Versatile Oxide Films Protect FeCrAl Alloys Under Normal Operation and Accident Conditions in Light Water Power Reactors

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The US has currently a fleet of 99 nuclear power light water reactors which generate approximately 20% of the electricity consumed in the country. Near 90% of the reactors are at least 30 years old. There are incentives to make the existing reactors safer by using accident tolerant fuels (ATF). Compared to the standard UO$_2$–zirconium-based system, ATF need to tolerate loss of active cooling in the core for a considerably longer time while maintaining or improving the fuel performance during normal operation conditions. Ferritic iron-chromium-aluminum (FeCrAl) alloys have been identified as an alternative to replace current zirconium alloys. They contain Fe (base) + 10–22 Cr + 4–6 Al and may contain smaller amounts of other elements such as molybdenum and traces of others. FeCrAl alloys offer outstanding resistance to attack by superheated steam by developing an alumina oxide on the surface in case of a loss of coolant accident like at Fukushima. FeCrAl alloys also perform well under normal operation conditions both in boiling water reactors and pressurized water reactors because they are protected by a thin oxide rich in chromium. Under normal operation condition, the key element is Cr and under accident conditions it is Al.

INTRODUCTION

Worldwide, the generation of electric power has several sources of energy that can be grouped as: (1) fossil fuels (coal, petroleum and natural gas), (2) nuclear and (3) renewable (wind, solar, hydroelectric, geothermal, biomass, etc.) sources. Figure 1 shows that the world energy consumption in the next two decades will be still dominated (~ 80%) by the burning of fossil fuels (liquid, gas and coal). Nuclear energy represents only 6% of the energy consumed worldwide. In the USA, 30% of the consumed natural gas is used to generate about 20% of electrical power produced in the country. On the other hand, 100% of the nuclear energy produced is used to generate electricity. This nuclear electricity also represents about 20% of all the electrical power produced in the country. The president of the Nuclear Energy Institute in the US said that one of her top priorities is to ensure the continuous “safe, reliable operation of the US commercial nuclear reactors” to avoid their premature retirement, which would be replaced by the burning of more natural gas, increasing the emissions of greenhouse gases. Currently, there are 99 operating power reactors in the US, 34 boiling water reactors (BWR) and 65 pressurized water reactors (PWR). Figure 2 shows in which year these power reactors were connected to the grid, affirming that 89% of the current reactors are at least 30 years old and 45% of the reactors are at least 40 years old. Only one new reactor was connected to the grid in the last 20 years. Currently, four new reactors are under construction in the USA. The existing nuclear power plants (NPP) in the USA are an aging infrastructure, which are investing a bare minimum in upgrades and maintenance. It is important to make these NPP safer in their operation for the remaining of their useful life.

After the Fukushima NPP black-out following the tsunami on March 2011, the international community has been dedicated to finding a fuel configuration that will be more resistant to a loss of coolant accident (LOCA) than the existing zirconium/UO$_2$ pellets. The massive building explosions at the
Fukushima plant were caused by the ignition of hydrogen gas formed by the fast-exothermic reaction of zirconium with water (or steam).

\[ \text{Zr} + 2\text{H}_2\text{O} = \text{ZrO}_2 + 2\text{H}_2 + \text{Heat} \] (1)

There is currently an international research effort to delay the generation of hydrogen gas and to expand the coping time allowed for quenching or cooling the reactor after a severe LOCA. There are several proposals on how to design an accident tolerant fuel (ATF) or an advanced technology fuel (ATF) that will permit for a safer operation of the existing light water reactors.\textsuperscript{5,6} The ATF proposals include the use of iron-chrome-aluminum (FeCrAl), silicon carbide (SiC), and coated zirconium or molybdenum (Mo) claddings. FeCrAl is the concept that would provide the fastest implementation for the safe operation of the remaining NPP for another 40 years.\textsuperscript{7} The proposed ATF configuration will use the current UO\textsubscript{2} fuel clad with a thin wall (less than 0.5 mm thick) FeCrAl tube. Eventually, other fuels such as uranium silicide may be industrially developed for the FeCrAl cladding.

**Oxidation Behavior of FeCrAl**

One of the requirements for ATF from the US Department of Energy is that the cladding must perform as well as or better than zirconium under normal operation conditions and much better than zirconium under severe accident conditions.\textsuperscript{8} That is, under normal operation conditions, the cladding must do well in high-purity water at 288°C for BWRs and at 330°C for PWRs. The water may contain dissolved hydrogen for corrosion control. Under accident conditions, the cladding must do well in superheated steam at 1200°C or higher temperatures. That is, under a LOCA situation, the cladding may not be able to oxidize through the wall for several hours, keeping its geometry intact for quenching operations with fresh water.

The composition of the FeCrAl alloys considered could cover the following range: Fe-based, plus 12–21 mass percent (% of Cr, plus 4–6% Al plus 2–3% Mo, with parts per million level of other elements such as yttrium, hafnium, zirconium, etc. Some FeCrAl alloys are made by powder metallurgy (e.g., APMT) and others by traditional melting, casting and forging (e.g., C26 M and Aluchrom 418 YH).
thicker dual layer oxide on the surface. As the immersion time progressed, the mass gain became smaller because of the formation of a protective inner layer of chromia. The external oxide may slowly dissolve in the oxygenated water. Under hydrogen conditions, the coupons lose initially some mass, but after a single layer chromia layer forms on the coupon surface, the mass loss is stabilized. Figure 4 shows comparatively the mass loss between the BWR (288°C) and PWR (330°C) hydrogenated environments. The mass loss was higher in the higher PWR temperature environment.

Figure 5 and Table II shows the characteristics of the oxides which form naturally on a FeCrAl APMT material exposed to simulated light water reactor environments. The chromium-rich oxides act as a barrier for the further dissolution of the material as shown by the low mass changes on the coupons (Figs. 3 and 4). The results presented here suggest that FeCrAl APMT has excellent environmental resistance characteristics under normal operation for both BWR and PWR coolants. There is no need to change the water chemistry of the coolants since FeCrAl is compatible with the current water chemistries in light water reactors.

FeCrAl Behavior Under LOCA and Severe Accident Conditions of NPP

The main reason FeCrAl alloys have been selected for ATF cladding is because they have a superior oxidation resistance in superheated steam in the event of LOCA or a severe accident. Chromium provides protection against oxidation in air or in steam to all stainless steels and FeCrAl alloys. As the temperature increases beyond 1000–1100°C, chromium does not effectively protect these alloys due to the evaporation of Cr₂O₃ in the environment. Beyond 1100°C, the alloys must contain approximately 4–6% Al to offer protection. The way the FeCrAl alloys work is by the initial formation of a Cr₂O₃ oxide on the surface. As the temperature increases, a continuous thin alumina layer (Al₂O₃) develops underneath the Cr₂O₃ film. Eventually,
Fig. 5. Composition of the oxide film formed on the surface of APMT exposed to 330°C and 288°C water with hydrogen and 288°C water with oxygen. Chromium oxide always develops.
the Cr₂O₃ oxide evaporates and the alumina layer protects the material up to its melting point of approximately 1500°C. The combined synergistic action between Cr and Al in FeCrAl at very high temperatures has been known for almost a century. Figure 6 shows that, when an FeCrAl APMT tube was exposed to steam at 1200°C for 4 h, it developed an approximately 1-μm-thick oxide which was pure alumina (it did not contain Cr, Fe or Mo). FeCrAl alloys such as APMT generally contain small amounts (less than 1%) of rare earth elements that may help peg the protective oxide to the alloy substrate. The FeCrAl producers such as Sandvik for APMT may sell the alloy in the pre-oxidized condition. Figure 6 shows that a pre-oxidized strip of APMT also contains an approximately 0.5-μm-thick alumina layer on the surface formed by exposure to air at 1050°C for 8 h. The oxide formed at 1050°C seems to contain residual amounts of Cr on its outer layer.

It was shown in this work that, when freshly fabricated specimens of FeCrAl alloys are exposed to ~300°C water, they develop a protective Cr₂O₃ oxide on the surface. Similarly, if these freshly fabricated specimens are exposed to >1000°C steam or air, they form a protective alumina on the surface. Figure 7 shows schematically the oxide development in both scenarios (normal operation or accident) described. A pre-oxidation treatment may be considered for FeCrAl tubes since the presence of an alumina layer will act as a barrier for tritium release into the coolant.

Two questions are repeatedly asked about the oxidation behavior of FeCrAl.

1. After the FeCrAl cladding is in high temperature water for several months, with a Cr oxide on its surface (Table II), would it be able to develop a protective alumina layer on the surface in the case of an accident?
2. If a pre-oxidation treatment is applied to the FeCrAl tube cladding, what happens to the alumina layer in the OD of the tube in contact with the water? Does it dissolve? Would a Cr oxide be able to develop on the surface (Table II) after the alumina dissolves?

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![Figure 6](image_url)

Fig. 6. Alumina oxide layer formed on FeCrAl alloys under exposure to temperatures higher than 1000°C.
|   | **Fe** | **Cr** | **Al** | **Mo** |
|---|---|---|---|---|
|   | ![Fe Image](image1) | ![Cr Image](image2) | ![Al Image](image3) | Not detected |
|   | ![Fe Image](image4) | ![Cr Image](image5) | ![Al Image](image6) | Not detected |

Fig. 6. continued
Question 1: Steam Oxidized APMT Tubes Exposed to High Temperature Water

Figure 8 shows the mass change (loss) as a function of immersion time for APMT tube specimens exposed to both BWR (288°C) and PWR (330°C) hydrogenated water. Each point is the average value for two or more specimens. The mass loss for the AR tube specimens was the same as for the AR flat specimens (Figs. 3 and 8). The tube specimens which were pre-oxidized (PO) in steam initially had a higher mass change (loss) than the tube specimens which were as-received (AR). However, once the alumina layer dissolved in the water, the mass loss of the PO specimens practically stopped, and the mass loss rate became the same as for the AR specimens. The surface area of each PO specimen tube in Fig. 8 was approximately 3.7 cm². When pre-oxidized in steam at 1200°C for 2 h, the tube specimens developed a 1-μm-thick alumina layer on the surface (Fig. 6). The volume of the alumina layer is \( 3.7 \text{ cm}^2 \times 0.0001 \text{ cm} = 0.00037 \text{ cm}^3 \) (volume of aluminum oxide). The mass of aluminum oxide is \( m = V \times \text{density aluminum} = 0.00037 \times 3.95 \text{ g/cm}^3 = 1.46 \text{ mg} \). Dividing the mass of 1.46 mg by the surface area of 3.7 cm², we get 0.4 mg/cm², which is exactly what the Fig. 8 plot shows as the mass loss of the PO tube specimens due to the dissolution of the alumina layer. This is a clear proof that the initial higher mass change is just the dissolution of the alumina, and then the dissolution stops because a \( \text{Cr}_2\text{O}_3 \) should develop on the surface.

Figure 9 shows the oxide composition formed on the surface of a APMT tube specimen after immersion in PWR hydrogenated water at 330°C for 284 days. These tube specimens were originally pre-oxidized in steam at 1200°C for 2 h, that is, they originally contained a 1-μm-thick alumina layer on the surface (Figs. 6 and 8). Figure 9 shows that the alumina layer from the surface of the pre-oxidized tube specimen dissolved in water and that a chromium-rich oxide developed, similarly as for a specimen which is as-prepared (not pre-oxidized) (Fig. 5).
Question 2: Water Oxidized APMT Tubes Exposed to Superheated Steam

Figure 10 shows the oxidation resistance for 4 h in superheated steam at 1200°C of the APMT tube specimens, both AR and pre-exposed to high-temperature water for 73 days. Figure 10 shows that specimens which were immersed in high-temperature water for 73 days had the same resistance to superheated steam as fresh AR specimens. That is, fuel rods clad with FeCrAl alloys in an operating plant will resist attack by superheated steam in the unlikely event of a LOCA. For both the AR tube specimens and for specimens pre-exposed to high-temperature water, the oxidation rate in superheated steam was in the order of 0.25–0.35 mg/cm².

Figure 11 shows schematically the evolution of the oxides on the surface of FeCrAl alloys from water to steam and steam to water. The alloy was designed with the versatility to develop a protective oxide in each situation.
DISCUSSION

Stainless steels (e.g., austenitic type 304 SS) contain at least 13% of chromium so they can offer passivity in most industrial applications. The long-lasting passivity is a slowing down of the corrosion rate of the steel by the formation of a thin protective and adherent layer of chromium oxide on the surface. The passivity provided by chromium oxide of the stainless steels and nickel-based alloys also applies to FeCrAl alloys in light water reactor environments at temperatures near 300°C.9 Nuclear power reactors have been successfully using stainless steels and nickel alloys with chromium in hot water for over 60 years. Chromium is the element that provides the resistance in hot water by the formation of a protective oxide film on the surface.6,9,10

Since 2012, it has been proposed to use ferritic FeCrAl alloys as an alternative for fuel cladding so they can provide resistance to a severe accident situation.11,12 Ferritic FeCrAl alloys have never been used in light water reactors before. The FeCrAl also contain chromium and it was expected and shown here and elsewhere that they will also develop a thin chromium oxide on the surface to protect against high-temperature water attack. FeCrAl alloys also have about 4–6% of aluminum which will “act” only in the case of a severe accident. If a plant never has an accident, the aluminum will just sit and wait. It does not participate in the passive film formed in high-temperature water. Aluminum may never be needed since, for the normal operation conditions, only the chromium is needed. However, we have also shown here that if an unlikely LOCA happens, the aluminum will form the protective alumina layer required to resist the attack by steam. Chromium can protect the cladding tubes against steam only until near 1100°C. At higher temperatures, aluminum offers the protection to the tubes until their melting point. The synergistic effect between Cr and Al in FeCrAl makes this material ideal for both situations, normal operation conditions and ready in the case of an accident condition.

![Fig. 10. Oxidation resistance of APMT tube specimens in superheated steam at 1200°C. The oxidation resistance is the same for as-prepared fresh tubes as for tubes that were exposed to water environments for up to 73 days. That is, in a case of an accident, a FeCrAl tube in the water will develop a protective alumina layer.](image)

![Fig. 11. Schematic representation of the oxides evolution on FeCrAl alloys under normal operation and accident conditions of a light water reactor.](image)
CONCLUSION

1. Iron-chrome-aluminum alloys (FeCrAl) contain both Cr and Al making it an ideal material to replace zirconium in light water reactors’ fuel claddings. Cr protects the alloy under normal operation conditions and Al protects the alloy at temperatures higher than 1100°C.

2. Under normal operation conditions, in a hydrogenated water environment, a thin layer of Cr oxide forms on the surface of the alloy slowing down the degradation process. Under oxidizing conditions, a dual layer may form. The inner layer is the protective layer of Cr oxide.

3. If the FeCrAl tube has an alumina layer on the surface, it will dissolve in high-temperature (~ 300°C) water and a protective Cr oxide will form in its place.

4. If a FeCrAl tube has a Cr oxide on the surface and it is exposed to accident steam conditions, the Cr oxide layer will evaporate and an alumina layer will form to protect the tube.

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REFERENCES

1. International Energy Outlook 2016. (US Energy Information Administration, 2016), https://www.eia.gov/todayinenergy/detail.php?id=26212. Retrieved 18 Aug 2017.
2. M. Korsnick, in press conference 20 Oct 2016 (Nuclear Energy Institute), www.nei.org/News-Media.
3. IAEA (International Atomic Energy Agency), PRIS (Power Reactor Information System), www.iaea.org.
4. J. Tollefson, Nature 537, 16 (2016).
5. C. Tang, M. Stueber, H.J. Seifert, and M. Steinbrueck, Corros. Rev. 35, 141 (2017).
6. F. Nagase, K. Sakamoto, and S. Yamashita, Corros. Rev. 35, 129 (2017).
7. R.B. Rebak, W.P. Gassmann, and K.A. Terrani, in Paper A0042, IAEA, Top Safe 2017 conference, 12–16 February 2017, Vienna, Austria.
8. S.M. Bragg-Sitton, M. Todosow, R. Montgomery, C.R. Staneck, R. Montgomery, and W.J. Carmack, Nucl. Technol. 195, 111 (2016).
9. K.A. Terrani, B.A. Pint, Y.-J. Kim, K.A. Unocic, Y. Yang, C.M. Silva, H.M. Meyer III, and R.B. Rebak, J. Nucl. Mater. 479, 36 (2016).
10. R.B. Rebak, M. Larsen, and Y.-J. Kim, Corros. Rev. 35, 177 (2017).
11. R.B. Rebak, Metall. Trans. E 2, 197 (2015).
12. K. Sakamoto, M. Hirai, S. Ukai, A. Kimura, A. Yamaji, K. Kusagaya, T. Kondo, and S. Yamashita, in WRFPreddit Conference (Jeju Island, September, 2017).