Review
Steam Oxidation of Silicon Carbide at High Temperatures for the Application as Accident Tolerant Fuel Cladding, an Overview

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Abstract: Since the nuclear accident at Fukushima Daiichi Nuclear Power Station in 2011, a considerable number of studies have been conducted to develop accident tolerant fuel (ATF) claddings for safety enhancement of light water reactors. Among many potential ATF claddings, silicon carbide is one of the most promising candidates with many superior features suitable for nuclear applications. In spite of many potential benefits of SiC cladding, there are some concerns over the oxidation/corrosion resistance of the cladding, especially at extreme temperatures (up to 2000 °C) in severe accidents. However, the study of SiC steam oxidation in conventional test facilities in water vapor atmospheres at temperatures above 1600 °C is very challenging. In recent years, several efforts have been made to modify existing or to develop new advanced test facilities to perform material oxidation tests in steam environments typical of severe accident conditions. In this article, the authors outline the features of SiC oxidation/corrosion at high temperatures, as well as the developments of advanced test facilities in their laboratories, and, finally, give some of the current advances in understanding based on recent data obtained from those advanced test facilities.

Keywords: SiC; ATF; fuel cladding; steam oxidation

1. Introduction
The severe accident in 2011 at Fukushima Daiichi Nuclear Power Station (FDNPS) has stagnated the usage of nuclear energy but has also accelerated the research and development for safety enhancement of light water reactors (LWRs). Soon after the occurrence of the Great East Japan earthquake, emergency scram of the FDNPS operating reactors was activated automatically as were the emergency power supplies to ensure fuel cooling operations. However, approximately 50 min after the initial earthquake, a huge tsunami struck the station, flooding the emergency power supplies (particularly diesel generators), which led to a loss of the reactor core cooling, resulted in lowered coolant level in the Reactor Pressure Vessel (RPV), and a temperature increase of fuel cores. Initially, this loss of coolant accident (LOCA) resulted in decay heat temperature increases of approximately 0.5 °C/s. When the fuel temperature attained around 1200–1400 °C, then a steam oxidation of zirconium (major component of fuel claddings and channel box) with steam accelerated, generating a huge amount of hydrogen and significant chemical heat to drive temperatures higher. Furthermore, boron carbide (B\textsubscript{4}C, as a neutron absorber) contained in the control blade could react not only interact with the stainless steel (lowering its melting point) but also react with steam and further accelerate the hydrogen and heat generation, as seen in the following equations [1]:

\[
\text{Zr} + 2 \text{H}_2\text{O} \rightarrow \text{ZrO}_2 + 2 \text{H}_2 (\Delta H = -6.4 \text{ MJ/kg-Zr}),
\]
B_4C + 8 H_2O → 2 B_2O_3 + CO_2 + 8 H_2 (ΔH = −15 MJ/kg-B_4C). \hspace{1cm} (2)

Once the exothermic reactions of Zry and B_4C in steam have started, they are hardly possible to mitigate or control. Besides the nuclear decay heat, these exothermic reactions became a significant heat source and result in local temperature escalations with prototypically 8 °C/s to values where there is significant degradation (melting/slumping) of the fuel. Once fuel cores have lost their original geometry and loss of coolant channels, the retrieval of heat removal capability is very difficult, even when the cooling system has been revived due to the blockage of the coolant channel. The water/steam supply needs to be able to cool large melted fuel masses of low surface area/volume ratios.

The meltdown sequence starts from the failure of control blades, which is initiated by the eutectic reaction between B_4C and SS at approximately 1250 °C [2–7]. This temperature is far below the melting points of these components. The B_4C-SS melt then attacks the zircaloy (Zry) channel boxes and penetrating them to reach the fuel rods. The degradation of the irradiated fuel rods, consisting of UO_2 fuel pellets in Zry cladding, starts with the liquefaction interaction between the partially oxidized inner Zry cladding (Zr(O)) and the UO_2 pellet at around 1900 °C [8–10], far below the melting allows the release of gaseous and volatile fission products (FPs) collected in the clad free space to the external steam atmosphere.

Concepts of accident tolerant fuel (ATF) had been studied to mitigate or block these chemical interactions by introducing advanced fuel and cladding materials replacing either UO_2 fuel and/or conventional Zry cladding [11]. Regarding the ATF-cladding, three major concepts have been proposed, such as silicon carbide [12,13], iron-based alloys [14–16], and coated zirconium-based claddings [17,18]. The motivation for transitioning away from the conventional Zry-cladding to the ATF-cladding alternatives is to delay or mitigate the chemical reactions and to reduce the amount of heat and hydrogen generation from the cladding steam oxidation of the. Each concept has a unique conceptual idea, based on the attributes of the base materials. For example, the concept of environmental barrier coatings of Zry is to delay or reduce the zirconium-steam interaction in the early stages. This concept is applicable to current nuclear safety technology and regulation as so-called “evolutionary” concept [11]. The idea of using iron-based alloys is to replace the major component of fuel cladding from zirconium to iron, which has “less heat generation capability”, hence mitigating the catastrophic temperature increase and H_2 generation in the early stage of accident. It also draws on the experience of rapid (fast neutron flux) reactors, where 316 stainless steels have been the standard cladding materials. Furthermore, Al and Cr as alloying elements should provide the capability to form protective oxide layers. The concept of replacing the major component of fuel cladding from zirconium to silicon carbide (SiC) is to decrease the heat and hydrogen generation capabilities, and to maintain the fuel integrity even at high temperatures, which could maintain the ability to suppress the release of fission products. However, in order to use a ceramic material as a fuel cladding, a significant improvement of LWR-relevant conventional technologies is needed (hence, SiC-cladding is a so-called “revolutionary” concept).

Among the potential ATF cladding materials, SiC seems to stand out in terms of oxidation resistance [19,20] and structural integrity at high temperatures [21,22], high radiation damage tolerance [23,24], high thermal conductivity [25], and high neutron economy [26]. In spite of these potential benefits of SiC cladding, however, there are some key issues that need to be addressed for the LWRs application, such as hydrothermal corrosion and fission product retention capability [23,27,28]. The present review attempts to outline: (1) features of high temperature SiC corrosion by comparing relevant reactions in steam to those in air or oxygen atmospheres, (2) the further needs for improved test facilities for SiC steam corrosion tests, and (3) the current knowledge based on recent data obtained at Japan Atomic Energy Agency ( JAESA) and Karlsruhe Institute of Technology (KIT).
2. Features of SiC Oxidation/Corrosion at High Temperatures in “Dry” and “Wet” Atmospheres and Remaining R&D Issues for ATF-Cladding

2.1. Oxidation/Corrosion of SiC at High Temperatures in “Dry” Atmospheres

As expected for high-temperature applications, such as components in gas turbine engines and LWRs, high-temperature oxidation of silicon carbide has been extensively studied in various environments and temperatures ranges. Oxidation of SiC at high temperatures in dry atmospheres, such as O$_2$-Ar, CO$_2$/CO environments, and technical helium gas as used in high-temperature gas-cooled reactors [29–33], can be classified into three features as active oxidation, passive oxidation and bubble formation [34–37]. At high oxygen potentials, the oxidation of SiC results in formation of a protective SiO$_2$ layer accompanying a mass gain, which is termed “passive oxidation” [34,38]:

$$\text{SiC}_\text{s}) + 3/2 \text{O}_2\text{(g)} = \text{SiO}_2\text{(s)} + \text{CO}_\text{2(g)}. \quad (3)$$

At lower oxygen potential, the oxidation of SiC at high temperatures induced a mass loss due to the formation of gaseous SiO, which is termed “active oxidation” [34,38]:

$$\text{SiC}_\text{s}) + 2 \text{SiO}_2\text{(s)} = 3 \text{SiO}_\text{2(g)} + \text{CO}_\text{2(g)}. \quad (4)$$

Figure 1 illustrates the expected weight change during the active and the passive oxidations which have been reported for Si and SiC [29,39,40]. In the passive regime, the chemical reaction, as shown in Equation (3), results in the formation of protective SiO$_2$, thus inducing a mass gain, whereas, in the active regime, the chemical reaction as shown in Equation (4) occurs, which causes the volatilization of the surface SiO$_2$ layer (formation of gaseous SiO) and leads to a mass loss. Reaction 4 also indicates there is always a SiO$_2$ surface layer to react with SiC substrate to form volatile SiO(g), and there is no SiC surface layer in contact with the atmosphere.

![Figure 1. Schematic diagram on weight change during active and passive oxidations of SiC according to oxygen partial pressure (drawn by the authors referring to data reported by Jacobson [39]).](image)

Oxidation mechanisms in active regime and the active-to-passive transitions can be classified into two types [40], as shown in Figure 2, according to the ambient environments. For example, Goto [38] observed a clear transition of active-to-passive of SiC in O$_2$-Ar environment at 1600 °C, in which the mass loss proceeds up to P$_{O2}$ = 146 Pa (volatilization of SiO), and, at P$_{O2}$ = 160 Pa, mass gain occurs (volatilization of SiO ceases and formation of SiO$_2$ as a non-volatile and protective layer dominates). The oxidation behavior of SiC in CO/CO$_2$ environment is a contrast to that in O$_2$-Ar atmosphere. In active oxidation regime of SiC, the active oxidation rate shows a maximum value at a certain value of P$_{CO2}$/P$_{CO}$ ratio. It is suggested that the vaporization of SiO and formation of SiO$_2$ in...
CO/CO₂ environment might proceed simultaneously (neither of them becomes dominant) during the active oxidation regime as the following reactions [37,40]:

\[
\text{SiC} (s) + 3 \text{CO}_2(g) = \text{SiO}_2(s) + 4 \text{CO}(g) \quad \text{(5)}
\]

\[
\text{SiO}_2(s) + \text{CO}(g) = \text{SiO}(g) + \text{CO}_2(g) \quad \text{(6)}
\]

**Figure 2.** Schematic diagram showing the differences of oxidation rate and active-to-passive transition of SiC oxidation in active regime in Ar-O₂ and CO-CO₂ atmospheres [40].

In other words, the high temperature oxidation behavior of SiC in dry air is fairly complicated and can vary according to the environment. Comprehensive reviews of SiC oxidation at high temperatures in various environments (dry and wet air) have been reported elsewhere [39,40].

### 2.2. Oxidation/Corrosion of SiC at High Temperatures in “Wet” Atmospheres

The steam oxidation of SiC at high temperatures can be categorized into two different processes, known as SiO₂ formation and its volatilization, as shown in Figure 3. At high temperatures, SiC reacts with water vapor to form SiO₂ on the sample surface via the following reaction:

\[
\text{SiC}(s) + 3 \text{H}_2\text{O}(g) = \text{SiO}_2 + 3 \text{H}_2\text{O}(g) + \text{CO}(g) \quad \text{(7)}
\]

The formation of SiO₂ is the diffusion rate-controlled process, which depends on the diffusion of oxidizing species through the oxide layer, thus obeying parabolic laws. The kinetics due to the formation of the oxide layer is described by:

\[
\frac{dx_1}{dt} = \frac{k_p}{2x} \quad \text{(8)}
\]

where \(x_1\) can be the mass gain or the thickness growth of the oxide scale. \(k_p\) is the parabolic rate constant and \(t\) the time. The driving force of this process is the difference of steam partial pressure at the top surface and the interface of SiO₂-SiC. It means that the parabolic rate constant is proportional to the steam partial pressure on the top surface [41]:

\[
k_p \propto P(\text{H}_2\text{O})^n \quad \text{(9)}
\]

where \(P(\text{H}_2\text{O})\) is water vapor partial pressure, and \(n\) the power law exponent. On the surface, the oxide scale reacts with water vapor at high temperatures and undergoes volatilization via, e.g., the following reaction:

\[
\text{SiO}_2(s) + 2\text{H}_2\text{O}(g) = \text{Si(OH)}_4(g) \quad \text{(10)}
\]
The volatilization of SiO$_2$ depends on the chemical reaction on the surface, and transport of gaseous products away from the surface, thus obeying linear laws. The change due to the volatilization of SiO$_2$ is described as:

$$\frac{dx_2}{dt} = k_l$$

where $x_2$ can be mass loss or thickness reduction of the SiO$_2$ scale. $k_l$ is the linear rate constant and $t$ the time. According to Opila et al. [42], the linear constant $k_l$ is proportional to the velocity ($v_o$) and total pressure $P_{total}$, as shown in the following equation:

$$k_l \propto v_o^{1/2} \frac{P(H_2O)^2}{P_{total}^{1/2}}.$$  

As steam oxidation of SiC at high temperatures is a combination of the SiO$_2$ formation and its volatilization, the mass/thickness evolution of the oxide scale follows the paralinear behavior [42,43] as shown in Equation (13).

$$\frac{dx}{dt} = \frac{dx_1 - dx_2}{dt} = \frac{k_P}{2x} - k_l.$$  

The oxidation behavior of SiC in steam at high temperatures obeys paralinear kinetics, which can be different from the oxidation behavior of SiC in dry air, as shown in Figure 4. Oxidation mechanism of SiC in dry air can be either active or passive oxidation, depending on oxygen potential. In wet atmosphere, the formation of SiO$_2$ at the SiC-SiO$_2$ interface and the volatilization of SiO$_2$ on the top surface happens simultaneously, which results in paralinear kinetics. Therefore, the high temperature oxidation of SiC in steam environments should be discussed separately, especially for applications in LWRs.

![Figure 3](image3.png)

**Figure 3.** Illustration of SiC oxidation in steam at high temperatures.

![Figure 4](image4.png)

**Figure 4.** Schematic image on oxidation kinetics of SiC at high temperatures in dry and wet atmospheres.
2.3. Oxidation Kinetics of SiC at High Temperatures and R&D Issues for ATF-Cladding

Oxidation behavior of SiC at high temperatures in wet air, such as H₂O/Ar [44–47] and H₂O/O₂ [30,45,48], have been reported. One has to recognize an important difference in the steam oxidation of SiC in “wet” atmospheres, in which the formation of both a condensed SiO₂ oxide and volatile phases, such as Si(OH)₄, is induced at the same time. While the formation of SiO₂ scale is a diffusion rate-controlled process, which obeys a parabolic law, the formation of volatile products depends on mass transport of the volatile products away from the surface, which yields linear kinetics [49,50].

To obtain 𝑘ₚ and 𝑘ₗ values based on Equation (13), one of the two terms (𝑘ₚ, 𝑘ₗ) needs to be known. For example, Opila et al. [42] conducted a separate experiment to directly obtain the linear volatilization rate (𝑘ₗ) of SiO₂ by exposing a fused quartz coupon to a 50%O₂/50%H₂O gas mixture using the thermogravimetric analysis (TGA) procedure. A similar solution has been proposed by Terrani et al. [13], in which electrically fused silica specimens were exposed in 100%H₂O using TGA and high temperature furnace (HTF) to obtain the volatilization rate 𝑘ₗ. The investigation conditions of the volatilization rate (𝑘ₗ) of SiO₂ obtained by various authors are shown in Table 1. Experimental results of the investigations for linear volatilization rate of SiO₂ reported by Opila [42] and Terrani [13] are summarized in Figure 5. After obtaining the volatilization rate 𝑘ₗ of SiO₂, the parabolic oxidation rate 𝑘ₚ can be determined by Equation (11) and the given experimental data (specifically, the mass change and investigation duration). Another method to obtain 𝑘ₚ and 𝑘ₗ together is a curve fitting technique (nonlinear least-squares analysis) of experimental data. Observing the paralinear kinetics of SiC, as well as the volatilization of fused silica, Opila concluded that the above techniques give similar results for the linear volatilization of SiC in water vapor [42].

Table 1. Investigation conditions of volatilization rate of SiO₂ reported by various authors.

| Author [Ref. No.] | Method | Specimen | Atmosphere          | Velocity, cm/s | Temperature (°C) |
|-------------------|--------|----------|---------------------|----------------|-----------------|
| Opila [42]        | TGA    | SiO₂     | 50% H₂O/O₂          | 4.4            | 1200–1400       |
| Opila [42]        | TGA    | Preoxidized SiC | 50% H₂O/O₂ | 4.4            | 1200–1400       |
| Terrani [13]      | TGA    | SiO₂     | 100% H₂O           | 1.3–1.6        | 1200–1500       |
| Terrani [13]      | HTF    | SiO₂     | 100% H₂O           | 18–138         | 1400–1600       |
| Pham [47]         | LAHF   | SiC      | 97% H₂O/Ar         | 1200           | 1400–1800       |

Figure 5. Summary of experimental results for linear volatilization rate of SiO₂ reported by Opila [42], Terrani [13], and Pham [47].
As the parabolic oxidation rate constant $k_p$ is proportional to the steam partial pressure, various authors have reported the oxidation kinetics of SiC in different environments. Table 2 briefly summarizes the investigation conditions of major studies for oxidation kinetics of CVD-SiC at high temperature in various steam partial pressures [13,42,45,47,48,51]. Representative results of these studies are presented in an Arrhenius-type diagram (Figure 6), which describes the relationship between logarithm of parabolic rate constant and reciprocal of temperature. The parabolic oxidation rate constant of SiC at a certain temperature, as shown in Figure 6, can vary, depending on many factors, such as the steam partial pressure and examination methods. The slopes (the activation energies) of the lines in Figure 6 give information about the diffusing species, which controls the process. In general, the activation energy for the permeation of a molecular species is less than that for the diffusion of a charged species, such as OH$^-$ and O$^{2-}$. For example, Deal [52] concluded that the activation energies for the oxidation of silicon in dry oxygen and water vapor by permeation mechanism to be 119 and 68 kJ/mol, respectively. Narushima [48] reported that the activation energy for SiC oxidation in 10 vol% H$_2$O/O$_2$ was 397 kJ/mol and stated that O$^{2-}$ was the diffusing species. In another study, Doremus [53] discussed that the diffusion of charged oxygen species is expected to have an activation energy magnitude similar to the bond energies of Si-O and Si-OH bonds (approximately 377 kJ/mol). More details of discussion on the activation energy for oxidation of SiC at high temperatures can be found elsewhere [45].

Table 2. Investigation conditions of major studies for oxidation kinetics of SiC in steam.

| Author [Ref. No.] | Method | Specimen | Atmosphere | Temperature (°C) |
|-------------------|--------|----------|------------|-----------------|
| Narushima [48]   | TGA    | CVD-SiC  | 10% H$_2$O/O$_2$ | 1550–1650       |
| Opila [51]       | TGA    | CVD-SiC  | 10% H$_2$O/O$_2$ | 1200–1400       |
| Opila [45]       | HTF    | CVD-SiC  | 10% H$_2$O/O$_2$ | 1100–1400       |
| Opila [45]       | HTF    | CVD-SiC  | 25% H$_2$O/O$_2$ | 1100–1400       |
| Opila [45]       | HTF    | CVD-SiC  | 50% H$_2$O/O$_2$ | 1100–1400       |
| Opila [45]       | HTF    | CVD-SiC  | 50% H$_2$O/O$_2$ | 1200–1400       |
| Opila [45]       | HTF    | CVD-SiC  | 70% H$_2$O/O$_2$ | 1100–1400       |
| Opila [45]       | HTF    | CVD-SiC  | 90% H$_2$O/O$_2$ | 1100–1400       |
| Terrani [13]     | TGA    | CVD-SiC  | 100% H$_2$O | 1200–1500       |
| Terrani [13]     | HTF    | CVD-SiC  | 100% H$_2$O | 1400–1600       |
| Pham [47]        | LAHF   | Monolithic SiC | 97% H$_2$O/Ar | 1400–1800       |

Figure 6. Major reports on the parabolic oxidation rate, $k_p$, of SiC in steam at high temperatures.
To investigate the influence of steam partial pressure on the parabolic oxidation rate of SiC (as seen in Equation (9)), Opila conducted oxidation tests of CVD-SiC at temperatures of 1100–1400 °C in H2O/O2 gas mixtures with compositions of 10–90 vol% water vapor at a total pressure of 1 bar [45]. The results of the investigation are presented in Figure 7, where the parabolic oxidation rate constant increased with the increase of steam partial pressure. A range of possible activation energies of 28–156 kJ/mol has been reported, which indicated that the molecular permeation mechanism is more dominant than a diffusion mechanism that involves a charged species.

Impurities from environment can also affect the oxidation of SiC in steam at high temperatures. Opila [51] investigated the oxidation kinetics of CVD-SiC at 1200–1400 °C in 10 vol% H2O/O2 mixture using TGA with either pure Al2O3 or fused quartz reaction tubes. Results of the investigation (Figure 8) indicated that the parabolic oxidation rate of SiC samples tested in Al2O3 reaction tube was about one order of magnitude higher than that of SiC samples tested in quartz tube. It was found that the steam oxidation of SiC in Al2O3 reaction tube led to sodium and aluminum contamination of the oxide scale. The sodium and aluminum might be transported from the Al2O3 reaction tube to the SiC sample surface in the gas phase to form less protective oxide scales.

**Figure 7.** Influence of steam partial pressure on parabolic oxidation rate constant reported by Opila [45].

**Figure 8.** Effects of sodium and aluminum from the reaction tube (alumina tube) on parabolic oxidation rate of SiC in steam reported by Opila [51].
Due to the characteristics of the applications of SiC, most of the oxidation studies of SiC focused at temperatures ranging from 1000 to 1600 °C. Although the normal cladding temperature of cladding is approximately 300 °C, SiC ATF claddings may suffer temperatures over 2000 °C in the event of severe accidents [54]. It is necessary to verify the performance of SiC cladding, as well as other potential candidates for ATF claddings under severe accident conditions, above at least 1600 °C in steam. However, the oxidation kinetics of SiC is still insufficient studied due to the lack of test facilities capable of operating in steam at temperatures above 1600 °C.

3. Advanced Test Facilities for ATF-Claddings

Steam oxidation tests at temperatures above 1600 °C are very challenging as major refractory lining materials, such as alumina and even zirconia, in conventional test facilities are at their limits. In recent years, several efforts have been made to modify existing or to develop new test facilities [46,47] to study SiC under extreme conditions. To perform the oxidation tests at temperatures higher than 1600 °C, the QUENCH-SR facility with utilization of induction heating was developed at Karlsruhe Institute of Technology [55]. First oxidation and quench tests of SiC at temperatures up to 2000 °C in steam atmosphere were reported by Avincola et al. [46]. Under the same motivation, JAEA has also developed a new test facility (Laser Heating Facility: LAHF) with utilization of laser heating for oxidation tests at extreme temperatures [47]. In this section, we introduce these two facilities.

3.1. Laser Heating Facility (LAHF)

A schematic diagram of the laser heating facility is shown in Figure 9. This apparatus was designed for heat-treatment tests of small samples (less than 20 mm in diameter) in controlled environments (including vacuum) at temperatures beyond 1600 °C. The main unit includes a laser head, vacuum pump, gas line heaters, flow rate controllers, and airtight test vessel which is equipped with two radiation pyrometers, a W-Re 5-26 thermocouple, and a video camera. Sample holders are made of zirconia. The laser light transmission, pyrometer temperature measurement, and in situ observation were made via five quartz windows (synthetic silica) located in the vessel wall. To avoid water vapor condensation, the test vessel, including gas pipes and quartz windows, are heated up to 200 °C using resistance heating. The flow rates can be controlled in the range of 0.1–3.0 g/min (water vapor), 0.1–5.0 L/min (= 0.18–8.9 g/min) (Ar), and 0.06–3.14 g/min (hydrogen), respectively. The 2500 W laser module allows the sample to be heated at the rate of 1000 °C/s. Parameters, such as laser beam power, sample temperatures, and gas flow rates, are monitored in a control system and recorded by a datalogger. After passing through a steam trap, outlet gases, such as hydrogen and carbon monoxide, are sampled and analyzed using a gas chromatograph.

![Figure 9. Schematic of the laser heating facility (LAHF) [47].](image-url)
Figure 10 shows experimental data of a preliminary investigation for steam oxidation of CVD-SiC/SiC composites under 1 bar pressure, in a stream of 50% H\textsubscript{2}O/Ar gas mixture, and at temperatures up to 2200 °C with 0.1 °C/s heating rate. In the temperature range of 200 to 1650 °C, the LAHF was automatically controlled by feedback of the pyrometer detecting sample temperature at the periphery. Above 1650 °C, the LAHF was manually controlled because the feedbacks from pyrometers are no longer reliable (pyrometer measurement is affected by the change of sample surface and the formation of gaseous products during the test). Using the laser current output and the immediately preceding temperature trends (before the interference mentioned above), the surface temperature of the sample at the center is estimated. An example is shown in Figure 10a–d. Hydrogen production during the test can be divided into 3 stages (below 1800 °C, 1800–2000 °C, and above 2000 °C). Below 1800 °C (points a and b): before the appearance of bubbles, the amount of H\textsubscript{2} produced during the test was less than 10 L/h/m\textsuperscript{2}SiC, indicating the excellent oxidation resistance in steam. In the temperature range of 1800–2000 °C (point c), the formation of bubbles on sample surface led to the degradation of SiC oxidation resistance in steam and an acceleration of H\textsubscript{2} production (to 700~800 L/h/m\textsuperscript{2}SiC). Above 2000 °C (point d), a significant increase of H\textsubscript{2} production (up to 1000 L/h/m\textsuperscript{2}SiC) was observed. However, the causes of this final increase are not fully understood.

3.2. QUENCH-SR Facility

The QUENCH-SR (single rod) facility at KIT Karlsruhe with inductive heating and the possibility of quenching of the samples by water is applied for steam oxidation tests to very high temperatures. The samples are enclosed in a quartz glass tube allowing observation of the tested sample by video recording. A water-cooled copper coil surrounding the quartz glass tube generates a magnetic field inducing eddy currents in an electrically conductive sample. The power is provided by a high frequency (HF) generator with a power of 20 kW working at a frequency up to 700 kHz. The temperature is measured and controlled via a two-color pyrometer (type IGAR 12-LO MB22) with a measuring range of 500 to 2200 °C working at the wavelengths 1.28 µm and 1.65 µm. Additionally, thermocouples can be attached on the surface of the samples. Gas supply is controlled by a Bronkhorst® gas flow controller, water flow controller, and a CEM (controlled evaporator and mixer). Usually, there is 40 L/h (max. 100 L/h) argon flow through the facility throughout all tests, and 60 g/h (max. 75 g/h) steam is injected during the oxidation phases resulting in a steam concentration of 65 vol%.

For metallic samples, the heating power is directly produced in the sample. SiC samples, however, are filled with graphite (used as a susceptor for the HF inductive heating purpose [46]) and tightly sealed with end caps at both sides to avoid interaction of the graphite with the oxidizing steam atmosphere. This heating method results in uniform and prototypical heat generation for cladding tube samples as in fuel rods, where the heat is also produced by the UO\textsubscript{2} fuel within the cladding. Figure 11 provides some illustrations of the facility and typical samples.

A number of experiments with SiC f-SiC ceramic-matrix composite (CMC) cladding tubes from various manufactures have been conducted already with this facility. Figure 12 provides, as one example, results of a transient test with such a cladding tube segment.
glass tube generates a magnetic field inducing eddy currents in an electrically conductive sample. The power is provided by a high frequency (HF) generator with a power of 20 kW working at a frequency up to 700 kHz. The temperature is measured and controlled via a two-color pyrometer (type IGAR 12-LO MB22) with a measuring range of 500 to 2200 °C working at the wavelengths 1.28 µm and 1.65 µm. Additionally, thermocouples can be attached on the surface of the samples. Gas supply is controlled by a Bronkhorst® gas flow controller, water flow controller, and a CEM (controlled evaporator and mixer). Usually, there is 40 L/h (max. 100 L/h) argon flow through the facility throughout all tests, and 60 g/h (max. 75 g/h) steam is injected during the oxidation phases resulting in a steam concentration of 65 vol%.

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A number of experiments with SiCf-SiC ceramic-matrix composite (CMC) cladding tubes from various manufactures have been conducted already with this facility. Figure 12 provides, as one example, results of a transient test with such a cladding tube segment from 1400 °C to 2000 °C [46], which correspond very well to the results obtained with smaller CVD-SiC samples in the LAHF facility. According to the gas release data (Figure 12b), which are a measure for the oxidation kinetics, a protective silica scale should have formed...
immediately after start of steam injection. The minimum of gas release rates at approximately 1600 °C, which is an indication of the beginning of volatilization from the oxide scale. From approximately 1750–1800 °C, in addition to the baseline, huge peaks of H₂, CO, and CO₂ releases are observed as large bubbles burst, as described above. Smoke (SiO-OH vapor) is also formed at temperatures above 1900 °C. The test was terminated at approximately 2000 °C by flooding a quartz glass cylinder with water from below. The sample was superficially degraded, but it survived these extreme conditions without breaking.

![Bubbles](image1)

![Si-O-OH smoke](image2)

(a) Video snapshots taken at temperatures between 1700 °C and 2000 °C

(b) Temperature and gas release during this ramp test

Figure 12. Results of a steam oxidation ramp test with a SiC cladding tube segment at temperatures between 1400 °C and 2000 °C [46].
4. Recent Findings in Advanced Test Facilities

4.1. Oxidation Kinetics of SiC in Steam at Temperatures Up to 1800 °C

With the development of LAHF, oxidation tests of monolithic SiC under a pressure of 1 bar, in a stream of 97% H₂O/Ar gas mixture, at a temperature range of 1400–1800 °C, were made possible and reported in the previous study [47]. Up to 1800 °C, the SiC samples underwent a mass loss process, which obeyed paralinear kinetics. Based on the mass change data, the parabolic oxidation rate, $k_p$ (see Figure 6), and the linear volatilization rate, $k_l$ (see Figure 5), were calculated. Comparison of parabolic oxidation rate obtained from this investigation is compared with the other work in Figure 6. The parabolic oxidation rate, $k_p$, obtained by using LAHF on monolithic SiC under 97% H₂O/Ar gas (1 bar), up to 1800 °C seemed to have the same activation energy (approximately 96 kJ/mol) with the one reported by Opila [45], which was conducted by using fused quartz tube furnace. In another study, in which steam oxidation tests of SiC was conducted in alumina tubes, Terrani [13] reported an activation energy of parabolic oxidation rate of 238 kJ/mol. As reported elsewhere [42,51], the use of alumina tube with water vapor substantially increased the parabolic oxidation rate $k_p$ of SiC due to the residual volatile sodium impurities. Because of the characteristics of LAHF facility (only the SiC sample was heated up to high temperature, the influence of the impurities from the environment was reduced. Therefore, the data of oxidation kinetics at high temperatures obtained from the LAHF facility best fit with the data reported by Opila [42,45], which were obtained in a high purity quartz tube. In summary, the study of SiC steam oxidation by using the LAHF is the first study that ranges up to 1800 °C. However, SiC oxidation kinetics above 1800 °C need further investigation as the SiC properties particularly viscosity and volatility are clearly changing the SiO₂ scale in this range.

4.2. Oxide Scale and Bubble Formation

The oxidation resistance of SiC at high temperatures relies on the formation of a protective SiO₂ scale. The influence of even small amounts (ppm levels) of impurities, such as alkali metals and aluminum, on oxidation kinetics of SiC has been reported in various studies [42,57,58]. These impurities may come from various sources, such as reaction tubes, residual additives/impurities from the tested sample, or the oxidizing atmosphere. The impurities will modify the SiO₂ melting point, but also its microstructure and properties, and enhance the diffusion of oxidizing species through the scale. However, from the elastic behavior observed in the LAHF and QUENCH-ATF experiments, the SiO₂ layer formed at high temperatures on the SiC is likely to be in a glassy (rather than a crystalline) form. The glassy form has SiO₂ tetrahedra chains with -O- bridging links, has a so-called glass transition point followed by a softening range. The chain length determines the transition point and softening range. In glass-making, small additions of alkali metal oxides are known reduce the chain lengths in silica melts and lower their viscosity [59]. Thus, amorphous/glassy phases formed between the impurities and silica may also be more permeable and volatile than the pure silica.

Bubble formation of SiC in steam has been reported in a number of studies [13,45–47]. Avincola [46] discussed three criteria, which have to be fulfilled for the formation of bubbles: (1) the formation of a dense, complete silica scale, (2) the presence of a higher gas pressure at the SiC/SiO₂ interface, and (3) sufficiently low viscosity of the silica scale. The impurities have been demonstrated to be able to significantly affect the viscosity of silica [60–62]. It means the less impurities the SiO₂ layer has, the higher temperature the bubble phenomenon occurs at and may explain the formation of bubbles over a wide range of temperatures with various bubble sizes.

Pressure may also influence the formation of SiO₂ bubbles. Currently, most of the oxidation studies for SiC at high temperature in steam were conducted at atmospheric pressure. High pressure environments in LWRs (60 bar before any depressurization phenomena) might suppress the formation of SiO₂ bubble at high temperatures. Nevertheless,
it remains necessary to conduct oxidation tests of SiC in high-pressure steam. Investigating the effect of controlled impurities on SiO$_2$ layer properties is a further aspect for testing.

4.3. Tests with Prototypical SiC-Based Cladding Tubes

More prototypical tests in the QUENCH-SR rig have confirmed the results obtained in the LAHF facility. The transient tests showed the same temperature ranges for stable and protective SiO$_2$ oxide scale formation, as well as for the development of bubbles and increasing volatilization of the silica scale in this temperature range. Isothermal tests confirmed the stability of such SiC-based cladding for a few hours up to 1700 °C. Longer term testing is also necessary to determine these limits. A rapidly enhanced oxidation kinetics is always observed when the superficial monolithic SiC environmental barrier layer (or seal-coat) failed, which gave access of the oxidizing atmosphere to the fiber-matrix composite [63]. The main reason for this dramatically increased oxidation kinetics, then, is the much higher accessible surface of the CMC system and the graphite-based fiber-matrix interface, which easily oxidizes. The temperature and time of CMC failure is mainly determined by the thickness and quality, respectively, of the external monolithic SiC layer. Most samples remained intact even during water quenching from temperatures of 1700 °C to 2000 °C.

5. Summary and Perspective

The steam oxidation of SiC at high temperatures and advanced facilities for accident tolerant fuel (ATF) cladding testing have been reviewed. In spite of many potential benefits, such as superior high-temperature properties and excellent irradiation resistance, there remain key issues, such as steam oxidation at extreme temperatures (1800–2000 °C), that needed to be addressed for LWR safety issues. Due to the lack of appropriate test facilities, study of oxidation behavior of SiC in steam at temperatures above 1600 °C was very challenging. Recent developments using remote heating and sensing (eddy current heating, laser heating, and pyrometry) have enabled oxidation studies to be carried out for SiC in steam up to 2000 °C. A summary of these developments and experiments are presented as follows:

- Development of non-contact heating and measurement techniques provide unique tools for studying of materials behavior in severe environments, which were not possible before. Further common features include localized heating of small sample masses allowing for high heating rates and avoiding contamination other materials of the facility, such as the support plate. All this occurs under fixed atmospheric conditions.
- Investigations conducted using the LAHF successfully yielded the SiC steam oxidation kinetics up to 1800 °C. The LAHF data at 1400–1800 °C are in good agreement with the data obtained at 1200–1400 °C by Opila. It indicated a similar SiC steam oxidation mechanism existed up to 1800 °C as was valid at 1400 °C. In other words, the excellent oxidation resistance of SiC in steam (producing relatively less heat and hydrogen gas) was maintained at temperatures up to 1800 °C. Investigation of SiC steam oxidation in conducted with the QUENCH-SR facility also confirmed the excellent performance of SiC, as prototypical SiC cladding samples remained intact even after quenching with water from 2000 °C.
- Future works: In spite of these recent efforts, further investigations of SiC steam oxidation in for a wider range of conditions should be carried out. For example, recent studies on steam oxidation of SiC were conducted in very clean environments. It is, therefore, necessary to conduct high temperature SiC steam oxidation tests under various impure environments, such as Cl$^{-}$, SO$_4^{2-}$, Ca$^{2+}$, or Na$^+$-containing steam (from sea water). Tests under high pressure conditions of SiC steam oxidation of at high temperatures or pressure cycles should also be conducted to reflect the pressurization / depressurization cycles during reactor accidents. Furthermore, this new data of high-temperature SiC steam oxidation kinetics needs to be tested to the highest available temperatures and to the longest times possible particularly for
operating fuel temperatures cycling in larger-scale experiments as planned in the QUENCH bundle facility at KIT. This is expected in the framework of the forthcoming OECD-NEA Joint Undertaking QUENCH-ATF [64].

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