POINT DEFECT EFFECTS ON HOT CORROSION
OF ZIRCONIA-BASED COATINGS

R. Reidy, D-H. Kim, and G. Simkovich

Depart of Materials Science and Engineering
The Pennsylvania State University
University Park, Pennsylvania 16802

Reproduction in whole or in part is permitted for any purpose of the United States Government. Distribution of this document is unlimited.
PENN STATE
College of Earth and Mineral Sciences

Undergraduate Majors
Ceramic Science and Engineering, Fuel Science, Metals Science and Engineering, Polymer Science;
Mineral Economics; Mining Engineering, Petroleum and Natural Gas Engineering;
Earth Sciences, Geosciences; Geography; Meteorology.

Graduate Programs and Fields of Research
Ceramic Science and Engineering, Fuel Science, Metals Science and Engineering, Polymer Science;
Mineral Economics; Mining Engineering, Mineral Processing, Petroleum and Natural Gas
Engineering;
Geochemistry and Mineralogy, Geology, Geophysics; Geography; Meteorology.

Universitywide Interdisciplinary Graduate Programs Involving EMS Faculty and Students
Earth Sciences, Ecology, Environmental Pollution Control Engineering, Mineral Engineering

Associate Degree Programs
Metallurgical Engineering Technology (Shenango Valley Campus).

Interdisciplinary Research Groups Centered in the College
C. Drew Stahl Center for Advanced Oil Recovery, Center for Advanced Materials. Coal Research
Section, Earth System Science Center, Mining and Mineral Resources Research Institute, Ore
Deposits Research Group.

Analytical and Characterization Laboratories (Mineral Constitution Laboratories)
Services available include: classical chemical analysis of metals and silicate and carbonate rocks;
X-ray diffraction and fluorescence; electron microscopy and diffraction; electron microprobe
analysis; atomic absorption analysis; spectrochemical analysis; surface analysis by secondary ion
mass spectrometry (SIMS); and scanning electron microscopy (SEM).
Thermal barrier coatings are vulnerable to certain types of hot corrosion: sulfidation and vanadic attack. Stabilized zirconia, an often used thermal barrier coating, is degraded by dissolution of the stabilizing component (e.g. Y$_2$O$_3$). To obtain the dissolution of the stabilizing components, mass transport in the coating must occur. The presence of point defects in a crystalline solid greatly affect the transport properties in that solid. The nature and concentration of these defects can be altered which, in turn, can impart large changes in the transport properties of a material (e.g. ionic conductivity and diffusion). In this study, we are determining the defect structures of yttria and ceria-stabilized zirconium oxides. Using electrical conductivity measurements, the activation energies of yttria-stabilized zirconia have been examined as a function of frequency, composition, and oxygen activity.

(see back)
Conductivity measurements on ceria-stabilized zirconia have shown anomalous results. At low oxygen activities, ZrO$_2$–CeO$_2$ samples did not reach equilibrium. The cause for this phenomenon has yet to be determined.

The transport properties of molten Na$_2$SO$_4$ have been investigated to aid to further understanding hot corrosion processes at 1173 K. An A.C. impedance technique for the total electrical conductivity, the potentiostatic polarization method for ionic transport numbers, and the steady state polarization method of Wagner and Hebb for the electronic conductivity were employed as a function of Na$_2$O activity in the melt by controlling the gas atmosphere.
ABSTRACT

Thermal barrier coatings are vulnerable to certain types of hot corrosion: sulfidation and vanadic attack. Stabilized zirconia, an often used thermal barrier coating, is degraded by dissolution of the stabilizing component (e.g. $\text{Y}_2\text{O}_3$). To obtain the dissolution of the stabilizing components, mass transport in the coating must occur. The presence of point defects in a crystalline solid greatly affect the transport properties in that solid. The nature and concentration of these defects can be altered which, in turn, can impart large changes in the transport properties of a material (e.g. ionic conductivity and diffusion). In this study, we are determining the defect structures of yttria and ceria-stabilized zirconium oxides. Using electrical conductivity measurements, the activation energies of yttria-stabilized zirconia have been examined as a function of frequency, composition, and oxygen activity. Conductivity measurements on ceria-stabilized zirconia have shown anomalous results. At low oxygen activities, $\text{ZrO}_2$-$\text{CeO}_2$ samples did not reach equilibrium. The cause for this phenomenon has yet to be determined.

The transport properties of molten $\text{Na}_2\text{SO}_4$ have been investigated to aid to further understanding hot corrosion processes at 1173 K. An A.C. impedance technique for the total electrical conductivity, the potentiostatic polarization method for ionic transport numbers, and the steady state polarization method of Wagner and Hebb for the electronic conductivity were employed as a function of $\text{Na}_2\text{O}$ activity in the melt by controlling the gas atmosphere.
INTRODUCTION

Due to the severity of their operating environments, marine turbines are subject to a variety of corrosive mechanisms. One of the most formidable forms of attack is hot corrosion, a chemical interaction leading to a breakdown of a protective oxide scale.(1) There are two dominant types of hot corrosion: sulfidation and vanadium attack.

In sulfidation, NaCl reacts with sulfur found in the fuel to form Na$_2$SO$_4$. The sodium sulfate reacts with the protective oxide scale resulting in scale failure.(1,2) A large body of work details the roles of Na$_2$SO$_4$ and NaCl in the destruction of the oxide scale.(1-8)

Vanadium enters the turbine as a fuel impurity in organic and inorganic forms. These compounds react with oxygen to form oxides (VO, VO$_2$, VO$_3$, and V$_2$O$_5$) and hydroxides (V$_2$O$_7$H$_4$, VO(OH)$_3$, and VO$_2$(OH)$_2$).(9) These gases either condense onto the turbine blades and react with sodium compounds to form sodium vanadates as in equations 1-3),

\[
2\text{VO(OH)}_3(g) = \text{V}_2\text{O}_5(l) + 3\text{H}_2\text{O}(g) \quad 1)
\]
\[
\text{V}_2\text{O}_5(g) = \text{V}_2\text{O}_5(l) \quad 2)
\]
\[
\text{V}_2\text{O}_5(l) + \text{Na}_2\text{O}(l) = 2\text{NaVO}_3(l) \quad 3)
\]

or react with sodium compounds in the gas phase forming sodium vanadates which condense onto the turbine blades as in equation 4).

\[
\text{VO(OH)}_3(g) + \text{NaOH}(g) = \text{NaVO}_3(l) + 2\text{H}_2\text{O}(g) \quad 4)
\]

In addition to Na$_2$SO$_4$, Luthra and Spacil found both V$_2$O$_5$ and NaVO$_3$ in the condensate; therefore, it is difficult to determine which compound is more destructive to the oxide.(9) Further, one can not ignore the effects of Na$_2$SO$_4$ in a separate or combined attack on the oxide scale.
The mechanisms by which the sulfur or vanadium compounds disrupt protective oxide scales of nickel-based superalloys in marine turbine hot zones are detailed in several literature surveys. (1, 7, 10) Bornstein and DeCrescente (2) and Jones (8) describe a general mechanism referred to as the salt fluxing or acid-base reaction model. In sulfidation, Bornstein explains that the oxide scales are insoluble in stoichiometric sodium sulfate, but due to the reaction,

\[ \text{Na}_2\text{SO}_4(\text{l}) = \text{Na}_2\text{O} + \text{SO}_3 \]

(molten salt) (base) (acid)

the oxide may dissolve in the base as an anion or in the acid as a cation. (1)

From the Lux-Flood theory (11), the activities of the acidic and basic components in the molten salt are fixed by the dissociation constant of the reversible reaction 5). (8) In vanadium attack, a similar dissolution mechanism utilizes the Lewis definition of acids (electron acceptors) and bases (electron donors). Jones describes the ability of vanadium pentoxide to behave as a Lewis acid and react with a protective oxide. (8)

\[ \text{Y}_2\text{O}_3(\text{s}) + \text{V}_2\text{O}_5(\text{l}) = 2\text{YVO}_4(\text{l}) \]

(base) (acid) (salt)

This dissolution mechanism involves the transport of electrons from the stabilizing component to the corrosive \( \text{V}_2\text{O}_5 \). A greater understanding of this transport phenomenon could yield the ability to retard or halt this dissolution.

**Defect Chemistry**

Rarely in nature does one find a perfect crystal. Instead, a number of disruptions occur in the periodic array of atoms. There are four general types of structural imperfections: point defects, line defects, plane defects, and
volume defects. In crystalline solids, the nature and concentration of point defects greatly affect transport properties. The type and concentration of point defects in an oxide is often dependent on the temperature, pressure, and chemical potentials of the oxide components. Atomic defects (vacancies, interstitials, impurities, or misplaced atoms) affect solid state diffusion and non-stoichiometry of compounds. Changes in these properties, in turn, alter reaction rates, ionic conductivities, and sintering rates. Electronic defects (electrons and electron holes) determine electrical conductivities as well as other properties. Therefore, a detailed understanding of the point defect behavior of a compound would provide great insight or even allow one to predict the transport phenomena in that material. Additionally, one may alter the transport properties of a compound by changing the nature or concentration of the point defects within it.

The effects of point defects on hydrometallurgical processes, such as flotation and dissolution, have been the subject of many studies. In lead sulfide (PbS) which shows p-type conduction at high $S_2$ partial pressures and n-type at low $S_2$ partial pressures, additions of $Ag_2S$ impart more p-type behavior while $Bi_2S_3$ doping creates a larger n-type contribution. When the reaction rate is controlled by the interfacial reaction, Simkovich and Wagner found that the dissolution rates in nitric acid of PbS containing $Ag_2S$ are greater than both pure PbS and $Bi_2S_3$-doped PbS. Similar effects of dopant-induced changes in dissolution rates in NiO (17) and ZnO (18) have been documented. Some alterations in the defect structure of a base material which result in changes in dissolution behavior are an increased concentration of electron holes in Li$_2$O-doped NiO (19) and an increase in zinc interstitials in Li$_2$O-doped ZnO (20). Simkovich and Wagner detail other systems and the mechanisms by which changes in defect concentration directly affect dissolution rates.
Thermal Barrier Coatings

In recent years, efforts have been made to apply protective ceramic coatings to turbine blades to improve resistance to hot corrosion as well as to increase engine operating temperatures. These higher operating temperatures are attained because these coatings have high melting points and low thermal conductivities, thus protecting the more temperature sensitive alloy beneath. These materials are referred to as thermal barrier coatings.

With additions of various stabilizers, zirconia has been used as a thermal barrier coating. Due to a relatively high coefficient of thermal expansion (as compared to other ceramics), $\text{ZrO}_2$ minimizes thermal expansion differences between metal substrate and coating. However, the primary reason for considering stabilized $\text{ZrO}_2$ as a thermal barrier coating is its resistance to many forms of corrosion.

Pure zirconium oxide is a polymorphic compound. Three different stable forms of zirconia exist: cubic (fluorite structure, 2370 - 2680°C), tetragonal (1170 - 2370°C), and monoclinic (below 1170°C). The phase transition from tetragonal to monoclinic has caused considerable concern. Early studies have determined that the transformation is martensitic and does not occur at a fixed temperature, but over a temperature range, and involves a large (about 9%) volume expansion. This volume increase causes cracking in the $\text{ZrO}_2$ when cooled below the transition; therefore, traditional cooling methods to below 1170°C can result in crumbling of the zirconia.

The addition of $\text{CaO}$, $\text{MgO}$, $\text{Y}_2\text{O}_3$, and $\text{CeO}_2$ to zirconia lowers the transition temperatures of both solid state phase transformations. Stubican and Hellmann have reviewed the binary oxide phase diagrams and have shown a partially stabilized form (a mixture of cubic and tetragonal or monoclinic phases) and a fully stabilized zirconia (cubic phase). Both the cubic and mixed phases are stable at room temperature.

Wagner demonstrated that stabilized zirconia contains oxygen ion vacancies. In calcia-stabilized zirconia, Hund discovered that calcium
and zirconium ions are distributed statistically in the cation sites while an oxygen vacancy concentration equal to the calcium dopant concentration provides electroneutrality. Using x-ray diffraction methods, Tien and Subbarao have determined similar defect behavior in other stabilized zirconias. Other investigators have concluded that the defect structure and concentration are fixed by dopant content. Alcock has found that the electronic defect structure of stabilized zirconia is affected by changes in atmosphere and temperature.

The primary thrust of this work is to determine the effect of point defects on the mechanisms and transport properties involved in the hot corrosion of zirconia coatings. It is prudent to briefly review the recent efforts to study these mechanisms by other methods. The sulfidation and vanadium attack studies by Jones and Williams demonstrated that ZrO₂ is substantially more resistant to hot corrosion than the dopants, Y₂O₃, CeO₂, and HfO₂. In other work, Jones discusses the degradation of the coating with respect to the acid-base reaction model. The V₂O₅ reacts with the Y₂O₃ in the zirconia and forms a yttrium vanadate (equation 5). The removal of yttria destabilizes the zirconia allowing the more voluminous monoclinic phase to form and leading to surface cracking of the coating. As the protective scale is thermally cycled, the cracks propagate and extend into the coating causing spallation. This mechanism has been put forth by a number of other investigators.

Also included in this report is a summary of recently completed work on transport properties in molten sodium sulfate which was in part supported by this contract. The transport properties of a Na₂SO₄ melt has been investigated to aid in further understanding hot corrosion processes at 1173 K. An A. C. impedance technique for the total electrical conductivity, the potentiostatic polarization method for ionic transport numbers, and the steady state polarization method of Wagner and Hebb for the electronic conductivity were carried out as a function of Na₂O activity in the melt by controlling the gas atmosphere.
EXPERIMENTAL

Electrical and Ionic Conductivity

The electrical conductivity will be measured by a digital bridge using a two-point probe or a four-wire technique. The conductivity measurements will be taken at a number of temperatures (700-1100°C) and oxygen partial pressures. In addition, an Arrhenius plot of conductivity and temperature can estimate the activation energy of conduction. By definition, conductivity is the product of the charge, the concentration of charge carriers, and the mobilities of those carriers. In an electrical conductivity measurement, the carriers can be either electronic or ionic.

To isolate the ionic contribution, potentiostatic polarization techniques can estimate the ionic transport numbers (the fraction of the total conductivity which is carried by ions).

\[
\Delta G = \Delta G^\circ + RT \ln Q
\]

\[
= (G^\circ_{\text{NO}} + RT \ln a_{\text{NO}}^\circ) + (G^\circ_{\text{O}_2} + RT/2 \ln a_{\text{O}_2})
\]

\[
- (G^\circ_{\text{NO}} + RT \ln a_{\text{NO}}^\circ) - (G^\circ_{\text{O}_2} + RT/2 \ln a_{\text{O}_2})
\]

\[
= RT \ln (a_{\text{NO}}^\circ/a_{\text{NO}}^\circ) + RT/2 \ln (P_{\text{O}_2}/P_{\text{O}_2}^\circ)
\]

\[
\Delta G = -nFE
\]

\[
= -NFE = RT \ln (a_{\text{NO}}^\circ/a_{\text{NO}}^\circ) + RT/2 \ln (P_{\text{O}_2}/P_{\text{O}_2}^\circ)
\]

therefore,

\[
E = (RT/nF) \ln (a_{\text{NO}}^\circ/a_{\text{NO}}^\circ) + (RT/2nF) \ln (P_{\text{O}_2}/P_{\text{O}_2}^\circ)
\]
Eq. 11 applies when all current is carried by ionic species or vacancies. However, if any electronic transport exists, then the right side of Eq. 11 must be multiplied by $t_i$, the ion transport number.

$$E = t_i \left[ \frac{RT}{N_f} \ln \left( \frac{a_{Mo}'}{a_{Mo}''} \right) + \frac{RT}{2nF} \ln \left( \frac{P_{O_2}''}{P_{O_2}'} \right) \right]$$ \quad 12)

Although the metal oxide activities are not known, they can be assumed to be nearly equal in a small oxygen potential gradient. Consequently,

$$a_{Mo}' \approx a_{Mo}''$$

$$E = E_{meas} = t_i \left[ \frac{RT}{2nF} \ln \left( \frac{P_{O_2}''}{P_{O_2}'} \right) \right]$$ \quad 13)

and

$$E = E_{max} = \frac{RT}{2nF} \ln \left( \frac{P_{O_2}''}{P_{O_2}'} \right)$$ \quad 14)

$E_{max}$ is calculated from experimental conditions, and $E_{meas}$ is the measured emf. The value of $t_i$ is the ratio of these results.

$$t_i \approx \frac{E_{meas}}{E_{max}}$$ \quad 15)

Wagner-Hebb Polarization Experiments

The electronic contributions will be determined by Wagner-Hebb type polarization method.\(^{(30-31)}\) This cell can be schematically represented as:

```
(negative) | (positive)
Zr | Stabilized | inert
electrode | Zirconia | electrode
```

The appropriate choice of electrodes enables the suppression of either electronic or ionic migration.\(^{(51,52)}\) The partial conductivity of the electronic species can be determined from the ratio of the current density and the potential gradient.\(^{(53,54)}\) Under steady state conditions, Wagner has derived equations for the total electronic current through the polarization cell.

$$I_{elec} = I_e + I_0$$ \quad 16)

$$= \frac{RTA}{LF} \left( \sigma_e [1 - \exp(-EF/RT)] \right)$$
Dividing eq. 17 by \[ [1 - \exp(-\frac{EF}{RT})] \] and rearranging yields:

\[
I_{\text{elec}} \cdot \frac{RTA/LF}{[1 - \exp(-\frac{EF}{RT})]} = \sigma_e^0 + \sigma_e^0 \exp(\frac{EF}{RT})
\]

Plotting the left side of eq. 18 versus \( \exp(\frac{EF}{RT}) \), the resulting line has \( \sigma_e^0 \) as its slope and \( \sigma_e^0 \) as its intercept.\(^{(53,54)} \) These quantities combined with the total electrical conductivity allow determination of the electronic transport numbers.

RESULTS AND DISCUSSION

Materials

In order to accurately describe the point defect structure of a material, the effects of macroscopic flaws such as pores and cracks should be reduced or, if possible, eliminated. As stated earlier, most commercial zirconia powders that were available did not fully densify during sintering. After a number of attempts, only one commercial powder had the necessary purity and could be fully densified (achieve densities 95% of theoretical values). Samples fabricated from other powders gave unreliable results. In order to control densities, composition, and purity, powders were synthesized by coprecipitation reactions. Using a protocol adapted from the work by Ciftcioglu and Mayo \(^{(48)} \), nano-crystalline powders were prepared. By controlling sintering temperatures and times, one should be able to fix grain size in samples. At present, such control has proved difficult. As was shown in Fig. 1, the electrical properties of these ultra-fine grained samples seemed very different from larger grained samples despite having the same composition. A coprecipitation procedure for CeO\(_2\)-ZrO\(_2\) by Duh et al. \(^{(49)} \) may also be adapted to yield nano-crystalline powders.
Electrical Measurements

Electrical conductivity measurements for 3 and 8 mole percent yttria-stabilized zirconia have provided some interesting results. As a function of temperature, conductivity behaved in accordance with the literature values (the activation energies were approximately 1 eV)(50)(Fig.2). However, as a function of oxygen activity, the results did not follow behavior described by other researchers. Previous work has found for a number of compositions in yttria-stabilized zirconia that oxygen does not significantly affect conductivity except at very low oxygen partial pressures.(50) Our results have shown conductivity to increase as a function of $P_{O_2}$ although only slightly (Fig.3). This behavior has been reproduced in a number of experiments.

Conductivity measurements on ceria-stabilized zirconia have also shown anomalous results. At low oxygen activities, $\text{ZrO}_2-\text{CeO}_2$ samples did not reach equilibrium. After well over 100 hours, the electrical conductivity continued to increase with no hint of leveling off (Fig.4). The cause for this phenomenon has yet to be determined. However, high temperature x-ray diffraction studies are currently underway to discover the effect of these low oxygen activities on the $\text{ZrO}_2-\text{CeO}_2$ structure.

$\text{Na}_2\text{SO}_4$ Studies

A large thrust of this experimental program was to obtain the transport properties in the aggressive molten salt $\text{Na}_2\text{SO}_4$. Table I list such information obtained from this study. The total electrical conductivity determined by utilizing an A.C. impedance technique indicates that pure $\text{Na}_2\text{SO}_4$ melt had a somewhat low total conductivity as compared to most ionic melts.

The cationic transport numbers of a pure $\text{Na}_2\text{SO}_4$ melt were obtained by utilizing the potentiostatic polarization technique. The results indicate that a pure $\text{Na}_2\text{SO}_4$ melt is a cationic conductor over a wide range of $\text{Na}_2\text{O}$ activities.
at 1173 K. From the above results, the diffusivity of cations, one of the important kinetic parameters, can be estimated as $8.1 \times 10^{-6}$ cm$^2$/sec.

The Wagner-Hebb type polarization experiments provided the electron and electron hole conductivities in a pure Na$_2$SO$_4$ melt at 1173 K. From these measurements along with total electrical conductivity data, the transport numbers of electrons, $t_e$, and electron holes, $t_+$, were calculated. Such experimental investigations show that the electronic conduction occurs primarily via the transport of electrons as indicated in Table I.

### Table I
Transport Properties of a Pure Na$_2$SO$_4$ Melt

<table>
<thead>
<tr>
<th>Ionic Transport Properties</th>
<th>Electronic Transport Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\sigma_{\text{total}} = 0.232$ (n-cm)$^{-1}$</td>
<td>$\sigma_e = 1.638 \times 10^{-3}$ (n-cm)$^{-1}$</td>
</tr>
<tr>
<td>$t_{\text{Na}^+} = 0.9838$</td>
<td>$t_+ = 7.055 \times 10^{-3}$</td>
</tr>
<tr>
<td>$D_{\text{Na}^+} = 8.083 \times 10^{-6}$ (cm$^2$/sec)</td>
<td>$t_e = 5.76 \times 10^{-5}$</td>
</tr>
<tr>
<td>$\mu_{\text{Na}^+} = 7.996 \times 10^{-5}$ (cm$^2$/volt-sec)</td>
<td>$D_e = 3.15 \times 10^{-3}$ (cm$^2$/sec)</td>
</tr>
<tr>
<td>$C_{\text{Na}^+} = 0.0294$ (mole/cm$^3$)</td>
<td>$\mu_e = 0.031$ (cm$^2$/volt-sec)</td>
</tr>
<tr>
<td></td>
<td>$C_e = 5.462 \times 10^{-7}$ (mole/cm$^3$)</td>
</tr>
</tbody>
</table>

The diffusion coefficient of electrons which are the predominant minor defects in a Na$_2$SO$_4$ melt, was determined by the voltage relaxation method suggested by Weiss (49). Its average value is $3.15 (\pm 0.17) \times 10^{-3}$ cm$^2$/sec. The corresponding drift mobility of electrons is $0.031$ cm$^2$/volt-sec which is identical to that in Na-NaCl melts at 1166 K with 0.81 mole percent of sodium.
The effects of Cr$_2$O$_3$, Al$_2$O$_3$, and SiO$_2$ on total conductivity as well as the electronic conductivity were examined by the A.C. impedance technique and Wagner-Hebb type polarization experiments. The introduction of Cr$_2$O$_3$ at concentration levels of $10^{-3}$, $10^{-2}$, $10^{-1}$, and 1 mole percent has significant effects on the total electrical conductivity i.e., as the amount of Cr$_2$O$_3$ in the melt increases the total electrical conductivity decreases. Additionally, the dissolved chromia decreases electron conductivities and increases electron hole conductivities. The addition of Al$_2$O$_3$ into the melt does not produce significant changes in total electrical conductivities. However, the Al$_2$O$_3$ lowers electron conductivities tremendously and increases electron hole conductivities slightly. The melts containing silica show small changes in total electrical conductivities but massive decrements in electron conductivities.

The oxidation and/or hot corrosion rate is often considered as a product of total conductivity, sum of ionic transport numbers, and the transport number of major electronic species. The results observed in the present investigation indicate the sum of ionic transport numbers is nearly unity; therefore, the product of total conductivity and the transport number of the major electronic species is considered. Such considerations are described in Fig.5 as a function of mole percent of oxides in the melt. The value of total conductivity x transport number of major electronic species for a pure Na$_2$SO$_4$ is $1.64 \times 10^{-3}$. As indicated in this figure the reaction rate is decreased as a function of mole percent of each oxide in the melt. For Cr$_2$O$_3$ and Al$_2$O$_3$ melts, it levels off at about $10^{-1}$ mole percent while SiO$_2$ melts do not level off even in supersaturated melts. It is clear from this figure that the addition of silica has the utmost effect on decreasing reaction rates among those protective oxides tested.
CONCLUSIONS AND SUMMARY

To obtain the dissolution of the stabilizing components, mass transport in the coating must occur. The presence of point defects in a crystalline solid greatly effect the transport properties in that solid. The nature and concentration of these defects can be altered which, in turn, can impart large changes in the transport properties of a material (eg. ionic conductivity and diffusion). In this study, we are determining the defect structures of yttria and ceria-stabilized zirconium oxides. Using electrical conductivity measurements, the activation energies of yttria-stabilized zirconia have been examined as a function of frequency, composition, and oxygen activity. Conductivity measurements on ceria-stabilized zirconia have shown anomalous results. At low oxygen activities, \( \text{ZrO}_2-\text{CeO}_2 \) samples did not reach equilibrium. The cause for this phenomenon has yet to be determined. However, high temperature x-ray diffraction studies are currently underway to discover the effect of these low oxygen activities on the \( \text{ZrO}_2-\text{CeO}_2 \) structure.

It was observed that the total electrical conductivity of pure \( \text{Na}_2\text{SO}_4 \) was of the order of \( 2.32 \times 10^{-1} \) (ohm-cm)\(^{-1} \) and varied only slightly with changes in the activities of \( \text{Na}_2\text{O} \). The cationic transport number of a pure \( \text{Na}_2\text{SO}_4 \) melt was found to be about 0.984 at 1173 K and was almost a constant as a function of \( \text{Na}_2\text{O} \). The partial conductivity of electrons in \( \text{Na}_2\text{SO}_4 \) was about two orders of magnitude less than the total electrical conductivity. Thus, the transport number of electrons, \( t_e \), is of the order of \( 10^{-3} \) in a pure \( \text{Na}_2\text{SO}_4 \) melt at 1173 K. The self-diffusion coefficient of \( \text{Na}^+ \) ions was estimated utilizing the above experimental data and its average value was \( 8.1 \times 10^{-6} \) cm\(^2\)/sec.

The transient relaxation method was employed to determine the diffusion coefficient of electrons in a pure \( \text{Na}_2\text{SO}_4 \) melt and its value was about \( 3.15 \times 10^{-3} \) cm\(^2\)/sec at 1173 K.
The introduction of the protective oxides such as $\text{Cr}_2\text{O}_3$, $\text{Al}_2\text{O}_3$, and $\text{SiO}_2$ into a $\text{Na}_2\text{SO}_4$ melt has significant changes in transport properties of the melt. These oxides decrease the electron conductivities and increase the electron hole conductivities as compared to pure $\text{Na}_2\text{SO}_4$. 
REFERENCES

1. Bornstein, N. S., Literature Review of Inhibition of Vanadate Attack, ONR N00014-88-M-0013, June 1988

2. Bornstein, N. S. and M.A. DeCrescente, Corrosion, 26, 7, 209-14, July 1988


7. Shores, D., D. McKee, and T. Kerr, Sodium Chloride Induced Hot Corrosion: Literature Survey and Preliminary Experiments, Contract 7A003-CIP105 (EPNHT05), April 1976


10. Fitzer, E. and J. Schwab, Corrosion, 12, 49-54, 1956


33. Wagner, C., Naturwissenschaften, 31, 265-68, 1943


Figure 1 AC conductivity (1kHz) vs temperature for two different grain sizes.
Figure 2  AC conductivity (1kHz) for 8wt% Y₂O₃-ZrO₂ as a function of temperature

activation energy = 1.008eV from slope
Figure 3  AC conductivity (1kHz) for 3 mole% $\text{Y}_2\text{O}_3$-$\text{ZrO}_2$ as a function of oxygen partial pressure.
Figure 4  AC conductivity (100 hz) vs time for 26 wt% CeO$_2$-ZrO$_2$ at 700°C
Figure 5  Average value of total conductivity times transport number of electronic species as a function of mole% of oxides added in the melt at 1173 K