Calcium-Magnesium-Aluminosilicate (CMAS) Reactions and Degradation Mechanisms of Advanced Environmental Barrier Coatings

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Abstract

The thermochemical reactions between calcium-magnesium-aluminosilicate- (CMAS-) based road sand and several advanced turbine engine environmental barrier coating (EBC) materials were studied. The phase stability, reaction kinetics and degradation mechanisms of rare earth (RE)-silicates Yb₂SiO₅, Y₂Si₂O₇, and RE-oxide doped HfO₂ and ZrO₂ under the CMAS infiltration condition at 1500 °C were investigated, and the microstructure and phase characteristics of CMAS-EBC specimens were examined using Scanning Electron Microscopy (SEM) and X-ray Diffraction (XRD). Experimental results showed that the CMAS dissolved RE-silicates to form crystalline, highly non-stoichiometric apatite phases, and in particular attacking the silicate grain boundaries. Cross-section images show that the CMAS reacted with specimens and deeply penetrated into the EBC grain boundaries and formed extensive low-melting eutectic phases, causing grain boundary recession with increasing testing time in the silicate materials. The preliminary results also showed that CMAS reactions also formed low melting grain boundary phases in the higher concentration RE-oxide doped HfO₂ systems. The effect of the test temperature on CMAS reactions of the EBC materials will also be discussed. The faster diffusion exhibited by apatite and RE-doped oxide phases and the formation of extensive grain boundary low-melting phases may limit the CMAS resistance of some of the environmental barrier coatings at high temperatures.

Introduction

The ever-present needs for developing higher efficiency high performance turbine engines have created demand for new hot-section component materials that are lighter and more temperature capable. Current Ni-based superalloys are reaching the upper limit of their temperature capabilities, and so silicon carbide (SiC) fiber-reinforced SiC/SiC ceramic matrix composites (CMCs) have been envisioned as alternative next generation turbine engine hot-section materials (Ref. 1). Silicon-based ceramics and composites are desirable because they have low density, superior strength and oxidation resistance in high temperature oxidizing environments. However, a major issue with Si-based materials is their vulnerability to corrosion and performance degradation in combustion water vapor containing environments due to volatilization of the protective silica (SiO₂) scales, leading to recession of the ceramics and strength degradation under combined thermo-mechanical loading conditions. An environmental barrier coating (EBC) can be used to prevent the SiC/SiC CMC from reacting with water vapor (Refs. 1 to 3). Advanced HfO₂, ZrO₂, and rare earth silicate (RE silicate) EBC systems have also been developed, achieving significantly improved temperature capability and environmental protection of SiC/SiC CMCs (Refs. 4 to 6).

The significantly higher operating temperatures envisioned above 1500 °C for next generation turbine engine hot-section components impose additional material design constraints and raise serious component environmental durability issues. Many common silicon-containing sand dust and volcano ash materials can melt at approximately 1200 °C. During engine operation, these materials deposit on the EBC and form glassy melts of calcium-magnesium aluminosilicate (CMAS), which can significantly reduce the

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silicon-based EBC temperature capability. This molten CMAS interacts chemically and mechanically with the EBC to cause premature coating failure. Thus when designing EBC materials, it is critical to understand the high-temperature interactions between the coating materials and CMAS in order to create CMAS-resistant coatings. Research work has been done to determine the mechanisms by which CMAS can cause failure and performance degradations in yttria stabilized zirconia thermal barrier coatings (Refs. 7 and 8) and early generation EBCs (Refs. 9 to 12).

The objective of the present paper is to investigate the stability of candidate rare earth silicates and rare earth oxide-doped HfO₂ and ZrO₂ environmental barrier coatings in contact with CMAS at high temperature. The emphasis is placed on the reactions and degradation mechanisms between CMAS and EBC materials particularly at 1500 °C, and compared with the results at 1300 °C. The current work is thus aiming at providing information on the CMAS resistance of RE silicates and RE oxide-doped HfO₂ and ZrO₂ systems.

**Experimental Materials and Methods**

The two primary EBC materials investigated in this study were ytterbium monosilicate (Yb₂SiO₅) and yttrium disilicate (Y₂Si₂O₇). These materials were NASA-designed and processed in the form of plasma-homogenized hollow-sphere (HOSP) powders to NASA powder specifications by Sulzer Metco (US), Westbury, New York, USA. In addition to the RE silicates, HOSP NASA four-component HfO₂-23.9mol%(Gd,Yb,Y)₂O₃ and six-component ZrO₂-4.4mol%(Y, Gd, Yb)₂O₃-2.6mol%(TiO₂+Ta₂O₅) compositions (Ref. 6) were also investigated.

Two types of samples were produced for this study. For the first kind, EBC powders were sintered using a hot press technique at 1500 °C and 103 MPa load in a vacuum environment to create dense, monolithic slabs ~4 mm thick or discs ~2 mm thick, which were then cut into several specimens. Each sample was prepared for CMAS exposure by polishing the surface of interest using 320-grit SiC paper and cleaning the surface with ethanol. To provide a baseline understanding for material behavior at 1500 °C, one specimen of each material was heat treated in air for 50 hr at 1500 °C. A model CMAS frit containing 35mol%CaO-10mol%MgO-7mol%Al₂O₃-48mol%SiO₂ and minor amounts of Fe₂O₃ and NiO was received from GE Aviation, Evendale, Ohio, USA. The CMAS frit was mixed with ethanol to form a paste and placed on each specimen surface to give a CMAS surface concentration of ~40 mg/cm². The specimens were placed in an alumina crucible and heat treated in air for 5, 50, 100, 150, or 200 h at 1500 °C. Every 50 h, the furnace was cooled to room temperature and the samples were removed and photographed. Because of observed volatility of CMAS and the reaction products, for samples tested longer than 50 h, an additional 40 mg/cm² CMAS was applied to the sample surface every 50 h. Compared to a single application of CMAS, this testing procedure better simulates engine operation where the source of CMAS is continuously available. To provide a better understanding of the CMAS-EBC reaction, additional tests were conducted for both of the RE-silicate materials at 1300 °C for 50 h.

The second type of sample was produced by mechanically mixing 25 and 30 wt% CMAS frit with Yb₂SiO₅ and Y₂Si₂O₇ EBC powders, respectively. The CMAS amounts were adjusted to reflect a higher concentration that would occur in a thin turbine coating system. The mixtures were sintered via hot press at 1050 °C and 103 MPa load in vacuum using a SiC/SiC CMC as a backing to produce two 1 mm thick RE-silicate + CMAS blend coating specimens onto CMC substrates. It was thought that these samples would more accurately represent the original CMAS composition in the closed, high pressure turbine section environment because the CMAS would volatilize less and thus react more completely with the EBCs when conducting the tests at such high temperatures.
Following testing, specimens were examined using x-ray diffraction (XRD) to determine phase structure using Cu Kα radiation with a wavelength of 1.541 Å. Each specimen was then sectioned using a low-speed diamond saw. One half of the specimen was mounted and polished using standard metallographic techniques and examined using a field-emission scanning electron microscope (SEM). The other half was reserved for examination of the surface with electron microscopy. Qualitative chemical compositions were obtained from surfaces and cross-sections using electron dispersive spectroscopy (EDS).

Experimental Results

Yb$_2$SiO$_5$-CMAS Surface Reactions

Figure 1 shows cross-sectional images of the Yb$_2$SiO$_5$ in the as-sintered condition and after 50 h of heat treatment in air at 1500 °C. There was little change in microstructure observed over time.

Figure 2 shows images of Yb$_2$SiO$_5$ before and after CMAS reactions. The dark areas in the 5 h tested specimen (Fig. 2(b)) are Yb$_2$SiO$_5$-CMAS reaction products. There is visibly much less of this reacted material on the 50 and 200 h tested specimens, suggesting that the CMAS components may be highly volatile at 1500 °C and atmospheric pressure in air.

SEM images of Yb$_2$SiO$_5$ reacted with CMAS for 50 h are shown in Figure 3. EDS was used to qualitatively determine the elemental composition in each region of the micrographs. CMAS reacted with the Yb$_2$SiO$_5$ to form a Ca-Yb-Si-containing layer with an average thickness of ~60 to 100 μm. The surface of this reaction layer (Fig. 3(b)) consisted of roughly hexagonal-shaped grains with an average diameter of 17 μm. Figure 4 shows cross-sectional images of Yb$_2$SiO$_5$ reacted with CMAS for 200 h. The heavily reacted surface layer had the same approximate composition, crystal structure, and average thickness as in Figure 3(a). The enlarged regions in Figures 4(b) and (d) have eutectic microstructures, indicating that they solidified from a low-melting liquid due to the higher CMAS content. EDS shows that the lighter phase of this eutectic is rich in Al, Mg, Si, and Yb, while the dark phase is rich in Al, Ca, Mg, and Si. As apparent from Figure 4(c), these low-melting particles are located between grains, providing evidence for preferential CMAS attack at grain boundaries. These particles were found to penetrate the entire specimen cross-section, which has a bulk thickness of ~4 mm.

XRD was used to determine that the reaction layers in Figures 3(a) and 4(a) had a crystal structure similar to calcium-ytterbium-silicate-oxyapatite (Ca$_2$Yb$_8$(SiO$_4$)$_6$O$_2$), a variation of the mineral apatite. The x-ray scan of the 200 h-reacted Yb$_2$SiO$_5$ is compared to a known apatite structure in Figure 5. The Al, Yb, Mg, Ca-containing grain boundary phases may be too small in quantity and/or preferred orientation with glass phases to contribute significantly to the diffraction pattern.

In order to better understand the grain size behavior, volatility, and surface melting of rare earth silicates, CMAS surface reaction treatments were carried out for 5 h at 1500 °C and 50 h at 1300 °C in air, and compared to 50 and 200 h treatments at 1500 °C. The XRD profile of the 5 h reacted specimen was found to be almost identical to the 200 h reacted sample in Figure 5, indicating that these were Ca$_2$Yb$_8$(SiO$_4$)$_6$O$_2$ grains. The surface of this specimen (Fig. 6(a)) has roughly hexagonal grains similar to those in the 50 h CMAS-reacted specimen (Fig. 3(b)). This suggests that the initially fast CMAS reaction kinetics and less further degradation of the specimens due to the CMAS volatility at the high temperatures and therefore the loss of available CMAS for reactions. The apatite grains appear to grow from an amorphous, CMAS-rich melt matrix, demonstrating that the apatite surface layer forms from a CMAS-rich melt. In contrast, the surface of the 1300 °C reacted specimen is glassy, with few notable microstructural features. This indicates significantly less reactions and slower kinetics at the lower testing temperatures.
**Y$_2$Si$_2$O$_7$-CMAS Surface Reactions**

Figure 7 shows images of Y$_2$Si$_2$O$_7$ specimens before and after reacting with CMAS for various times at 1500 °C. The same issue of CMAS volatility at atmospheric pressure observed in Yb$_2$SiO$_5$ was also observed for this material at such a high test temperature. Figure 8 shows SEM images of Y$_2$Si$_2$O$_7$ reacted with CMAS for 50 h. The region shown in Figure 8(a) represents a portion of the ~1 mm thick well-melted layer in contact with high CMAS concentration, which has been converted from Y$_2$Si$_2$O$_7$ to a Ca-Si-Y-rich reaction product with isolated Si-rich regions. The surface of reaction layer (Fig. 8(b)) consists of angular, needle-like grains. SEM micrographs of Y$_2$Si$_2$O$_7$ reacted with CMAS for 200 h are shown in Figure 9. The light colored phase shown in Figure 9(a) is a portion of the ~1 mm thick melted layer and is rich in Ca, Si, and Y. The darker phases located between the lighter grains are rich in Al, Si, and Y and can be found throughout the entire ~4 mm thick cross-section. Unlike the Yb$_2$SiO$_5$-CMAS reaction products, these grain boundary phases are not visible in the surface images (Fig. 9(b)), possibly due to even higher volatility in the Y, Al, Ca, Mg, Si-containing phases. XRD identifies the Ca-containing Y$_2$Si$_2$O$_7$-CMAS reaction product as calcium-yttrium-silicate-oxyapatite (Ca$_2$Y$_8$(SiO$_4$)$_6$O$_2$), as shown in Figure 10. This material has a very similar structure to the Yb$_2$SiO$_5$-CMAS reaction layer.

It can be seen from Figure 11(a) that the Y$_2$Si$_2$O$_7$ reacted with CMAS for 5 h at 1500 °C has a surface microstructure nearly identical to the 50 h CMAS reacted specimen in Figure 8(b) with the same amount CMAS applied. As with Yb$_2$SiO$_5$, the result suggests that the reaction occurred fast in the initial a few hours, and the reaction and CMAS loss due to its volatility prevented further significant more reactions in the test condition. The surface of the 1300 °C reacted sample in Figure 11(b) contains apatite grains of similar shape to the 1500 °C CMAS-reacted specimens, while un-reacted glass phase CMAS seemed still present. The XRD profile of this specimen is nearly identical to Figure 10, showing the crystalline calcium-yttrium-silicate-oxyapatite (Ca$_2$Y$_8$(SiO$_4$)$_6$O$_2$) phase.

**RE-Silicate + CMAS Blend**

The hot-pressed rare earth silicate-CMAS blends are shown before and after heat treatment in Figure 12. Before heat treatment, both specimens appear dense and have a mottled appearance. Heat treatment at 1500 °C results in complete melting of both samples. This melt partially infiltrated the porous alumina crucible to penetrate and react with a portion of the ceramic brick beneath it. The experiments approximately determined upper temperature limits of the coating materials in contact with CMAS, and showing some extensive melting of the Yb$_2$SiO$_5$ and Y$_2$Si$_2$O$_7$ specimens at the high 25 to 30 wt% CMAS concentrations at 1500 °C.

**RE-Oxide Doped HfO$_2$- and ZrO$_2$-CMAS Surface Reactions**

Images of the RE-oxide doped HfO$_2$ reacted with CMAS for 100 h are shown in Figure 13. Based on initial examinations, the large, light colored grains contain HfO$_2$ with a much lower RE-oxide concentration than the original as-processed coating material. The darker grain boundary phases are rich in both CMAS elements and RE elements. These RE-rich grain boundary phases penetrated through the entire specimen, which had a bulk thickness of ~2 mm.

Figure 14 shows images of the six-component ZrO$_2$ reacted with CMAS for 100 h. The EDS analysis indicates that the large, white grains visible in the cross-section and surface images contain mainly ZrO$_2$. The dark grain boundary regions appear to be rich in Si and Zr. No Ca, Mg, or Al was detected in the sample cross-section of surface, possibly due to high volatility of the unreacted CMAS components.
Discussion

RE-Silicates

The main mechanism of CMAS penetration in the low CMAS concentration surface reaction tests appears to be grain boundary attack. As mentioned in more detail later, some surface melting may have occurred at relatively short CMAS exposure times during which CMAS concentration was high. High CMAS volatility likely reduced the CMAS surface concentration at longer exposure times. In the Yb$_2$SiO$_5$ specimens, the widening of the grain boundary regions containing low-melting eutectics with testing time implies continuing dissolution of the rare earth (Yb) and recession of the grain surfaces even when the overall CMAS surface concentration has decreased due to CMAS volatilization. In both materials, low melting grain boundary phases were observed throughout the bulk thickness of the 200 h reacted specimens, demonstrating overall poor CMAS resistance, particularly along the grain boundaries, at the test temperatures. Because EBC layers generally have a thickness no greater than 200 μm, CMAS would penetrate the entire RE silicate EBC top coat quickly, causing the EBC to fail and also exposing the intermediate and bond coat layers to CMAS and other environment attack.

Y$_2$Si$_2$O$_7$ appears to have suffered a more severe reaction, possibly due to its higher SiO$_2$ concentration. The needle-like grains in the surface images (Figs. 8(b), 9(b), and 11(a)) of Y$_2$Si$_2$O$_7$ are formed from a liquid phase, indicating at least the surface of the EBC experiences localized melting at 1500 °C. This also appears to be the case for Yb$_2$SiO$_5$, as the 5 h CMAS experiments confirmed localized melting of the Yb$_2$SiO$_5$ and Y$_2$Si$_2$O$_7$ surfaces in contact with high CMAS concentrations. However, more work is under way to explain the difference in grain morphology and grain growth kinetics. The Y$_2$Si$_2$O$_7$-CMAS reaction kinetics appeared faster than the Yb$_2$SiO$_5$-CMAS kinetics, as evidenced by the fully formed apatite grains on the 5 h reacted Y$_2$Si$_2$O$_7$ specimen and the partially formed apatite grains in the 5 h reacted Yb$_2$SiO$_5$ specimen. CMAS attack on Y$_2$Si$_2$O$_7$ appears to be a significant problem even at 1300 °C, as demonstrated by the extensive localized melting and formation of Y-containing apatite grains on the specimen surface (Fig. 11(b)). Conversely, Yb$_2$SiO$_5$ does not display the same aggressive CMAS attack at 1300 °C, which can be seen the lack of a crystalline apatite reaction product (Fig. 6(b)) during the short CMAS exposure.

The CMAS volatility observed in the CMAS surface reaction tests may be unrealistic, as the effective compositions of a localized CMAS source may change rapidly in the 1500 °C air furnace test conditions and no constant CMAS concentration is maintained during the given testing cycle times. The closed turbine section environment and continuous deposition of CMAS during the engine operation would lessen CMAS volatility, effectively raising the CMAS concentration on the EBC surface and thus likely promoting even more severe CMAS infiltration and attack. In this way, the rare earth silicate-CMAS blend experiments represent a more realistic scenario and help determine the upper temperature limit of the CMAS concentrations. In these tests, the CMAS dissolved the entire EBC specimen, including partially the SiC/SiC CMC substrates. This is an extreme demonstration of the possible catastrophic effects of CMAS on EBCs and turbine section components. More work must be done to determine the upper CMAS concentration limits that RE-silicates and other candidate EBC materials can endure for their designed lifetimes without catastrophic failure.

The RE silicate-CMAS reaction layers observed in the coating experiments suggest very fast transport of CMAS into the bulk grains of the specimens via a solid state diffusion process. An effort was made to explain this fast diffusion by examining the stoichiometry of the apatite reaction layers. An atomic number-absorption-fluorescence (ZAF) technique was used to produce approximate semi-quantitative molar percentages of oxides from the EDS data collected from the Yb$_2$SiO$_5$-CMAS and Y$_2$Si$_2$O$_7$-CMAS reacted specimens. These molar percentages were then used to determine the ratio of alkaline earth oxides (AE oxides) to RE oxides in the apatite reaction products. This ratio is shown plotted as a function of CMAS reaction time in Figure 15. Stoichiometric apatite has an AE oxide-RE oxide ratio of 0.5. From the composition standpoint, the non-stoichiometric Ca$_2$Yb$_8$(SiO$_4$)$_6$O$_2$ appears to be Yb$_2$O$_3$-lean (with a more oxygen vacant sublattice), while the Ca$_2$Y$_8$(SiO$_4$)$_6$O$_2$ appears Y$_2$O$_3$-rich (with an AE vacant sublattice).
The modified phase diagram shown in Figure 16 indicates the shifted compositions of the two reaction products. The non-stoichiometric materials contain high concentration anion and cation vacancies and defects, which may substantially increase the rate of CMAS element diffusion in the solid material.

**RE-Oxide Doped HfO₂ and ZrO₂**

The doped HfO₂ and ZrO₂ specimens generally exhibited less melting as compared to RE-silicates thus may have showed better CMAS phase stability than the RE-silicates. In the case of high rare earth element doped HfO₂, the CMAS appears to penetrate the specimen at grain boundaries and maybe forming higher melting RE-aluminosilicate phases. The low concentration of RE elements in the HfO₂ bulk grains indicates that these elements also substantially dissolve into the CMAS grain boundary melts to form the grain boundary reaction phases. Unlike the rare earth silicates, however, the doped HfO₂ seems to react with CMAS very little but seemed also recessed due to loss of RE elements.

The doped ZrO₂ had much lower dopant concentrations and exhibited the least reaction and maybe the best CMAS resistance of the materials studied. The lack of Ca, Mg, and Al in the bulk specimen suggests that the unreacted CMAS volatilized during the course of each 50 h heat treatment. Experiments involving higher CMAS concentrations will have to be done to fully understand the interaction between CMAS and this six-component ZrO₂ composition.

**Summary and Conclusions**

The thermochemical reactions at 1500 °C between calcium-magnesium-aluminosilicate- (CMAS) and several EBC materials were studied. After 50 h, Yb₂SiO₅ and Y₂Si₂O₇ developed a locally melted, Ca- and RE-containing reaction layer from which non-stoichiometric apatite-like grains precipitated. CMAS penetrated the specimen preferentially at grain boundaries and dissolved the EBC material to form low-melting grain boundary pockets. After 200 h, these low melting phases extended through the entire specimen, indicating that CMAS penetrated the bulk thickness of ~4 mm. Y₂Si₂O₇ appears to react more quickly with CMAS forming even lower melting phases than Yb₂SiO₅. RE-silicate + 25 to 30 wt% CMAS specimens were fully melted after 50 h heat treatment at 1500 °C, indicating that there is an upper limit to the allowable CMAS concentration in these materials. RE-oxide doped HfO₂ and ZrO₂ demonstrated better CMAS reaction and phase stability than the RE-silicates, although appears that high RE-oxide dopant concentration still facilitated severe grain boundary CMAS attack in HfO₂ and the grain boundary surface recession. Low rare earth dopant ZrO₂ appeared relatively stable and unreacted under the experimental conditions, and the effect of transition metal dopants Ta and Ti on CMAS resistance is a future topic to be studied. Higher CMAS exposure and some controlled coating experiments must be carried out to fully understand the EBC material’s CMAS-resistance capabilities.

**References**


Figure 1.—Sintered and as-processed Yb2SiO5. Cross-section SEM Back-Scattered Electron (BSE) images of Yb2SiO5 (a) as-sintered; and (b) after 50 h heat treatment at 1500 °C in air.
Figure 2.—Yb$_2$SiO$_5$ reacted with CMAS. Macro-photographs of Yb$_2$SiO$_5$ specimens (a) as-sintered; and reacted with CMAS for (b) 5 h, (c) 50 h, and (d) 200 h. There is visibly much less of the reacted material on the 50 h and 200 h tested specimens, suggesting that the CMAS components may be highly volatile at 1500 °C in atmospheric pressure in air.

Figure 3.—SEM micrographs of Yb$_2$SiO$_5$ reacted with CMAS for 50 h. (a) cross-section; (b) surface BSE images of Yb$_2$SiO$_5$ reacted with CMAS for 50 h at 1500 °C; (c) and (d) EDS of regions I and II, respectively.
Figure 4.—SEM images of Yb$_2$SiO$_5$ reacted with CMAS for 200 h. (a), (b), (c), and (d) low and high magnification surface BSE images of Yb$_2$SiO$_5$ reacted with CMAS for 200 h at 1500 °C; and (e) and (h) EDS of regions I-IV, respectively.
Figure 5.—XRD of Yb$_2$SiO$_5$-CMAS reaction layer. The pattern for Ca$_2$Yb$_8$(SiO$_4$)$_6$O$_2$ matches well with the observed x-ray scan for the Yb$_2$SiO$_5$-CMAS reaction layer.

Figure 6.—Yb$_2$SiO$_5$ surface reacted with CMAS for (a) 5 h at 1500 °C and (b) 50 h at 1300 °C.
Figure 7.—Photographs of Y$_2$Si$_2$O$_7$ specimens reacted with CMAS for various time at 1500 °C. (a) as-sintered; and (b) reacted with CMAS for 5 h, (c) 50 h, and (d) 200 h. Due to the volatility of the CMAS and its melts at high temperature, the amount of CMAS for reactions was not sustainable for the long-term testing at 1500 °C in atmospheric pressure in air.

Figure 8.—Y$_2$Si$_2$O$_7$ reacted with CMAS for 50 h at 1500 °C. (a) A cross-section SEM BSE image of the Y$_2$Si$_2$O$_7$; (b) A surface SEM BSE image of the Y$_2$Si$_2$O$_7$; (c) and (d) EDS analyses of regions I and II in the cross-section surface, respectively.
Figure 9.—Y$_2$Si$_2$O$_7$ reacted with CMAS for 200 h at 1500 °C. (a) A cross-section SEM BSE image of the Y$_2$Si$_2$O$_7$; (b) A surface SEM BSE image of the Y$_2$Si$_2$O$_7$; (c) and (d) EDS analyses of regions I and II in the cross-section surface, respectively.

Figure 10.—XRD of Y$_2$Si$_2$O$_7$-CMAS reaction product. The pattern for Ca$_2$Y$_8$(SiO$_4$)$_3$O$_2$ matches.
Figure 11.—$Y_2Si_2O_7$ reacted with CMAS for (a) 5 h at 1500 °C and (b) 50 h at 1300 °C.

Figure 12.—RE-silicate + CMAS mixtures. Digital photographs of (a) Yb$_2$SiO$_5$ + 25 wt%CMAS as-sintered; (b) Y$_2$Si$_2$O$_7$ + 30 wt% CMAS as-sintered; (c) Yb$_2$SiO$_5$ + 25 wt% CMAS heat treated for 50 h at 1500 °C; and (d) Y$_2$Si$_2$O$_7$ + 30 wt% CMAS heat treated for 50 h at 1500 °C.
Figure 13.—HfO$_2$ reacted with CMAS for 100 h. (a) and (b) Low and high magnification cross-sectional images of RE-oxide doped HfO$_2$ reacted with CMAS for 100 h at 1500 °C; (c) and (d) Low and high magnification surface BSE images of RE-oxide doped HfO$_2$ reacted with CMAS for 100 h at 1500 °C. Similar to the rare earth silicate cases, the high concentration rare earth doped HfO$_2$ had relatively substantial grain boundary recessions and CMAS reaction phases.
Figure 14.—ZrO$_2$ reacted with CMAS for 100 h at 1500 °C. (a) A cross-sectional image of six-component RE oxide-doped ZrO$_2$ reacted with CMAS; (b) A surface BSE image of six-component RE oxide-doped ZrO$_2$ reacted with CMAS.

Figure 15.—Stoichiometry of CMAS reaction layers. The molar ratio of alkaline earth-oxides (AEO) CaO+MgO to rare earth-oxides (RE$_2$O$_3$) Yb$_2$O$_3$ and Yb$_2$O$_3$ in the Yb$_2$SiO$_5$-CMAS and Y$_2$Si$_2$O$_7$-CMAS reaction layers as a function of time. The average and stoichiometric molar ratios are indicated as horizontal lines.
Figure 16.—AEO-RE₂O₃-SiO₂ phase diagram. Redrawn ternary phase diagram indicating shifted non-stoichiometric apatite compositions. The letters A and B indicate the compositions of CMAS-reacted Yb₂SiO₅ and Y₂Si₂O₇, respectively.
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Experimental results showed that the CMAS dissolved RE-silicates to form crystalline, highly non-stoichiometric apatite phases, and in particular attacking the silicate grain boundaries. Cross-section images show that the CMAS reacted with specimens and deeply penetrated into the EBC grain boundaries and formed extensive low-melting eutectic phases, causing grain boundary recession with increasing testing time in the silicate materials. The preliminary results also showed that CMAS reactions also formed low melting grain boundary phases in the higher concentration RE-oxide doped HfO2 systems. The effect of the test temperature on CMAS reactions of the EBC materials will also be discussed. The faster diffusion exhibited by apatite and RE-doped oxide phases and the formation of extensive grain boundary low-melting phases may limit the CMAS resistance of some of the environmental barrier coatings at high temperatures.