I. Introduction

Non-oxide ceramics such as silicon carbide (SiC) and silicon nitride (Si₃N₄) are promising materials for a wide range of high temperature applications. These include such diverse applications as components for heat engines, high temperature electronics, and re-entry shields for space vehicles. Table I lists a number of selected applications. Most of the emphasis here will be on SiC and Si₃N₄. Where appropriate, other non-oxide materials such as aluminum nitride (AlN) and boron nitride (BN) will be discussed. Proposed materials include both monolithic ceramics and composites. Composites are treated in more detail elsewhere in this volume, however, many of the oxidation/corrosion reactions discussed here can be extended to composites. In application these materials will be exposed to a wide variety of environments. Table I also lists reactive components of these environments.
It is well-known that SiC and Si₃N₄ retain their strength to high temperatures. Thus these materials have been proposed for a variety of hot-gas-path components in combustion applications. These include heat exchanger tubes, combustor liners, and porous filters for coal combustion products. All combustion gases contain CO₂, CO, H₂, H₂O, O₂, and N₂. The exact gas composition is dependent on the fuel to air ratio or equivalence ratio. (Equivalence ratio (EQ) is a fuel-to-air ratio, with total hydrocarbon content normalized to the amount of O₂ and defined by EQ=1 for complete combustion to CO₂ and H₂O). Figure 1 is a plot of equilibrium gas composition vs. equivalence ratio. Note that as a general rule, all combustion atmospheres are about 10% water vapor and 10% CO₂. The amounts of CO, H₂, and O₂ are highly dependent on equivalence ratio.

Other proposed applications for SiC include high temperature semi-conductors and for AIN include electronic substrates. In these situations, high temperature oxidation behavior is a prime issue. Re-entry shields have long been a useful application of ceramic materials—here the environment is a complex mixture of atoms and molecules. Finally a growing area is the application of ceramics as pump components and ball bearings. The environment depends on the fluid of interest, but is generally high temperature/high pressure water.
In this chapter we will discuss the interaction of non-oxide ceramics with some representative environments. We begin with pure oxidation which is important for nearly all the applications and then proceed to the more system-specific environments such as molten salts and high pressure water.

II. High Temperature Oxidation

Figure 2 summarizes oxidation behavior for a range of non-oxide ceramics (14). These data are presented in terms of recession in one hundred hours. Clearly long-term operation requires very low recession. The key oxidation reactions can be summarized as follows:

1. Silica formers—SiC, Si₃N₄, MoSi₂

   \[ \text{MSi} + \frac{3}{2} \text{O}_2(g) = \text{SiO}_2 + \text{MO} \]  \hspace{1cm} (1)

   The oxidation of SiC and Si₃N₄ is described in detail in the following section.

2. Alumina formers—primarily alloys, but also AlN

   \[ 2 \text{Al} + \frac{3}{2} \text{O}_2(g) = \text{Al}_2\text{O}_3 \]  \hspace{1cm} (2)

   The underline indicates aluminum is at less than unity activity, as would be found in an alloy.

   \[ 2 \text{AlN} + \frac{3}{2} \text{O}_2(g) = \text{Al}_2\text{O}_3 + \text{N}_2(g) \]  \hspace{1cm} (3)

   The oxidation of AlN is discussed more fully in the following section.
3. Borides

\[ 2 \text{MB} + \frac{5}{2} \text{O}_2(g) = \text{B}_2\text{O}_3(l) + 2\text{MO} \]  \hspace{1cm} (4)

4. Carbides

\[ \text{MC} + \text{O}_2(g) = \text{MO} + \text{CO}(g) \]  \hspace{1cm} (5)

5. Nitrides

\[ \text{MN} + \text{O}_2(g) = \text{MO} + \text{NO}(g) \]  \hspace{1cm} (6)

or

\[ 2 \text{MN} + \text{O}_2(g) = 2 \text{MO} + \text{N}_2(g) \]  \hspace{1cm} (7)

Most of these materials oxidize according to a parabolic rate law, but some oxidize according to linear kinetics. Hence Figure 2 gives oxidation kinetics in terms of recession to account for both kinetic laws. The reader is referred to (14) and references contained therein for more details on the kinetics of these reactions.

The important point from Figure 2 is that only materials which form protective silica or alumina films are useful for long times in oxidizing environments. Most of the borides, carbides, and nitrides do not form protective metal oxide scales. Hence the focus of this chapter will be on silica forming ceramics, SiC and Si$_3$N$_4$, ...
and the alumina forming ceramic, AlN. We shall also include a brief discussion of BN, which is an important material in many areas of technology.

II.A. Oxidation of SiC and Si$_3$N$_4$

II.A.1. Oxygen Transport in Silica

The low oxidation rates of silica-forming ceramics are due to remarkably low oxygen transport rates in silica. Thus we begin with a brief discussion of oxygen transport in silica. Amorphous silica, found for short-time oxidation or in very clean systems, consists of a random network of Si-O tetrahedra as shown schematically in Figure 3a (15). Cristobalite, the crystalline form of silica most often observed after long-term oxidation or oxidation in less clean environments, is composed of the same Si-O tetrahedra but arranged in an orderly structure as shown in Figure 3b.

Oxygen transport through these structures occurs by two different mechanisms. First, molecular oxygen can move through the interstices between the tetrahedra by a permeation mechanism. Alternatively, ionic oxygen can move from network site to network site by a bond breaking exchange mechanism. The permeation mechanism requires less activation energy since bond breaking is not required. Permeation through cristobalite is expected to be slower than through
amorphous silica since the regular structure is restricted to six member oxygen rings whereas the irregular structure found in amorphous silica allows for seven and eight member rings (15).

Permeation rates of molecular oxygen in amorphous silica have been determined (16), however, the corresponding rates in cristobalite have not been measured. The measurements are difficult since bulk cristobalite is not available and must be nucleated and characterized at temperature. In addition, the presence of short circuit transport paths along cristobalite grain boundaries and cracks formed from the β to α cristobalite phase transformation upon cooling make the measurement of intrinsic oxygen transport rates in cristobalite impossible.

II.A.2. Oxidation of silica forming ceramics—Experimental Techniques

Oxidation kinetics for these materials are determined by measuring either weight changes using a sensitive thermogravimetric balance as shown in Figure 4 or oxide thickness changes using optical techniques. An example of weight change versus time for SiC oxidation in dry oxygen is shown in Figure 5. Since the oxidation rate is controlled by diffusion of oxygen through the silica scale, the weight change decreases with time according to a parabolic law and is described by the parabolic rate constant, $k_p$. Accurate determinations of oxidation kinetics are more difficult for silica forming materials than for other metal-oxide forming
materials because silica grows at such a slow rate, and because silica growth kinetics are affected by even small amounts of impurities in the sample or oxidation environment (17). A comparison of the parabolic oxidation rates for silicon, SiC, and Si₃N₄ as a function of temperature is shown in Figure 6. Similarities and differences in the oxidation kinetics of these materials are discussed in the following sections.

It is instructive to begin with the relatively simple case of silicon oxidation. We will then focus on very pure SiC and Si₃N₄ in order to understand the fundamental oxidation behavior of these materials. These pure materials are also important in microelectronic applications.

II.A.3. Oxidation of Silicon

Silicon oxidation has been studied in great detail due to its application in microelectronic devices. The model of Deal and Grove (18) has been used to successfully describe silicon oxidation behavior at all but the shortest times. This model considers two possible rate limiting steps for the oxidation of silicon. First the reaction of oxygen with silicon according to a linear rate law controls the oxidation rate at short times or for thin scales. Second, transport of oxygen through the growing silica scale controls the oxidation rate according to a parabolic rate law for long times and for thick scales. These two rate laws are combined into a single expression that is valid for all times:
where \( X_0 \) is the oxide thickness, \( t \) is time, \( \tau \) is the offset time which corrects for the presence of an initial oxide layer, \( B \) is the parabolic rate constant and \( B/A \) is the linear rate constant. At short times this expression simplifies to the linear rate law \( x_0 = (t+\tau)B/A \) while at long times the expression simplifies to the parabolic law \( x_0^2 = Bt \). For long-term applications of silica forming materials, the parabolic law usually describes the oxidation kinetics adequately.

Deal and Grove have derived the parabolic rate constant in terms of the properties of silica, i.e.:

\[
B = 2D_{\text{eff}}C^*/N
\]  

(9)

and

\[
C^* = kP^n
\]  

(10)

where \( D_{\text{eff}} \) is the effective diffusivity of oxidant in silica, \( C^* \) is the equilibrium concentration of oxidant in the oxide, \( N \) is the number of oxidant molecules incorporated into a unit volume of the oxide layer, \( P \) is the oxidant pressure, \( n \) is a power law exponent, and \( k \) is a constant. The parabolic rate constant was found to vary linearly with the oxidant pressure for both oxygen and water vapor, i.e., \( n=1 \), indicating that the oxidant did not dissociate and molecular permeation through the silica is the rate limiting step for oxidation in the parabolic regime.

II.A.4. Oxidation of SiC
The oxidation of SiC is similar to the oxidation of silicon since a silica scale forms in both cases. In this case, however, the gases generated, as shown in the following oxidation reactions, will cause some differences:

$$\text{SiC} + \frac{3}{2}\text{O}_2(g) = \text{SiO}_2 + \text{CO}(g) \quad (11)$$

$$\text{SiC} + 2\text{O}_2(g) = \text{SiO}_2 + \text{CO}_2(g) \quad (12)$$

The linear reaction rate is different since the oxidation of C to CO(g) or CO$_2$(g) occurs in addition to the oxidation of Si. The parabolic oxidation rate could be different if the outward transport of CO or CO$_2$, rather than the inward transport of oxygen, limited the oxidation rate. It has been shown that the oxidation rates of SiC are about a factor of two slower than silicon (19-21) due to the extra consumption of oxygen in the reaction with C, as predicted (22). In addition, the activation energy for oxidation (20) is nearly identical to that of silicon (18) (see Figure 6) and the permeation of molecular oxygen through silica (16). Finally, the oxidation rate of SiC is found to depend on the oxygen partial pressure (23). Therefore, it is generally agreed that oxygen transport inward is the rate limiting step for parabolic oxidation of SiC. This issue has been discussed more fully in other references (24, 25).

The dependence of the parabolic rate constant on the oxygen partial pressure, given by the power law exponent, $n$, in Equation 10 above, gives information about the type of oxygen transport occurring in the silica. For $n=1$ molecular permeation of oxygen occurs. For $n<1$, some dissociation of oxygen into a
charged species occurs and network diffusion of oxygen by a bond breaking process is likely. Zheng et al. (23) have determined the power law exponent for the oxidation of SiC to vary between 0.6 and 0.3 at temperatures from 1200 to 1500°C. This implies that some combination of permeation and network diffusion limits the oxygen transport through the silica scale grown on SiC, with network diffusion increasing with temperature. This is confirmed by 18O tracer diffusion studies which show isotope exchange with network oxygen becomes increasingly important as the temperature is increased (26). However, the similar activation energies for molecular oxygen permeation of silica (16) and for oxidation of SiC up to temperatures of 1500°C (20) indicates that oxygen transport is dominated by the permeation mechanism.

The effect of crystallization of silica on SiC oxidation is a complex topic. Some investigators have used crystallization to explain observed rate changes and/or activation energy changes. However, current evidence indicates that many of these deviations in rate law and activation energies can be explained by impurities in either the environment or the SiC (17, 20, 27, 28). Ogbuji (29) has shown that silica scales fully crystallized during argon anneals at 1300°C do result in slower oxidation rates by about a factor of 30, but that fully crystalline scales are not found in actual practice since amorphous silica is continually formed during the course of SiC oxidation.

II.A.5. Oxidation of Si₃N₄
Like SiC, the oxidation of Si$_3$N$_4$ results in the formation of a silica layer and the generation of gaseous products according to the following simplified reaction:

$$Si_3N_4 + 3O_2(g) = 3SiO_2 + 2N_2(g) \quad (13)$$

An additional complication, however, is the formation of a suboxide layer of amorphous silicon oxynitride of variable stoichiometry (30, 31). The oxidation reaction can be written as (32):

$$Si_3N_4(1-x) O_{6x} + 3 \delta x O_2(g) = Si_3N_4(1-x-\delta x) O_{6(\delta x+\epsilon)} + 2 \delta x N_2(g) \quad (14)$$

for $x$ varying between 0 and 1. Measurements of the oxidation kinetics of Si$_3$N$_4$ have shown they are parabolic, but significantly slower than oxidation of both silicon (33) and SiC (20) to temperatures of 1500°C. Pure Si$_3$N$_4$ is the slowest oxidizing material known today. The higher activation energy for Si$_3$N$_4$ reflects the additional energy required in the breaking of bonds for the nitrogen-oxygen substitution reaction. The oxidation rates were found to be dependent on the oxygen partial pressure, but independent of the nitrogen partial pressure (33). In this case, the oxidation reaction is limited by oxygen transport and reaction in the oxynitride layer (32) rather than by oxygen transport in silica.

II.A.6. Oxidation of Additive-Containing Materials
The above discussions have considered only very pure materials. SiC and Si$_3$N$_4$ materials used for structural applications often contain additives to aid sintering. These additives affect long-term oxidation behavior. First, additives can diffuse into the silica scale during oxidation (34-38). Impurities present in silica increase oxygen transport rates by modification of the silica network structure and thereby increase oxidation rates of the silica-forming material (39). Second, additives such as Y$_2$O$_3$, La$_2$O$_3$, MgO, and CeO$_2$ present in significant amounts (4-10%) will diffuse to the silica surface and form discrete particles of silicates often of the M$_2$Si$_2$O$_7$ phase where M=Y, La, Ce. In these cases, the oxidation kinetics are still parabolic to times as long as 1000h (D.S. Fox, personal communication, 1997), but limited by the outward diffusion of metal cations to the matrix/oxide interface (34,35). In addition, for silicon nitride ceramics, the presence of impurities (40) and additives (41) prevent the formation of the silicon oxynitride inner layer, thus, oxidation rates of Si$_3$N$_4$ in practical applications are never as low as those found for pure CVD Si$_3$N$_4$. As additive levels and impurities increase further, the oxidation kinetics often do not follow simple parabolic rate laws. These more complex oxidation kinetics have been modeled in various ways (42,43). Finally, for systems containing large amounts of additives, >10%, at intermediate temperatures 1000-1200°C, reactions to form silicates occur in the grain boundaries with a volume expansion large enough to cause disintegration of the ceramic (44).
II.A.7. Thermal Cycling Effects

Long-term applications of structural ceramics, such as heat engines, require thermal cycling. Thermal cycling is a concern for several reasons. First, upon cooling, cristobalite undergoes the $\beta$ to $\alpha$ phase transformation with an accompanying 3% volume contraction (45). This volume change causes cracks to form in cristobalite upon cooling. Second, the thermal mismatch between SiC or Si$_3$N$_4$ and cristobalite, shown in Figure 7, results in tensile stresses in the oxide layer upon cooling. Cracks in the oxide may then form. However, cyclic oxidation tests at 1300°C in 5-hour cycles for 1000 hours have shown little deleterious effects on the oxidation kinetics when measured by weight change (46), as shown in Figure 8. One possible explanation is that upon reheating, stress is relieved and the cracks heal. Note that the tensile stresses formed in these ceramic/oxide systems contrast with those in superalloy/oxide systems where compressive stresses form in the scale on cooling and oxide spallation is typically observed.

II.B. Oxidation of other non-oxide Ceramics
BN is a useful crucible material in a vacuum or inert atmosphere. However, in oxygen it readily forms a liquid $B_2O_3$ scale which is not protective (47). Furthermore this $B_2O_3$ scale readily reacts with water vapor and forms stable volatile $H$-$B$-$O$($g$) species, such as $HBO_2(g)$, $H_3BO_3(g)$, and $H_3B_3O_6(g)$. These species have strongly negative free energies of formation and form even with ppm levels of water vapor. A similar situation exists with TiB$_2$, which is an attractive structural material due to its high strength (48).

AlN is currently a candidate for electronic substrates due to its high thermal conductivity (9). As an alumina former, it is expected to exhibit slow oxidation kinetics. However it appears to oxidize quite rapidly above about 1000°C, with rates quite dependent on grain size, porosity, the presence of second phases, and impurity content (49). In many instances, linear kinetics have been reported, in contrast to alumina formation on metal alloys (50). In addition, it is well established that water vapor in the oxidizing stream leads to extensive attack (51). The reasons for this extensive alumina formation in dry and wet oxygen are not clear. Clearly, nitrogen must escape through the alumina scale, which may lead to micropore formation. It has been suggested that the injection of $N_3^-$ leads to excess vacancy formation in the alumina and more rapid diffusion rates (52). In addition, the CTE mismatch between AlN and $Al_2O_3$ may lead to scale cracking (53).
III. Complex Environments

In most applications, ceramic materials are subjected to more aggressive environments than high temperature oxygen alone. These are outlined in Table I. As in the case of pure oxidation, most of the available data on interactions of non-oxide ceramics in complex gas mixtures are for SiC and Si$_3$N$_4$.

III.A. Water Vapor

There is general agreement that water vapor enhances the oxidation rate of silicon, SiC and Si$_3$N$_4$. There is a large amount of disagreement as to the magnitude of this effect for SiC and Si$_3$N$_4$. This disagreement arises in part from the complex effects water vapor has on the growing silica scale. These effects include enhanced impurity transport to the silica scale in water vapor containing atmospheres, enhanced solubility of water in the silica scale, alterations in the silica scale viscosity and structure, and finally, formation of volatile silicon hydroxide and oxyhydroxide species. Each one of these topics will be discussed separately below. Again, references to the oxidation behavior of silicon in water vapor are made for comparison.

Oxidation of silicon by water vapor occurs by the following reaction:
\[ \text{Si} + 2\text{H}_2\text{O}(g) = \text{SiO}_2 + 2\text{H}_2(g) \]  

(15)

The enhanced growth rates of \( \text{SiO}_2 \) by this reaction, relative to dry oxygen, has commercial application in the more rapid fabrication of microelectronic devices.

Oxidation of \( \text{SiC} \) by water vapor occurs by the reactions (54):

\[
\begin{align*}
\text{T}>1400K & \quad \text{SiC} + 3\text{H}_2\text{O}(g) = \text{SiO}_2 + \text{CO}(g) + 3\text{H}_2(g) \\
\text{T}<1400K & \quad \text{SiC} + 2\text{H}_2\text{O}(g) = \text{SiO}_2 + \text{CH}_4(g)
\end{align*}
\]

(16)

and (17)

and for \( \text{Si}_3\text{N}_4 \) by the following reaction:

\[
\text{Si}_3\text{N}_4 + 6\text{H}_2\text{O}(g) = 3\text{SiO}_2 + 2\text{N}_2(g) + 6\text{H}_2(g)
\]

(18)

Note that in each case the solid product \( \text{SiO}_2 \) is formed with the generation of additional gaseous products.

III.A.1. Impurity Transport

Impurities normally found in water vapor-containing environment form M-OH(g) species, where M is the impurity element such as Na, K, Fe, etc. Since M-OH(g) species are so thermodynamically stable, the quantity of impurities transported to a silica scale forming on \( \text{SiC} \) or \( \text{Si}_3\text{N}_4 \) in a water-vapor containing environment is increased. This increased contamination of the silica scale results in faster transport rates of oxidant through the scale, and thus increased oxidation rates of \( \text{SiC} \) or \( \text{Si}_3\text{N}_4 \). This effect has been identified for both Na- (55) and K-contamination (56) during the oxidation of \( \text{SiC} \).

III.A.2. Enhanced solubility of water in the silica scale
Deal and Grove (18) have shown that the parabolic oxidation rate of silicon in water vapor is increased by about an order of magnitude over the rate found in dry oxygen. This is explained by examination of Equation 9. While the diffusivity of water vapor in silica is almost two orders of magnitude slower than molecular oxygen, the solubility of water is nearly three orders of magnitude larger than oxygen. The net result is that parabolic oxidation rates of silicon in water vapor are more than one order of magnitude larger than those observed in dry oxygen. This explanation is directly applicable to the discussion for SiC and Si$_3$N$_4$ since silica is the oxidation product for these materials. This enhancement in parabolic oxidation rate constants has been observed for SiC (55,57) and in some cases for Si$_3$N$_4$ (58).

Deal and Grove (18) have also demonstrated that the parabolic oxidation rate has a power law dependence on the water vapor partial pressure, with the power law exponent, $n$ (Equation 10), equal to one, indicating that molecular water diffusion is the rate controlling step in the oxidation of silicon. Power law exponents for the oxidation of SiC of 0.67 (59) and of about 0.5 (54) have been obtained. This indicates dissociation of water into a singly charged species is probable. The discrepancy between the findings for silicon and SiC is unexpected since the transport properties of silica should be independent of the substrate material. A complex relationship between parabolic rate constant and
water vapor partial pressure has been observed for Si₃N₄ (58), but this may be explained by silica volatility, as described in section III.A.4.

III.A.3. Alterations in the structure of silica

It has been shown that the viscosity of amorphous silica decreases as the hydroxyl content increases (60). Hydroxyl groups are effective in breaking Si-O-Si bonds in amorphous silica. It is suggested that the resulting silica allows the more rapid permeation of molecular oxygen (57). The reduced viscosity of amorphous silica in conjunction with increased amounts of gaseous products results in the formation of bubbles in the scale formed on SiC (54) as shown in Figure 9. These bubbles, in turn, create shorter transport paths for oxidant to the SiC/SiO₂ interface by decreasing the effective oxide thickness, thereby increasing the oxidation rate. In contrast, bubbles are not observed in the scales formed on SiC in dry oxygen or on silicon in wet oxygen.

III.A.4. Silica scale volatility

Water vapor reacts with the silica scale formed on SiC and Si₃N₄ to form volatile hydroxide and oxyhydroxide species by the following reactions:

\[
\begin{align*}
\text{SiO}_2 + 2\text{H}_2\text{O}(g) &= \text{Si}({OH})_4(g) \\
\text{SiO}_2 + \text{H}_2\text{O}(g) &= \text{SiO}({OH})_2(g)
\end{align*}
\]
These volatile silicon hydroxides and oxyhydroxides have been identified experimentally using the transpiration technique (61) as well as mass spectrometry (62). The reaction of water vapor with SiC or Si$_3$N$_4$ involves the oxidation reaction (Equations 16-18) and the simultaneous linear volatilization reaction (Equations 19-20) resulting in overall paralinear kinetics (63). Paralinear kinetics, as measured by weight change, are shown in Figure 10. At long times, the weight change and recession can be approximated by the linear volatilization rate alone. At this time a steady state is achieved, the silica scale is consumed at the same rate it is formed leaving a constant oxide thickness. The volatilization rate of silica is controlled by transport of the volatile species through a gaseous boundary layer (63). This boundary layer controlled volatility rate can be expressed in terms of the application conditions such as gas velocity, total pressure, and water vapor partial pressure. For Si(OH)$_4$(g) formation, the volatility rate has the following dependences:

$$k_i \propto \frac{v^{1/2}P(H_2O)^2}{P_{total}^{1/2}}$$

(21)

where $k_i$ is the linear volatility rate, $v$ is the gas velocity and $P$ is pressure. Thus, in high pressure and high velocity applications, such as a gas turbine engine, the recession of a SiC or Si$_3$N$_4$ component by the volatility mechanism can be significant.

In summary, water vapor has deleterious effects on the durability of silica formers through a number of different mechanisms: increased transport of impurities to the oxide surface, increased oxide formation rates due to impurity
effects, enhanced solubility of water vapor, short circuit paths for oxidant transport, increased permeability of the oxidant, and finally consumption of the silica scale and component recession due to silica volatility.

III.B. Carbon Dioxide

Although CO$_2$ is a major component of combustion gases (Figure 1), there are only a few limited oxidation studies of SiC in CO$_2$ (64-67). It has been found that the oxidation rates of SiC in CO$_2$ are less than those in oxygen. Because the oxide growth rates are so low it is difficult to determine whether the kinetics are linear, parabolic, or more complex. The oxidation weight gain for SiC in CO$_2$ is shown compared to that observed in oxygen and a 50% water vapor/oxygen mixture in Figure 11 (67). Thus, in a complex combustion environment, the effects of CO$_2$ as an oxidant are negligible.

III.C. Effects of low P(O$_2$), Reducing Gases, H$_2$S, Cl$_2$

III.C.1. SiO(g) Formation

A unique, but very important issue with silica formers is the highly stable volatile sub-oxide, SiO(g). Consider the free energy of formation at 1500 K:

\[ \frac{1}{2} \text{Si} + \frac{1}{2} \text{O}_2(g) = \frac{1}{2} \text{SiO}_2(s) \quad \Delta G = -322 \text{ kJ/mol} \quad (22) \]

\[ \text{Si} + \frac{1}{2} \text{O}_2(g) = \text{SiO}(g) \quad \Delta G = -227 \text{ kJ/mol} \quad (23) \]
Note these equations are normalized to one mole of oxygen atoms. The free energies of formations are close, indicating SiO(g) can readily form.

There are two conditions which lead to SiO(g) formation (68): active oxidation and oxidation in mixed oxidizing/reducing gases. In the active oxidation case, the partial pressure of oxygen is too low to form a stable SiO$_2$ film, but sufficient to form SiO(g). This can occur in certain heat treating environments (69). Let us begin with a bare SiC surface. As the partial pressure of oxygen is increased, SiO(g) will form in increasing quantities. Then SiO(g) formation will stop and a stable SiO$_2$ film will form. This is the active-to-passive transition. Wagner has derived this for pure silicon (70). His results can be easily extended to SiC and Si$_3$N$_4$ (71). The active-to-passive transition occurs when sufficient SiO(g), via Equation 23, is generated to satisfy the SiC/SiO$_2$ condition for equilibrium. There is some controversy about the exact equilibrium condition, but reasonable agreement with measurements is obtained from the following equilibria:

$$2 \text{SiC} + \text{SiO}_2 = 3 \text{Si} + 2 \text{CO(g)}$$ (24)

The transition P(O$_2$) for active-to-passive oxidation is calculated based on equilibrium conditions and diffusion through the gas boundary layer to the sample. Calculated and measured active-to-passive transitions (72-74) are shown in Figure 12.
Now consider the case of passive-to-active transition. Beginning with a stable SiO$_2$ film on SiC, as the partial pressure of oxygen is lowered the film will become unstable and SiO$_2$ will react to form SiO(g). The transition pressure is calculated from the decomposition of the protective SiO$_2$ film (70):

\[
\text{SiO}_2(s) = \text{SiO}(g) + \frac{1}{2} \text{O}_2(g)
\]  

(25)

The calculated passive-to-active line is shown in Figure 12—note it is several orders of magnitude lower than the active-to-passive transition.

A second route to SiO(g) formation occurs in a mixture of oxidizing and reducing gases (68). These can be present in a fuel-rich combustion situation with appreciable amounts of CO$_2$, CO, H$_2$O and H$_2$. It has also been observed in laboratory experiments with CO/CO$_2$ (73) and H$_2$/H$_2$O (72). Thus a sequence of reactions might be:

\[
\begin{align*}
\text{SiC} + 3 \text{H}_2\text{O}(g) &= \text{SiO}_2 + \text{CO}(g) + 3 \text{H}_2(g) \\
\text{SiO}_2 + \text{H}_2(g) &= \text{SiO}(g) + \text{H}_2\text{O}(g)
\end{align*}
\]

(26) \hspace{1cm} (27)

This leads to paralinear kinetics—simultaneous oxide growth and linear reduction kinetics.

III.C.2 Effects of H$_2$S and Cl$_2$
Hydrogen sulfide is found in coal gasifiers and various other petrochemical environments (5). At low oxygen potentials, when SiO\textsubscript{2} does not form, both SiO(g) and SiS(g) may form:

\[ \text{Si}_3\text{N}_4 + 3 \text{H}_2\text{O}(g) = 3 \text{SiO}(g) + 3 \text{H}_2(g) + 2 \text{N}_2(g) \]  
(28)

\[ \text{Si}_3\text{N}_4 + 3 \text{H}_2\text{S}(g) = 3 \text{SiS}(g) + 3 \text{H}_2(g) + 2 \text{N}_2(g) \]  
(29)

These reactions can lead to material consumption (5), however when the oxygen potential is high enough to form SiO\textsubscript{2}, the corrosion rate drops substantially.

Chlorine, which may be found in certain chemical process environments, also leads to volatile products (75). In the case of SiC, reaction occurs with the silicon, leading to formation of various silicon chlorides with residual carbon. As the oxygen potential is increased, attack becomes less severe (75).

Current interest focuses on chlorine as HCl in waste incineration applications (76). Here the situation is quite complex involving not only HCl, but also a range of deposits.

III.D. Oxidation in the Presence of Impurities and Deposits
III.D.1. Low Levels of Na and K

Small amounts of impurities have several possible effects on the structure and thus the transport properties of silica. First, impurities can nucleate the formation of cristobalite at temperatures and times where amorphous silica would be expected. Second, impurities can modify the silica network as shown in Figure 3c. Breaking up the network tends to increase the oxygen transport rate through silica. Small amounts of alkali metals act as network modifiers and can increase oxidation rates an order of magnitude (56, 77, 78).

III.D.2. Na₂SO₄ Deposits

As impurity levels increase, actual deposits may form. The chemistry of these depends on application, as shown in Table I. Corrosive deposits have long been known to be an important issue for metals and alloys in various high temperature applications (79). Here we shall focus on sodium sulfate deposits on silica. Many of the general principles described for this system apply to the other cases.

It is difficult to simulate the effect of an actual deposit. Simple laboratory experiments involve a one-time deposition of a salt on a test coupon, followed by
a heat treatment. This type of experiment has the advantage of precise control of experimental variables such as temperature and gas composition. However, the actual situation involves a continuous deposition process. This can be done in a laboratory furnace. However, this can be accomplished more effectively by seeding a flame in a burner rig, as illustrated in Figure 13. Figure 14 is a comparison of SiC treated in a burner without salt and with a flame seeded with NaCl. Note the extensive corrosion in the latter case.

The formation of Na$_2$SO$_4$ occurs when ingested NaCl reacts with sulfur impurities in the fuel (80, 81):

$$2 \text{NaCl(v)} + \text{SO}_3(g) + \text{H}_2\text{O(g)} = \text{Na}_2\text{SO}_4(l) + 2 \text{HCl(g)} \quad (30)$$

The source of NaCl depends on the application—it may be from a marine environment in the case of a heat engine, or from the process chemicals in the case of an industrial furnace. Under the appropriate conditions, Na$_2$SO$_4$ forms as a condensed phase, depositing on parts. Figure 15 shows the calculated dew points for Na$_2$SO$_4$ deposition (81). The rates of deposition are also critical and have been treated in detail (82).

The most useful interpretation of this process is with the acid/base theory of molten salts. Na$_2$SO$_4$ decomposes to Na$_2$O, which is the key reactant:
salt solution models (84) and leads to Mg and Ca silicates. Impure fuels lead to
vanadate deposits. $V_2O_5$ is an acidic molten salt and thus does not react with
$SiO_2$. However it is reported to accelerate $Si_3N_4$ oxidation, possibly due to the
solubility of $SiO_2$ in $V_2O_5$ which may be further enhanced by the presence of a
$Y_2O_3$ in $Si_3N_4$ (85).

A more complex case is that of a molten slag, which may contain up to ten
different oxides. Here it more difficult to define basicity, however a useful index
has been the weight percent ratio of basic to acid oxides (86). These are the
basic and acidic oxides encountered:

**Basic Oxides:** $Na_2O$, $K_2O$, $MgO$, $CaO$, $Fe_2O_3$

**Acidic Oxides:** $SiO_2$, $Al_2O_3$, $TiO_2$

As in the $Na_2SO_4$ case, the basic slag tends to cause more rapid material
degradation than the acidic slag. Under some conditions, metal silicides may
form with a basic coal slag (4).

**III.E. High Temperature Effects**
Silica melts at 1723°C. Transport rates are very high in a liquid scale and rapid oxidation occurs, as shown in Figure 17. Some of the bubbles in this sample are from escape of CO(g), but some are also from the interaction of SiC and SiO₂ at high temperatures:

\[
\text{SiC}(s) + 2 \text{SiO}_2(s) = 3 \text{SiO}(g) + \text{CO}(g) \quad (34)
\]

Figure 18a is a plot of total pressures generated by this reaction (87). Note that for carbon saturated SiC the pressures are much higher, leading to a lower upper use temperature. An analogous situation exists for Si₃N₄, but based on the Si₂N₂O interaction with SiO₂:

\[
\text{Si₂N₂O}(s) + \text{SiO}_2(s) = 3 \text{SiO}(g) + \text{N}_2(g) \quad (35)
\]

The total pressure is shown in Figure 18b.

IV. Refractory Oxide Coatings

One possible solution to the preceding issues of volatility and molten salt corrosion is the use of refractory oxide coatings on SiC and Si₃N₄. Refractory oxides are generally more chemically inert and this approach offers the possibility of combining the benefits of both materials. Table II lists some coefficients of thermal expansion (CTEs) for SiC, Si₃N₄, and refractory oxides. There is a good match between SiC and mullite. Mullite coatings on SiC were
first developed at Solar Turbines (San Diego, CA) and further developed at NASA Lewis (88, 89). The critical processing issues are surface roughening of the SiC for adherence and application of a fully crystalline mullite coating. A mullite coating on a SiC coupon is shown in Figure 19. There are clear advantages with mullite coatings in Na$_2$O-induced corrosion due to formation of high melting sodium-alumino-silicates, as opposed to lower melting sodium silicates. However the chemical activity of silica in mullite is only about 0.4, so reactions which volatilize SiO$_2$ can still occur readily.

It may be possible to apply other refractory oxides to SiC. As Table II shows, the problem of CTEs must be considered. One approach is graded coatings from SiC to a refractory oxide such as alumina or zirconia.

V. Hydrothermal Corrosion of SiC and Si$_3$N$_4$

Hydrothermal corrosion involves the attack by water at high temperatures and high pressures. Typical conditions are shown in Table I. These conditions are often near the critical point of water. Table I also shows some potential applications where these environments may be encountered. Another promising application of SiC and Si$_3$N$_4$ is a vessel for oxidation of waste materials to form safe products using super-critical water (90).
The hydrothermal corrosion of SiC may form silica by the following reactions (91, 92):

\[ \text{SiC} + 2 \text{H}_2\text{O}(g) = \text{SiO}_2 + \text{CH}_4(g) \]  
\[ \text{SiC} + 4 \text{H}_2\text{O}(g) = \text{SiO}_2 + \text{CO}_2(g) + 4 \text{H}_2(g) \]

Given nearly equal amounts of SiC and water vapor at high temperatures, free carbon can form (92, 93):

\[ \text{SiC} + 2 \text{H}_2\text{O}(g) = \text{SiO}_2 + \text{C} + 2 \text{H}_2(g) \]

This provides a method for synthesis of carbon films. However, in most applications there is excess water and SiO$_2$ is formed as a protective film. The hydrothermal corrosion of Si$_3$N$_4$ is similar and occurs by:

\[ \text{Si}_3\text{N}_4 + 6 \text{H}_2\text{O}(g) = 3 \text{SiO}_2 + 4 \text{NH}_3(g) \]

At temperatures above 430°C, NH$_3$(g) decomposes to N$_2$(g) and H$_2$(g).

In both cases, the silica is eventually dissolved by the high pressure water as:

\[ \text{SiO}_2 + \text{OH}^- = \text{HSiO}_3^- \]
In the case of Si$_3$N$_4$, this leads to attack of the grain boundary phases and dislodging of the grains (94). Methodical studies have been done on the effect of additives in Si$_3$N$_4$ on its hydrothermal corrosion behavior (13, 94). The morphology of attack has been correlated with the type of additive and its resistance to hydrothermal attack.

Ceramics which form oxides other than silica behave differently under hydrothermal conditions. AlN forms AlOOH (95). Other ceramics such as TiC and ZrC which do not form protective oxides at high temperatures also do not form protective oxides under hydrothermal conditions (91).

**VI. Liquid Metals**

Ceramics have long been used as containers for processing of liquid metals. More recent applications involve use of liquid metals as a heat transfer medium. A wide range of non-oxide ceramics have been considered for these applications (12, 96, 97). Dissolution of the ceramic and capillary action are two important issues in liquid metal corrosion (12). Dissolution can be diffusion controlled or interface controlled. Local variations in dissolution due to phase and structural differences can lead to surface roughening. Capillary action leads to internal attack of porous ceramics. More important it can lead to substantial grain
boundary attack, depending on the relation of the solid/liquid interfacial energy and the grain boundary energy (12).

VII. Effect of Oxidation/Corrosion on Mechanical Properties

It is evident from current materials research that we cannot view one phenomenon such as oxidation or corrosion alone, but rather must view its effect on the entire system. Nearly all the applications listed in Table I utilize the ceramic in some type of load-bearing situation. A critical question that arises, is how oxidation and corrosion alter the ability of the ceramic to bear a load (98-112). Again, the emphasis here is on SiC and Si$_3$N$_4$.

Two recent reviews summarize the critical issues involved in corrosion/mechanical property interactions (98, 99). Many of the studies in this area deal with the change in mechanical properties after corrosion. However these processes often act together. The oxidation rate for various types of Si$_3$N$_4$ is increased by the application of either compressive or tensile stresses (100,101).
It is well established that short-term oxidation tends to blunt cracks and may actually increase strength (103). Long-term isothermal oxidation (5000 h) for SiC containing B, C, and Si leads to strength increases whereas the same treatment for SiC containing Al₂O₃ and WC leads to strength decreases (104). Little strength degradation of a variety of commercial SiC and Si₃N₄ materials was observed for 3500h burner rig tests with 12 min. cycles (105). Recent research (106) shows the effects of water vapor on strength are complex, however, in some cases the rapid oxidation rates lead to flaw healing. In the case of stress rupture for Si₃N₄, water vapor has little effect (107).

Molten salt corrosion (108-110) leads to extensive pitting and strength reduction. Figure 20 is a fracture origin due to a corrosion pit from molten salt attack (108). This indicates the need for tougher, more flaw resistant ceramics. In the case of Si₃N₄ it has been shown that corrosion reactions often effect the grain boundary phase. Various mechanical properties are often related to the stability of this phase. Compositional changes in this grain boundary phase may lower the threshold stress intensity for crack growth and also increase the creep rate.

Corrosion reactions at a growing crack tip are quite important as evidenced by moisture-assisted crack growth of ceramics (111). Henager and Jones (112) have shown that the presence of Na₂SO₄ increases the slow crack growth velocity by a factor of two over air at 1573 K.
Similar effects occur in the case of hydrothermal corrosion. Pitting and dislodging of grains in various types of Si₃N₄ leads to substantial strength reductions. Analogous to the high temperature situation, changes in the grain boundary phase of Si₃N₄ lead to changes in the mechanical behavior. Weakening and dissolution of this phase leads to substantial strength reductions. Similarly, liquid metal attack of grain boundaries will lead to strength reductions (96).

VIII. Conclusions

Non-oxide ceramics are a promising class of materials for a wide range of applications at high temperatures. Before application, their interactions with the high temperature environment must be well-understood. The silica forming ceramics (SiC and Si₃N₄) exhibit the best oxidation behavior and the focus of this chapter is on them. However some information is included on AlN and BN which show promise for several specific applications.

High-temperature oxidation is critical to most applications. SiC oxidizes with the diffusion of oxygen inward through the growing silica scale being rate limiting.
The oxidation of Si$_3$N$_4$ is more complex—here the oxynitride scale plays a role. Most engineering ceramics contain additives, which tend to enhance oxidation.

Actual applications involve complex gas mixtures. The effects of water vapor, carbon dioxide, low oxidant pressures, and mixed oxidizing/reducing gases are discussed. Water vapor enhances oxidation, whereas carbon dioxide is a less effective oxidant. Low oxidant pressures and mixed oxidizing/reducing gases lead to SiO(g) formation.

Corrosive deposits are also encountered in some applications. These include sodium sulfates, vanadate, and slags. Deposit-induced corrosion can be described with the acid/base theory of oxides. Since silica is an acidic oxide, it is readily attacked by a basic molten salt.

These problems may be minimized with the application of a refractory oxide coating. Mullite shows a good thermal expansion match to SiC and is a promising starting point. It may be possible to develop other more refractory oxide coatings.
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These problems may be minimized with the application of a refractory oxide coating. Mullite shows a good thermal expansion match to SiC and is a promising starting point. It may be possible to develop other more refractory oxide coatings.
Some proposed applications involve hydrothermal conditions—water at high temperatures and pressures. Hydrothermal corrosion is analogous to high temperature oxidation in many ways. Ceramics which form protective high temperature oxides also form protective hydrothermal oxides. The form of corrosive attack is also similar.

Finally the interaction of corrosive attack and degradation of ceramic properties is briefly discussed. This is essential to understand before these materials can be applied. Microstructural changes such as pitting and grain boundary attack lead to a concurrent loss of mechanical integrity. The development of tougher ceramics and coatings are necessary to minimize this problem.
<table>
<thead>
<tr>
<th>Application</th>
<th>Non-Oxide Ceramic Composites</th>
<th>Environment</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Turbine engine components</td>
<td>SIC, Si₃N₄ Composites</td>
<td>Combustion gases, deposits: Na, Mg, Ca sulfate, sodium vanadates</td>
<td>1</td>
</tr>
<tr>
<td>Combustor liners</td>
<td>SIC, Si₃N₄ Composites</td>
<td>Combustion gases</td>
<td>2</td>
</tr>
<tr>
<td>Bladed &amp; Vanes</td>
<td>SIC, Si₃N₄ Composites</td>
<td>Combustion gases, various deposits</td>
<td>3</td>
</tr>
<tr>
<td>Pistons</td>
<td>SIC, Si₃N₄ Composites</td>
<td>Combustion gases, slag</td>
<td>4</td>
</tr>
<tr>
<td>Valves</td>
<td>SIC, Si₃N₄ Composites</td>
<td>Various gases including air, H₂S, HCl</td>
<td>5</td>
</tr>
<tr>
<td>Industrial Furnaces</td>
<td>SIC</td>
<td>Reduced pressure N₂O₅, CO₂, N₂, Air</td>
<td>6</td>
</tr>
<tr>
<td>Heat Exchangers</td>
<td>SIC</td>
<td>Air</td>
<td>7</td>
</tr>
<tr>
<td>Particulate filters</td>
<td>SIC</td>
<td>Fiber coatings: reduced pressure</td>
<td>8</td>
</tr>
<tr>
<td>Coal Gasifiers, Waste</td>
<td>SIC</td>
<td>Vacuum, inert gases</td>
<td>9</td>
</tr>
<tr>
<td>Incinerators</td>
<td>SIC</td>
<td>Fiber coatings: reduced pressure</td>
<td>10</td>
</tr>
<tr>
<td>Re-entry Shields</td>
<td>SIC</td>
<td>Vacuum or inert gas and</td>
<td>11</td>
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<tr>
<td>High Temperature Semi-</td>
<td>SIC</td>
<td>Liquid metal containers</td>
<td>12</td>
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<tr>
<td>Conductors</td>
<td>AIN</td>
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<td>Electronic Substrates</td>
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<td>Liquid metal containers</td>
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<tr>
<td>Processing Heat transfer</td>
<td>and nitrides</td>
<td>liquid metals</td>
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<td>-------------------------</td>
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</tr>
<tr>
<td>Pump bearings, Cooling lines for nuclear reactors</td>
<td>SiC, Si$_3$N$_4$</td>
<td>300-600</td>
<td>High Pressure Fluids, 10 - 100 atm</td>
</tr>
</tbody>
</table>

Table I. Selected applications which are proposed for non-oxide ceramics.
<table>
<thead>
<tr>
<th>Material</th>
<th>CTE ($10^{-6}$ K$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiC</td>
<td>5.2</td>
</tr>
<tr>
<td>Si$_3$N$_4$</td>
<td>3.2</td>
</tr>
<tr>
<td>Mullite (3Al$_2$O$_3$·2SiO$_2$)</td>
<td>5.4</td>
</tr>
<tr>
<td>Alumina (Al$_2$O$_3$)</td>
<td>9</td>
</tr>
<tr>
<td>Partially stabilized zirconia</td>
<td>10</td>
</tr>
<tr>
<td>(0.08 Y$_2$O$_3$·ZrO$_2$)</td>
<td></td>
</tr>
</tbody>
</table>

Table II. Coefficients of thermal expansion of SiC and several refractory oxides (89).
References


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Figure Captions

1. Equilibrium gas composition vs. equivalence ratio (24).
2. Recession due to oxidation for selected materials, with protective oxide scale indicated (14).
3. Structure of SiO₂ (a) Amorphous (b) Crystalline (c) With sodium cations (Adapted from (15))
4. Schematic of electro-balance and vertical tube furnace used for oxidation studies.
5. Oxidation of SiC at 1300 C in 1 atm oxygen, showing parabolic kinetics (20).
6. Arrhenius plot of log kp vs 1/T for Si, CVD SiC, CVD Si₃N₄, and Si₃N₄ with additives obtained in dry oxygen(18, 20, 35).
7. Thermal expansion of SiC, Si₃N₄, amorphous SiO₂, and crystalline SiO₂ as a function of temperature (24).
8. Cyclic oxidation kinetics of SiC and Si₃N₄ (46).
9. Bubbles formed in SiO₂ due to oxidation of SiC in 90% water vapor/10% O₂ at 1200°C (54).
10. Schematic showing components of paralinear kinetics (63).
11. Comparison of SiC oxidation in 50% H₂O/O₂, O₂, and CO₂ at 1200°C (67).
12. Calculated active/passive and passive/active transitions and measured active/passive transitions (72-74).
13. Schematic of jet fuel burner used for corrosion studies.
14. Optical micrographs of sintered SiC coupons with carbon and boron additives, oxidized in a burner rig at 1000°C (a) 46 h with no sodium (b) 13.5 h with sodium chloride seeded flame (24).

15. Calculated dew points for Na$_2$SO$_4$ deposition (81).

16. Quartz coupons treated in burner rig for 1 h with 2 ppm sodium (as NaCl) seeded flame: (a) No. 2 diesel fuel (0.5% sulfur), 5 h (b) Jet A fuel (0.05% sulfur) (81).

17. CVD SiC oxidized for 1 h at 1800°C (Courtesy of D. Fox, NASA Lewis).

18(a). Total pressure [P(SiO) + P(CO)] generated from SiC/SiO$_2$ interactions.

and (b). Total pressure [P(SiO) + P(N$_2$)] generated from Si$_2$N$_2$O/SiO$_2$ interactions (24).

19. Micrograph of polished cross-section of mullite on SiC (88).

20. Fracture origin for sintered $\alpha$-SiC reacted with Na$_2$SO$_4$ for 48h at 1000°C (108). The area noted in (b) is enlarged in (c). (c) Preferential grain boundary attack is observed at arrows.
Figure 1.

(a) Pressure, bar

(b) Adiabatic flame temperature, K

E-10922 Jacobson 9pt/100% JJ from E-6965 Fig 3.
Figure 2.
Figure 4.

E-10922 Jacobson 9pt/100% JJ from E-6965 Fig 6.
Figure 5.

Specific weight change, mg/cm²

Specific weight change², mg²/cm⁴

\[ k_p = 2.3 \times 10^{-4} \, \text{mg}^2/\text{cm}^4 \, \text{hr} \]
Figure 6.

- **Si**, 119 kJ/mol, extrapolated from (18)
- CVD SiC, 118 kJ/mol (20)
- CVD Si$_3$N$_4$, 363 kJ/mol (20)
- Si$_3$N$_4$, 6 w/o Y$_2$O$_3$, 450 kJ/mol (35)
Crystalline SiO$_2$,
SiC, Si$_3$N$_4$,
Amorphous SiO$_2$.

Figure 7.
Figure 8.

- CVD SiC
- CVD Si$_3$N$_4$
- NT 154 Si$_3$N$_4$

Specific weight change, mg/cm$^2$

Time, hr
Figure 9.
Figure 10.

Solid scale weight

Net weight

Volatile weight

$k_p = 3 \times 10^{-3} \text{ mg}^2/\text{cm}^4 \text{ h}$

$k_f = 4 \times 10^{-3} \text{ mg}^2/\text{cm}^2 \text{ h}$

Specific weight change, mg/cm$^2$

Time, hr
Figure 11.

50% H₂O/50% O₂

O₂

CO₂

Specific weight change, mg/cm²

Time, hr

E-10922 Jacobson 9pt/100% JJ from (author's files)
Figure 12.

Measured active to passive

Calculated active to passive

Calculated passive to active

- $H_2O/H_2 (72)$
- $CO_2/CO (73)$
- $O_2/Ar (74)$
Combustion temperature probe
Exit nozzle
Preheated-air temperature probe
Jet-A fuel supply line
Air inlet
Combustion chamber
Salt solution injector/aspirator
Ignitor
Burner pressure tap
Test specimen

Figure 13.
Figure 14.
Figure 15. Dewpoint temperature, K vs. Pressure, bar for various S content, percent and Na content, ppm. The graph shows the transitions from liquid to solid phases. Data sources include E-10922 Jacobson 9pt/100% JJ from E-6965 Fig 24. or CD-89-40832.
Figure 16.

E-10922 Jacobson 9pt/100% JJ from E-6965 Fig 28.
Figure 17.
Figure 18.

(a) SiC/C/SiO₂

(b) Si₂N₂O/Si/SiO₂

Figure 18.