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Article

# Steam Oxidation of Aluminide-Coated and Uncoated TP347HFG Stainless Steel under Atmospheric and Ultra-Supercritical Steam Conditions at 700 °C

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Abstract: The efficiency of ultra-supercritical (USC) steam power plants is limited by the materials properties, in particular, the steam oxidation resistance of the currently used steels at temperatures higher than 600 °C. Under these conditions, steam oxidation results in the development of thick oxide scales which spall and can accumulate in tube bends leading to blockage, overheating and premature creep rupture, as well as erosion of downstream components such as steam valves and turbine blades. Most published work related to oxidation testing is carried out at atmospheric pressure, with significantly less testing of austenitic steels in supercritical steam, and rarely including protective coatings. Indeed, the effect of high-pressure steam in the oxidation process is not quite understood at present. This paper covers a comparison of the behaviour of TP347HFG stainless steel at 700 °C under atmospheric pressure and 25 MPa, with and without slurry-applied diffusion aluminide coatings. The results show a very protective behaviour of the aluminide coatings, which develop a very thin Al-rich protective oxide, and no significant difference between the two environments. In contrast, the uncoated steel exhibited a different behaviour. Indeed, under atmospheric pressure after 3000 h, very thin scales, rich in Cr and not surpassing 5 to 10 µm in thickness, covered the samples along with some much thicker Fe-rich oxide nodules (up to 150 µm). However, under 25 MPa, a thick multilayer scale with a non-homogeneous thickness oscillating between 10 to 120  $\mu$ m was present. A microstructural investigation was undertaken on the oxidised uncoated and coated substrates. The results suggest that pressure increases the oxidation rate of the chromia former steels but that the oxidation mechanism remains the same. A mechanism is proposed, including early detachment of the outer growing scales under supercritical pressure.

Keywords: ultra-supercritical steam; coating; oxidation; aluminide; austenitic steel

## 1. Introduction

The efficiency of ultra-supercritical (USC) steam power plants is currently limited, in part, because of insufficient steam oxidation resistance at temperatures >600 °C. Indeed, increasing the operating temperature poses a challenge in the selection of materials without significantly increasing costs [1]. In these power plants, the materials are subjected to very severe environments including pressures of 25 MPa and higher. Steam oxidation takes place under these conditions, generating very thick oxide scales, which spall and accumulate in tube bends leading to blockage, overheating and premature creep rupture, as well as erosion of the main steam valves and turbine blades.



Supercritical water (SCW) is also key for other power generation processes such as SCW-cooled Generation IV nuclear reactors [2]. In this case the conditions are similar but within the 300–650 °C temperature range. Moreover, SCW is also being considered for heavy oil (asphaltene and bitumen) refining because of the increasing demand of this resource [3]. In the corresponding plants, temperatures can reach values higher than 600 °C and the materials are in general austenitic steels or Ni-base alloys, although Ti- and Zr base materials are also being considered.

In order to validate new alloys or coatings to be used in plants, expensive and uncontrolled conditions testing must be currently carried out in operating plants. More experimental data are needed as a better understanding of supercritical conditions effects as well as the possibility of establishing correlations between laboratory supercritical and atmospheric steam rigs would be very useful, since plant testing is not always available and high-pressure testing facilities are more expensive and challenging to operate.

Few studies have compared the behaviour of materials exposed to real operating conditions in power plants with that observed in the laboratory under atmospheric pressure steam and the results are dissimilar in some cases. For instance, Knödler and Strauß compared several ferritic/martensiticand Ni-based alloys tested at: (a) 630 and 725 °C under high velocity steam at 190 bar in a plant, and (b) at 650 and 750 °C in the lab under low velocity steam at atmospheric pressure [4]. They found that ferritic/martensitic steels yielded similar results under both conditions, indicating that the steam pressure and velocity did not influence the scale growth. On the other hand, Ni-base alloys showed considerable higher oxidation rates in the plant tests. According to the authors, this may be due to the higher pressure and/or the high frequency of thermal cycling which occurs in the plant. Other factors such as different steam compositions (pH, oxygen content, etc.,) might also play a role. In contrast, Holcomb found different behaviours when he studied several steels and Ni alloys at 267 bar of slowly flowing steam and 670 °C in a pilot test in the laboratory, and compared them with the observed behaviour under atmospheric pressure [5]. In this case, all studied alloys showed corrosion rates orders of magnitude higher as well as higher thickness scales after 293 h of exposure to steam under supercritical conditions. In particular, fine-grained austenitic steel TP304H showed a corrosion rate four orders of magnitude higher at 267 bar. Indeed, the oxide scale was significantly thicker with an outer Fe oxide layer and an inner Fe-Cr spinel, whereas at 1 bar very thin pure chromia formed according to the author.

Some other studies related to testing austenitic steels under ultra-supercritical steam can be found in the open literature at temperatures ranging from 600–800 °C, and in general, at higher pressure, higher oxidation rates are observed [6–11]. At lower temperatures (<700 °C), a thin protective Cr-rich oxide accompanied by the presence of deep and large oxide nodules was observed at relative short. At higher temperatures and/or longer exposure times, dual-layer oxides developed, with an outer layer containing Fe<sub>2</sub>O<sub>3</sub> and Fe<sub>3</sub>O<sub>4</sub>, while the inner layer was composed of (Fe,Cr)<sub>3</sub>O<sub>4</sub>.

Considerable efforts have been dedicated to the development of coatings for super-critical power plants [12–19] but little work has been published regarding the behaviour and coatings under high pressure steam. Coatings such as MCrAlY overlays (M = Ni, Fe) [20,21] ceramics such as Al<sub>2</sub>O<sub>3</sub> [22], and diffusion aluminides deposited by slurry application [23–25] or by pack cementation [26] have been studied over the past few years and in most cases these coatings displayed a substantial improvement in terms of high pressure steam oxidation resistance. For instance, specimens of P92, both uncoated and coated with a slurry aluminide were exposed to real operating conditions in a bypass of a steam power plant and the results were compared to those obtained in the laboratory under atmospheric steam [23]. The authors found that under 180 bar and at 640 °C uncoated P92 exhibited earlier oxide scale spallation, faster oxide scales growth and a higher degree of porosity than the samples exposed in the laboratory under atmospheric pressure at 650 °C. In contrast, the coatings were very protective after at least 21,700 h of exposure under both conditions, but evidence of faster coating degradation was found for the field-tested specimen, as a significantly thicker protective scale was present under these conditions. As the plant was subjected to many temperature cycles, it is possible that protective

oxide spallation was intensified, consuming Al to reform it and therefore, Al depletion at the surface may occur faster than under isothermal conditions. Alternatively, high pressure may also have an effect on the protective oxide growth.

When examining these results as a whole, in particular in the few cases on which testing under atmospheric and high pressures is compared, no total agreement is found. Nevertheless, a general tendency can be observed, in particular regarding austenitic steels that develop protective oxides under atmospheric steam with isolated thick un-protective oxide nodules. On the other hand, thick, continuous multi-layered oxides are observed under supercritical conditions even after relatively shorter exposures. Regarding coatings, mostly alumina formers have been examined under supercritical conditions and they remain stable and protective despite the fact that some differences have been detected as for instance by Boulesteix and coworkers [25] who observed different oxide scale morphologies, as whisker- and plate-like features were detected under flowing atmospheric steam and high stagnant steam pressure respectively. In any case, the dissimilar results obtained by the different authors could be probably attributed to different testing conditions such as linear velocities, water quality and compositions, etc.

In the present work, a study has been carried out to compare the behaviour of slurry aluminide coatings deposited on SS TP347HFG under supercritical (25 MPa) as well as atmospheric flowing steam at 700 °C. The uncoated material that is a chromia former, was also tested and the evolution of the corresponding oxides as well as that of the coating microstructure was studied up to 3000 h. The aim was to gain further understanding on the effects of steam pressure on the oxide morphologies and the oxidation rates on both Cr and Al protective oxide former materials. As a result of this investigation, a steam oxidation mechanism for austenitic steels is proposed.

### 2. Experimental Part

## 2.1. Materials

Sample coupons (20 mm  $\times$  10 mm  $\times$  3 mm) of TP347HFG were machined from the wall thickness of a cold finished seamless tube obtained from Sumitomo Metal Industries LTD (Tokyo, Japan). The actual and nominal compositions are shown in Table 1. The samples were ground with 1200 Emery paper prior to testing. Al powder (particle size 4–6  $\mu$ m) was obtained from Bende-Lutz Werker GmbH.

Chemical Composition (wt.%)		С	Si	Mn	Р	S	Cr	Ni	Nb + Ta	Fe
Test material		0.09	0.36	1.4	0.02	0.01	18.3	11.4	0.9	balance
Nominal	min	0.06	-	-	-	-	17.0	9.0	$8 \times C$	balance
	Max <sup>(1)</sup>	0.10	0.75	2.00	0.04	0.03	20.0	13.0	1.0	balance
(1) TP347HFG (VdTÜV WB 547).										

Table 1. Composition of TP347HFG stainless steel from Sumitomo Metal Industries LTD (Tokyo, Japan).

#### 2.2. Coating Deposition

The Al slurry is a Cr VI free, water-based proprietary composition paint, containing spherical Al particles (nominal average  $\emptyset$ : 5 µm), a binder constituted by a mixture of non-toxic inorganic compounds developed by INTA and water as solvent. The mixture was stirred by means of a magnetic bar prior to coating and the sample's surface was ground and solvent degreased. The slurry was applied at room temperature by spraying with a Sagola spray gun (spray distance: 250 mm; pressure: 2 bar), and the coated samples were left to dry under laboratory air and then subjected to a diffusion heat treatment performed under vacuum at 700 °C for 1 h followed by 5 h at 900 °C. Finally, slight cleaning with a Scotch Brite sponge was undertaken to remove undiffused slurry residues (bisque). More insight into the deposition procedure can be found elsewhere [27].

## 2.3. Atmospheric Pressure Steam Oxidation Testing

The schematics of the closed loop laboratory tubular rig employed at INTA are shown elsewhere [28]. Prior to testing, laboratory air is displaced from the chamber by means of  $N_2$  which is kept flowing at 150 L/min while heating up to the test temperature (approximately at 600 °C/h). Once the test temperature is achieved (700 °C), the  $N_2$  flow is closed and pure steam is introduced at a linear velocity of 8 cm/s. Water used to produce steam is constantly bubbled with  $N_2$  and the pH is maintained at 7. To carry out weight measurements or to remove samples for metallographic analysis, the furnace is cooled to ~300 °C under  $N_2$  and the specimens are subsequently removed. The reheat cycle is also carried out under  $N_2$ . At least three samples per exposed time where weighted.

#### 2.4. High Pressure Steam Oxidation Testing

The high-pressure steam test was carried out at VTT in a custom built recirculating supercritical water autoclave at 700 °C/25 MPa with 150 ppb dissolved oxygen at the inlet flow and a pH of 7, measured at room temperature in the low-pressure part of the recirculation loop. The test environment was the same during all heating and cooling cycles, except when the autoclave was opened for sample retrieval at room temperature. At least three samples per exposed time where weighted.

The heating rate at the start of testing sequences was 20 °C/min up to 500 °C and 7 °C/min above this level, and the cooling rate at the end of the testing sequence was 20 °C/min. The schematics as well as further details of the test rig can be found elsewhere [24]. The flow rate during the exposure was around 7 mL/min (of water) resulting in a refresh time of the autoclave volume roughly every 1 h. Specimens were hung on the specimen holder rack using electrically insulating ZrO<sub>2</sub> rings.

#### 2.5. Characterisation

The coatings were characterised by light microscopy (Leica DV 6) and field emission gun scanning electron microscopy (FESEM) employing a ThermoScientific APREO C-LV (ThermoFisher Scientific, Brno, Czech Republic) microscope equipped with an energy-dispersive X ray microanalysis system (EDX) from Aztec Oxford. Cross section observations were obtained by cutting the specimens with a diamond wheel ATM 92002401, mounting them in conductive resin Konductomet (Buehler, Lake Bluff, IL, USA) and polishing with silice coloidal MarterPolish2 (Buehler, Lake Bluff, IL, USA). The samples were not subjected to etching. Thicknesses were determined by a minimum of 20 measurements and their averages are reported in the text. Phase composition was examined by X-ray diffraction (XRD) in a Philips X'Pert equipment (Panalytical, Almello, Holland) using the Cu K $\alpha$  line 0.15418 nm.

## 3. Results

## 3.1. Initial Coating Microstructure

Slurries are suspensions of Al particles in a binder and are applied as any paint. After drying and curing, a layer of agglomerated Al particles is obtained and when this layer is subjected to appropriate heat treatment, Al melts and partially dissolves the substrate forming intermetallics, which solidify and later evolve depending on the heat treatment length and temperature. In the present case, a heat treatment consisting of 1 h at 700 °C followed by 5 h at 900 °C under vacuum was applied and a three-zone coating microstructure measuring ~100  $\mu$ m was obtained as shown in Figure 1. The element matrix shows a decreasing Al content from the surface towards the substrate and the presence of "islands" of Al richer phases as well as a Cr-rich phase within the most external layer (zone 1). According to the composition measured by EDS as well as the XRD pattern (Figure 2), the matrix in zone 1 (Figure 3) corresponds to Fe<sub>2</sub>Al<sub>5</sub> (Al: 50.9; Fe: 38.7 in wt.%) with some Cr and Ni (4.6 and 3.2 wt.%). The Al-poorer zones correspond to FeAl (Al: 33.0; Fe: 49.8 in wt.%) with Ni (6.2 wt.%) and Cr (8.3 wt.%), and the Cr-rich phase to Cr<sub>2</sub>Al<sub>13</sub> (Al: 64.8; Cr: 15.0 in wt.%). Some peaks matching NiAl could also be observed in the XRD pattern. In addition, submicron scale precipitates rich in Nb (~45 wt.%) can also be detected throughout the coating and can be attributed to Nb carbides and

carbonitrides which are present in alloy TP347HFG [29]. Indeed, low intensity peaks corresponding to Nb (C, N) could also be identified in the XRD pattern. The intermediate layer composition (zone 2 in Figure 1) corresponds again to FeAl (Al: 34.1; Fe: 49.8 in wt.%) with some Ni (6.8 wt.%) as well as dissolved Cr (9.4 wt.%) and at the interface with the third and last zone there is a Ni rich thin layer (Figure 1) with Al: 34.2; Fe: 30.4; Ni: 25.1 in wt.%. In the interdiffusion layer (zone 3 in Figure 1), the Al content is reduced to 15 wt.% near the interface with zone 2 and gradually decreases to 3 wt.% at the interface with the substrate. Likely, this phase is a solid solution of Ni, Cr and Al on Fe. Within this inner layer, some darker grey features can be found, which are richer in Al and Ni than the

surrounding areas (Figure 4). The same morphological features were observed by Yang et al. who recently produced an aluminide coating on Super 304 H alloy with a different heat treatment than ours and a different overall morphology [19]. However, within the interdiffusion zone (zone 3 in our case), they were able to identify these features (called "butterflies" in their case) as NiAl, by electron diffraction, in agreement with our results. Dark AlN precipitates are also present in this zone in our case, resulting from N present in the alloy, as it reacts with incoming Al. This zone is also rich in NbC precipitates as shown in the corresponding mapping in Figure 4.



**Figure 1.** FESEM cross section SE image and EDS element map of a slurry aluminide coating deposited on HT347HFG steel after being heat treated for 1 h at 700 °C and 2 h at 900 °C.



**Figure 2.** XRD pattern of a slurry aluminide coating deposited on HT347HFG steel after being heat treated for 1 h at 700 °C and 2 h at 900 °C.



**Figure 3.** FESEM cross section and element map of the external layer (zone 1 in Figure 1) of the slurry aluminide coating deposited on HT347HFG steel.



**Figure 4.** FESEM cross section of the most internal layer (zone 3 in Figure 1) of the slurry aluminide coating deposited on HT347HFG steel.

#### 3.2. Steam Oxidation Test Results

Figure 5 illustrates the mass variation of all tested samples at 700 °C. Under atmospheric steam pressure, no significant weight differences between the coated and uncoated specimens could be observed, except perhaps between the samples exposed for 3000 h, as the coated specimens exhibited lower weight gain. In contrast, the uncoated substrate experienced considerable weight loss after 300 h of exposure under supercritical conditions, and then the samples began to gain weight up to more than 1.5 mg/cm<sup>2</sup> after 3000 h. This was explained by evidence of substantial spallation of the 300 h specimens, as shown in Figure 6, where images of the specimens before and after 300 h of exposure to supercritical steam are presented. After longer exposure, the scale regrew but there is also evidence of a significantly smaller degree of spallation as observed on the sample exposed for 3000 h in Figure 6. A similar behaviour was observed by Tepylo and coworkers [26] and also by Chen and collaborators [11] who pointed out that this is not often observed weight loss for TP347H after 1500 h at 650 °C, whereas in the data showed by Tepylo, at 700 °C significant weight loss was observed for 304 SS after 1000 h.



Figure 5. Mass variation of all tested specimens as a function of time exposed to steam at 700 °C.



Figure 6. HT347HFG specimens before and after exposure to steam at 25 MPa at 700 °C.

Unfortunately, intermediate weights were not recorded for the coated specimens tested under supercritical conditions but after 3000 h the corresponding specimens gained less than 0.2 mg/cm<sup>2</sup>.

## 3.2.1. Uncoated TP347HFG

In Figure 7, the cross sections of the uncoated specimens exposed to steam both at atmospheric and supercritical pressures are shown for all the test intervals. Under atmospheric pressure, TP347HFG is mostly un-attacked by steam oxidation, but exhibits oxide nodules which appear to grow in width and thickness as the exposure time increases, reaching up to 150  $\mu$ m in some nodules after 3000 h. In contrast, under supercritical steam an irregular double layer scale develops and grows up to 120  $\mu$ m after 3000 h. The apparent lower thickness of the oxides developed under supercritical steam compared to that of the oxide nodules formed under atmospheric steam, may be caused by the already mentioned initial oxide spallation experienced under supercritical conditions. In fact, on the sample exposed for 300 h, only some residues of the outermost layer appear on some areas of the shown image.



**Figure 7.** Light microscope cross section of HT347HFG specimens exposed to steam at 700 °C as a function of time and pressure.

The XRD patterns of the samples tested for 3000 h under atmospheric and supercritical conditions are shown in Figure 8a,b respectively. Under atmospheric pressure, the pattern corresponds mostly to the substrate but there are low intensity peaks matching  $Fe_2O_3$  and  $(Fe,Cr)_3O_4$ , whereas under supercritical steam, no substrate peaks could be detected and high intensity peaks corresponding to  $Fe_2O_3$  and  $(Fe,Cr)_3O_4$  were present. Spinel  $Fe_3O_4$  could also be present, as the XRD peaks corresponding to this phase overlap with those of  $(Fe,Cr)_3O_4$ . This is coherent with the findings of several authors [10,11] and is also supported by the microscopy investigation carried out by light and FESEM microscopy in this work as shown below.



**Figure 8.** XRD patterns of HT347HFG steel exposed to steam at 700 °C under: (**a**) atmospheric pressure and (**b**) 25 MPa.

### 3.2.2. Microstructure of the Oxide Scales Developed under Atmospheric Steam

A  $\sim$ 5 µm oxide covers most of the surface of the samples exposed to atmospheric pressure, allowing the peaks of the substrate to be picked up by XRD. In Figure 9, a FESEM cross section of this sample is shown, exhibiting one of the nodules as well as the protective oxide which covers most of the sample surface. The thin protective oxide is rich in Cr, Fe and Mn and is fractured on most of the examined surface, likely because of the metallographic preparation, and therefore appears to be brittle. A higher magnification image of this  $4-5 \mu m$  oxide is shown in Figure 10 at a section on which it had not fractured. It appears to be a single layer, but when examining the EDS element map, different compositions are detected as a function of the layer depth, and a very thin Si-rich oxide is also present at the scale-substrate interface. It has been found that in the presence of enough Si, this very thin oxide forms and behaves as a diffusion barrier improving the oxidation resistance [30]. Indeed, Li and coworkers also observed this layer by TEM microscopy on SS310 exposed to atmospheric steam pressure at 625 °C [6]. Right on top of this layer, a thin Cr (23.7 in wt.%) and Mn (4.5 in wt.%) containing oxide measuring less than 1 µm is observed, and is in turn covered by an Fe-rich thicker oxide containing Mn (4.1 in wt.%) and very little Cr (3.5 in wt.%). Behnamian et al. [10] found a similar oxide microstructure when exposing 316 SS to supercritical steam at 25 MPa and 800 °C for only 12 h, but for TP347HFG only a single composition layer was found which they attributed to (Fe,Mn)Cr<sub>2</sub>O<sub>4</sub>. The elemental analysis of the external layer in our case is compatible with  $(Fe_{,}Mn)_{3}O_{4}$ , whereas the inner layer is likely (Cr,Mn)<sub>2</sub>O<sub>3</sub>. This was also proposed by Yang and coworkers to form on 304H after 1000 h under flowing steam at 650 °C [19], and by Li et al. when exposing SS310 to supercritical (29 MPa) steam at  $625 \,^{\circ}$ C [6]. The lower Cr content of the outer zone of the scale may be caused by Cr evaporation as proposed by other authors [8]. Under this scale a Cr depletion substrate zone was observed, with only 10.5 wt.% Cr as opposed to 18 wt.% in the bulk alloy.



Figure 9. FESEM cross section and EDS element maps of HT347HFG exposed to steam at 700 °C under atmospheric pressure after 3000 h.



**Figure 10.** FESEM cross section of the thin oxide formed on HT347HFG exposed to steam at 700  $^{\circ}$ C under atmospheric pressure after 3000 h.

The oxidation nodule shown in Figure 9 presents a complex oxide morphology with an external Fe-rich layer corresponding to Fe<sub>2</sub>O<sub>3</sub> according to EDS and XRD and an internal complex Cr-rich zone. It has already been accepted by many authors that this two-layered oxide results from Fe outwards diffusion forming the outer layer, and oxidising species inwards diffusion resulting in Cr-rich spinel formation. The external oxide exhibits a high degree of porosity near the interface with the internal oxide (see also Figure 7). In addition, this outer portion of the nodule appears to grow on top of a thin Cr-, Fe- and Ni-rich oxide which corresponds to the protective oxide which covers most of the sample surface. Underneath, an internal oxidation zone, characterised by intergranular and lath oxidation patterns is observed. In fact, the original substrate grains appear to be surrounded by a Cr- and Mn-rich oxide, and inside the grains, preferential lath boundary oxidation takes place, as exhibited in the EDS element map shown in Figure 11. This confirms that oxygen inwards diffusion is faster along the grain and lath boundaries. Indeed, within the shown grain (Figure 11), metallic un-oxidised zones can still be observed inside the grain. Lath preferential oxidation was also observed by Bischoff and coworkers on ferritic steels [31] and similar microstructures were observed by Li [8] Yang [19] and Tepylo [26] and coworkers on 304 H stainless steel and by Zhu et al. on TP347HFG stainless steel [32] after steam oxidation. In agreement with Chen and coworkers' findings [11], the formation of Cr-rich layers in this inner oxide leads to Cr depletion and indeed, on the areas surrounding the nodules, Cr depletion can be observed on the element map showed in Figure 9. According to EDS analysis, Cr content of 14.1 wt.% was measured, which is lower than that found in the bulk material (18 wt.%).



**Figure 11.** FESEM cross section of internal oxidation zoned formed on HT347HFG exposed to steam at 700 °C under atmospheric pressure after 3000 h.

The observed porosity between the inner and outer scales, may lead to oxide detachment as for instance in one of the nodules shown in Figure 7 on the specimen exposed to atmospheric steam for 300 h. The outer layer is formed by Fe outwards diffusion and the thin protective layer located at the interface between the outer and inner oxide may hinder Fe outwards diffusion and moreover, the inner spinel oxide appears to be depleted in Fe. For the generation of two layered oxides, Rouillard and coworkers proposed that Fe atoms' outward diffusion caused vacancies that coalesce forming pores at the interface with the inner growing oxide [33,34]. The stable Cr-rich oxides present on the inner scale may also slow further Fe diffusion from the substrate.

Regarding the causes of oxide nodule growth on this Nb containing steel, recent studies verified that the oxidation of Nb-rich phases, or dissolved Nb in the matrix could promote the failure of protective Cr-rich scales when the austenitic stainless steels are exposed to high temperatures. Nb oxidation and the generation of stresses leading to cracking and pitting corrosion [35] could cause oxidation through cracks formed in the protective oxide, resulting in the development of oxide nodules such as the one exhibited in Figure 9. For instance, Behnamian found pits related to Nb when exposing SS 347 H to supercritical steam at 25 MPa at 800 °C for only 12 h [10]. In addition, as mentioned earlier, in the present study the protective scale appears to be brittle and prone to cracking so temperature cycles could also result in provoking crack development due to mismatch between the coefficient of thermal expansion of the oxide and the substrate. An alternative explanation to the origin of oxide nodules can be found in the investigations carried out by Halvarsson and coworkers based on Cr evaporation on stainless steel 304 [36]. They suggested that the oxide formed on top of grain boundaries, but due to Cr species evaporation, surface areas away from the grain boundaries cannot maintain a Cr-rich protective scale for long exposures.

## 3.2.3. Oxide Scales Developed under Supercritical Steam

The cross section of the sample exposed for 3000 h to supercritical steam is shown in Figure 12. As mentioned, the thickness of this rather wavy layer is not uniform, suggesting that it develops as a result of the coalescence of nodules as observed in the samples exposed to atmospheric steam (see Figure 7). The top oxide contains mostly Fe with very low amounts of Mn and Ni as well, but no Cr. The average Fe content in the oxide is of 64.5 wt.%. The XRD pattern shown in Figure 8b is indicative of Fe<sub>2</sub>O<sub>3</sub>, although Fe<sub>3</sub>O<sub>4</sub> could likewise be present as the corresponding peaks overlap with (Fe, Cr)<sub>2</sub>O<sub>3</sub> also present in the XRD pattern. In addition, different colorations can be observed within this top oxide in Figures 7 and 12. The inner layer is rather complex as in the case of the nodules formed under atmospheric pressure, and exhibits a lower Fe content which varies from 12 to 34 wt.%. It also contains

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very rich Cr zones as for instance at the interface with the substrate, as shown in the corresponding element map in Figure 12. At the bottom of the oxide, a grain surrounded by a Cr-rich layer can be observed in analogy with the microstructure seen in the nodule formed under atmospheric steam (Figure 9), which confirms that the oxidation process initiates at the grain boundaries. Surrounding the nodule and the grains, Cr depletion zones can also be observed with for instance a Cr content of 11.1 wt.% at the bottom of the nodule in the substrate zone immediately adjacent. The Cr-rich thin layer found at the interface between the outer and inner scales in the sample exposed to atmospheric pressure after 3000 h, was not found in the case of the specimen exposed to supercritical steam as it probably delaminates along with the outer growing Fe<sub>2</sub>O<sub>3</sub> scale.



**Figure 12.** FESEM cross section and EDS element maps of HT347HFG exposed to steam at 700 °C 25 MPa after 3000 h.

As mentioned earlier, delamination of the oxide formed under supercritical conditions takes place mostly at the initial stages of the process and later, regrowth of the scale supersedes the weight loss due to spallation. Tepylo and collaborators also observed oxide spallation on 304 stainless steel at but they only reported the weight loss after 1000 h so there is no indication on when the spallation took place [26]. On the other hand, Chen reported weight loss only at 650 °C (not at lower temperatures) and beginning after 1500 h [11]. The optical micrographs shown in Figure 7 indicate that it is the outer layer (Fe<sub>2</sub>O<sub>3</sub>) that mostly spalls at the beginning, and then it reforms and spallation continues with exposure time but very likely at a lesser rate.

## 3.2.4. Slurry-Aluminised TP347HFG

As shown in Figure 5, the aluminised specimens gained very little weight after 3000 h at 700 °C under both atmospheric and supercritical steam. These results contrast with those obtained by Tepylo

and coworkers [26] who observed slight weight loss when exposing pack cementation-deposited aluminide coatings on SS 304 to supercritical steam at 700 °C. The authors however, indicated that the coatings exhibited defects that led to substrate oxidation.

Figure 13 shows the coating microstructure evolution as a function of time for the specimens exposed to atmospheric steam. No degradation by oxidation is apparent and the main significant changes are the transformation of the outer layer (zone 1) and the increase in thickness of the interdiffusion zone (zone 3), no doubt caused by coating-substrate interdiffusion. The Al content at the surface diminished from an initial 50.9 wt.% corresponding to the Fe<sub>2</sub>Al<sub>5</sub> intermetallic phase to 25 wt.% in the sample exposed for 3000 h indicating that the Fe<sub>2</sub>Al<sub>5</sub> has transformed into FeAl. Since there is no weight loss nor evidences of protective scale spallation, it can be concluded that the transformation into to a lower Al containing intermetallic phase is caused by Al inwards diffusion, as further supported by the increase in thickness of zone 3 (interdiffusion zone). The presence of Kirkendall porosity between zones 2 and 3 was also observed, as a result of relatively faster outwards diffusion of the elements of the substrate than Al inwards diffusion.



**Figure 13.** FESEM cross sections of aluminide coated HT347HFG exposed to atmospheric pressure steam at 700 °C as a function of time (hours).

Some cracks perpendicular to the samples' surface that are typical of Fe aluminide coatings also appeared. Fe-Al intermetallics tend to be brittle and the thermal cycles associated with the test interruptions to extract samples are likely behind this. Slurry aluminide coatings deposited on ferritic steels develop cracks after the initial diffusion heat treatment. It has been shown that these cracks do not propagate into the substrate, nor allow steam to reach the substrate even after more than 70,000 h of exposure to steam under cyclic conditions at 650 °C as they self-heal by developing protective  $Al_2O_3$  on their surfaces [17].

Figure 14 shows the cross sections of the samples exposed to both atmospheric and supercritical conditions for 3000 h at 700 °C. No substantial difference between the two can be appreciated regarding both microstructure and composition. Both samples exhibit a two-zone coating with similar thickness and the surface Al content of both samples is ~25–28 wt.% and it is kept constant as a function of depth, up to the interface with the interdiffusion layer (zone 3).



**Figure 14.** XRD pattern of slurry-aluminised HT347HFG steel exposed to steam at 700 °C for 3000 h under: (**a**) atmospheric pressure and (**b**) 25 MPa including a higher magnification of the low intensity peaks.

The XRD patterns of these two samples (Figure 15) display the FeAl phase, confirming the full transformation of the initial Fe<sub>2</sub>Al<sub>5</sub> phase present in zone 1 (see Figure 1) to FeAl. Again, although some Al could have been "spent" in building a protective Al<sub>2</sub>O<sub>3</sub> scale, most is probably lost from the surface due to interdiffusion between Al from the coating moving inwards and Fe from the substrate moving outwards. This explains the growth in thickness of the interdiffusion zone that has gone from an initial ~33 µm to ~70 µm after 3000 h. The XRD pattern of the sample exposed to supercritical pressure exhibits very low intensity peaks corresponding to  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>. An Al-rich oxide scale with approximately the same thickness (~2–5 µm) is present on the samples exposed to both conditions (Figure 15a,b), although XRD peaks corresponding to  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> could not be detected on the sample exposed to atmospheric steam. The element mapping of the oxide formed on the samples exposed to atmospheric steam is shown in Figure 16 confirming that the oxide is Al<sub>2</sub>O<sub>3</sub>. The same element distribution is observed on the mapping of the corresponding supercritical steam sample (not shown). In earlier work carried out in our laboratory [37], we found that under atmospheric steam at 650 °C,  $\chi$ -Al<sub>2</sub>O<sub>3</sub> forms on slurry aluminide coatings at the initial stages but that it slowly transforms into  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>.



**Figure 15.** XRD pattern of slurry-aluminised HT347HFG steel exposed to steam at 700 °C for 3000 h under: (**a**) atmospheric pressure and (**b**) 25 MPa.



**Figure 16.** FESEM cross sections of the scale found at the surface of aluminide-coated HT347HFG exposed to steam at 700 °C, at atmospheric pressure and at 25 MPa.

Figure 17 shows a high magnification FESEM image of the interdiffusion zone in the sample exposed to atmospheric steam for 3000 h. The same features than those observed in the coating prior to exposure are observed, but after 3000 h the NiAl precipitates appear more elongated, and aligned perpendicularly to the substrate which may be explained for a preferential growth as the Al diffuses inwards.



**Figure 17.** FESEM cross section of the interdiffusion layer (zone 3) of the slurry aluminide coating deposited on HT347HFG steel after 3000 h of exposure to atmospheric steam at 700 °C.

## 4. Discussion

The results of the present work confirm those obtained by most other authors, in that the steam oxidation rate exhibited by austenitic steels is higher under supercritical conditions than under atmospheric pressure steam. Some different causes of this steam pressure-related increase in the oxidation rate have been proposed. These include: (a) increased solid state diffusion within the scales,

(b) increased hydrogen content in the alloy, (c) a higher content of oxidising species that can absorb on the materials surface increasing the partial oxygen pressure at the interface of the oxide scale and steam and (d) higher rate of Cr evaporation [5,38–40].

Observation of the different type of scales and also their evolution with exposure time leads to propose the following oxidation mechanism (Figure 18):

- (1) Initially, under both atmospheric and supercritical conditions, a thin Cr-rich oxide develops the surface of TP347HFG. Such an oxide was the only one observed by Behnamian and workers when exposing several austenitic steels to supercritical steam at 25 MPa and 800 °C [10] after only 12 h of exposure. Cr, Mn and Fe containing oxides with a thickness of 1 μm were observed along with some nodules as in the samples exposed to low atmospheric pressure in our case. Li and collaborators also found a similar Cr-, Mn- and Fe-rich oxide when exposing SS 310 to 29 MPa at 625 °C for 1000 h [6].
- (2) Thick unprotective nodules develop because of either the presence of Nb-rich carbides in the surface and/or to localised Cr evaporation, and grow both in thickness and sideways at faster oxidation rate under high pressure, as observed by many other authors. Cracks developed within the protective layer could also lead to nodule formation as the Cr-depleted area under scale may not provide sufficient Cr to reform the protective scales. Once the nodules begin to form, the outer oxide ( $Fe_2O_3$ ) grows in a radial direction form the defect originating the nodule, on top of the thin protective scale.
- (3) At high pressure, the oxide grows at faster rates as mentioned, and massive spallation occurs at earlier stages than under atmospheric pressure. Spallation is possibly caused by pore coalescence at the interface between the outer growing hematite layer and the inner growing Fe, Cr spinel. Pore development could be caused by the presence of the thin protective oxide which may act as a barrier to Fe outwards diffusion. Stresses generated by radial growth of hematite on top of the original Cr-rich protective thin oxide could also result in the outer oxide detachment. When the top oxide spalls it may carry the Cr-rich layer placed directly underneath.
- (4) After spallation, new oxide will grow on top of the inner spinel scale by Fe outward diffusion but at slower rates because of the somewhat protective role of the inner Cr-rich spinel and therefore, overall scale growth will not be as fast as during the initial oxidation stages.
- (5) Under higher steam pressure, the growth of these nodules appears to be significantly faster so they end up coalescing after only 300 h.



Figure 18. Proposed mechanism for steam for steam oxidation of HT347HFG at 700 °C.

In contrast, the slurry aluminide coating appears to behave in the same manner under both atmospheric and supercritical steam, as it develops a protective  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> scale which does not seem to be influenced by pressure up to at least 3000 h. The principal degradation mechanism in this case could be coating-substrate interdiffusion as the concentration of Al at the surface diminishes as function of exposure time at 700 °C. However, up to 3000 h the content of Al is high enough to sustain and maintain the scale, and since diffusion follows a parabolic law, the time required to reach a content of Al lower than the critical amount needed to sustain and reform the scale is expected to be high. Moreover, the relatively high content of Cr exhibited by these coatings may contribute to reduce the Al critical content. The appearance of cracks in the coatings both exposed to atmospheric and supercritical conditions, was not surprising given the brittleness of the Fe aluminide intermetallics. However, these cracks do not propagate into the substrate, at least within the tested timeframe.

In an early study, aluminide coatings deposited on ferritic steel P92 [23], a thicker protective oxide was developed on exposure to 180 bar in the bypass of an operating turbine for 21,700 h at 640 °C, when compared to a similar exposure in the laboratory. Two possible reasons were suggested to explain this behaviour: (1) a higher degree of spallation related to relatively more frequent thermal cycles in the plant and (2) a faster growth and/or spallation/reforming cycles of the protective Al oxide scale caused by higher steam pressures. The as deposited coating had cracks perpendicular to the surface, and in the sample exposed to high pressure steam, the cracks widened to a larger extent but again did not cause substrate oxidation. In contrast, Boulesteix and collaborators found different scale morphologies when exposing aluminide coatings to steam, as under atmospheric pressure, whiskers where observed on the surface of an aluminide coating whereas at 300 bar of stagnant steam there were platelets and clusters after 5000 h [25]. Although not shown, in the present work we did not observe differences on the cluster-like surface morphology of the coating exposed to both conditions. When comparing these results with those obtained in the present work, we found no evident differences in the behaviour of the coating as a function of pressure. This could be due to a shorter exposure in our case, but also to the fact that our experiment was carried out under flowing steam (Boulesteix high pressure experiment was carried out under stagnant steam [25]) and with significantly less cycles as those experienced in the plant bypass experiment.

## 5. Conclusions

Uncoated and aluminide-coated TP347HFG were exposed to steam at 700 °C under atmospheric and supercritical conditions up 3000 h. Under supercritical conditions the oxidation rate was higher for the uncoated steel and early massive oxide spallation took place. The results indicate that both under atmospheric and high-pressure steam, a thin protective Cr-rich oxide develops initially on the surface of the steel but oxidation nodules begin to form and grow both in depth and sideways until nodule coalescence takes place. The nodules may form due to: (a) Protective oxide cracking due to thermal cycles; (b) the presence of Nb-rich precipitates at the surface as they oxidise generating stresses and causing pit formation; and (c) localised Cr depletion away from grain boundaries. After 3000 h under atmospheric pressure, the protective oxide is still partially present but under supercritical conditions, at exposure times lower than 300 h the nodules have fully coalesced. The nodules exhibit an outer growing  $Fe_2O_3$  scale on top of the thin Cr-rich oxide which in turn covers an inner growing Fe, Cr, Mn spinel. This inner oxidation process begins at the grain boundaries where Cr-rich oxide is formed. We propose that early massive spallation occurs under supercritical steam due to the higher oxidation rate and the presence of the thin protective scale under the outer growing  $Fe_2O_3$  scale which contributes to generate porosity and/or stresses. Under atmospheric steam the same process occurs but at a much lower rate as after 3000 h, extensive areas of the steel surface were still covered by the protective scale as already mentioned.

In contrast, slurry-deposited aluminide coatings protect TP347HFG from oxidation under both conditions by developing a thin protective  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> scale. In this case, no differences in behaviour were perceived under the two examined conditions, indicating that up to 3000 h,  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> is very stable and

grows slowly, independently of the steam pressure. The main coating degradation mechanisms are temperature-driven interdiffusion with the substrate, the development of Kirkendall porosity at the interface with the interdiffusion zone and the development of thickness through cracks that stop at the interface with the interdiffusion zone. None of these three degradation modes led to substrate attack up to the experiment exposure timeframe (3000 h) at 700 °C.

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