Microstructural Regulation, Oxidation Resistance and Mechanical Properties of C\textsubscript{f}/SiC/SiHfBOC Composites prepared by chemical vapor infiltration with precursor infiltration pyrolysis

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Abstract:
To further improve the oxidation resistance of PDC (Polymer Derived Ceramic) composites in harsh environments, C\textsubscript{f}/SiC/SiHfBOC composites were prepared by CVI (Chemical Vapor Infiltration) and PIP (Precursor Impregnation Pyrolysis) methods. The weight retention change, mechanical properties, and microstructure of C\textsubscript{f}/SiC/SiHfBOC before and after oxidation in air were studied in detail. Microscopic analysis showed that only the interface between the ceramic and fibers were oxidized to some extent and hafnium had been enriched on the composite surface, after oxidation at different oxidation environments. After C\textsubscript{f}/SiC/SiHfBOC composites oxidized at 1500 °C for 60 min, it was mainly determined by the HfO\textsubscript{2} and HfSiO\textsubscript{4} phase. Moreover, the weight retention ratio and compressive strength of the C\textsubscript{f}/SiC/SiHfBOC composites are 83.97 % and 23.88 ± 3.11 MPa, respectively. It indicates that the C\textsubscript{f}/SiC/SiHfBOC composites can be used for a long time in the oxidation environment at 1500 °C.

Keywords: C\textsubscript{f}/SiC/SiHfBOC composites, Precursor infiltration pyrolysis method, Mechanical properties, High-temperature oxidation resistance

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1. Introduction

With the emergence of hypersonic spacecraft, the requirements of thermal protection materials such as lightweight, strengthening-toughening, and oxidation resistance of thermal protection materials are increasing in the combustion chamber and tail nozzle of aerospace vehicles [1-3]. The thermal protection materials represented by ceramic thermal insulation tiles are difficult to meet the requirements of increasing the service environment temperature of hypersonic spacecraft [4-7].

In addition, ceramic matrix composites have attracted much attention due to their excellent thermal stability and high compressive strength [8]. However, the intrinsic brittleness and poor thermal shock resistance of ultra-high temperature seriously restrict the wide application of ceramic thermal protection materials [9,10]. The high-performance silicon-based precursor ceramics have excellent thermodynamic stability. SiOC based precursor ceramics have been applied in the hot end components of aerospace vehicles. However, the strong carbothermal reduction occurs in the environment above 1300 °C, which seriously restricts the development of silicon-based precursor ceramics. It is found that the high-temperature stability of SiOC ceramics can be effectively improved by doping boron, nitrogen, zirconium, hafnium, and other elements. At present, a variety of precursor ceramics have been synthesized, such as SiBOC [11], SiZrOC [12], SiHfOC [12], SiAlOC [13], SiBCN [14], SiHfBOC [3], SiBCNZr [14], etc.

Continuous carbon fiber (or carbon fiber perform) reinforced ceramic matrix composites have excellent properties, such as low density [15], high toughness [16], high strength, and high reusability [17], especially the toughening property [18], which helps to solve the inherent brittleness of ceramic materials [19]. It not only improves the thermal shock resistance of ceramic materials but also maintains the inherent high-temperature stability and low thermal expansion coefficient of ceramic materials [8]. Therefore, continuous carbon fiber reinforced ceramic matrix composites has been widely studied in recent years.

However, due to the low reaction activity of carbon fiber and the damage of carbon fiber in a high-temperature oxidation environment, the interface bonding performance between continuous fiber-reinforced phase and the ceramic matrix will be reduced, and the excellent characteristics of
continuous carbon fiber cannot be brought into full play. The coating will reduce the area of carbon fiber exposed to oxygen atmosphere, thus improving the oxidation resistance of the composites. Therefore, Chemical vapor deposition (CVD), chemical vapor infiltration (CVI), and hydrothermal methods can be used to prepare coatings on carbon fiber surface [1,17,20]. The types of coatings can include C coating [21], BN coating [22], SiC coating [23], etc. These coatings can not only change the surface roughness of carbon fiber but also improve the interface bonding performance between ceramic matrix and carbon fiber reinforcement. In the process of ceramic pyrolysis, the coating can also protect carbon fibers by inhibiting the surface damage of carbon fiber.

The precursor solution of SiHfBOC was prepared in the previous work. Therefore, in this paper, SiC coating was prepared on the surface of carbon fiber by CVI, and then SiHfBOC precursor solution was ultrasonically impregnated into the framework of carbon fiber preform coated with SiC coating by PIP. After solvothermal reaction and high-temperature pyrolysis, Cf/SiC/SiHfBOC ceramic matrix composites were prepared. Their micromorphology and phase evolution were analyzed. Additionally, the strengthening-toughening mechanism and oxidation resistance of Cf/SiC/SiHfBOC ceramic matrix composites were also evaluated.

2. Experimental

2.1 Materials

The commercially available methyltriethoxysilane (MTES, CH₃Si(OCH₂CH₃)₃, 99 % purity, purchased from Aladdin Co. Ltd., Shanghai, China), boric acid (B(OH)₃ 98 % purity, purchased from Harbin Chemical Reagent Factory Co. Ltd., Harbin, China), tetrachloride hafnium (HfCl₄, 99.5 % purity, purchased from Aladdin Co. Ltd., Shanghai, China), T700 PAN-based carbon fibers (C₇ preform, purchased from Yixing Tianniao High Technology Co. Ltd., Jiangsu, China) and Methyl trichlorosilane and Hydrogen (MTS and CH₃SiCl₃H₂, 99.99 % purity, purchased from Institute of metal research, Chinese Academy of Sciences, Shenyang, China) were used here as received.

2.2 Materials synthesis and processing

In our previous study, the preparation method of the SiHfBOC precursor solution was reported [3]. At the same time, SiC coating was prepared on the surface of carbon fiber by CVI method to protect the carbon fiber and improve its bonding strength with the SiHfBOC precursor solution. After
degumming, the C$_f$ preform (64 mm × 64 mm × 25 mm) was placed in chemical vapor infiltration (CVI) equipment. The MTS mass fraction of the precursor gas mixture was 40 wt.%, the gas flow rate was 40 ml/min, and the deposition temperature was 1000 °C. Afterward, the SiHfBOC precursor solution was ultrasonically impregnated into the C$_f$ preform coated with SiC coating. The precursor solution of SiHfBOC was impregnated into SiC/C$_f$ preform by ultrasonic and then put into the reactor. After a solvothermal reaction at 120 °C for 720 min, the sample was further pyrolyzed in a tubular furnace. Finally, C$_f$/SiC/SiHfBOC composites were prepared by multiple PIP cycle times.

![Illustration of the preparation process of C$_f$/SiC/SiHfBOC composites.](image)

**2.3 Characterization**

This experiment used an X-ray diffractometer (XRD) of the type Empyrean Sharp (Panalytical, Netherlands) equipped with monochromatic Mo Kα radiation at a scan speed of 10 °/min in the 2-theta range of 10-90 °. It was used for crystal identification, phase identification, and quantitative analysis. It has high reception efficiency, sensitivity, and precision. The surface morphology C$_f$, C$_f$/SiC, and C$_f$/SiC/SiHfBOC composites were analyzed by SEM. Scanning electron microscopy (SEM) was performed on a HELIOS NanoLab 600i (FEI, USA) with energy dispersive spectroscopy (EDS). Secondary electron image resolution: 0.9 nm (15 kV), 1.4 nm (1 kV), magnification: 40-600000; acceleration voltage: 0.5-30 kV; ion imaging resolution: 4 nm (30 kV). Acceleration voltage: 0.5-30 kV. The density and porosity of C$_f$ preform, C$_f$/SiC preform and C$_f$/SiC/SiHfBOC composites
were measured and calculated by Archimedes drainage method. The oxidation process of the thermogravimetric analyzer was simulated by a muffle furnace.

2.4 Static oxidation tests and mechanics performance testing

The static oxidation resistance test was carried out in a muffle furnace. The C$_f$/SiC/SiHfBOC composites with the dimension of 10 mm × 10 mm × 10 mm were placed in a corundum crucible, and then the crucible was placed in a muffle furnace at the target temperature. After oxidation at the set oxidation temperature for a period, the oxidized samples were taken out and cooled naturally at room temperature. The weight of samples before and after the oxidation test was recorded by analytical balance. The calculation formula of oxidation weight retention ratio (W%) was shown in Eqs (1):

$$W\% = \frac{m_0 - m_1}{m_0} \times 100\%$$

Where W% is oxidation weight retention ratio of sample; $m_0$, $m_1$ is the weight of the sample before and after oxidation, respectively. The oxidation weight retention ratio is the average of three samples. Finally, the static oxidation resistance of C$_f$/SiC/SiHfBOC composites was analyzed according to formula (1) and SEM, EDS, and XRD results.

The flexural strength and compressive strength of C$_f$/SiC/SiHfBOC composites were tested by three-point bending using DCS-250kN electronic universal material testing machine. The prepared samples of flexural strength were processed into 3 mm × 4 mm × 22 mm, and the samples were ground and polished to eliminate the error. The effective span was set at 16 mm and the loading ratio of indenter was set at 0.5 mm/min. The loading direction was divided into x/y direction and z direction, which were perpendicular to and parallel to the carbon fiber layer respectively. The samples of compressive strength were processed into 10 mm × 10 mm × 10 mm. The loading ratio of the indenter and loading direction were the same as above. The calculation formula of flexural strength ($\sigma_f$) and compressive strength ($\sigma_c$) was shown in formula (2) and (3), respectively [24]:

$$\sigma_f = \frac{3PL}{2bh^2W}$$

$$\sigma_c = \frac{P}{hW}$$
Where $P_f$ and $P_c$ are maximum loads during the test; $L$ is span; $h$ is sample height; $W$ is sample width; $\sigma_f$, $\sigma_c$ are flexural and compressive strength, respectively. The flexural and compressive strength is the average of three samples. Then the strength retention ratio was calculated by the ratio of compressive strength after and before the static oxidation test. And the strengthening and toughening mechanism of C/SiC/SiHfBOC composites were analyzed based on the fracture images observed by SEM.

3. Results and discussion

3.1 Microstructure regulation of C/SiC/SiHfBOC composites

The XRD patterns and microstructure of C$_f$ and SiC/C$_f$ are shown in Fig. 2. The amorphous diffraction peak of C was detected near $2\theta=25.5$° in untreated C$_f$ samples. In addition to the amorphous diffraction peaks of C, the diffraction peaks of SiC at 35.6°, 60.1°, and 71.9° were also found in the XRD patterns of SiC/C$_f$ samples [23,25,26], which further indicated the existence of SiC coating. It can be seen from Figs. 2 (b)-2 (d) that the carbon fiber bundles are evenly arranged. The diameter of carbon fiber monofilament is about 6-7 μm. SiC coating uniformly covers carbon fiber, and the thickness of the SiC coating is about 300 nm.
Archimedes drainage method was used to test the bulk density and porosity of C\textsubscript{f}/SiC/SiHfBOC composites under different PIP cycle times. The specific results are shown in Table 1. With the increase of PIP cycle times, the density of C\textsubscript{f}/SiC/SiHfBOC composites increase gradually, while the porosity decreases gradually. The density of untreated C\textsubscript{f} preform is 0.35 g/cm\textsuperscript{3}, and its porosity is 85.73 %. The density of C\textsubscript{f}/SiC preform is 0.50 g/cm\textsuperscript{3}, and its porosity is 74.86 %. After the third PIP cycle times, the density of C\textsubscript{f}/SiC/SiHfBOC composite was 1.02 g/cm\textsuperscript{3}, and its porosity was 47.98 %. The density of C\textsubscript{f}/SiC/SiHfBOC composites increased by about 0.12 g/cm\textsuperscript{3} after the third to fifth PIP cycle times, and its density growth rate gradually slowed down. After the seventh PIP cycle times, the density of C\textsubscript{f}/SiC/SiHfBOC composites increased only by 0.03 g/cm\textsuperscript{3}, so the density of C\textsubscript{f}/SiC/SiHfBOC composites has reached the upper limit.

| Sample          | Cycle times of PIP | Density (g/cm\textsuperscript{3}) | Porosity (%) |
|-----------------|--------------------|------------------------------------|--------------|
| C\textsubscript{f} | 0                  | 0.35                               | 85.73        |
| C\textsubscript{f}/SiC | 0            | 0.50                               | 74.86        |
| C\textsubscript{f}/SiC/SiHfBOC-3 | 3            | 1.02                               | 47.98        |
| C\textsubscript{f}/SiC/SiHfBOC-4 | 4            | 1.15                               | 41.35        |
| C\textsubscript{f}/SiC/SiHfBOC-5 | 5            | 1.27                               | 35.23        |
| C\textsubscript{f}/SiC/SiHfBOC-6 | 6            | 1.37                               | 30.13        |
| C\textsubscript{f}/SiC/SiHfBOC-7 | 7            | 1.40                               | 28.60        |

Figure 3 are SEM images of C\textsubscript{f}/SiC/SiHfBOC composites after different PIP cycle times. Combined with Table 1, the density of the composite increases with the increase of PIP cycle times. In Fig. 3 (a), the C\textsubscript{f}/SiC/SiHfBOC-4 was prepared after four PIP cycle times, with a density of 1.15 g/cm\textsuperscript{3} and a porosity of 41.35 %. Therefore, a great number of pores can be observed in the fiber preform body, and many carbon fibers have not been filled by SiHfBOC ceramics. The C\textsubscript{f}/SiC/SiHfBOC-5 was prepared after five PIP cycle times, with a density of 1.27 g/cm\textsuperscript{3} and a porosity of 36.84 % (Fig. 3 (b)). However, there are still many holes in the composite, and it is observed that the carbon fiber not coated by SiHfBOC ceramic has decreased significantly. The C\textsubscript{f}/SiC/SiHfBOC-6 was prepared after
six PIP cycle times. Its density is increased to 1.37 g/cm$^3$, and the porosity is 33.44 % (Fig. 3(c)). It is observed that the pores in the composite have been further reduced, and most of the carbon fibers are covered by SiHfBOC ceramics, and the internal filling is relatively complete. The C/SiC/SiHfBOC-7 was prepared after seven PIP cycle times as in Fig. 3 (d), and the surface of the carbon fiber in the composite material is completely wrapped by SiHfBOC ceramic. The density of C/SiC/SiHfBOC composites increased was not obvious, which was 1.40 g/cm$^3$, and its porosity was 28.87 %. SiHf-BOC ceramics covered most of the carbon fibers, that was, the composites were filled with SiHfBOC ceramic matrix. There are only a few holes in the composite, and the maximum pore width is about 3 μm. With the increase of PIP cycle times, the spacing of fibers is uniform, and the surface pores are gradually reduced.

Fig. 3 SEM of C/SiC/SiHfBOC composites after different PIP cycle times: (a) 4 PIP cycle times, (b) 5 PIP cycle times, (c) 6 PIP cycle times, (d) 7 PIP cycle times

It is found that the microstructure and density of the composites can be controlled by the different PIP cycle times, and the macro and micropores in the composites are greatly reduced after seven times of impregnation and pyrolysis.
3.2 Study and analysis of oxidation resistance of C/SiC/SiHfBOC composites

The oxidation behavior of C/SiC/SiHfBOC composites at different PIP cycle times are evaluated by XRD and SEM. As shown in Fig. 4, the XRD phase diagram of C/SiC/SiHfBOC-7 composite is oxidized at 1100 °C for 10 min. It indicates that not only the diffraction peaks of SiC and HfO$_2$ are detected [27], but also the diffraction peaks of HfSiO$_4$ are detected at 20.0 °, 27.1 °, 44.0 °, and 53.7 ° after oxidation test [28].

![XRD pattern of C/SiC/SiHfBOC-7 composites before and after oxidation](image)

The SiHfBOC ceramic content in the sample was less (Figs. 5 (a) and (b)), and foamy oxidation products were formed after oxidation at 1100 °C for 10 min. In Fig. 5 (c), filamentous oxidation products and obvious pores were observed. The pores on the surface of the samples decrease obviously when PIP cycle times reach 7 times (Fig.5(d)). From the local enlarged Fig. 5 (e), the surface of carbon fiber in the composite is completely wrapped by SiHfBOC ceramic, and the SiHfBOC ceramic on the surface of carbon fiber is oxidized by O$_2$ to form particles with different volumes. In Fig. 5, the C/SiC/SiHfBOC composites were gradually filled and the surface was gradually dense after oxidation, as the increase of PIP cycle times. There were filamentous oxidation products, and the pores were also more obvious (Fig. 5 (c)). The pores on the surface of C/SiC/SiHfBOC-7 were significantly reduced after oxidation. From Fig. 5 (e), the SiHfBOC ceramic on the surface of the carbon fiber are oxidized.
to form particles with different volumes. Fig. 5 (e) shows the weight retention ratio of C/SiC/SiHfBOC composites with different PIP cycle times after oxidation. The weight retention ratio of the composite material increases from 93.43 % to 98.75 %, when the PIP cycle times increase from 4 to 7. Combined with Fig. 5 and Table 1, it can be found that the pore of the C/SiC/SiHfBOC composites surfaces gradually decreases with the increase of PIP cycle times. At the same time, the contact area between carbon fiber and oxygen is gradually decreasing, so the oxidation resistance of the composite material is also improved.

Fig. 5 SEM images and weight retention ratio of C/SiC/SiHfBOC composites with different densities oxidized at 1100 ℃ for 10 min: (a) 4 PIP cycle times, (b) 5 PIP cycle times, (c) 6 PIP cycle times, (d), (e) 7 PIP cycle times, (f) weight retention ratio

Figure 6 exhibits the SEM image and EDS surface scanning analysis of the C/SiC/SiHfBOC - 7 after oxidation at 1100 ℃ for 10 min. In Figs. 6 (a) and (g), there are many spherical particles attached to the top of the carbon fiber in C/SiC/SiHfBOC composites, which are similar in volume and evenly distributed. The EDS surface scanning results of the oxidized composites are shown in Fig. 6 (b-f), from which the distribution position and uniformity of Si, Hf, B, O, and C on the z-direction surface of the oxidized C/SiC/SiHfBOC composites can be seen. Combined with the atomic content
ratio in Fig. 6 (h), the Si, Hf, O, and C are the most abundant elements, while the B element is relatively low. This is mainly due to the B element escaping in the form of B$_2$O$_3$ during the oxidation process.

![SEM images and EDS diagram of composite materials after oxidation at 1100 °C for 10 min](image)

Fig. 6 SEM images and EDS diagram of composite materials after oxidation at 1100 °C for 10 min

The C$_x$/SiC/SiHfBOC-7 samples were subjected to a static oxidation treatment in a muffle furnace at 1200 °C, 1300 °C, 1400 °C, and 1500 °C for 10 min, respectively. The micromorphology of samples after static oxidation treatment at different temperatures are shown in Fig. 7. There are bare SiC/C$_t$ and large pores on the surface of the samples after oxidation at 1200 °C (Figs. 7 (a) and 7(e)), which further forms oxygen channels to accelerate the internal oxidation of the composite. The surface of the sample with the same density is gradually filled and coated by the molten oxide layer (Figs. 7 (b)-7 (d) and 7 (f)-7 (h)), with the increase of oxidation temperature, so its pores gradually decrease. After oxidation at 1500 °C for 10 min, a dense oxide film has been formed on the surface of the sample.
Fig. 7 SEM images of Cf/SiC/SiHfBOC composites after 10 min of oxidation at different oxidation temperatures:

(a, e) 1200 °C, (b, f) 1300 °C, (e, g) 1400 °C, (d, h) 1500 °C

Fig. 8 shows the XRD pattern and weight retention ratio of the Cf/SiC/SiHfBOC composites after oxidation at 1200 °C, 1300 °C, 1400 °C, and 1500 °C for 10 min. The oxidized sample is mainly composed of SiC, m-HfO₂, and HfSiO₄, where the diffraction peaks of SiC and part of the diffraction peaks of HfSiO₄ overlap (Fig. 8(a)). When the oxidation temperature is 1200 °C, the m-HfO₂ in the sample is the main crystalline phase. As the oxidation temperature increases, it is found that the diffraction peak of m-HfO₂ gradually decreases, and the diffraction peak of HfSiO₄ gradually increases. This indicates that m-HfO₂ and SiO₂ in Cf/SiC/SiHfBOC composites are more likely to react to form HfSiO₄ in an oxidizing environment with higher temperatures. The weight retention ratio of the composite material decreases from 96.05 % to 94.85 % (Fig. 8(b)), when the oxidation temperature increases from 1200 °C to 1500 °C. With the increase of oxidation temperature, the weight retention ratio of Cf/SiC/SiHfBOC composites gradually decreases, but the decreasing range is small. It shows that as the oxidation temperature increases, the surface dense oxide film of the sample reacts with oxygen gradually becomes severe, so the oxidation resistance of the Cf/SiC/SiHfBOC composites at high-temperature also decreases.
Fig. 8 XRD pattern and weight retention ratio of C/rSiC/SiHfBOC composites after 10 min oxidation at different oxidation temperatures: (a) XRD pattern, (b) weight retention ratio

The micromorphology of the C/rSiC/SiHfBOC-7 composites after static oxidation treatment at 1500 °C in a muffle furnace for 30 min, 60 min, 90 min, and 120 min is shown in Fig. 9. As the oxidation time increases, the surface of the composite material becomes denser and the pores are gradually filled. In Figs. 9 (a) and 9 (e), the sample oxidized at 1500 °C for 30 min has no obvious pores, but the surface is rough. In Figs. 9 (b) and 9 (f), the sample oxidized at 1500 °C for 60 min has some small pores, but the surface roughness decreases. In Figs. 9 (c) and 9 (g), the pores on the surface of the sample after oxidizing at 1500 °C for 90 minutes are the largest, and the distribution is uneven, but the surface is relatively flat. In Figs. 9 (d) and 9 (h), the pores on the surface of the sample oxidized at 1500 °C for 120 min are smaller and the surface is the flattest. The results show that a large amount of CO₂, CO, B₂O₃, and other gases escape during the oxidation process of C/rSiC/SiHfBOC-7 composites, so many pores are formed on the surface of the samples. At the same time, molten SiO₂ and borosilicate glass phases are produced under high-temperature oxidation conditions to fill the surface, thereby gradually smoothing and compacting.
Fig. 9 SEM images of C/SiC/SiHfBOC composites after different oxidation time at 1500 °C:
(a, e) 30 min, (b, f) 60 min, (c, g) 90 min, (d, h) 120 min

Figure 10 (a) exhibits the XRD pattern of the C/SiC/SiHfBOC-7 composites after oxidation at 1500 °C for 30 min, 60 min, 90 min, and 120 min. The diffraction peaks of crystalline SiO$_2$ were observed at 22.0° in the XRD curves of samples oxidized for 60 min, 90 min, and 120 min. Meanwhile, with the increase of oxidation time, the diffraction peaks of crystalline SiO$_2$ and m-HfO$_2$ gradually decreases, while the diffraction peaks of HfSiO$_4$ gradually increase. It shows that the chemical reaction of crystalline SiO$_2$ and m-HfO$_2$ is easier to produce HfSiO$_4$ under the condition of long-time high-temperature oxidation. The oxidation products of C/SiC/SiHfBOC-7 composites are mainly composed of HfSiO$_4$, m-HfO$_2$, SiO$_2$, and SiC. In Fig. 10 (b), the weight retention of C/SiC/SiHfBOC-7 composites first increase, then decreases, and then increases with the extension of oxidation time, but it generally shows a decreasing trend. On the whole, the weight reduction of C/SiC/SiHfBOC-7 composites is very small. According to the analysis in Fig. 9, the C/SiC/SiHfBOC-7 composites is seriously oxidized with the prolongation of oxidation time, high-temperature oxidation resistance is gradually weakened. It also indicates that C/SiC/SiHfBOC-7 composites still have a high weight retention ratio after long time oxidation at 1500 °C. However, the weight retention ratio of C/SiC/SiHfBOC-7 composites after oxidation at 1500 °C for 120 min is roughly the same as that after oxidation for 90 min, indicating that the weight change of the composites after oxidation for 90 min reaches a dynamic equilibrium.
The main phases of the C/SiC/SiHfBOC composites prepared in this paper mainly include C, SiC, HfO$_2$, BC$_x$O$_{3-x}$, B(SiO)$_3$, and SiO$_2$C$_y$, according to the above data and reference [2,14,15,28-32]. Therefore, the high-temperature oxidation mechanism of C/SiC/SiHfBOC composites are analyzed, and the possible chemical reaction formulas under high-temperature oxidation conditions are as follows:

\[
C_f + O_2 \rightarrow CO(g) \quad (4)
\]

\[
C_f + O_2 \rightarrow CO_2(g) \quad (5)
\]

\[
B_2O_3(l) \rightarrow B_2O_3(g) \quad (6)
\]

\[
SiO_xC_y(s) + O_2(g) \rightarrow SiO_2(s) + SiC(s) \quad (7)
\]

\[
BC_xO_{3-x} + O_2(g) \rightarrow B_2O_3(g) + CO_2(g) \quad (8)
\]

\[
B(SiO)_3(s) + O_2(g) \rightarrow B_2O_3(g) + SiO_2(s) \quad (9)
\]

\[
SiC(s) + O_2 \rightarrow SiO_2(s) + CO_2(g) \quad (10)
\]

\[
SiC(s) + O_2 \rightarrow SiO_2(s) + CO(g) \quad (11)
\]

\[
HfO_2(s) + SiO_2(s) \rightarrow HfSiO_4(s) \quad (12)
\]

Comparing the Gibbs free energy of each phase composition in C/SiC/SiHfBOC composites, it is found that the Gibbs free energy of the reaction between SiC and O$_2$ is the lowest [33,34]. Combined
with the morphology, XRD phase diagram, and the above chemical reaction formula under different oxidation conditions, the oxidation behavior can be divided into three parts:

1. The oxidation temperature rises from room temperature to 1100 °C. Due to the difference in thermal expansion coefficients of each phase’s composition and the volatilization of gaseous components, it can be seen from the SEM that there are many cracks and pores on the surface of the composites. Although the generated B$_2$O$_3$ has a certain fluidity in this temperature range, it can protect the composites to a certain extent. However, due to the low B element content in the material, the defects of the surface of the composites cannot be healed. When the temperature continues to rise, the generated B$_2$O$_3$ gradually begins to volatilize. The process continues from low temperature to high-temperature, while a small amount of HfSiO$_4$ is generated.

2. The oxidation temperature rises from 1200 °C to 1500 °C. The SiO$_2$ in the composites begin to melt, so the fluidity of the surface of the composites is greatly improved. The cracks on the surface of the composites are gradually healed, leaving only a few holes. At this time, the oxygen channel is gradually blocked, so that oxygen atoms can no longer enter the interior of the composites to react. Therefore, the thickness of the oxide layer decreases, and the generation of HfSiO$_4$ gradually increases with increasing temperature.

3. After oxidation at 1500 °C for 30 min, there are many pores on the surface of the composites. After 60 minutes of oxidation, the amorphous SiO$_2$ is oxidized to crystalline SiO$_2$. However, with the extension of the oxidation time, the peak intensity of crystalline SiO$_2$ gradually weakens, while the content of HfSiO$_4$ gradually increased.

### 3.3 Characterization of mechanical properties and analysis of strengthening-toughening of C$_f$/SiC/SiHfBOC composites

Figure 11 shows the compressive strength of C$_f$/SiC/SiHfBOC composites in x/y and z directions under different PIP cycle times. The mechanical properties of the C$_f$/SiC/SiHfBOC composites were improved with the increase of PIP cycle times. The compressive strength of samples after three PIP cycle times in the x/y direction and z-direction is 31.32 ± 6.12 MPa and 9.31 ± 1.23 MPa respectively. After five PIP cycle times, the compressive strength of C$_f$/SiC/SiHfBOC-5 was 71.97 ± 8.97 MPa in the x/y direction, and 23.92 ± 4.11 MPa in the z-direction. The density of C$_f$/SiC/SiHfBOC-7 was the
highest after seven PIP cycle times, and the compressive strength of C/SiC/SiHfBOC-7 increased slightly in the x/y direction and z-direction to 77.56 ± 8.56 MPa and 40.03 ± 5.48 MPa, respectively. The increase of compressive strength decreases with the increase of PIP cycle times. As shown in Fig. 11, the compressive strength of C/SiC/SiHfBOC composites in x/y direction and z-direction is positively correlated with PIP cycle times of the sample. Besides, the compressive property of the samples with the same PIP cycle times in the x/y direction is higher than that in the z-direction. The main reason is that the fiber arrangement mode of the three-dimensional carbon fiber preform is that the two-dimensional carbon fiber cloth is arranged along the z-direction, and the z-direction is the needling process.

![Graph showing compressive strength vs PIP cycle times](image)

**Fig. 11** Compressive strength of C/SiC/SiHfBOC composites with different densities

In Fig. 12, the compressive strength-strain curves of C/SiC/SiHfBOC composites consist of three stages: (1) In the elastic strain stage, the whole structure of the sample is not damaged until the compressive strength reaches the maximum value, and the stress-strain curve appears inflection point. Then the internal structure of C/SiC/SiHfBOC composite began to destroy. The initial corresponding slope is the elastic modulus of the composites, that is the compressive modulus; (2) The yield stage of C/SiC/SiHfBOC composites are also called the strain strengthening stage, where irreversible inelastic deformation occurs; (3) In the destruction stage, the composite material is seriously damaged and the structure collapses.
The compressive strength-strain curve in the x/y direction is obvious changed (Fig. 12 (a)), with the increase of PIP cycle times of C/SiC/SiHfBOC composites. It can be seen from the first elastic strain stage that the compressive strength reaches the maximum value when the PIP cycle times is 6, and the composite itself has a large deformation. There are only two stages in the compressive strength-strain curve of C/SiC/SiHfBOC composites in the z-direction (Fig. 12 (b)), namely, elastic stage and yield stage. The reason is that only part of the fiber bundle breaks in the z-direction of the sample under the external load, while the fiber cloth layers perpendicular to the z-direction start to stack without being damaged, so the stress-strain curve of the composite rises continuously.

As shown in Fig. 13 (a), the x/y and z flexural strengths of C/SiC/SiHfBOC-3 composites are $43.12 \pm 4.18$ MPa and $4.93 \pm 0.85$ Mpa, respectively. The flexural strength of C/SiC/SiHfBOC-5 composites reaches $51.27 \pm 4.55$ MPa in the x/y direction and $6.84 \pm 1.44$ MPa in the z-direction after five PIP cycle times. The flexural strength of C/SiC/SiHfBOC in the x/y and z directions is $59.94 \pm 4.24$ MPa and $12.47 \pm 2.17$ MPa after seven PIP cycle times, respectively. Figure 13 (b) shows the flexural strength-displacement curves of C/SiC/SiHfBOC composite after oxidation at five different PIP cycle times. The three stages of flexural strength-displacement curves in the x/y direction of C/SiC/SiHfBOC composite are obvious with the increase of oxidation temperature. It can be seen from the first elastic stage that the flexural strength of composites reaches the maximum value when the PIP cycle times are the maximum, and the composites themself have a large deformation.
Fig. 13 Flexural strength and flexural strength-displacement curves of C/SiC/SiHfBOC composites with different PIP cycle times in the x/y direction and z direction: (a) flexural strength, (b) flexural strength-displacement curves

The fracture morphology of C/SiC/SiHfBOC composites is shown in Fig. 14. In this work, to overcome the intrinsic brittleness of ceramics, continuous carbon fiber was selected as the second phase to toughen. The continuous carbon fibers added to the SiHfBOC ceramic matrix are arranged orderly in the SiHfBOC ceramic matrix, which can greatly improve the flexural strength of C/SiC/SiHfBOC composites. During elastic deformation stage process, the continuous carbon fiber absorbs a lot of energy, and most of the mechanical energy absorbed is converted into surface energy, which is mismatched with the ceramic matrix and begins to debond [35]. In Fig. 14 (b), as the load continues to increase, the crack interface between continuous carbon fiber and ceramic matrix begins to loosen, resulting in stress relaxation. The ceramic matrix is damaged and then spalling occurs. The continuous carbon fiber is exposed, which is called the fiber pull-out phenomenon. When the carbon fiber is pulled out, it will continue to absorb external energy, and absorb the most energy. The continuous carbon fiber begins to fracture (Fig. 14 (c)). In the process of carbon fiber fracture, when the crack extends to the surface of carbon fiber, the carbon fiber will continue to absorb energy and deform, until the critical point of carbon fiber fracture, it cannot continue to absorb energy, and finally fracture. After the fracture, the carbon fiber will recover to the original state, and then release the energy absorbed before. It is indicated that the strengthening and toughening mechanism of C/SiC/SiHfBOC composites are mainly composed of fiber debonding, fiber pulling out, and fiber
fracture. The debonding and pullout of continuous carbon fibers play an important role in the failure of C/SiC/SiHfBOC ceramic matrix composites during flexural fracture experiments.

![Fracture morphology of C/SiC/SiHfBOC composites: (a) fiber debonding, (b) fiber pulling out, (c) fiber fracture](image)

Fig. 14 Fracture morphology of C/SiC/SiHfBOC composites: (a) fiber debonding, (b) fiber pulling out, (c) fiber fracture

The compressive strength of C/SiC/SiHfBOC composites in x/y direction has three stages, which are representative of research. Therefore, the analysis of oxidative compressive strength in x/y direction is taken as the research object. The compressive strength and compressive strength-strain curves of C/SiC/SiHfBOC-7 composites after oxidation at five oxidation temperatures are shown in Fig. 15. The compressive strength in the x/y direction of C/SiC/SiHfBOC-7 composites after oxidation at 1100 °C, 1200 °C, 1300 °C, 1400 °C and 1500 °C for 10 min is 75.16 ± 8.17 MPa, 64.93 ± 7.22 MPa, 60.53 ± 7.01 MPa, 55.19 ± 5.25 MPa, 39.59 ± 4.78 Mpa (Fig. 15 (a)), respectively. The results show that the compressive strength of C/SiC/SiHfBOC composites decreases with the increase of oxidation temperature. The compressive strength of C/SiC/SiHfBOC composites decreases with the increase of oxidation temperature. The compressive strength decreased to 96.90 % after oxidation at 1100 °C and 51.03 % after oxidation at 1500 °C., but it still has high mechanical properties after a short time of high-temperature oxidation. Three stages of the stress-strain curve in the x/y direction can be observed (Fig. 15 (b)). In the first elastic stage of all composites, the compressive strength is the largest and the deformation is the smallest, when the oxidation temperature is 1100 °C.
Fig. 15 Compression strength and compressive strength-strain curves of C/SiC/SiHfBOC composites in the x/y direction at different oxidation temperatures: (a) compression strength, (b) compressive strength-strain curves

Figure 16 exhibits the compressive strength and compressive strength-strain curves of C/SiC/SiHfBOC-7 composite after four oxidation time at 1500 °C. The compressive strength of C/SiC/SiHfBOC-7 composite in the x/y direction is 30.38 ± 3.55 MPa, 23.88 ± 3.11 MPa, 18.68 ± 2.34 MPa and 16.50 ± 2.08 MPa after oxidation at 1500 °C for 30 min, 60 min, 90 min, and 120 min, respectively. The compressive strength decreased to 39.17 % after 30 min oxidation, 24.08 % after 90 min oxidation, and 21.27 % after 120 min oxidation. The results show that the compressive strength of C/SiC/SiHfBOC-7 composites decreases with the increase of oxidation time, and the compressive strength changes little after oxidation at 1500 °C for 90 min, and 120 min. Fig. 16 (b) shows the stress-strain curves of C/SiC/SiHfBOC-7 composites after oxidation at 1500 °C for a different time. Three stages of the stress-strain curve in the x/y direction can be seen (Fig. 16 (b)), with the increase of oxidation time. From the first elastic stage, when the oxidation time is 30 min, the compressive strength reaches the maximum value, and the sample itself has a large deformation. Moreover, the strain range of yield point in the first stage is between 2 % and 7 %. After reaching the yield point, stress yield begins to occur.
4 Conclusion

In this paper, the on-demand preparation of Cf/SiC/SiHfBOC composites is realized, to control the microstructure and properties of the composites. When the PIP cycle times increased from 3 to 7, the density and porosity of the composites increased from 1.02 g/cm³ and 52.53 % to 1.40 g/cm³ and 28.87 %, respectively. And the compressive strength in the x/y and z directions increased from 31.32 ± 6.12 MPa and 9.31 ± 1.23 MPa to 77.56 ± 8.56 MPa and 40.03 ± 5.48 MPa, respectively. The strengthening and toughening methods mainly include fiber debonding, fiber pulling out, and fiber breaking. The products of Cf/SiC/SiHfBOC composites after the static oxidation test mainly include CO₂, CO, B₂O₃, SiO₂, HfSiO₄, etc. According to the analysis of compressive stress-strain curve, oxidation weight retention rate, and oxidation surface SEM, it was found that the weight retention and compressive strength of Cf/SiC/SiHfBOC-7 sample were 83.97 % and 23.88 ± 3.11 MPa respectively after being oxidized at 1500 °C for 60 min, which further proved that the Cf/SiC/SiHfBOC composites could be used in oxidation environment at 1500 °C for a long time.

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