Simulated fireside corrosion of T91 in oxy-combustion systems with an emphasis on coal/biomass environments

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ABSTRACT

Oxy-combustion is the burning of a fuel in oxygen rather than air for the ease of capture of CO₂ for reuse or sequestration. Corrosion issues associated with the change in heat exchanger tube operating environment (replacement of most of the N₂ with CO₂ and potentially higher SOₓ levels) from air- to oxy-combustion were examined. The ferritic-martensitic alloy T91 was used in accelerated fireside corrosion tests using several different gas compositions and ash deposit overcoats to simulate air-fired, oxy-fired coal, and oxy-fired co-fired coal/biomass conditions. Initial corrosion was observed after 240 h of exposure by examining cross-sections with retained ash. Metal section losses were determined after exposures of up to 1440 h at 600–700°C. Severe corrosion was observed, and a corrosion response with respect to ash deposit chemistry was observed. Corrosion response differences with respect to gas phase chemistry were minimal. Alloy-oxide scale-ash morphologies were consistent with oxide fluxing mechanisms.

INTRODUCTION

Research into improved materials systems and associated manufacturing and reliability issues is a major part of initiatives to economically reduce greenhouse gas emissions (primarily CO₂). The UK–US Memorandum of Understanding on Energy R&D between the United Kingdom Department of Climate Change (now Department of Business, Energy and Industrial Strategy) and the United States Department of Energy has provided the opportunity for collaboration between UK and US organisations in advanced materials for power generation [1]. Under this agreement, the organisations involved collaborate and share the results of Government or self-funded projects to maximise the return and benefit for the participating organisations and more rapidly advance underlying mechanistic understanding and materials development. Using this framework, a task addressing boiler fireside corrosion under oxy-fuel conditions was initiated. The focus was on the effects of deposit and gas-phase chemistry on the potential corrosion mechanisms.

Oxy-combustion-based power generation combusts coal in relatively pure oxygen (O₂) and re-cycled flue gases to enable the capture of carbon dioxide (CO₂) from the flue gas stream [2]. A typical oxy-combustion process consists of an air separation unit (ASU) that produces nearly pure O₂ (≥95 vol %) to combust coal in an oxy-combustor, with the combusted gas stream exchanging heat with a steam cycle to generate electric power. The oxy-combustion process performance is limited by (a) the efficiency of the ASU, which can impose a significant parasitic load, and (b) the recycling of flue gas required to limit temperatures within the oxy-combustor and maintain the mass flow of the combusted gas stream (especially in systems intended for retrofitting).

Most schemes to limit temperatures within the oxy-combustor involve recirculating a fraction of the CO₂-rich flue gas back into the oxy-combustor. If this is done with a gas stream taken before flue gas desulphurisation (FGD), then the oxy-combustor must operate with much higher levels of sulphur oxides (SOₓ), which increase corrosion concerns. This is termed hot-gas-recycle (HGR). If recirculation is done after FGD, then there will be a higher efficiency loss, but SOₓ levels will be relatively low and similar to air-combustion. The recirculation path after FGD could go directly back into the oxy-combustor or go through a water removal step to reduce its water content. These recirculation paths are shown schematically in Figure 1 as recirculation paths A, B, and C. There are three major areas of corrosion concern within the boiler: water wall corrosion, superheater (SH)/reheater (RH) corrosion, and dew point corrosion. The research presented here targets SH/RH conditions.

The primary fireside corrosion mechanism of SH/RH boiler tubing relies on the formation of liquid sulphates that result from reactions between deposited ash...
(containing relatively high melting point (Na,K)$_2$SO$_4$) and SO$_2$ with iron oxides either from the alloy scale or from the deposited ash [3]. The lowest melting point sulphates within these types of deposits are alkali iron trisulphates – although the more general description of a (Na,K)$_2$SO$_4$-Fe$_2$(SO$_4$)$_3$ solution being the corrosive liquid deposit is likely to be more accurate [3]. Rapid corrosion occurs after the molten phase contacts the alloy, either through flaws in the oxide scale or after the oxide scale dissolves ('fluxes') into the molten phase. The fluxing reactions can be either acidic or basic, depending on the local chemistry of the liquid phase and oxide scale [4]. This is analogous to type II hot corrosion of turbine blades, where rapid corrosion occurs after an incubation period and is characterised by an uneven and pitted surface. High Cr content alloys, up to 22 wt %, are beneficial in reducing fireside boiler corrosion [5].

The reported effects of oxy-combustion on fireside corrosion have been mixed for Fe9Cr ferritic-martensitic steels [5–22]. Hjörnhede et al. [6] examined oxyfuel pilot plant exposures of Fe9Cr steels at Vattenfall and found no significant differences between oxy-fired and air-fired ash deposits, corrosion rates, or carburisation. This was not expected as SO$_2$ levels were a factor of 4 greater for oxy-combustion than for air-combustion. Chandra et al. [7] found that 9–12 Cr ferritic-martensitic steels in oxy-combustion environments (CO$_2$, 27% N$_2$, 2% O$_2$, 1% SO$_2$) were susceptible to carburisation. Stein-Brzozowska et al. [8,9] found fireside corrosion for Fe9Cr steels in oxy-combustion environments to increase with increasing SO$_2$ and increasing temperature (to 650°C). Robertson et al. [5] found corrosion rates for T91 to be the same, or somewhat higher, in air-combustion than in oxy-combustion (for the same SO$_2$ content). They concluded that conventional materials used in subcritical and supercritical air-combustion boilers would be suitable for oxy-combustion with similar SO$_2$ levels. It should be noted that observing carburisation generally requires etching, and not all studies performed this step.

Holcomb et al. [10] found for short times (100 h), with a deposit of K$_2$SO$_4$, Na$_2$SO$_4$, and Fe$_3$O$_5$ in a 1.5:1.5:1.0 molar ratio, that there was an apparent threshold in SO$_2$(g) levels (between 10$^{-3}$ and 10$^{-5}$ atm), below which corrosion was much reduced. Section loss measured after 240 h for T91, with a deposit of 40% Al$_2$O$_3$, 30% SiO$_2$, 30% Fe$_2$O$_3$, 5% Na$_2$SO$_4$ and 5% K$_2$SO$_4$ (mol%), showed more metal loss for oxy-combustion gases with high H$_2$O contents (for both low and high SO$_2$ contents) than for air-combustion or low H$_2$O/low SO$_2$ oxy-combustion gases. The present work presented here includes an update of this work, and these longer exposure times will be shown to change the metal loss response with gas composition.

Combining co-firing of fuels and oxy-firing systems is an approach to develop efficient power plants while reducing CO$_2$ emissions [11–14]. Biomass is classed as carbon neutral fuel, and so has zero net CO$_2$ emissions, and the use of oxy-firing technologies for fuel combustion facilitates the post-combustion capture of CO$_2$ [15–17]. It is believed that co-firing biomass such as cereal co-product (CCP) with a UK coal will release elevated levels of Cl and alkali metal (K), with a possible formation of KCl [16,18] on the heat exchanger surfaces. At present time alkali salts have been in focus extensively as well as chlorine-induced high-temperature corrosion caused by alkali chlorides.

Figure 1. Schematic of an oxy-combustion coal boiler showing major areas of corrosion concern and three different possible flue gas recirculation paths.
Much attention to the subject is given in the literature [21,22]. Similarly, oxy-firing of such fuel blends will result in elevated levels of CO$_2$, H$_2$O and corrosive gases (HCl and SO$_2$) [14]. Both approaches (co-firing and oxy-firing) will change the combustion gas chemistry considerably and raise concerns for the life of the SH/RH tubes. Therefore, retrofitting existing power plants into co-firing units or oxy-firing without sufficient experimental data carries substantial risk.

In this paper, degradation of T91 (~9% Cr) in simulated oxy-firing conditions is presented. Tests conducted at 600°C focused on deposit chemistry effects under combined co-firing coal/biomass and oxy-combustion conditions. Tests conducted at 700°C focused on gas-phase chemistry effects under oxy-combustion conditions.

**Experimental procedures**

The tests were conducted at 600°C and 700°C, with different heats of T91 used at each temperature. These steels have a metastable-tempered martensite structure that is quite persistent at boiler temperatures; ferrite is the stable phase. Exposure temperatures of 700°C and above are higher than the ferritic-martensitic steels would see in steam boiler service as a metal temperature. However, actual service conditions are not isothermal and ash and outer oxide layers on the fireside of the boiler tube would be expected to be at higher temperatures than the alloy.

The alloy compositions are given in Table 1. The composition of the heat used at 600°C was measured by energy dispersive x-ray analysis (EDX). The composition of the heat used at 700°C was measured by wavelength dispersive x-ray fluorescence (WDXRF) with a Rigaku ZSX Primus II. The exceptions were for C and N, where nominal compositions are listed.

Test coupons were machined from commercial tubes. For the 600°C tests, the coupons were curved tube segments (~50° tube arcs), with dimensions approximately 15 mm long with a 4 mm wall thickness. All surfaces were ground to ISO/FEPA 600 grit surface (25.8 µm particle diameter, roughness average ($R_a$) of <0.4 µm). For the 700°C tests, the coupons were flat, with dimensions approximately 19 × 13 × 3 mm. All surfaces were ground to CAMI 600 grit surface (16.0 µm particle diameter, $R_a$ of 0.13 µm).

**600°C test procedures (tests focused on deposit effects in coal/biomass co-firing)**

| Alloy | $T$, °C | Fe | Cr | Ni | Mo | C | N | Si | Al | Mn | V | Cu |
|-------|---------|----|----|----|----|---|---|----|----|----|---|---|
| T91   | 600     | Bal| 8.36| 0.21| 0.90| 0.10| 0.05| 0.12| 0.02| 0.45| 0.17|
| T91   | 700     | Bal| 8.48| 0.16| 0.99| 0.10| 0.05| 0.35| 0.41| 0.20| 0.18|

A vertical alumina-lined controlled atmosphere furnace was used for these tests. This system can expose up to 24 samples in alumina crucibles held by an alumina frame. The test was run in a series of five 200 h segments (to give a total exposure time of 1000 h) using the well-established ‘deposit recoat’ technique [18,23].

The test exposure conditions were established by a detailed study of the potential gas compositions and deposit compositions/fluxes that could be produced if an oxy-fired system was fired using a common UK coal (Daw Mill) and biomass product (cereal co-product, CCP) for use in UK power stations [16]. The gas composition predicted for the use of 80 wt % of the UK coal and 20% of the CCP combined with HGR and O$_2$ (at 95 vol % purity) were calculated using spreadsheet-based models that have been validated using pilot plant data [13,16]. These gas compositions were simplified (to one composition, 59% CO$_2$, 5.2% N$_2$, 31% H$_2$O, 4% O$_2$, 0.626% SO$_2$ and 0.17% HCl) to include their key active components for corrosion testing in SH/RH environments – shown in Table 2, in the Oxy-Combustion ‘Co-Fired’ 600°C column. The Oxy-Combustion ‘Co-Fired’ gas mixture was achieved by combining three pre-mixed gas bottles (HCl-N$_2$, CO$_2$, SO$_2$, CO$_2$, O$_2$, and CO$_2$) using mass flow controllers. Deionised water was added with a peristaltic pump.

The chemical compositions of the deposits used in this portion of the study are shown in Table 3 (D0-D5). Deposit D6 was only used in the 700°C tests that were focused on gas composition effects and is described later. At the start of the test, these deposits were applied to the convex surfaces of the samples, using a paintbrush, to give a coverage of ~20 mg/cm$^2$. After each 200 h cycle, the samples were unloaded from the furnace and weighed: first in their individual crucibles (together with any spalled deposit/oxide) and then without the crucible (and any spall) to enable the calculation of gross and net mass changes (to assist with monitoring the progress of the test). Following these weight measurements, the samples were each recoated with a further ~20 mg/cm$^2$ of deposit before being reloaded into their crucibles and furnace.

The deposits can be characterised as [24]:

- D0: no deposit (as a reference)
- D1: a deposit composition that is widely used in screening tests. At test conditions, it generates an aggressive alkali–iron trisulphate that has been frequently associated with fireside
The presence of CaO in D2-D4 and CaCO\(_3\) in D5 can lead to the formation of alkaline calcium trisulphates instead of alkaline iron trisulphates and are generally less corrosive [24,26].

Following their exposure, samples were carefully mounted using jigs and a low shrinkage epoxy resin 50% filled with ballotini (to further reduce shrinkage). These mounted samples were then sectioned (using a CBN cutting wheel and oil-based lubricant) perpendicular to their lengths. The cross-sections were prepared using standard SiC grinding papers and then diamond pastes using oil-based lubricants.

After preparation, the cross-sections were measured using a calibrated x-y stage on a Leitz optical microscope linked to an image analyser system to generate accurate measurements of the edge of the remaining metal and the depth of any internal damage. These measurements were compared to the pre-exposure sample dimensions (collected using a digital micrometre, with a resolution of 1 \(\mu\)m) to produce distributions of the changes in metal thickness (with and without any internal damage) that had been caused by the exposures. Detailed descriptions of this procedure for dimensional metrology were reported previously [e.g. 24].

The microstructure of the deposits and corrosion products were characterised using an FEI environmental scanning electron microscope (ESEM) with Oxford Instruments energy dispersive X-ray analysis (EDX).

Additional details of the procedures used for the 600°C exposures can be found in references [23, 24 and 27].

### 700°C test procedures (tests focused on gas composition effects in air- and oxy-firing)

The tests were conducted in horizontal tube furnaces, with the samples placed face down in alumina boats. Synthetic ash was used to cover the samples. The components of the ash were ball-milled for good mixing. The specimens were buried with a 3 mm cover of ash, which was then lightly compacted using a mass equivalent to 40 g/cm\(^2\). The composition of the ash used at 700°C (D6) is given in Table 3. Compared to the other deposits, D6 used Al\(_2\)O\(_3\) and SiO\(_2\) instead of Kaolinite for the (expected) inert portion of the ash. Deposit D6 is similar to D1 in that it does not contain KCl, CaO or CaCO\(_3\), but it has a much smaller amount of sulphates. It should be less of an accelerated test than the use of D1.

The specimens were exposed for either 240 h (ash-retained exposures), or for 1440 h in 240 h test segments (section loss exposures) in the four gas mixtures associated with either air-combustion or oxy-combustion gas mixtures as shown in Table 2. The shorthand for the four cases are ‘air’ for air-combustion, ‘FGD <H\(_2\)O’ for oxy-combustion after flue gas desulphurisation (FGD) and water removal in the flue gas recirculation stream, ‘FGD’ after desulphurisation in the flue gas...

### Table 2. Gas phase compositions, vol %.

| Gas   | Oxy-Combustion 'Co-Fired' | Air-Combustion 'Air' | Oxy-Combustion 'FGD <H\(_2\)O' | Oxy-Combustion 'FGD' | Oxy-Combustion 'HGR' |
|-------|--------------------------|---------------------|-------------------------------|---------------------|---------------------|
| T, °C | 600                      | 700                 | 700                           | 700                 | 700                 |
| CO\(_2\) | 59.0                     | 14                  | 80.2                          | 69.2                | 68.6                |
| N\(_2\) | 5.2                      | 74.2                | 8.0                           | 8.0                 | 8.0                 |
| H\(_2\)O | 31.0                     | 9.0                 | 9.0                           | 20.0                | 20.0                |
| O\(_2\) | 4.0                      | 2.5                 | 2.5                           | 2.5                 | 2.5                 |
| SO\(_2\) | 0.626                    | 0.3                 | 0.3                           | 0.3                 | 0.9                 |
| HCl  | 0.170                    |                     |                               |                     |                     |

### Table 3. Chemical composition (mol %) of the deposits used.

| ID   | T, °C | Kaolinite\(^a\) | Al\(_2\)O\(_3\) | SiO\(_2\) | Fe\(_2\)O\(_3\) | Na\(_2\)SO\(_4\) | K\(_2\)SO\(_4\) | KCl | CaO | CaCO\(_3\) |
|------|------|-----------------|----------------|----------|----------------|-----------------|----------------|-----|-----|-----------|
| D0   | 600  |                 |                |          |                |                 |                |     |     |           |
| D1   | 600  |                 |                |          |                |                 |                |     |     |           |
| D2   | 600  | 57.0            |                |          |                |                 |                |     |     |           |
| D3   | 600  | 57.0            |                |          |                |                 |                |     |     |           |
| D4   | 600  | 53.0            |                |          |                |                 |                |     |     |           |
| D5   | 600  | 57.0            |                |          |                |                 |                |     |     |           |
| D6   | 700  |                 |                |          |                |                 |                |     |     |           |

\(^a\)Al\(_2\)O\(_3\) · 2SiO\(_2\) · 2H\(_2\)O

corrosion of SH/RH in coal-fired power stations [13,17,25]. This deposit causes accelerated fireside corrosion damage.

- **D2**: a simplified alkali–iron trisulphate composition (without K) diluted with kaolinite (Al\(_2\)O\(_3\) · 2SiO\(_2\) · 2H\(_2\)O), Fe\(_2\)O\(_3\) and CaO to represent the minerals often found in coals.
- **D3**: similar to D2, but with 5% of the Na\(_2\)SO\(_4\) replaced with K\(_2\)SO\(_4\).
- **D4**: similar to D3, but with the K present as KCl to investigate one potential effect of biomass co-firing.
- **D5**: similar to D2, but with CaO replaced by CaCO\(_3\).
recirculation stream, and 'HGR' for the hot-gas-recycle case (no purification of the recirculation stream). The four gas mixtures were achieved by combining pure gases using mass flow controllers. Deionised water was added with a syringe pump. The gas mixtures flowed at a rate of 25 cm/min (at 700°C) first past a Pt/Rh mesh and then over the boat with samples and ash. The Pt/Rh mesh was to catalyse SO$_3$ formation from SO$_2$ and O$_2$. Oxides in the ash, whether in laboratory ash or in a boiler, also catalyse the formation of SO$_3$.

**Ash-retained exposures**

After the test, the entire alumina boat (with samples and intact ash) was embedded in epoxy and cross-sections were cut and examined. This allowed observation of the alloy-scale-ash largely intact after exposure. The sectioning step used water for lubricant and cooling, so water-soluble species containing Na and K may have been lost. Examinations were by polarised light (PL) and differential interference contrast (DIC) with a Leitz metallurgical microscope. Additional examinations were done using field-emission scanning electron microscopy (SEM) with both a JEOL 7000F equipped with Oxford energy dispersive x-ray (EDX) and wavelength dispersive spectrometer (WDS) detectors and an FEI Quanta 600 equipped with an Oxford X-Max 80 EDX.

**Section loss exposures**

Exposures were done in 240 h increments on triplicate samples. After each exposure, the ash was removed, and the samples gently cleaned in isopropyl alcohol. Mass changes measurements were made, but since significant ash remained on the surface, these measurements were of minimal value. Light microscopy macro images were made of each sample. One of each triplicate or duplicate sample was cut for cross-section analysis after 240 and 480 h. All the samples (the enact sample(s) and the remaining portion of the cut sample) of each alloy were exposed to the next 240 h exposure. Care was taken to ensure that the same side was facing upwards on both exposures. After the final exposure period, all the samples were cross-sectioned for analysis. Section loss measurements were made by comparing pre-test thickness measurements from 15 separate locations on the sample with image analysis done on high-contrast light microscopy images of cross-sections after the test. The image analysis consisted of measuring the metal thickness at many regularly spaced locations (typically 50 locations) along the width of the sample. Measurements near the edges of the cross-section were not included in the analysis. Similar section loss measurements were also made for alloys in the 24 h ash-retained exposures.

**Results and discussion**

**600°C tests (tests focused on deposit effects in coal/biomass co-firing)**

Figure 2 shows the time dependence of the mass change data for the different deposits. After 1000-h exposure, these data ranked the mass gains for the T91 samples with different deposits (from lowest to highest) as D0, D2, D3, D5, D4 and D1. As expected, the sample without deposit (D0) had the lowest mass change, and the most aggressive deposit (D1) produced the highest mass change. However, more detailed interpretation of such data is limited by deposit/corrosion product spallation. Dimensional metrology needs to be used to obtain a better dataset for the metal damage generated by fireside corrosion. The pre- and post-exposure dimensional metrology processes are used to produce datasets of changes in metal thickness as a function of position around the samples. These datasets were then converted into metal losses and corrected for calibration differences between the pre- and post-exposure metrology processes (using data from reference samples), before being reordered into distributions of metal loss (from least to greatest). These processed data are plotted in Figure 3 as metal loss (with units of µm) as a function of cumulative normal probability (with units of standard deviations).

These data show that the lowest metal losses were obtained from samples with no deposit (D0) and with deposit D2; with the samples still being in the incubation stage of fireside corrosion. The highest metal losses were obtained with deposit D1 (which had been anticipated to be the most aggressive deposit); with the sample having been in the propagation stage of fireside corrosion for most of its exposure. Intermediate metal losses were obtained for deposits D3-D5; with samples...
having transitioned from incubation to propagation damage during the course of their exposures.

ESEM/EDX analyses were performed after 1000 h of exposure of samples exposed with deposit D1 and without deposit (D0); backscattered ESEM images are shown in Figure 4. Without a deposit (D0), the scale was ~90 µm thick and had an outer layer of Fe$_3$O$_4$ (with some Cr) and an inner layer of Fe-Cr spinel (with some S). In contrast, with deposit D1 the scale was much thicker (~300–340 µm), with a surface deposit/corrosion product mix of ~300 µm. The inner oxide layer was an Fe-Cr spinel with S near the alloy and Fe$_3$O$_4$ with Cr, S, Na, and K below a surface deposit/corrosion product mix.

Based on the research reported elsewhere [e.g. 24, 25, 28], it is suggested that the compounds applied in deposit D1 have reacted with each other and the gaseous environment to form alkali-iron sulphates that have melted below the exposure temperature for this test and enabled the propagation of fireside corrosion to start quickly. In contrast, the initial deposit D2, with the addition of an alumina-silicate mineral and CaO as well as the removal of K$_2$SO$_4$, resulted in a surface condition that did not cause fireside corrosion during this exposure (due to its melting point being higher than the exposure temperature). Deposits D3-D5, with varying initial levels of Na$_2$SO$_4$, K$_2$SO$_4$, KCl, CaO and CaCO$_3$, resulted in surface conditions that enabled the transition between the incubation and propagation stages of hot corrosion to occur during these exposures. This was due to reactions between the complex initial deposits and gaseous environments generating mixtures with melting points that have moved below the test temperature, enabling the fireside corrosion fluxing reactions to start.

700°C tests (tests focused on gas composition effects in air- and oxy-firing)

Ash-retained exposures

Significant corrosion was observed after exposure. Figure 5 shows T91 with retained ash after 240 h exposure in an oxy-combustion ‘FGD’ gas (CO$_2$ + 8% N$_2$ + 20% H$_2$O + 2.5% O$_2$ + 0.3% SO$_2$) with a D6 ash cover. The oxide scale consists of an inner scale containing (Fe,Cr) oxide and an outer scale consisting of Fe oxide. This is typical of 9Cr steel oxidation. Sulphur was found within the inner (Fe,Cr) oxide scale. Above the outer oxide scale is an oxide deposition zone, where dissolved oxide, from a fluxing reaction, has deposited within the ash cover. Above the oxide deposition zone is an ash-affected zone, which by PL has a different colouration than the remainder of the ash cover. Examination of the ash-affected zone at higher magnification, Figure 5(d), shows needle-shaped Fe oxides growing within the pores in the ash. These needles are not present further away from the alloy. The needles appear to be the location of a reaction front where iron oxides precipitate, in a non-protective morphology, after being dissolved at the scale-ash interface. This is consistent with a fluxing-type model of hot corrosion [4].

Figure 6 shows edge effects for T91 after exposure for 240 h at 700°C with a D6 ash and an oxy-combustion FGD <H$_2$O gas (CO$_2$ + 8% N$_2$ + 9% H$_2$O + 2.5% O$_2$ + 0.3% SO$_2$). The oxide deposition region is much thicker near the edges of the sample. There is visible section loss on the right-hand edge of the sample. However, the inner and outer oxide scales...
have similar thicknesses, regardless of distance from an edge.

**Section loss exposures**

An example of the remaining thickness measurements taken on cross-sections after a test is in Figure 7(a) for T91 after 1440 h at 700°C in the oxy-combustion HGR gas with D6 ash. Each measurement was taken at regular intervals, approximately 250 µm apart from each other. Normally five images for each specimen, covering the entire cross-section, are used, generating about 50 measurements per specimen. The difference between these measurements and the initial thickness is used to determine the metal loss. Metal loss data were arranged from smallest to largest and put into cumulative probability plots. Figure 7(b) shows a cumulative probability plot for the entire T91 sample shown in Figure 7(a). There is a good agreement with a normal distribution, which is associated with general corrosion. The cumulative normal distribution is determined from the mean and variance of the metal loss data, using the Excel Normdist function – it is not a fit of the individual data points.

Even though corrosion coupons were placed in the alumina boat, and covered with ash on one side, corrosion damage was typically observed on both sides of the sample.

Figure 8 shows the mean metal section loss for T91 after exposure under a D6 ash and with four different gas compositions. Several observations can be made. First, there was little difference in mean metal section loss (720–820 µm) after 1440 h of exposure. Second, the initial differences after 240 h, noted in an earlier publication [10], are no longer apparent after longer exposures. Finally, the overall section loss kinetics appeared linear, and the total section losses substantial.

In similar tests (alloy T92 in with deposit D1 for 1000 h) Hussain [29] showed metal loss differences at...
700°C of ~1100 μm for an HGR case and ~850 μm for an air-fired case. The major differences in test procedures were the much higher SO₄ levels in the Hussain result, but with the use of much thinner deposits (20 mg/cm² as opposed to ~3 mm of ash cover). The section loss results in Figure 8 could be a consequence of thicker ash cover masking some of the gas composition differences.

Conclusions

This paper reported the collaborative results on examining the corrosion resistance of T91 in a variety of oxy-fuel combustion environments. Tests conducted at 600°C focused on deposit chemistry effects under combined co-firing coal/biomass and oxy-combustion conditions. Tests conducted at 700°C focused on gas-phase chemistry effects under oxy-combustion conditions.

The exposures at 600°C (tests focused on deposit effects in coal/biomass co-firing) showed:

- Substantial metal losses (30–120 µm) were observed for all 1000 h exposures.
- Under deposit D1 the formation alkali iron sulphates enabled a relatively high rate of metal loss (the highest of the five deposits used at 600°C).
- Lower corrosion damage was observed with deposits D3–D5, which appeared to have delayed the transition to the propagation stage of fireside corrosion.
- Under deposit D2, the metal losses were similar to those observed with no deposit, with the sample still being in the incubation stage of fireside corrosion.

The exposures at 700°C (tests focused on gas composition effects in air- and oxy-firing) showed:

- Substantial metal losses (720–820 µm) were observed for all 1440 h exposures.
- The oxidation kinetics were linear.
- Iron oxide morphology, within the ash at the oxide scale-ash interface, was consistent with a fluxing (dissolving and later non-protective precipitation) mechanism.

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