Article
High-Temperature Oxidation of Cr-Coated Resistance Upset Welds Made from E110 Alloy

Dmitri Sidelev *, Sergey Ruchkin and Egor Kashkarov

School of Nuclear Science and Engineering, National Research Tomsk Polytechnic University, 634050 Tomsk, Russia; ruchkin.2010.2010@mail.ru (S.R.); egor_kashkarov@mail.ru (E.K.)
* Correspondence: sidelevdv@tpu.ru; Tel.: +7-3822-70-17-77 (ext. 2518)

Abstract: The resistance upset welds (RUW) made from E110 alloy without and with Cr coatings were oxidized in air atmosphere at 1100 °C for 2, 10 and 30 min. The cross-section microstructure, elemental composition and hardness were studied before and after oxidation using optical and scanning electron microscopy, and indentations in welding region. The RUW welding does not noticeably change oxidation kinetics of E110 alloy. The most crucial effect has surface non-regularities formed after welding, which prevent uniform coating deposition on full surface of welded cladding tube and end plug. Cr coating deposition can strongly reduce oxidation of welded E110 alloy, while additional post-processing treatment should be applied to improve surface morphology after RUW welding. Several suggestions favorable to development of ATF Zr-based claddings using Cr coating deposition on welded nuclear rods were discussed.

Keywords: Zr welds; high-temperature oxidation; nuclear fuel claddings; zirconium-based alloys; magnetron sputtering; chromium coatings

1. Introduction

Currently, accident tolerant fuel (ATF) is being developed to improve safety of light or pressurized water reactors [1,2]. The short-term ATF strategy is aimed to improve safety of already produced Zr-based alloy claddings by deposition of protective coatings. A lot of deposition methods such as magnetron sputtering, cold spraying, laser cladding, arc evaporation and various coating materials (Cr, FeCrAl, CrN, etc.) are still being studied to develop ATF claddings resistant under conditions simulating normal and accidental operation of nuclear reactors [2–9]. Nevertheless, the last published data showed that Zr-based alloys can be protected by magnetron deposited Cr coating during high temperature (HT) oxidation for a long-term period [9]. Such an approach seems to be one of the best prospects for short-term development of ATF. The high potential of Cr coatings can be also confirmed by the participation of industrial/research institutes such as CEA (France), VNIINM (Russia), KIT (Germany) and others [7,9–11] in several full-scale tests of ATF Zr-based claddings with UO2 fuel in nuclear reactors.

A nuclear fuel rod consists of uranium-based fuel placed in Zr cladding tube (~Ø9 mm and wall thickness ~0.5 mm) with two end plugs. Welding is necessarily applied to join end plugs to a cladding tube at both sides to ensure leak tightness of fuel rods. Different welding methods such as electron-beam welding (EBW), laser beam welding (LBW), tungsten inert gas welding (TIG) and resistance upset-butt welding (RUW) can be used. Therefore, the special attention of ATF development should be devoted to estimate oxidation behavior of Zr-based alloy welds especially in the loss of coolant accident conditions (LOCA). Oxidation tests of EBW, LBW, TIG and RUW welds of Zr-based alloys were studied in several works [12–17]. Vandegrift et al. concluded the most protective behavior of EBW welds of Zr-1Nb alloy in comparison with TIG welds and/or made from Zircalloys (Zry-3 and Zry-4) at HT isothermal and transient conditions in both dry and
humid air [12]. Laser processing of Zr-based alloys usually reduces their corrosion resistance in autoclave tests [14,16,18]. Segregation of Nb, Fe or Sn at grain boundaries results in secondary phases precipitations such as Zr(Fe,Cr)2 which affect the oxidation kinetics of Zr-based alloy welds [17,19]. Additional post-processing can be applied to improve homogeneity of weld microstructure and distribution of secondary phases. The heat treatment at 577 °C for 5 h in vacuum resulted in good ductility and strength of LBW welds similar to bulk Zr-based alloys [20]. Corrosion resistance of EBW welds can be improved using electron beam thermocycling treatment coupled with annealing at 580 °C and mechanical treatment/surface etching [21]. Amorphization of LBW welds made from Zry-4 under low dose Kr+ irradiation was used to improve their corrosion resistance [22]. Using a distributed heat source had a positive effect on structure and corrosion resistance of EBW welds made of Zr-2.5%Nb alloy [23]. Despite special post-treatment processing, the welds made from Zr-based alloys demonstrate poor corrosion resistance. Coating deposition was applied to protect welds from Al-, Mg-based alloys and steels and considered only for relatively low temperatures [24–26]. Zirconia has a non-protective behavior during HT oxidation; therefore, additional approaches are developing to enhance corrosion resistance of welds for application in ATF nuclear fuel materials. Currently, there are no literature or experimental data concerning the oxidation behavior of RUW welds made from Zr-Nb alloys under high temperatures (800 °C and more). The behavior of coated Zr-Nb welds under HT oxidation has not yet been studied.

The aim of this study is to determine the role of Cr coating deposition on oxidation behavior of resistance upset welds made from E110 alloy. The RUW welds were selected as such type of welding is currently used in technological process of nuclear fuel production in Russia (Rosatom, PJSC (NCCP), Novosibirsk, Russia). Therefore, the samples of RUW welds without and with 10 µm thick Cr coatings were prepared and their oxidation behavior was analyzed.

2. Materials and Methods
2.1. Sample Preparation

The E110 (0.9–1.1 wt.% Nb, 0.06–0.1 wt.% O, Zr balance) alloy samples with RUW welds were prepared by PJSC (NCCP), their geometry and outer view are shown in Figure 1. Chromium coatings were deposited onto the welded samples by using the vacuum installation equipped with multi-cathode magnetron sputtering systems. The detailed description of the installation is presented in previous study [27]. Firstly, the samples were rinsed with acetone in an ultrasonic bath and dried by compressed air for 2 min. Then, the samples were fixed to the substrate-holder and placed in vacuum chamber, which was evacuated up to the base pressure of 2 × 10⁻³ Pa. The special substrate-holders were used to deposit Cr coating on outer surface of the samples (the inner surface remained uncoated). Prior to deposition, the samples were etched by Ar⁺ ions using an ion source with closed electron drift for 20 min. The following etching parameters were used: Ar pressure of 0.15 Pa, accelerated voltage of 2.5 kV and ion current of 40 mA. The multi-cathode magnetron sputtering system with three disk Cr (99.95%, Ø90 mm) targets was used for deposition. The temperature of the samples was measured and controlled using an Optris CT Laser 3MH1CF4 (Optris GmbH, Berlin, Germany) infrared pyrometer (spectral range—2.3 µm) during coating deposition. According to our previous results [28], thick and dense Cr coating can protect Zr-Nb alloy from oxidation at high temperatures during a long-term period. Similar results were also shown in the literature [7,9]. Thus, the deposition mode of the Cr coating was selected based on the previous studies of Cr coating deposition on flat samples made from Zr-Nb alloy and its oxidation performance. The deposition parameters are shown in Table 1. Two types of the coated samples were obtained. The first type of the samples had a Cr coating on full outer surface, while the second type—only on a cladding surface (end plug and RUW weld were uncoated). The appearance of the samples is shown in Figure 1.
Table 1. Deposition mode.

| Q, W/cm² | P, Pa | t, min | U_{sub}, V | j_{sub}, mA/cm² | T_{sub}, °C | h, µm |
|----------|-------|--------|------------|-----------------|------------|-------|
| 39.1     | 0.25  | 192    | -100       | 20              | <390       | 9.8 ± 0.4 |

Note: Q—target power density, P—operation pressure, t—deposition time, U_{sub}—substrate bias voltage, j_{sub}—ion current density on the substrate, T_{sub}—substrate temperature, h—coating thickness.

2.2. Oxidation Test

An atmospheric furnace ATS 3210 (Applied Test Systems Inc., Butler, PA, USA) was used to perform HT oxidation tests at 1100 °C for different isothermal periods. The furnace was pre-heated up to 500 °C and the samples were loaded in the chamber. The alumina tube holders were used. The samples were heated up to 1100 °C with the rate of ~21° C/min and then were oxidized for 2, 10 and 30 min. After HT oxidation, the chamber was opened and the samples were naturally cooled to room temperature (~22 °C/min). The appearance of the samples in the furnace before and after oxidation is shown in Figure 1. The samples were necessarily photographed before and after each oxidation test.

![Figure 1](image)

**Figure 1.** The photographs of the samples (a) before and (b) after oxidation at 1100 °C for 2 min: left sample—partially Cr-coated; central sample—uncoated; right sample—fully Cr-coated.

2.3. Sample Characterization

The mass gain of the samples was measured using an analytical balance machine Sartorius CP124S (Goettingen, Germany) with an accuracy of 10^{-4} g. The thickness of the as-deposited Cr coating and elemental composition of the samples after oxidation were analyzed using scanning electron microscopy (SEM) with energy-dispersive X-ray spectroscopy (EDS) detector (LEO EVO 50, Zeiss, Oberkochen, Germany). To control thickness of as-deposited Cr coatings and their microstructure, polished Si (110) substrate was also used in the deposition experiment. Figure 2 shows the cross-section of Cr coating deposited on Si substrate. The Cr coating had dense and columnar microstructure. The thickness of the coating was uniform and equal to 9.8 ± 0.4 µm.

![Figure 2](image)

**Figure 2.** Cross-section of Cr coating deposited on Si substrate.

The microstructure of the as-received and oxidized samples were studied by an optical microscopy using AXIOVERT 200MAT (Zeiss, Göttingen, Germany). Hardness was measured on the cross-section of the samples with a load of 0.5 N using Vickers micro-hardness tester Pruftechnik KB-30S (Hochdorf-Assenheim, Germany). The Vickers pyramid from diamond was used as the indenter. The hardness measurements were performed in a welding region before and after the oxidation tests. The distance between indentations was equal to ~100 µm (it was higher than 3d, where d—averaged length of diagonals). The dwell time was 15 s. In total, 36 indentations were done for each sample, and the distribution of hardness in a welding region was plotted.
3. Results

3.1. As-Received Samples

Figure 3 presents the SEM images of the cross-section of the as-received RUW weld made from E110 alloy.

There was no difference in microstructure of welding region and end plug or cladding tube. The microstructure of the weld region consists of non-equiaxed coarse grains with the average size of ~10–12 µm. The elemental analysis by EDS technique showed uniform distribution of Zr and Nb over welding region without any Nb precipitations. Similar microstructure behavior of Zr-Nb and Zr-Sn alloys were also found after EBW welding in [29]. Most likely that the temperature during welding process was enough for dissolution any precipitates in zirconium matrix.
3.2. Weight Gain Measurements

Table 2 shows the outer view of the samples after oxidation. White-colored film was grown on the surface of the uncoated sample after oxidation indicating the formation of zirconia scale. The coated sample had dark-green color that is usually observed for chromia scale. The sample with partially Cr coating had a sharp border of zirconia and chromia that corresponds to the initial position of uncoated and Cr-coated surfaces. Moreover, local oxidation (white-colored area) was found in a burr of RUW welds for the fully Cr-coated samples. These regions are additionally highlighted by red arrows in Table 2 for each sample (after 2, 10, and 30 min oxidation). The samples had similar outer view after 10 min oxidation, but swelling of all samples in bottom part (in region of cladding tube) was found. This effect was caused by oxidation of the inner uncoated area of the tube. The changes of sample geometry were observed in other studies [4], where an inner surface of the tubes was uncoated. The severe degradation of the uncoated sample was found after 30 min oxidation test. The failure of this sample in bottom part was due to its geometry where a thin wall of a cladding tube rapid oxidized on a three sides (outer, internal and face surfaces). At the same time, swelling of the Cr-coated samples in bottom became more pronounced.

| Title                | Uncoated | Partially Cr-coated | Fully Cr-coated |
|----------------------|----------|---------------------|-----------------|
| 2 min oxidation      | ![Image](image1) | ![Image](image2) | ![Image](image3) |
| 10 min oxidation     | ![Image](image4) | ![Image](image5) | ![Image](image6) |
| 30 min oxidation     | ![Image](image7) | ![Image](image8) | ![Image](image9) |

Due to complex geometry of the samples, mass gain after oxidation was calculated and compared only to the uncoated and fully Cr-coated samples to estimate protective properties of Cr coating (Figure 4) using the next equation:

$$\text{Mass gain} = \frac{m_{1100} - m_0}{m_0} \times 100\%$$

(1)

where $m_{1100}$—mass of the sample after oxidation at 1100 °C, $m_0$—initial mass of the sample.

Figure 4 shows higher mass gain of the uncoated samples than for the Cr-coated samples. Moreover, the mass gain increased with the change of oxidation time from 2 to 30 min. However, the Cr-coated sample had lower mass gain after 30 min oxidation (2.0%) than that after 10 min (2.3%). Such result can be caused by spalling of zirconia from the inner surface of this sample during oxidation test. In common, the obtained results present the decrease in oxidation rate of Cr-coated samples by ~2.5–5.0 times in comparison with the uncoated ones for the considered oxidation conditions. The significant decrease in mass gain of Cr-coated Zr-based alloys under HT oxidation was found in other studies: in 9 times in air at 1100 °C for 20 min [27], in 8–11 times in a steam at 1100 °C for 850 s [7], in 2–4 times in a steam at 1200 °C for 1500 s [9].
Figure 4. Mass gain of uncoated samples and with full Cr coating after oxidation at 1100 °C for 2, 10 and 30 min. (*) Spalling of zirconia from the inner surface of this sample.

3.3. Cross-Section Microstructure of the Samples

Figures 5 and 6 show cross-section microstructures of the uncoated and fully Cr-coated samples after oxidation at 1100 °C. The uncoated sample was predominantly oxidized on its outer surface, while lower oxidation was observed on the internal surface. The most pronounced and crucial oxidation was found in a weld burr region (up to 800 µm) and end plug (~1 mm). The thickness of oxide layers was less (50–500 µm) in other regions.

According to Figure 6, high oxidation was observed in a weld burr region of the uncoated samples. The thickness of oxide layers increased from 300 to 800 µm as the oxidation time was changed from 2 to 30 min. The high non-uniformity of the oxide layer formed on the uncoated sample can be additionally caused by “nitrogen effect” [30,31]. The latter strongly accelerates oxidation kinetics of Zr-based alloys due to ZrN→ZrO₂.
transformation accompanied by high volume expansion (~48%). The accelerated degradation of Zircalloy-4 and M5 alloys in air were also found in [31], where the significant role on oxidation rate and non-uniformity was belonged to re-oxidation process of ZrN to ZrO2. This was usually pronounced at temperatures higher than 800 °C.

In the case of fully Cr-coated samples, outer surface of the samples had protective chromia scale under the considered oxidation conditions excluding the region of weld burr. This effect was clearly seen in the appearance of the samples after oxidation tests (Table 2). For both types of the samples, the formation of different surface morphology in a weld burr region during welding should be taken into account, when performing the comparative analysis. Thus, it also can influence the oxidation behavior (see Figure 6b).

![Figure 6. Optical photographs of cross-section microstructures of the (a) uncoated and (b) Cr-coated samples in the region of RUW welds after oxidation at 1100 °C for 2, 10 and 30 min.](image)

The ZrN and ZrO2 grains were clearly seen in the uncoated regions of the samples at higher optical resolution (Figure 7). While the fully Cr-coated sample had the multi-layered structure of Cr2O3 oxide layer/residual Cr layer/Cr-Zr interdiffusion layer/E110 alloy which is typically observed in oxidized Cr-coated Zr-based alloys during protective scale (Figure 7c) [9,28]. The cross-section microstructure of the partial Cr-coated sample is presented in Figure 7b. The Cr coating was still adherent at the border of coated/uncoated regions of the sample even at 1100 °C up to 30 min. Moreover, oxide layer was predominantly grown in the depth, while less kinetics was observed in the side direction (underneath the Cr coating). It indicates high coating adhesion of Cr to Zr-based alloys, which is typical for magnetron sputtering technologies and in the case of using special pre-treatment procedures like ion cleaning, preliminary heating, etc. [32,33].
Figure 7. Optical images of cross-section microstructures of the (a) uncoated, (b) partially and (c) fully Cr-coated samples in the region of cladding tube after oxidation at 1100 °C for 30 min.

3.4. Elemental Composition of the Samples

Figure 8 shows elemental compositions over depth of the uncoated and Cr-coated samples after 30 min oxidation at 1100 °C. According to EDS line-scans, the uncoated sample was strongly oxidized. A lot of microcracks were found on its surface. The O and N signals were found at the depth of 500 µm and more in E110 alloy. At the same time, the penetration depth of oxygen and nitrogen was less ~5 µm in the case of the Cr-coated samples. It corresponds to one half of the thickness of the as-deposited Cr coatings. No sign of O or N were found in the E110 alloy, when the fully Cr-coated sample was studied. However, a thin (1.5–2.5 µm) diffusion layer of Cr and Zr was detected underneath the residual Cr layer.

Figure 8. EDS-line scans of the (a) uncoated and (b) Cr-coated samples after 30 min oxidation at 1100 °C. White dashed lines indicate the measurement position.

3.5. Hardness Distributions

Figure 9 shows hardness maps of the cross-section of the RUW welds made from E110 alloy before and after oxidation at 1100 °C for 2, 10 and 30 min. The hardness of the as-received sample in the welding region was uniform and equal to 185–215 HV, which corresponds to the value for bulk Zr-Nb alloys [34]. The increase in hardness up to ~300, 350 and 420 HV was found for the uncoated samples after 2, 10 and 30 min oxidation, respectively. The oxygen and nitrogen diffusion results in the increase in hardness of Zr-based alloy due to the formation of α-Zr(O) or α-Zr(N) phases under HT oxidation [35]. The hardness maps for Cr-coated welds demonstrate high protective properties of the coating. The average hardness values in the weld region are slightly decreased to ~145–190 HV after oxidation that can be caused by thermal treatment (grain growth).
4. Discussion

The oxidation behavior of the uncoated and Cr-coated RUW welds made from E110 alloy was studied. Oxidation tests were performed in air atmosphere at 1100 °C for 2, 10 and 30 min. The uncoated samples were strongly oxidized due to non-protective scale of zirconia at high temperatures. The presence of N in atmosphere accelerated oxidation rate of the samples and resulted in the non-uniform growth of oxide layer over the cross-section of the samples. Cracking and spallation of zirconia layers can cause breakaway oxidation of Zr-based alloys that was also found in other studies during high-temperature (800–1200 °C) oxidation, especially in nitrogen-content atmosphere [27,31,36]. Nevertheless, the comparison of oxide thickness in welding zone, end plug and cladding tube showed the predominantly oxidation in the region of weld burr.

The geometry of the samples can also have influence on oxidation rate in different regions of the uncoated samples. Zirconium oxidation is an exothermic reaction (ΔH = −584.5 kJ/mol at 1200 °C) [27] that can result in non-uniform heating of the samples during HT oxidation. Taking into account the equal (or similar) surface area of cladding tube and end plug, a thin-wall cladding tube can be heated up to higher temperatures than end plug having higher volume. This effect can also influence on oxidation kinetics of the investigated samples that was clearly observed in their appearance (Table 2).
The deposition of 10 µm-thick Cr coatings effectively prevented the oxidation of welded E110 alloy under the considered conditions. The formation of protective scale of chromia with the thickness of ~5 µm and Cr-Zr diffusion layer at the “Cr coating-E110 alloy” interface were found after the oxidation test at 1100 °C for 30 min. According to EDS line-scans and hardness maps (Figures 8 and 9), oxygen and nitrogen can diffuse to the welding region with the formation of hard and brittle solid solutions of α-Zr(O) and α-Zr(N). Therefore, the double increase (up to 420 HV) of hardness was found in the region of RUW weld for the uncoated samples after 30 min HT oxidation. The above-mentioned results showed enough coating thickness (10 µm) of Cr to protect the welded E110 alloy under 30 min oxidation at 1100 °C. Such or some higher coating thickness (10−15 µm) was also suggested as an optimum for steam oxidation at 1100−1200 °C by other authors [7,9]. Nevertheless, the Cr-Zr interdiffusion can be a problem of coated nuclear rods at HT oxidation conditions, so a coating thickness can be increased. The protective coating can be consumed by diffusion and dissolution of Cr in Zr-based alloys at high temperatures [10]. Moreover, the melting temperature of liquid eutectic Cr2Zr phase is in the range of 1305−1332 °C [9,37–39], so surface of Cr-coated cladding can melt that results in loss of protectiveness and cause oxidation of zirconium. Thus, a barrier sublayer (e.g., Mo, Ta, etc.) should be additionally applied for prevention of Cr-Zr interdiffusion at temperatures higher 1300 °C [39,40].

Despite high protectiveness of the Cr coatings presented in Section 3.3., the local oxidation can occur in the region of weld burr (Figure 10). Thus, the additional post-treatment of the RUW welds should be needed to decrease surface roughness of weld burr and uniform coating deposition. For this purpose, mechanical processing, chemical etching [21] or ion-plasma treatment [41] can be considered.

**Figure 10.** Oxidation of Cr-coated RUW weld in a weld burr region.

It should be noted that lower oxidation kinetics of E110 alloy in the border of uncoated and Cr-coated regions also like as lower oxide thickness in the defect region of weld burr in the case of the fully Cr-coated samples. The oxide layer predominantly formed into the depth of E110 alloy than in side direction (under the protective coating). However, the deposition of Cr coating onto full surface of the samples seems to be better than partially coating deposition (only on cladding tube). The obtained results can be used for development of coating technology for ATF Zr-based claddings.

5. Conclusions

The oxidation behavior of RUW-welded samples made from E110 alloy with and without Cr coatings was investigated. The comparative analysis of their cross-section microstructures, elemental distributions over a depth, hardness before and after oxidation in air at 1100 °C revealed the following conclusions.
1. The RUW welding of E110 alloy does not change oxidation kinetics at high temperatures. The surface non-regularities in the region of weld burr have the most pronounced effect on oxidation rate.

2. The Cr coatings can protect RUW-welded samples from oxidation at high temperatures. Partial coating deposition on Zr claddings, especially in the case of uncoated weld, leads to uneven oxidation and material degradation in the area of the end plug/Zr cladding tube. Therefore, the coating deposition on the full surface of nuclear fuel rods after welding of end plugs and cladding seems to be advantageous for development of ATF Zr-based claddings.

3. Additional post-treatment procedures like mechanical processing, chemical etching or ion-plasma treatment should be considered for surface smoothing of nuclear fuel rod after RUW welding.

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