Sintering and Creep Behavior of Plasma-Sprayed Zirconia and Hafnia Based Thermal Barrier Coatings

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SINTERING AND CREEP BEHAVIOR OF PLASMA-SPRAYED ZIRCONIA AND HAFNIA BASED THERMAL BARRIER COATINGS

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ABSTRACT

The sintering and creep of plasma-sprayed ceramic thermal barrier coatings under high temperature conditions are complex phenomena. Changes in thermomechanical and thermophysical properties and in the stress response of these coating systems as a result of the sintering and creep processes are detrimental to coating thermal fatigue resistance and performance. In this paper, the sintering characteristics of ZrO2-8wt%Y2O3, ZrO2-25wt%CeO2-2.5wt%Y2O3, ZrO2-6w%NiO-9wt%Y2O3, ZrO2-6wt%Sc2O3-2wt%Y2O3 and HfO2-27wt%Y2O3 coating materials were investigated using dilatometry. It was found that the HfO2-Y2O3 and baseline ZrO2-Y2O3 exhibited the best sintering resistance, while the NiO-doped ZrO2-Y2O3 showed the highest shrinkage strain rates during the tests. Higher shrinkage strain rates of the coating materials were also observed when the specimens were tested in Ar+5%H2 as compared to in air. This phenomenon was attributed to an enhanced metal cation interstitial diffusion mechanism under the reducing conditions. It is proposed that increased chemical stability of coating materials will improve the material sintering resistance.

I. INTRODUCTION

Plasma-sprayed ceramic thermal barrier coatings are being developed for advanced gas turbine and diesel engine applications to improve engine reliability and efficiency. Since these coatings are experiencing severe thermomechanical cycling during engine operation, it is especially challenging to develop coating systems with high reliability and durability. In particular, ceramic
coating sintering and creep at high temperature are among the most important issues for the development of advanced thermal barrier coatings, as has been recognized by many investigators [1-10]. The ceramic sintering and creep at high temperature can result in coating shrinkage and through-thickness cracking during cooling, thereby further accelerating the coating failure process. Sintering-segmentation-enhanced delamination can be an important failure mechanism for a thermal barrier coating system, due to stress concentration from the through-thickness cracks, and increased coating elastic modulus from the sintering densification process. The increase in coating thermal conductivity is also detrimental to coating performance. Research efforts involving various techniques have also been made in characterizing the ceramic coating sintering and creep behavior at high temperature and under temperature gradients simulating those encountered in the engine [1, 2, 7, 11-13].

The sintering and creep of plasma-sprayed, porous and microcracked ceramic thermal barrier coatings are complex phenomena. The early work by Firestone et al [1, 2] indicated that the ceramic creep appeared to be a thermally activated process, with the ceramic splat-sliding being an important creep deformation mechanism. More recently, it has been reported that the ceramic thermal barrier coatings can sinter and creep significantly under compressive stress states at relatively low temperatures [11, 13, 14]. The "creep" of plasma-sprayed ZrO2-8wt%Y2O3 at room temperature has also been observed at a tensile stress of 7.4 MPa [15]. A mechanism-based model has been proposed to describe the densification and deformation occurring in thermal barrier coatings at temperature by taking into account the thermally and stress activated diffusion, and the mechanical compacting processes [13]. The dopants in the ceramic coatings can significantly modify the point defect and microstructures in the bulk, at splat-grain boundaries and microcrack surfaces of the materials, thereby can significantly affect these sintering and creep processes. A better understanding of the dopant effects will help to develop future advanced, sintering/creep resistant "superalloy-type" ceramic coatings.

The purpose of this paper is to investigate sintering kinetics of several zirconia and hafnia based ceramic coating materials. The ceramic materials investigated include: (a) ZrO2-8wt%Y2O3, a NASA-Lewis Research Center reference (or baseline) material; (b) ZrO2-25wt%CeO2-2.5wt%Y2O3, a commercially available coating material developed for hot corrosion resistance; (c) HfO2-27wt%Y2O3, a potential new coating material developed at NASA for high temperature stability [16]; (d) ZrO2-6wt%NiO-9wt%Y2O3, a NiO-doped ZrO2-Y2O3 coating material reported to suppress the tetragonal-monoclinic phase transformation [17]; (e) ZrO2-6wt%Sc2O3-
2wt%Y₂O₃, developed for improved hot corrosion resistance [18]. The coating sintering and creep mechanisms and dopant effect on coating sintering rates are discussed based on experimental observations and possible defect reactions.

II. EXPERIMENTAL MATERIALS AND METHODS

The five ceramic coating materials mentioned above, ZrO₂-8wt%Y₂O₃, ZrO₂-25wt%CeO₂-2.5wt%Y₂O₃, ZrO₂-6wt%NiO-9wt%Y₂O₃, ZrO₂-6wt%Sc₂O₃-2wt%Y₂O₃ and HfO₂-27wt%Y₂O₃, were chosen for this study. The actual compositions of these materials were close to their nominal compositions. Each of the above materials was prepared by sintering and crushing except for ZrO₂-25CeO₂-2.5wt%Y₂O₃ which was spray dried and plasma spheroidized. A single set of standard plasma-spray parameters was used for each material. The powders with an average particle size of 60 μm of these coating materials were first plasma-sprayed onto 3 mm diameter graphite cylindrical bars, using the plasma spray conditions described previously [19]. The coating thickness was about 0.76 mm, and porosity was about 10%. The graphite bars were then slowly burnt off at 600°C for 6 hours in a furnace in air. The hollow ceramic cylinders were cut into 25.4 mm dilatometer specimens.

Ceramic sintering experiments were carried out in air and in Ar+5%H₂ within the temperature range of 900 to 1400°C, using a UNITHERM™ high temperature dilatometer system shown in Figure 1. Since the push rod in the dilatometer exerts a certain amount of force (measured at approximately 450g using a spring device) on the specimen, a uniaxial stress of

![Fig. 1 Schematic diagram showing the ceramic sintering experiment using dilatomentry.](image-url)
approximately 0.5 MPa was acting on the specimen during the entire sintering test. Therefore, this experiment can also be considered a low constant-stress creep test for the ceramic materials. During the sintering/creep experiments at various test temperatures, all specimens were heated in at a rate of 5°C/minute and held at the given test temperature for 15 hours, and then cooled down at a rate of 5°C/minute to room temperature. Specimen shrinkage during the heating/cooling cycles was continuously recorded in a computer system. Surface morphology changes of the specimens due to the sintering process were examined using a scanning electron microscope (SEM). Phase structures of the specimens before and after dilatometer sintering tests were also examined by X-ray diffractometry with Cu $K_\alpha$ radiation.

III. EXPERIMENTAL RESULTS

Figure 2 shows thermal expansion (shrinkage) results for the coating materials during the sintering experiments at various temperatures measured by the dilatometry technique. Sintering shrinkage were observed for all materials when the specimens were held at temperature for 15 hours. The shrinkage strains increased with increasing temperature. It can be seen that the HfO$_2$-27wt% Y$_2$O$_3$ showed the best sintering resistance. In contrast, CeO$_2$, Sc$_2$O$_3$, and NiO-doped ZrO$_2$-Y$_2$O$_3$ materials exhibit significant sintering shrinkage. Below the temperature of 900°C, no significant shrinkage strains were detected for the given test time. Figure 3 illustrates the sintering shrinkage strains occurring at the isothermal sintering stages as a function of temperature.

The sintering rates of the ceramic materials at the isothermal stages change with time, especially at the early sintering time period. As shown in examples in Figure 4 (a) and (b), faster shrinkage rates were observed initially, however, relatively constant rates were observed for longer sintering times. At 1400°C as shown in Figure 4 (c), the "steady state" sintering rates for ZrO$_2$-Y$_2$O$_3$, ZrO$_2$-CeO$_2$-Y$_2$O$_3$, ZrO$_2$-Sc$_2$O$_3$-Y$_2$O$_3$, ZrO$_2$-NiO-Y$_2$O$_3$, and HfO$_2$-Y$_2$O$_3$ are $2.6 \times 10^{-8}$/sec., $3.8 \times 10^{-8}$/sec., $4.2 \times 10^{-8}$/sec., $8.5 \times 10^{-8}$/sec., and $6.4 \times 10^{-9}$/sec., respectively. Figure 4 (d) shows that for ZrO$_2$-NiO-Y$_2$O$_3$, the second cycle resulted in further shrinkage of the specimen at 1200°C.

Figure 5 shows the sintering shrinkage kinetics of plasma-sprayed ZrO$_2$-8wt%Y$_2$O$_3$ at 1200°C in air and in Ar+5%H$_2$. It can be seen that when the specimen was tested in a reducing atmosphere, a faster sintering shrinkage rate was observed. In addition, the ceramic coating material turned black after this Ar+H$_2$ treatment. Increased sintering rates and darkened color were observed for all other materials under the reduced oxygen partial pressure condition. This may imply that the more defective structure of the materials due to Ar+H$_2$ sintering would increase the
Fig. 2 Thermal expansion and sintering shrinkage response for the coating materials during the dilatometry sintering experiments at various temperatures. (a) ZrO$_2$-8wt%Y$_2$O$_3$; (b) ZrO$_2$-25wt%CeO$_2$-2.5wt%Y$_2$O$_3$; (c) ZrO$_2$-6wt%Sc$_2$O$_3$-2wt%Y$_2$O$_3$; (d) ZrO$_2$-6wt%NiO-9wt%Y$_2$O$_3$; (e) HfO$_2$-27wt%Y$_2$O$_3$. 
minority defect transport especially at the internal surfaces and grain boundaries, thus resulting in a faster sintering rate.

Figure 6 illustrates some examples of the X-ray diffraction spectra for ZrO$_2$-Y$_2$O$_3$, ZrO$_2$-CeO$_2$-Y$_2$O$_3$, HfO$_2$-Y$_2$O$_3$ and ZrO$_2$-Sc$_2$O$_3$-Y$_2$O$_3$ after 15 hours sintering at 1200°C in air. From X-ray diffraction experiments, it was found that the baseline ZrO$_2$-8wt%Y$_2$O$_3$ primarily consisted of tetragonal $t'$ phase. The CeO$_2$-doped ZrO$_2$-Y$_2$O$_3$ also showed significant amount of $t'$ phase, however, the possibility that the cubic $c$ phase might also be increased as compared to the baseline material requires further study. Due to the high concentration of yttria dopant, HfO$_2$-27wt%Y$_2$O$_3$ had a fully stabilized cubic $c$ phase. No appreciable monoclinic phase was observed in these three materials. Heat treatments related to the sintering experiments under various temperature and oxygen pressure conditions did not measurably alter the phase structures of these materials. The as-sprayed ZrO$_2$-Sc$_2$O$_3$-Y$_2$O$_3$ material showed tetragonal $t'$ phase and an increased amount of the monoclinic $m$ phase. The monoclinic phase in the Sc$_2$O$_3$-doped materials increased after the sintering tests.
Fig. 4 Sintering behavior of the ceramic materials at the isothermal stages. (a) and (b) The sintering strains as a function of time and temperature for ZrO2-8wt%Y2O3 and ZrO2-25wt%CeO2-2.5wt%Y2O3, respectively; (c) Steady state creep rates for the ceramic materials at 1400°C; (d) Sintering shrinkage of ZrO2-6wt%NiO-9wt%Y2O3 at 1200°C under two temperature cycles.
Fig. 5  Sintering shrinkage kinetics of plasma-sprayed ZrO₂-8wt%Y₂O₃ at 1200°C in air and in Ar+5%H₂.

The NiO-doped ZrO₂-Y₂O₃ showed more complex phase structures. As shown in Figure 6, the majority phase in ZrO₂-NiO-Y₂O₃ was the cubic \( c \) phase, instead of tetragonal phase in the baseline material. However, the monoclinic \( m \) phase was also present in this material. Because of the limited solubility of NiO in ZrO₂-Y₂O₃ (about 3 mol% at 1600°C\textsuperscript{[17]}), NiO phase was observed in the as-sprayed and air-sintered specimens. In the Ar+H₂ sintered specimens, however, a Ni phase was present because of the reduction of NiO.

Surface microstructure changes were also observed after the sintering experiments. Certain regions showed more noticeable sintering densification and grain growth as compared to other regions, indicating there were some heterogeneities in the observed sintering phenomena. Figure 7 shows micrographs of ceramic surfaces of the ZrO₂-8wt%Y₂O₃ coating material before and after the dilatometry sintering at 1200°C. It can be seen that sintering which occurred could result in microcrack healing and material densification, accompanying with substantial grain growth in some regions.
Fig. 6. X-ray diffraction spectra of the plasma-sprayed ceramic coating materials. (a) Diffraction spectra of ZrO$_2$-Y$_2$O$_3$, ZrO$_2$-CeO$_2$-Y$_2$O$_3$, ZrO$_2$-Sc$_2$O$_3$-Y$_2$O$_3$, and HfO$_2$-Y$_2$O$_3$ after 1200°C sintering in air and 1200°C sintering in Ar. (b) Diffraction spectra of ZrO$_2$-NiO-Y$_2$O$_3$ under the as-sprayed condition and after 1200°C sintering in Ar.
Fig. 7 Surface micrographs of the ZrO$_2$-8wt%Y$_2$O$_3$ ceramic coating material before and after the dilatometry sintering at 1200°C for 15 hours. (a) Before the sintering test; (b) After the sintering test.

IV. DISCUSSION

The sintering and low-stress creep characteristics of the ceramic coating materials, determined by the dilatometer technique, are similar to the creep behavior of plasma-sprayed coatings obtained from high temperature mechanical creep tests [1, 12, 20] and the laser
sintering/creep test [13]. The fast initial creep rate and low creep activation energy have been attributed to mechanical sliding, fast surface and grain boundary diffusion, and temperature and stress gradient enhanced transport in the porous and weak ceramic coatings [13]. Figure 8 illustrates the creep rates of the plasma-sprayed ZrO$_2$-8wt%Y$_2$O$_3$ as a function of stress and temperature determined by the laser sintering technique [13]. It can be seen that with higher compressive stresses acting on the coating, a long primary creep stage and substantial sintering/creep rates can be observed at much lower temperatures. In the high temperature, low stress sintering/creep tests by the dilatometer technique, mechanical sliding becomes less predominant, and a nearly "steady-state" creep region has been reached in a relatively short period of time. Diffusion-related processes become more important mechanisms for the low stress sintering and creep deformation. The observed grain growth phenomena also suggest the complex diffusion occurring during the dilatometer sintering test.

Fig. 8 The coating creep rates of plasma-sprayed ZrO$_2$-8wt%Y$_2$O$_3$, determined by laser high heat flux sintering/creep technique, as a function of stress and temperature.
Creep deformation of ceramic coating materials require diffusion of the cations and anions in these materials. The creep rate in ceramics is therefore determined by the diffusion of the slowest species, diffusing along the fastest path. In yttria-stabilized zirconia, the majority defect types are oxygen vacancies and yttrium aliovalent dopants at normal cation sites. The possible minority defects are zirconia interstitials, zirconium vacancies, and yttrium interstitials, and the zirconium and yttrium cation transport is confirmed to be the slowest process in yttria-stabilized single crystals [21]. The defect reactions in the yttria-stabilized zirconia can be written according to Kröger-Vink notation [22] as

\[
\begin{align*}
Y_2O_3 &= 2Y'_{Zr} + 3O'^+ + V_O^- \quad (1) \\
O'^+ &= V_O^- + 2e^- + \frac{1}{2}O_2(g) \quad (2)
\end{align*}
\]

and

\[
\begin{align*}
2O'^+ + Zr'^+ &= Zr'^++ 4e^- + O_2(g) \quad (3) \\
\frac{1}{2}O_2(g) &= V'_{Zr} + 4h^+ \quad (4)
\end{align*}
\]

(for majority defects)

(for minority defects)

In the extrinsic region, the majority defect oxygen vacancy concentration \([V_O^-]\) is determined by the dopant yttria concentration \([Y'_{Zr}]\), which follows the electroneutrality condition

\[
[Y'_{Zr}] = 2[V_O^-]. \quad (5)
\]

At lower oxygen partial pressures in the intrinsic region where the electron conductivity becomes important, oxygen vacancies can be further introduced according to Equation (2), that is

\[
[V_O^-] n^2 = K_{V_O} P_{O_2}^{-1/2} \exp\left(-\frac{\Delta H_{V_O}}{RT}\right) \quad (6)
\]

where \(n\) is electron concentration, \(K_{V_O}\) is reaction constant, \(\Delta H_{V_O}\) is the enthalpy of formation of oxygen vacancies, \(R\) and \(T\) are gas constant and temperature, respectively. Metal interstitials can be an important defect type in the oxygen deficient oxide [23], and the zirconium interstitial concentration can be obtained from Equation (3)

\[
[Zr'^+ n]^4 = K_{Zr'^+ P_{O_2}^{-1}} \exp\left(-\frac{\Delta H_{Zr'^+}}{RT}\right) \quad (7)
\]

where \(K_{Zr'^+}\) is a constant, \(\Delta H_{Zr'^+}\) is the enthalpy of formation of zirconium interstitials. In this
intrinsic region, the electroneutrality can be expressed as

\[ n = 2\left[ V_O \right] + 4\left[ Zr_i^{\text{\textasciitilde}} \right]. \]  

(8)

By combining Equations (6)-(8), the oxygen vacancy and zirconia interstitial concentrations can be written as

\[
\left[ V_O \right] = \left( \frac{K_{V_O}}{4} \right)^{1/3} \left( \frac{p_{O_2}}{P_{O_2}} \right)^{-1/6} \exp\left( -\frac{\Delta H_{V_O}}{3RT} \right) 
\]

(9a)

\[
\left[ Zr_i^{\text{\textasciitilde}} \right] = \left( \frac{K_{Zr_i^{\text{\textasciitilde}}}}{(2K_{V_O})^{1/3}} \right) \left( \frac{p_{O_2}}{P_{O_2}} \right)^{-1/3} \exp\left( -\frac{3\Delta H_{Zr_i^{\text{\textasciitilde}}} - 4\Delta H_{V_O}}{3RT} \right) 
\]

(9b)

(when \( V_O^{\gg} > \left[ Zr_i^{\text{\textasciitilde}} \right] \))

\[
\left[ V_O \right] = \left( \frac{K_{V_O}}{(4K_{Zr_i^{\text{\textasciitilde}}})^{2/5}} \right) \left( \frac{p_{O_2}}{P_{O_2}} \right)^{-1/10} \exp\left( -\frac{5\Delta H_{V_O} - 2\Delta H_{Zr_i^{\text{\textasciitilde}}}}{5RT} \right) 
\]

(10a)

\[
\left[ Zr_i^{\text{\textasciitilde}} \right] = \left( \frac{K_{Zr_i^{\text{\textasciitilde}}}}{256} \right)^{1/5} \left( \frac{p_{O_2}}{P_{O_2}} \right)^{1/5} \exp\left( -\frac{\Delta H_{Zr_i^{\text{\textasciitilde}}}}{5RT} \right) 
\]

(10b)

(when \( V_O^{\ll} < \left[ Zr_i^{\text{\textasciitilde}} \right] \))

In the very high oxygen pressure region where the zirconium vacancies are predominant, the electroneutrality condition can be written as

\[ p = 4\left[ V_{Zr}^{\text{\textasciitilde}} \right] \]

(11)

where \( p \) is electron hole concentration. The zirconium vacancy concentration can be obtained from Equation (4) as

\[
\left[ V_{Zr}^{\text{\textasciitilde}} \right] = \left( \frac{K_{V_{Zr}^{\text{\textasciitilde}}}}{256} \right)^{1/5} \left( \frac{p_{O_2}}{P_{O_2}} \right)^{1/10} \exp\left( -\frac{\Delta H_{V_{Zr}^{\text{\textasciitilde}}}}{5RT} \right) 
\]

(12)

where \( K_{V_{Zr}^{\text{\textasciitilde}}} \) is a constant, \( \Delta H_{V_{Zr}^{\text{\textasciitilde}}} \) is the enthalpy of formation of zirconium vacancies. A Kröger-Vink diagram is constructed based on these defect reactions and Equations (5), (9), (10) and (12), as shown in Figure 9.

The increased sintering rate of ZrO\textsubscript{2}-8wt\% Y\textsubscript{2}O\textsubscript{3} at the reducing Ar+5\% H\textsubscript{2} atmosphere is probably related to the defect structure change in the oxide. From the proposed Kröger-Vink diagram shown in Figure 9, it can be seen that both concentrations of oxygen vacancies and zirconia interstitials increase with reducing partial pressure of oxygen, especially in the low oxygen activity region. At extremely low oxygen pressures, the metal cation interstitials can even become the dominant defect type. Therefore, it is possible that the highly defective oxide structures under
low oxygen pressures facilitates the metal cation interstitial formation, thus resulting in faster metal cation diffusion and the increased sintering rate. Thornton et al. [24] have also observed enhanced cerium migration and segregation in the ZrO$_2$-25wt%CeO$_2$-2.5wt%Y$_2$O$_3$ material under relatively moderate reducing conditions, further confirming the increased cation mobility in more oxygen-deficient oxide under the low oxygen activity conditions.

Since the ceramic sintering requires the transport of the minority cations, the stability of the ceramic materials (both dopants and base materials) will have influence on the sintering behavior. The present study has shown that there is a close relationship between the oxide chemical and phase stability and the sintering rate. Halfnia-based oxides have higher chemical stability, and lower oxygen partial pressures for the transition of ionic conductivity to electronic conductivity, as compared to zirconia-based oxides, therefore it is not surprised that the HfO$_2$-27wt%Y$_2$O$_3$ exhibited the lowest sintering rates. On the other hand, the CeO$_2$-doped ZrO$_2$ exhibited large electron contributions at even moderate temperatures and oxygen activities [25]. As shown in Figure 9, the increased region of electron conductivity implies an extended metal cation interstitial
region, in which the cation interstitial concentration is increased with decreasing oxygen partial pressure, and thus resulting in possible enhanced metal cation diffusion with reducing oxygen activity. Insufficient doping, as possibly occurred for Sc2O3-doped ZrO2-Y2O3, will have a similar effect on the metal cation diffusion. For the NiO-doped ZrO2-Y2O3, the observed high sintering rates may also be related to NiO segregation at the grain boundaries, which may act as a sintering agent. At lower oxygen partial pressures, the NiO reduction to metallic Ni, as observed in this experiment, can further enhance the sintering process. It is suggested that the chemical and phase stability of both the base oxides and dopant oxides is critical to the sintering and creep behavior of the ceramic materials.

V. CONCLUSIONS

1. Sintering shrinkage strains were observed at the isothermal stage for all ceramic coating materials tested in the dilatometer sintering experiments. The HfO2-27wt%Y2O3 and baseline ZrO2-8wt%Y2O3 exhibited the best sintering resistance, and NiO-doped ZrO2-Y2O3 showed the highest shrinkage strain rates during the tests.

2. The higher shrinkage strain rates of the coating materials were observed for the specimens tested in Ar+5%H2 as compared to those tested in air. This phenomenon was attributed to a proposed enhanced metal cation interstitial diffusion mechanism under the reducing conditions.

3. There was a close relationship between the observed sintering behavior and chemical and phase stability of the coating materials. Increased chemical stability of base oxides and dopants seems to improve materials phase stability at high temperature, and sintering/creep resistance. Insufficient doping and dopant-segregation-induced depletion will facilitate the sintering process.

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