polished to a 1 micron diamond paste finish, annealed in vacuum for 30 minutes at 1100°C followed by a 10 hour oxidation at 1100°C.
away from the lace covered areas have a needle like structure. A similar effect was observed at a higher temperature as shown in Figure 22. The sample in Figure 22a) was annealed for 30 minutes in vacuum at 1200 °C while the sample in Figure 22b) had not been given any vacuum anneal. Both the samples were polished by a 1 micron diamond paste and oxidised for 10 hours at 1200 °C. The formation of lace on the sample with vacuum anneal was much more dramatic. These observations suggest that vacuum annealing encourages the formation of lace at least at short oxidation times.

To study the effect of vacuum anneal on lacy structure for large oxidation times, two samples were oxidised at 1100 °C for 50 hours. Both samples were annealed in vacuum for 30 minutes at 1100 °C before oxidation. Figure 23 shows the microstructure of the oxide formed on the two samples. The sample in Figure 23a) has been polished with 1 micron diamond paste while the sample in Figure 23b) has been polished with 600 grit polishing paper. Both samples do not show lace. However, Figure 18 shows a dramatically different microstructure. The sample in Figure 18
22 a) Vacuum anneal at 1100°C for 30 minutes prior to oxidation.
22 b) No vacuum anneal.

FIGURE 22 - Effect of vacuum anneal on oxide microstructure at 1200°C. Both samples were polished to a 1 micron diamond paste finish and oxidised for 10 hours at 1200°C.
23 a) Microstructure of oxide at gas-oxide interface. The sample has been polished to a 1 micron diamond paste finish, annealed in vacuum for 30 minutes at 1100°C and oxidised for 50 hours at 1100°C.
23 b) Microstructure of oxide at gas-oxide interface. The sample has been polished by 600 grit polishing paper annealed at 1100°C for 30 minutes and oxidised for 50 hours at 1100°C.

FIGURE 23 - Microstructure of oxide after long time oxidation.
has also been annealed in vacuum for 30 minutes at 1100 °C, after which it was oxidised for 40 hours at 1100 °C. The specimen has been polished by 3 micron diamond paste. This difference in microstructure may be due to different alloy grain orientation.

In conclusion, some of the facts about lace and its structure are stated without an attempt to explain the reason or the mechanism of structure variation. Occurrence of lace seems to be sensitive to the vacuum anneal, at least at smaller times. The vacuum anneal encourages the formation of lace. The structure of the lace itself may vary from a cell-like structure in Figure 14b) to a spider-like structure in Figure 21. The cell size and shape of the lace is sensitive to the alloy grain orientation. The cell size of the lace may decrease with time under isothermal oxidation. The lace seems to have a radial texture, i.e., the ridges seem to form radially from a nucleus.

The formation of lace and its structure does not seem to be consistent and can vary from sample to sample. Its erratic behaviour is a good reason to refrain from speculating on its origin.
5.3 Sputter etching

An important point to consider is the quality of data obtained by SIMS. The question to be addressed is that if the alloy was to undergo planar oxidation in a manner described by the model, how accurate would be the tracer concentration profile obtained by SIMS. Data obtained by SIMS are in the form of ion counts versus sputtering time. Even though the counts are a strong function of the oxide matrix, the matrix dependency is removed by taking the ratio of O-18 counts to the total number of oxygen counts. However, the conversion of sputtering time to depth is valid only if the sputtering rate is a constant or a known function of time. The manner in which data is collected by SIMS is that the oxide is sputtered for a given length of time (1 sec in this case) and the counts reported is the average of the signal received over this period.

To understand how SIMS collects the data, consider a planar oxide with a true concentration profile \( C(y) \) across it as shown in Figure 24. The data is collected by sputtering for a time \( \Delta t \) and
true concentration profile \( C(y) \)

volume sputtered during a period \( \Delta t \)
at time \( t \)

\( \Delta y_1, \Delta y_2, \Delta y_n \)

gas

oxide

metal

sputtering direction

\( Y_n, Y_{n+1}, y \)

FIGURE 24 - Sputtering of oxide scale by SIMS.
averaging the number of counts collected over that time period. Let $\Delta y_1, \Delta y_2, \ldots, \Delta y_n$ be the thickness of oxide sputtered away after successive sputterings of time $\Delta t$. The sputtered surface is assumed to be flat. Then the concentration reported by SIMS at time $t$, $C(t)$, is given by:

$$C(t) = C\left(\frac{y_n + y_{n+1}}{2}\right) = \frac{1}{\Delta y_n} \int_{y_n}^{y_{n+1}} C \, dy$$

...(5.1)

If the sputtering rate is constant, then all the $\Delta y$'s are the same.

Figure 25a) shows a step function concentration profile and Figure 25b) shows how the SIMS data would appear for a constant sputtering rate when the data in the form of counts versus sputtering time is converted to a plot of concentration versus distance. The profile obtained would be a symmetric distortion of the true profile. However it would be area preserving, i.e., the area under the experimental concentration profile would be the same as that under the true concentration profile. Thus the mass balance ideas discussed in Chapter 2 would be valid and applicable to the observed profile. However, if the sputtering rate is not uniform, then the conversion of counts versus time
FIGURE 25 - Distortion of concentration profile due to sputtering by SIMS. a) Concentration profile. b) Output from SIMS.
to concentration versus distance by using Equation 4.2 would lead to an unsymmetrically distorted profile which would not be area preserving. In such a case the mass balance idea would not be applicable to the data.

A further complication occurs if the sputtered surface of the oxide is not planar but has instead a convoluted morphology as shown in Figure 26. Let \( C(y) \) be the true concentration profile of the tracer in the oxide. In this case for a sputtering time of \( \Delta t \) information is being collected from a range of depths \( y_1 \) to \( y_2 \) and the reported concentration is the average of the concentration over this depth range weighted by the amount of material removed from a given depth. Let \( dV(y) \) be the volume of oxide removed at depth \( y \) by sputtering for a time \( \Delta t \). The average concentration reported is given by:

\[
\bar{C} = \frac{\int_{y_1}^{y_2} C(z) \, dV(z)}{\int_{y_1}^{y_2} dV(z)} \quad \cdots (5.2)
\]

If the range of depths over which information is simultaneously received is too large it may distort the concentration profile to an extent that obtaining quantitative information from it is
FIGURE 26 - Effect of non-uniform sputter surface.
impossible.

Another major problem faced when a sputter etching technique is used on a polycrystalline oxide is the phenomenon of preferential sputtering. The grain boundary region may sputter at a faster rate than the grains. The sputtering rate of the grains themselves may be a function of their orientation. This means that the range of depths over which the signal is collected will vary across the sputter surface. Figure 27a) shows the oxide scale being made up of 2 regions A and B where B sputters twice as fast as A. A could represent the grain while B could represent the grain boundary or alternatively they could represent grains of fast and slow sputter orientation. Figure 27 shows the sputter surface after a sputtering time $\Delta t$. The reported concentration is the average of the concentration measured over the shaded area in Figure 27b). This preferential sputter etching can also severely limit the resolution of the data.

5.4 SIMS data
FIGURE 27 - Preferential sputtering of oxide by SIMS.
Figure 28 shows the total counts of oxygen plotted as a function of normalised scale depth for a sample polished by 1 micron diamond paste and oxidised successively in air and O-18 tracer at 1100°C for 20 hours and 16 hours, 25 minutes respectively. It is important to note that the oxygen counts drop off very gradually and there is no discontinuity indicating a sharp oxide-metal interface. This could be due to a combination of preferential sputtering and non-planar oxide growth. Figure 29 shows a SEM picture of the same crater. The metal grains and the regions of oxide can be seen side by side as marked by M and O respectively. The lacy structure is evident in the oxide. The apparent tracer concentration profile of the same sample is plotted in Figure 30. If there was a sharp interface, the tracer concentration profile would have fallen off rapidly once the metal was reached. However, not only does the concentration profile at the interface not fall off steeply but it has a lesser slope than the near surface profile. This indicates that the problems of non-planar oxide growth and preferential sputtering are too severe to apply our model to obtain quantitative data.
FIGURE 28 - Plot of total oxygen counts versus normalised depth.

0-16 oxidation time = 20 hours
0-18 oxidation time = 16 hours, 25 minutes
at 1100°C

sputtering time
\[ \frac{\text{total sputtering time}}{X 100 \text{ (normalised depth)}} \]
FIGURE 29 - Top view of the sputtered crater.
slope | A | > | slope | B |

0-16 oxidation time=20 hours
0-18 oxidation time=16 hours, 25 minutes
at 1100°C

FIGURE 30 - Tracer concentration profile from SIMS
5.5 Conclusions

The aim of this study was to develop the double oxidation tracer technique. The thrust of the work was to come up with a method of analysis of the tracer concentration profile such that valuable information could be obtained from it.

The mass balance ideas developed can be used to deduce the oxidation mechanism from the tracer distribution in the oxide scale, if the oxide grows in a planar fashion. For all planar growing scales, if a true tracer concentration profile can be obtained, the fraction of inward and outward growth can be deduced by mass balance.

An effort was then made to develop an analysis which could be used to derive quantitative diffusion data from the tracer concentration profile. The specific diffusion problem of tracer distribution in a polycrystalline oxide scale growing by inward oxygen grain boundary diffusion with exchange was solved using a model based on the Oishi-Ichimura combined lattice and boundary diffusion model. For uniform grains across the
scale, unique profiles could be predicted using a combination of \(D, D\) and \(r\). It was found that each \(b\) parameter had a different effect on the profile. The model could be generalized to accommodate any grain size distribution obtained in a real oxide scale. Careful consideration was also given to the factors that could distort the data.

Having developed the analysis, it was tested on the FeCrAl-Zr alloy. Based on the extensive studies on NiCrAl alloys and long time oxidation studies on FeCrAl alloys by Smialek, the FeCrAl-Zr alloy was expected to oxidise in a planar fashion. However it was found that the oxidation behaviour of FeCrAl-Zr alloy was extremely unpredictable and the microstructure of the oxide varied from sample to sample and even within the sample. The formation and changes in the lacey structure at the gas-oxide interface suggested that outward cation transport also occurred. So it was found that the FeCrAl-Zr alloy was not a good system to apply the model. The SIMS which had been very successfully used on amorphous materials led to preferential sputtering in the polycrystalline oxide scale and caused distortion of the tracer concentration.
profile. Thus no attempt was made to derive diffusion coefficients from the data obtained.

To exploit the analysis and quantitative modelling done in this thesis, it is necessary to apply the double oxidation technique to systems with typically planar growing scales. There is also a need to improve the profiling techniques on polycrystalline scales using proton activation or SIMS having uniform sputter etching.
CHAPTER 6

SUGGESTIONS FOR FUTURE WORK

1. The realistic oxide microstructure could be incorporated in the computer program by treating the problem of varying grain size and shape across the oxide scale. For oxide scales having columnar grains, the exchange process could be treated as diffusion into a cylinder. The only modification required would be to substitute Equation 3.7 by the corresponding equation of diffusion of tracer into a cylinder having a constant surface concentration.

2. The presence of precipitates of a second phase with different oxygen diffusivity could be incorporated in the program by considering the variation of lattice diffusivity across the oxide scale.

3. The problem of combined inward oxygen tracer diffusion with exchange and outward aluminum diffusion could be addressed.

4. The amount of O-16 in the old oxide undergoing direct surface exchange with the gas during tracer oxidation can be calculated. This would lead to a
modification of the mass balance ideas. However the amount of O-16 involved in direct exchange with the gas is not very significant (Appendix B).

5. A system with an inherently planar growing oxide scale should be chosen to apply this model. The analysis techniques developed could be applied to the existing tracer concentration data on NiO by Atkinson et. al.[51] and NiCrAl-Zr by Reddy [15].
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APPENDIX A

Listings of the Computer Program

dimension c(40), ci(40), ca(40), ti(40), cstrain(40)
open (unit=1, status='old', name='fpi.dat')
READ(1,*) concint
read(1,*) dop, sc
READ(1,*) W3
read(1,*) tstop, temp, tau, dl, sbwd, okp
READ(1,*) W4
read(1,*) ssize, dsb
open(unit=2, status='new', name='lpfl.obj')
open(unit=3, status='new', name='lrate.obj')
open(unit=4, status='new', name='sbc.obj')
write(2,3)
3 FORMAT(5X,6HTEMPE,10HO-18TIME,5HO-16TIME)
write(2,4)temp, tstop, tau
4 format(4X, F6.0, 4X, F6.0)
write(2,8)
8 FORMAT(5X, 14HGRAIN SIZE(M), 25HGB-DIFF
* COEFF (M**2/SEC), 21HLDIFF COEFF (M**2/SEC), 5X,5hsbwd)
write(2,9)ssize, dsb, dl, sbwd
9 format(5X, e11.3, 5X, e11.3, 13X, e11.3, 10X, e11.3)
write(2,10)
10 format(3X, 5H time, 4X, 1h1, 5X, 1h2, 5X, 1h4, 5X, 1h6, 5X, 1h8, 5X, 2h10, 4X, 2h12, 4X, 2h14, 4X, 2h16, 4X, 2h18, 4X, 2h20)
READ(1,*) W5
read(1,*) tinc, dtp

115
deltw = ((okp*tau)**0.5)/19.

\[ a = 4*dl/( ssize*sbwd) \]

\[ B = ((3.14/GSIZE)**2.)*DL*T \]

\[ \alpha = \frac{d*b*dop/(sc*temp*((0kF*(T+tau)**0.5))}{(Ssize*Sbwd)} \]

var = 0.

DO 12 N4 = 1, 20

\[ var = var + \exp(-b*(N4**2.)) \]

VAR4 = \exp(-B*(N4**2.))

IF (VAR4/VAR.LE.0.01) GO TO 100

12 continue

N4 = 1

beta = a * var

\[ x = (dSb/(deltw**2.))-(alpha/(2.*deltw)) \]

\[ y = (1./tinc)-(2*dSb/(deltw**2.)) \]

\[ z = (dSb/(deltw**2.))+(alpha/(2.*deltw)) \]

do 16 J = 2, nscount

\[ \text{ci}(j) = tinc*(x*c(j)+y*c(j)+z) \]

\[ \text{ci}(1) = 1.0*concint \]

\[ \text{ci}(40) = 0.0 \]

do 35 k = 1, nscount

if (\text{ci}(k).le.0.0) go to 35

\[ \text{ti}(k) = \text{ti}(k) + tinc \]

35 continue

if (t-tprint).le.20,17,17

17 continue

write(4,800) t, ci(2), ci(8), ci(16),

* ci(20), ci(21), ci(22), ci(23),
* ci(24), ci(25), ci(26), ci(27), ci(28)

800 format(1x,f7.0,12f5.2)

\[ b = ((3.14/ssize)**2.)*dl*ti(j) \]

\[ \text{call expcal(b, vop)} \]

\[ ca(j) = (1-((6.)/(3.14**2.))*vop)) \]

* (\text{ci}(j)-\text{csrain}(j))+\text{csrain}(j)

\[ ca(j) = (((\text{ssize}/2.0)**3.)* \]

* \text{ca}(j)+1.5*((\text{ssize}/2.)**2.)*

* sbwd*\text{ci}(j)/(((\text{ssize}/2.)+(sbwd/2.))**3.)

80 continue

write(2,18) t, ca(1), ca(2), ca(4),

* ca(6), ca(8), ca(10), ca(12), ca(14),

* ca(16), ca(18), ca(20)

18 format(1x,F7.0,11f6.2)

\[ \text{tprint} = \text{tprint} + dtpr \]

20 if (t-tstop).le.515,21,21

21 do 90 j = 1, nscount

\[ b = ((3.14/ssize)**2.)*dl*ti(j) \]

\[ \text{call expcal(b, vop)} \]
ca(J) = ((size/2.0)**3.)
* (ci(J) - c strain(J)) + c strain(J)
ca(J) = ((size/2.0)**2.)*
* sbwd * ci(J) / ((size/2.0) + (sbwd/2.0)**3.)
continue
write(2,30), ca(1), ca(2), ca(4),
* ca(6), ca(8), ca(10), ca(12), ca(14),
* ca(16), ca(18), ca(20)
30 FORMAT(1X,F7.0,11F6.2)
write(2,150) ci(20)
150 format(///f7.2)
area1=0.0
do 600 n=1,19
area1 = (ca(n) + ca(n+1))/2.0
area1 = area1 + area1
600 continue
area2=0.0
ncs=ncount-1
do 680 j=20,ncs
areaJ = 1.0 - ((ca(n) + ca(n+1))/2.0)
area2 = area2 + areaJ
680 continue
xinc = (area1 - area2)/(1.0 - ci(ncount))
xscale = xscale + xinc
write(3,610), xscale, xinc,
* ca(21), ca(22), ca(23), ca(24), ca(25),
* ca(26), ca(27), ca(28), ca(29)
610 format(1x,f7.0,2x,f6.2,1x,f5.2,1x,9f6.2)
stop
515 continue
tcounter = t + tau
if (tcounter - tsrate) 520, 530, 530
530 continue
do 580 j=1,ncount
b=((3.14/size)**2.)*dl*ti(j)
call expcal(b,vop)
ca(j) = (1-((6./(3.14**2.))*vop))
* (ci(j) - c strain(j)) + c strain(j)
ca(j) = ((size/2.0)**3.)*ca(j)
* +1.5*((size/2.)**2.)*
* sbwd * ci(j) / ((size/2.0) + (sbwd/2.0)**3.)
580 continue
write(2,640), ca(1), ca(2), ca(4),
* ca(6), ca(8), ca(10), ca(12), ca(14),
* ca(16), ca(18), ca(20)
640 format(1x,f7.0,11f6.2)
area1 = 0.0
do 500 n=1,19
area1 = (ca(n) + ca(n+1)) / 2.0
area1 = area1 + area1
continue
area2 = 0.0
jnsc = ncount - 1
do 670 j = 20, jnsc
area2 = 1.0 - ((ca(n) + ca(n+1)) / 2.0)
area2 = area2 + area2
continue
xinc = (area1 - area2) / (1.0 - ci(ncount))
xscale = xscale + xinc
write (3, 510) tscale, xinc,
* ca(21), ca(22), ca(23), ca(24), ca(25),
* ca(26), ca(27), ca(28), ca(29)
format (1x, f7.0, 2x, f6.2, 1x, f5.2, 9f6.2)
tsrage = ((tau**0.5) / 19.0 + (tsrate**0.5)**2.0)
ctrain(ncount+1) = ci(ncount)
ci(ncount+1) = ci(ncount)
ncount = ncount + 1
520 go to 24
24 DO 25 K = 1, 40
c(k) = ci(k)
continue
520 go to 26
end
subroutine expcal(b1, out)
out = 0.0
do 60 n = 1, 20
n2 = n**2
out = out + (exp(-b1*n2)/n2)
var2 = (out - (exp(-b1*n2)/n2))/out
if(var2 > 0.99) go to 70
continue
70 n = 1
end
APPENDIX B

Justification of Assumptions Made

While developing the mass balance ideas in Section 3.3, it is assumed that the amount of $O-16$ in the old scale involved in direct surface exchange with the tracer when the sample is oxidized in $O-18$ gas is negligible. The diffusion of $O-16$ atoms from the scale to the gas is due to the concentration gradient only and can be evaluated by setting the chemical potential gradient equal to zero. Area A in Figure 31 represents the amount of tracer retained in the old oxide due to inward grain boundary diffusion of tracer with exchange. Area B is the amount of $O-16$ in the old scale which undergoes direct surface exchange with a same amount of $O-18$ in the gas during the tracer oxidation. Area B is about 7% of area A. So the amount of tracer diffusing into the oxide due to direct surface exchange is not very significant.
FIGURE 31 - Comparison of direct surface exchange to total exchange.

0-16 oxidation time - 18 hours
0-18 oxidation time - 14 hours
at 1100°C

\[ D_b = 1.3 \times 10^{-16} \text{ m}^2/\text{sec} \]
\[ D = 10^{-20} \text{ m}^2/\text{sec} \]
grain size = 0.3 microns

(normalised depth \( \frac{Y}{X} \times 100 \))
The problem of tracer exchange between the grain boundaries and grains was solved by treating the grains as spherical sinks with a constant surface concentration $\phi_y$ (Section 3.5). This was done for mathematical convenience so that $\phi_y$ could be evaluated in Equation 3.8. However, $\phi_y$ is not a constant but is a function of time. However, the assumption of a constant grain surface tracer concentration is not very severe. As shown in Figure 32, the relatively high value of grain boundary diffusivity causes the grain surface tracer concentration to rapidly reach a near steady state value. There is only about an 8% variation in the grain surface tracer concentration when the grains are being filled with tracer due to exchange. Thus the assumption of a constant does not lead to a large error.
FIGURE 32 - The variation of grain surface and average grain concentration with time.

0-16 oxidation time-18 hours
0-18 oxidation time-14 hours
at 1100°C

\[ D_b = 1.3 \times 10^{-16} \text{ m}^2/\text{sec} \]
\[ D = 10^{-20} \text{ m}^2/\text{sec} \]

grain size = 0.3 microns
### Analysis Techniques for Tracer Studies of Oxidation

Soumendra Nath Basu

**Case Western Reserve University**  
Dept. of Metallurgy and Materials Science  
Cleveland, Ohio 44106

**National Aeronautics and Space Administration**  
Washington, D.C. 20546

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**Abstract**

Analysis techniques to obtain quantitative diffusion data from tracer concentration profiles were developed. Mass balance ideas were applied to determine the mechanism of oxide growth and to separate the fraction of inward and outward growth of oxide scales. The process of inward oxygen diffusion with exchange was theoretically modeled and the effect of lattice diffusivity, grain boundary diffusivity and grain size on the tracer concentration profile was studied. The development of the tracer concentration profile in a growing oxide scale was simulated. The double oxidation technique was applied to a FeCrAl-Zr alloy using O-18 as a tracer. SIMS was used to obtain the tracer concentration profile. The formation of lacy oxide on the alloy was discussed. Careful consideration was given to the quality of data required to obtain quantitative information.