The Corrosive Effect of Chlorine Containing Species on Waterwalls and Superheater Materials in Waste and Biomass-Fired Power Plants

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Department of Chemistry and Chemical Engineering
CHALMERS UNIVERSITY OF TECHNOLOGY
Gothenburg, Sweden 2016
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Cover: TG mass gain curves of T22 at 400 °C and an image of a cross section of T22 in the presence of KCl + HCl(g) after 1 hour of exposure at 400 °C.

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Abstract

To increase power generation from waste- and biomass-fired boilers, it is necessary to increase the temperature and pressure of the steam generated. This implies an increase in the temperature both of the waterwalls and of the steam superheaters in the boiler. Low-alloyed steels are preferred in the waterwalls due to low cost and excellent mechanical properties and weldability. The recent shift from fossil fuels (oil, coal and natural gas) to biofuels and waste for the generation of heat and power has introduced several challenges to the boilers. One is the rapid corrosion of superheaters and waterwalls. Many observations imply that the corrosivity of the fireside environment in boilers fired by biomass and waste is connected to the relatively high chlorine/sulphur - ratio in the flue gas. Hence, this thesis investigates the role of chlorine compounds on fireside corrosion of both water walls and superheaters.

The waterwall part of this thesis investigates the corrosive effect of alkali chlorides (KCl and NaCl), PbCl₂ and ZnCl₂, including the effect of KCl+ZnCl₂ and NaCl+ZnCl₂ mixtures. This work shows that all four salts accelerates the corrosion of waterwall steels in the temperature range of interest. A new mechanism describing alkali chloride-induced corrosion of low-alloyed steels is presented. The other part of the thesis investigates the influence of SO₂ on the corrosion of superheater steels in the presence of alkali chlorides in a laboratory setting. Also, complementary corrosion experiments were performed in a commercial waste-fired power plant. The results show that the corrosivity of chlorides in superheater deposits is mitigated by SO₂ in the flue gas. The effect is attributed to the conversion of KCl and NaCl into K₂SO₄ and Na₂SO₄, the two sulfates being relatively harmless from a corrosion point of view at the temperatures studied (525 °C and 600 °C). However, the laboratory study indicated that the simultaneous presence of KCl and SO₂ can cause steel grain boundary attack beneath the original KCl particle.

Keywords: Biomass, Waste, Combustion, Alkali-induced corrosion, Heavy metal chlorides, Sulfur recirculation, Waterwalls, Superheater corrosion.
List of publications
This thesis is based on the following appended papers.

Paper I
E. Larsson, T. Jonsson and J. Liske
The influence of small amounts of KCl(s), NaCl(s), ZnCl₂(s) and salt mixtures on the high temperature oxidation of a Fe-2.25Cr-1Mo steel at 400°C
Submitted to oxidation of metals

Paper II
E. Larsson, T. Jonsson, J. Liske and L-G. Johansson
The influence of KCl and HCl on the high temperature oxidation of a Fe-2.25Cr-1Mo steel at 400°C
Submitted to oxidation of metals

Paper III
A comparative study of the initial corrosion of KCl and PbCl₂ on a low alloyed steel
Accepted to Oxidation of Metals

Paper IV
A Laboratory Study of the In-situ Sulphation of Alkali Chloride-Rich Deposits
–Corrosion perspective
Energy Fuels, 2016, 30 (9), pp 7256–7267

Paper V
Sulfur recirculation for increased electricity production in Waste-to-Energy plants
Waste Management Volume 34, Issue 1, January 2014, Pages 67–78

Paper VI
E. Larsson, S. Andersson, J. Liske
The influence of sulfur recirculation on superheater corrosion rates, evaluated by a novel image analysis method combined with SEM
Manuscript
Statement of contribution

Paper I
I was the main author and carried out all exposures and all analysis.

Paper II
I was the main author and carried out all exposures and all analysis.

Paper III
I was the main author and carried out the KCl exposures and analysis, H. Gruber carried out the PbCl$_2$ exposures and analysis.

Paper IV
Performed BIB cross sections and participated in the writing.

Paper V
I assisted with the exposures in the boiler and carried out the SEM analysis and material loss measurement. I wrote the corrosion part of the paper together with Jesper Liske.

Paper VI
I was the main author. I assisted with the exposures in the boiler and carried out the SEM, XRD and IC analysis as well as material loss measurement.
Related papers and technical reports not included in the thesis

Froitzheim, J, Ravash, H, Larsson, E, et al.  
Investigation of Chromium Volatilization from FeCr Interconnects by a Denuder Technique  
JOURNAL OF THE ELECTROCHEMICAL SOCIETY  Volume: 157  Issue: 9  Pages: B1295-B1300  Published: 2010

“Evaluation of different fuel additives’ ability to master corrosion and deposition on steam superheaters in a waste fired CFB-boiler”  
KME report, 2010

”Measures for simultaneous minimization of alkali related operating problems, Phase 3”  
Värme forsk report, 2010

M. Gyllenhammar, S. Herstad Svärd, K. Davidsson, J. Liske, E. Larsson, T. Jonsson, D. Zhao  
”Additiv för att minska driftproblem vid rostförbränning”  
Waste refinery 2013

“Increased electrical efficiency and service life assessment of super heaters from combustion of difficult fuels”  
KME report 2014
Preface
This thesis summarizes the research that was performed within the Swedish competence center for High Temperature Corrosion (HTC), from 2010 to 2016. The work has been carried out Environmental Inorganic Chemistry at the Department of Chemistry and Chemical Engineering, Chalmers University of Technology, Göteborg, Sweden, under the supervision of Professor Jan-Erik Svensson. The work was funded by the High Temperature Corrosion Centre (HTC), Konsortiet för Materialteknik för termiska Energiprocesser (KME), Värmeforsk (VF), Waste Refinery (WR), Proenviro and the Swedish Energy Agency.
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<th>Description</th>
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<tr>
<td>BF</td>
<td>Bright field</td>
</tr>
<tr>
<td>BIB</td>
<td>Broad ion beam</td>
</tr>
<tr>
<td>BSE</td>
<td>Backscattered electron</td>
</tr>
<tr>
<td>EDX</td>
<td>Energy dispersive x-ray</td>
</tr>
<tr>
<td>FEG</td>
<td>Field emission gun</td>
</tr>
<tr>
<td>FIB</td>
<td>Focused ion beam</td>
</tr>
<tr>
<td>HAADF</td>
<td>High angle annular dark field</td>
</tr>
<tr>
<td>IC</td>
<td>Ion chromatography</td>
</tr>
<tr>
<td>SE</td>
<td>Secondary electron</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning electron microscope</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission electron microscope</td>
</tr>
<tr>
<td>XRD</td>
<td>X-ray diffraction</td>
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1 Introduction

Since oil and coal (fossil fuels) result in a net contribution of CO$_2$ to the atmosphere when combusted, it is believed to be problematic due to its contribution to the global warming effect. The projections are that the trend in increasing global temperatures will continue if drastic counter measures are not undertaken [1]. As a consequence there is a political driving force to reduce the dependence of fossil fuels for energy conversion. The Swedish government has decided to reduce the emissions of greenhouse gases by the year 2020 by 40 \% from 1990 emission levels and, additionally, the government has set the target for an increase in efficiency of energy use by 20 \%. In addition it has been decided that at least 50 \% of overall energy consumption should come from renewable sources [2]. In order to reach these goals more or less all power conversion sectors of society are effected. This thesis deals with the material challenges that occurs when shifting from fossil fuels to renewables such as biomass and waste in heat and power plants.

There are inherent problems that must be solved when shifting from fossil fuels to waste and biomass. One important issue is the high corrosion rates that occurs on heat exchange materials in the boilers, where heat is transferred from the combustion reaction to the steam system (superheaters) that powers a turbine, and to boil the water within the waterwall tubes. Biomass, and waste, upon combustion releases high levels of compounds such as hydrogen chloride (HCl(g)), alkali chlorides (NaCl, KCl) and heavy metal chlorides (ZnCl$_2$, PbCl$_2$) [3-11]. This makes the flue gas and deposits considerably more corrosive compared to when coal or oil are used as fuels. In order to limit the corrosion issues, plant operators can lower the steam temperature in the superheaters which decrease not only the material temperature but also the electrical efficiency of the process. It is also possible to use expensive heat exchange materials with high corrosion resistances. As both strategies result in a less cost effective process, understanding and mitigation of material degradation is of great importance.

Knowledge about superheater corrosion is relatively well established and is usually attributed to the presence of alkali chlorides. Therefore, research is at a stage where different mitigating techniques are being tested and elaborated upon. Research directed into waterwalls is at a stage where the main contributors regarding corrosion is being researched. To mitigate the corrosion in this region of the boiler, expensive materials are often welded onto existing tubes.
Chlorine-containing compounds are generally considered to account for the more corrosive environment when these alternative fuels are used [4, 5]. However the underlying mechanisms for this are not completely understood.

Certain species or mixtures can be present in the tube deposit during combustion of waste and biomass, with melting points below the local temperature of the waterwalls. Because of the lower material temperatures compared to at the superheaters, more corrosive species (i.e. heavy metal chlorides) can condense out on the heat exchanger surfaces. Substantially elevated corrosion rates in the presence of melts have been reported in the literature [12-14]. The low melting point has been attributed to the presence of heavy metal chlorides in flue gas deposits.

The aim of this thesis is to investigate the effect of chlorine containing compounds in different parts of a boiler i.e. at the waterwalls and on the superheater tubes. The waterwall research was carried out in well controlled laboratory exposures, in which T22 sample coupons were coated with KCl, NaCl and ZnCl₂ or with mixtures of ZnCl₂/(KCl or NaCl). Oxidation in the presence of PbCl₂ was also investigated. The combined effect of KCl and a gaseous chlorine containing specie HCl(g), which are normally present simultaneously during combustion were also assessed. The superheater research was conducted in a commercial waste fired boiler with the aim of evaluating the effectiveness of a novel corrosion mitigating technique, in which sulfur is recirculated back into the combustion area. The effect of in-situ sulfation of relatively large amounts of KCl towards a 304L steel was also investigated in a laboratory setting.
2 Steels

Steel is a generic name that covers many different alloys, however in its most basic form a steel is an interstitial solid solution of carbon in an iron matrix. The carbon concentration is typically within the range of ~ 0.01 – 1 wt. %. The addition of carbon increases the strength and toughness of the material drastically compared to pure iron [15].

Pure iron exists in two allotropes at atmospheric pressure, ferrite (α) and austenite (γ). At room temperature ferrite is the thermodynamically favored form in which the atoms are arranged in a body-centered cubic structure (bcc). If pure iron is heated, a phase transformation from ferrite to austenite occurs at 910 °C. In austenite the atoms have a face-centered cubic arrangement (fcc), see Figure 2.1 [16].

![Figure 2.1: Cubic unit cells. a) Body-centered cubic (bcc), b) face-centered cubic (fcc).](image)

Steels are polycrystalline materials consisting of grains with an approximate size range from 5 – 100 µm depending on the type of steel. Carbide particles are generally dispersed in the material. Both the grain size and distribution of carbides are important parameters to consider since these greatly influence the strength and toughness of the steel. This is done by carefully controlled heat treatments and cold working, which makes it possible to customize the desired properties [17].

The most important alloying element for corrosion resistance in steels is chromium. With a chromium content of approximately 10 wt. % a stainless steel is formed [15]. The corrosion resistance of stainless steels can be attributed to their ability to form a chromium rich oxide
scale \((\text{Fe}_{1-x}\text{Cr}_x)_2\text{O}_3\) that protects the material from further oxidation. Stainless steels can be in both ferritic and austenitic form, however austenitic stainless steels have superior corrosion resistance. Austenitic steels are more ductile but not as strong as ferritic steels [17]. To attain the austenite phase at room temperature, substantial amounts of austenite stabilizing elements such as nickel or manganese are required. This results in a considerably higher price compared to ferritic steels.

The low alloyed ferritic Fe-2.25Cr-1Mo steel (T22) that was used in the laboratory study contains approximately 2.2 wt. % chromium, while the 16Mo3 that was used in the field study contains no chromium at all. As a consequence these materials are not protected by a chromium-rich oxide. Instead low-alloyed steels rely on iron oxides as protection against further oxidation and are normally used when the environment is less corrosive.

Two stainless steels were investigated in this study, the alloy 304L was studied in the laboratory part and in the field part the highly alloyed stainless steel Sanicro 28 was studied. Sanicro 28 contains considerably higher concentrations of Cr and Ni than 304L and thus has a better corrosion resistance. The nickel based Inconel alloy 625 that was used in the field study forms a more or less pure chromium oxide since its Cr content is above wt. 20 % [18]. Material compositions are listed in Table 1.

Table 1. Elemental composition of the alloys investigated in the thesis in weight %.

<table>
<thead>
<tr>
<th>Alloy</th>
<th>% Fe</th>
<th>% Cr</th>
<th>% Ni</th>
<th>% Mo</th>
<th>% Si</th>
<th>% C</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe-2.25Cr-1Mo</td>
<td>96</td>
<td>2.2</td>
<td>-</td>
<td>0.9</td>
<td>0.3</td>
<td>0.1</td>
<td></td>
</tr>
<tr>
<td>16Mo3</td>
<td>99</td>
<td>-</td>
<td>-</td>
<td>0.3</td>
<td>0.3</td>
<td>0.12</td>
<td>0.6 % Mn</td>
</tr>
<tr>
<td>304L</td>
<td>68</td>
<td>19</td>
<td>10</td>
<td>0.5</td>
<td>0.6</td>
<td>&lt;0.03</td>
<td>1.4% Mn</td>
</tr>
<tr>
<td>Sanicro 28</td>
<td>35</td>
<td>27</td>
<td>31</td>
<td>3.5</td>
<td>0.7</td>
<td>&lt; 0.02</td>
<td>1 % Cu, 2% Mn</td>
</tr>
<tr>
<td>Inconel alloy 625</td>
<td>5</td>
<td>22</td>
<td>58</td>
<td>9</td>
<td>0.5</td>
<td>&lt; 0.10</td>
<td>0.5 % Mn, 3.15-4.15 Nb+Ta</td>
</tr>
</tbody>
</table>
3 Oxidation of metals

3.1 Oxide formation

A metal's resistance to corrosion depends on the properties of the oxide scale that is formed on the surface. For ultimate resistance the scale should act as a barrier to the surrounding environment, in other words, the scale ideally should be well adherent to the metal and have low diffusion coefficients for ions and atoms.

Almost all metals are thermodynamically unstable when exposed to oxygen, which means that an oxide layer rapidly forms on the metal surface. When a clean metal surface is exposed to oxygen, the oxygen is adsorbed to the surface and reduced by the metal to form a metal oxide (Me$_x$O$_y$) see reaction 3.1.

$$x\text{Me} + 0.5y\text{O}_2(g) \rightarrow \text{Me}_x\text{O}_y$$  \hspace{1cm} (3.1)

The oxidation process initiates with the adsorption of oxygen molecules on the metal surface, followed by a charge transfer and the formation of O$^{2-}$ (see Figure 3.1 a). The next step is the formation of individual oxide nuclei that grow laterally and eventually coalesce to cover the entire metal surface (b). These first two steps are very fast and normally not observable at room temperature. At this point, further oxide growth is controlled by the diffusion of reactants across the continuous oxide film. The diffusion rate through the oxide at room temperature is close to zero and the oxide growth is negligible. However, with increasing temperature, diffusion rates normally also tend to increase [18]. Thus, the high temperature performance of a metal is, to a great extent, determined by the diffusion properties and defect structure of its formed oxide.
3.2 Diffusion processes

For the growth of oxide scales, metal or oxygen ions must diffuse through the oxide layer. Transport of these ions depends on the presence of defects and imperfections in the structure of the oxide. Transport through metal oxides at high temperature, are either bulk diffusion or short-circuit diffusion [19].

Metal oxides are seldom stoichiometric. The non-stoichiometry of oxides affects the properties of the oxides, e.g. the electronic conductivity in the oxide that is very important in high temperature oxidation. Non-stoichiometric oxides have either an excess or a deficit in metal content, which leads to interstitial or vacant sites in the lattice. In an oxide with metal excess, interstitial ions dominate the defect structure. Metal excess can also be achieved by means of oxygen ion deficiency. In both of these instances, the defects have positive charges, and these are compensated by the formation of equivalent concentrations of negative electrons [18]. The charge is, thus, transferred by negative carriers and the semiconductor is classified as n-type, (see Figure 3.2). Positive (p-type) semiconductors have a metal deficit, either by the removal of metal ions or by the addition of oxygen ions. These defects will have negative effective charges that are compensated through the formation of positive electron holes [18].
3.2.1 Lattice diffusion

Lattice diffusion occurs through the movement of point defects, e.g. vacancies and interstitial ions. The most common diffusion mechanisms are vacancy diffusion, interstitial diffusion and the interstitialcy mechanism, (see Figure 3.3). Vacancy diffusion occurs when an ion in a normal lattice site jumps to an adjacent unoccupied site in the lattice, (a) in Figure 3.3. Interstitial diffusion refers to the movement of an ion situated in an interstitial site to a neighboring interstitial site, (b) in Figure 3.3. This mechanism is probable only when the interstitial ion is smaller than the ions in the normal lattice. The interstitialcy mechanism occurs when an interstitial ion moves to a normal lattice site, forcing the ion in that position to move to the next interstitial position, (c) in Figure 3.3. Vacancy diffusion is possible in crystals with Schottky defects, while interstitial movements of ions occur in crystals with Frenkel defects.

Figure 3.3: Diffusion mechanisms in a lattice: (a) vacancy mechanism; (b) interstitial mechanism; and (c) interstitialcy mechanism [19].
3.2.2 **Short-circuit diffusion**

Ion diffusion can also occur through short-circuit diffusion paths. Short-circuit diffusion take place in three-dimensional dislocations, e.g. along grain boundaries or cracks, which provide fast transport paths through the oxide. In steels the bulk consists of small grains that give rise to many of these dislocations. At low temperatures this is the most dominant diffusion process. With increasing temperature bulk diffusion becomes increasingly significant. The activation energies for short-circuit diffusion are lower than those for lattice diffusion (typically 50–70 %), which makes short-circuit diffusion dominant except at very high temperatures [18].

### 3.3 Kinetics of oxidation

The kinetics of oxygen uptake during high temperature oxidation, as illustrated by the mass gain curves in Figure 3.4, can follow several different growth models. The three most common rate laws are the linear, parabolic and logarithmic. All these models are idealized, and in practice, they often only describe part of the oxidation process [18].

![Mass gain curves for three common oxidation rate laws: linear oxidation, parabolic oxidation and logarithmic oxidation.](image)

3.3.1 **Linear rate law**

The linear rate law applies to situations in which the oxidation of a metal proceeds at a constant rate. Typically, the oxidation rate is determined by a phase boundary reaction, but there are also other processes that can exhibit linear kinetics. Oxide growth by the linear rate law is common...
in the early stages of oxidation before the oxide is thick enough to effectively separate the reactants. Linear oxidation can also occur if there is no protective oxide. The transport of ions and electrons will then be fast regardless of oxide thickness, and the oxidation rate will remain linear. In some cases, linear kinetics can occur after a period of parabolic behavior. This is termed breakaway corrosion or catastrophic corrosion and indicates that the oxide has lost its protective properties. Mathematically, the linear rate law is described as

\[ x = k_{\text{lin}} t \]  

(3.2)

where \( x \) is the oxide thickness, \( k_{\text{lin}} \) is the linear rate constant, and \( t \) is the exposure time [20].

### 3.3.2 Parabolic rate law

Most alloys and metals used at high temperatures follow the parabolic rate law. The rate of oxidation is determined by the transport of ions and electrons through the oxide scale. Consequently, the mass gain curve levels off as the oxide thickness increases. The transport is driven by chemical potential gradients between the oxide/gas interface and the metal/oxide interface.

The mathematical expression for parabolic growth was derived by Wagner in 1933 [21]. It can be represented as equation (3.3):

\[ x^2 = k_p t + C \]  

(3.3)

In this expression \( x \) is the oxide thickness, \( k_p \) is the parabolic rate constant, \( t \) is the exposure time and \( C \) is the integration constant. Wagner’s derivation of parabolic oxidation kinetics is highly idealized and based on the following assumptions [18, 20, 21]:

- The oxide layer is compact and perfectly adherent.
- Migration of ions or electrons is the rate determining process.
- Thermodynamic equilibrium is established at both the metal/oxide and oxide/gas interfaces.
- The oxide shows only small deviations from stoichiometry.
- Thermodynamic equilibrium is established locally throughout the oxide scale.
- Oxygen solubility in the metal can be neglected.

3.3.3 Combined rate laws

As mentioned in the linear rate law section, it is not uncommon that an oxidation process can follow different rate laws during different stages of oxidation when abrupt changes in the oxidation behavior occur as illustrated in Figure 3.5. This may occur when an initially protective slow growing oxide abruptly loses its protectiveness, an occurrence known as breakaway oxidation. The reason for such behavior may be, changes in the chemical composition of the scale with a dramatic increase of the diffusion rate and/or mechanical failure of the scale.

Figure 3.5: Example of oxidation according to a combined rate law, with breakaway oxidation after \( t^b \) (time for breakaway).

3.4 Mechanical properties of oxide scales

During the growth of oxide scales, stresses may arise, which can lead to mechanical failure of the scale. This exposes the underlying metal, and fast oxidation can occur. There are several reasons for stress build up, including the oxide growth mechanism, differences in volume between oxide and metal, thermal cycling, differences in thermal expansion coefficient. Upon relaxing the system is often deformed in some way. Stress relief can be divided into four categories: plastic deformation of the scale, detachment of the scale, cracking of the scale, and deformation of the metal substrate [18].
3.5 Corrosion products

The chemical reactions that occur when a metal is exposed to a corrosive environment at high temperatures usually result in the formation of various corrosion products. The most common products are oxides, as all metals used at high temperatures are thermodynamically unstable in the presence of oxygen. As mentioned earlier, the properties of the oxide determine the extent of corrosion. Other elements present in the system may form other compounds with the metal. The properties of these compounds and their interactions with the oxide may also influence further corrosion. The compounds most commonly encountered in bio-mass and waste combustion environments are, e.g. chlorides, sulfides, nitrides and carbides.

Wüstite, Fe$_{1-x}$O

Wüstite forms on pure iron above 570 °C (on iron alloys, higher temperatures are required). The oxide exhibit the rock salt crystal structure (see Figure 3.7), and, since iron can have more than one oxidation state, deviations from stoichiometry are common. The oxide is a metal deficit p-type semiconductor correctly described as Fe$_{1-x}$O, with $x$ from 0.05 to 0.15. It has a high cation vacancy concentration and, consequently, has high ion diffusion rates and is poorly-protective [18].

Magnetite, Fe$_3$O$_4$

Magnetite is less protective than hematite but not as poor as wüstite. The oxide contains both divalent and trivalent iron ions and has the inverse spinel crystal structure (B$_{tet}[A,B]_{oct}O_4$). This structure consists of a face-centred cubic array of oxygen anions. One eighth of the tetrahedral holes are occupied by half of the Fe$^{3+}$ ions, and all of the octahedral holes are occupied by the other half of the Fe$^{3+}$ ions and all the Fe$^{2+}$ ions, (see Figure 3.7) [18].

Hematite, $\alpha$-Fe$_2$O$_3$

Hematite is the most protective of the iron oxides. In dry atmospheres, it remains protective up to 500 °C. It has the corundum structure, which consists of a hexagonal array of oxygen anions, with trivalent metal ions in two thirds of the octahedral holes, (see Figure 3.7). Hematite has n-type conductivity at 650 – 800 °C and p-type conductivity at higher temperatures, but at the
lower temperatures, that are studied in this thesis, both cations and anions are assumed to be transported [18].

**Eskolaite, Cr$_2$O$_3$**

Eskolaite, or Cr$_2$O$_3$, is the most protective oxide of those formed on stainless steels and is the major reason for their high corrosion resistance. Its protectiveness can be attributed to a low level of non-stoichiometry and low diffusion rates of ions. Eskolaite is the only solid chromium oxide that is stable at high temperatures. It has the corundum structure and grows by transport of chromium cations [18].

**Corundum type solid solution (Fe,Cr)$_2$O$_3$**

Eskolaite, Cr$_2$O$_3$ is completely soluble in Fe$_2$O$_3$, which on stainless steels usually results in an oxide with a solid solution of both, i.e. (Fe$_{1-x}$Cr$_x$)$_2$O$_3$. The protectiveness of this type of oxide relies on its chromium content, and corrosion is caused by reactions that remove chromium from it, e.g. the formation of chromic acid [22-27] or chromates [28-33]

**Spinel type solid solution (Fe,Cr,Ni)$_3$O$_4$**

Oxides based on Fe, Cr and Ni may form solid solutions with a spinel-type structure, i.e. (Fe, Cr, Ni)$_3$O$_4$. Compared to magnetite, some of the Fe$^{3+}$ ions may be replaced with Cr$^{3+}$ ions, whereas some of the Fe$^{2+}$ ions may be replaced with Ni$^{2+}$ ions. The actual composition depends on the quantity of Fe, Cr and Ni, as well as the oxygen activity at the oxide/metal interface [18]. Spinel type oxides with other compositions may also form e.g. (Fe, Cr)$_3$O$_4$, which is formed in the inward growing part of the scale on T22 during oxidation. Because of the slow diffusion of Cr in spinel oxides (see Figure 3.6), Cr can act as a marker for the interface between inward/outward growing parts of the scale. This interface also marks the initial alloy surface.
Figure 3.6. Cation diffusion coefficients in magnetite at 1200 °C [34].

Figure 3.7: Crystal structures of metal oxides.

Transition metal chloride

When compounds such as KCl, NaCl, HCl, PbCl₂ are present in the flue gas, transition metal chlorides have been reported to form on metallic heat exchange surfaces. Depending on the alloying elements in the steel, various chlorides, such as FeCl₂, FeCl₃, CrCl₃ and NiCl₂ may form. The presence of transition metal chlorides is normally accompanied by severe corrosion attacks. Iron chlorides are commonly detected on low-alloyed steels in the presence of alkali or
heavy metal chlorides. The diffusion rates of ions in FeCl₂ are quite high. Consequently, iron chloride does not protect the metal. FeCl₂ is a good ion conductor but a poor electron conductor, which may block the electron transport to the surface of the oxide scale, thus disturbing the O₂ reduction on the surface. Iron chloride is volatile at high temperatures and, consequently, may remove material through evaporation. An example of this can be seen in Figure 3.8, which shows that chlorides have evaporated from the sample with subsequent condensation on the sample holder. Iron chlorides are also hygroscopic and may cause severe atmospheric corrosion when the metal is at room temperature in humid air after exposure.

Figure 3.8: Sample holder with condensed iron chloride from evaporation, which has reacted into iron oxide.
4 Fireside corrosion in waste and biomass fired boilers

Examples of biomass used in heat and power production are wood chips, straw, grass and bark. Municipal waste and industrial waste are also used for heat and power production. When combusted, these fuels release higher levels of corrosive compounds, than fossil fuels, such as oil and coal. Most important among the corrosive compounds are alkali salts (e.g. KCl and NaCl) and hydrogen chloride, HCl(g). The lower sulfur to chlorine ratios of waste and biomass are also suggested to make them more corrosive [35, 36]. Compounds that contain chlorine are generally considered to be the most corrosive constituents of the environment in the boiler [37]. Waste normally releases higher levels of chlorine-containing compounds than biomass, as e.g. plastics are common in waste. However, some biofuels, such as straw and grasses, release high amounts of alkali chlorides when combusted. The concentrations of HCl(g) in the flue gas are commonly in the range from 300 ppm to 1000 ppm in waste fired boilers and from 5 ppm to 100 ppm in biomass fired boilers. The KCl + NaCl concentrations in the flue gas in biomass combustion range from ~ 5 to 50 ppm and < 120 ppm for waste combustion (see Table 2 for comparisons).

Table 2. Approximate flue gas composition in coal, biomass and waste [38]

<table>
<thead>
<tr>
<th>Fuel</th>
<th>O₂ %</th>
<th>H₂O %</th>
<th>SO₂ ppm</th>
<th>HCl ppm</th>
<th>KCl+NaCl ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coal</td>
<td>~ 4-5</td>
<td>~ 4-16</td>
<td>~400-1200</td>
<td>~10-50</td>
<td>-</td>
</tr>
<tr>
<td>Biomass</td>
<td>~ 5-10</td>
<td>~ 10-20</td>
<td>~0-70</td>
<td>~25-1000</td>
<td>~5-50</td>
</tr>
<tr>
<td>Waste</td>
<td>~ 5-11</td>
<td>~ 10-20</td>
<td>~0-150</td>
<td>~250-1300</td>
<td>&lt;120</td>
</tr>
</tbody>
</table>

4.1 Fuel types

Several varieties of biomass categories are used for heat and power production. This section focuses on the fuels with the highest concentrations of heavy metal chlorides, which have been identified as a contributor to elevated corrosion rates of boiler waterwalls [39, 40].
4.1.1 Municipal solid Waste (MSW)

Waste is characterized as a biofuel and usually contains higher levels of chlorine and alkali than ordinary biomass such as peat and forest residues. The main contributor of chlorine in MSW has been found to be PVC plastics [41]. There are two main reasons why MSW is used as a fuel. The first one is that it is a simple way to reduce the amount of waste that is deposited in landfills. The motivation for minimizing the amount of waste that end up in landfills has grown due to increased deposition fees in Sweden and the western world. The second reason is that by combusting waste, the intrinsic heat content is recovered and can be transformed into electricity and district heating. When waste was introduced as a fuel source for energy conversion, it was discovered that the resulting flue gas was far more corrosive than in conventional biomass combustion. This caused severe material degradation, with very high corrosion rates, where tube failures occurred after only a few months of operation [42]. Because of the increase in corrosion rates, boilers were forced to operate at lower steam temperatures, with low electrical efficiency as a result.

4.1.2 Recovered waste wood (RWW)

Recovered waste wood is considered to be a renewable a CO$_2$ neutral fuel. Apart from renewability it is fairly inexpensive compared to virgin wood. RWW is highly heterogeneous, originating from a variety of sources. The main sources comes from construction and demolition, MSW and industrial waste. Typical constituents are demolition wood, packaging material and used wood from industrial and residential activities. Since the wood has been used, it is likely to contain non-wood contaminants, such as plastics, paint, glue and chemicals for wood treatment. Paints are the primary source of zinc and lead contaminants (as pigments), which are expected to increase the corrosion rate of boiler water walls. For an elemental composition see Table 3.

Table 3. Average fuel composition of 13 RWW samples [39].

<table>
<thead>
<tr>
<th>S [wt% ds]</th>
<th>Cl [wt% ds]</th>
<th>K [mg/kg ds]</th>
<th>Na [mg/kg ds]</th>
<th>Pb [mg/kg ds]</th>
<th>Zn [mg/kg ds]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.07</td>
<td>0.07</td>
<td>870</td>
<td>830</td>
<td>160</td>
<td>1800</td>
</tr>
</tbody>
</table>
4.2 Combustion systems

The often high levels of alkali, chlorine and water content has forced boiler operators to develop the combustion techniques in aspect of temperature, steam pressure and boiler design. The most frequently used combustion techniques are grate firing, bubbling fluidized bed and circulating fluidized bed. To mitigate the corrosion attack, boilers are constructed with the aim of placing the heat exchanger with the lowest metal temperature in contact with the warmest flue gas. Consequently, the material temperature of the waterwalls is lower than that of the superheaters, since the gas temperature is higher in the furnace where the waterwalls are located. In the event of highly corrosive fuels, boilers are usually designed to have a radiation section without heat exchange tubes, in order to lower the flue gas temperature before it interacts with the superheaters.

4.2.1 Grate

In the group of grate fired boilers, some types can be distinguished based on if the grate is stationary sloping, travelling or vibrating see Figure 4.1. Depending on size, the grate is either air- or water-cooled. Grate firing is mainly used for solid fuels in small- to medium-sized units. The main fuels are bark, straw and municipal solid waste. Grate firing cannot combust fuels where the humidity exceeds 60 %, since the fuel is dried, devolatilized and the remaining carbon is burned as it is transported over the grate. If the fuel humidity is too high, incomplete combustion can occur [43].
Figure 4.1: Illustration of a grate boiler [43].

4.2.2 Bubbling fluidized bed (BFB)

A bubbling fluidized bed is filled with sand particles with diameters ranging from 0.5 to 1.2 mm. Air is introduced through nozzles at the bottom of the bed (see Figure 4.2). When the velocity of the air is sufficient, the bed becomes fluidized, and the combustion takes place in the bed. The bed temperature is typically between 750-900 °C. A general feature of fluidized bed systems is that they are flexible in relation to the homogeneity of the fuel to be fired which makes them suitable for co-firing. A drawback of this combustion technique, is that the fuel must be processed before combustion since the fuel particles must be relatively small in order to enable fluidization [44].
4.2.3 Circulating fluidized bed (CFB)

Compared to a BFB, the sand particles in a circulating fluidized bed are smaller, usually less than 0.5 mm in diameter, and the fluidizing air velocity is higher. As a consequence, the sand is forced upwards out from the bed by the air flow. A cyclone separates the sand from the flue gas and the sand are then recycled back into the bed via a loop seal. Similar to the BFB, the CFB system is fuel-flexible. Another advantage of the CFB is that the carbon burnout efficiency is very high. A drawback of the system is the risk of bed agglomeration if the alkali content in the fuel is too high [46]. A schematic illustration of a CFB system is shown in Figure 4.3.
4.3 Superheater corrosion

The use of cheaper fuels for energy production has resulted in significant material degradation of heat exchanger surfaces. The two foremost areas in boilers where high corrosion rates have been identified are on furnace water walls (discussed in the next section) and on superheater tubes. Estimating tube lifetimes is not an easy task since the chemistry involved is quite complex and several corrosion mechanisms are probably at play simultaneously. The complex chemistry depends on the type of fuel that is combusted. As mentioned earlier, MSW and RWW are highly heterogeneous compared to fossil fuels, with high levels of elements that are known to play a role in the degradation of superheater materials see Figure 4.4.
The tube material is usually steels or alloys with varying levels of chromium. For simplicity the alloys can be divided into categories according to the formed oxide (see Table 4). The first category is low-alloyed steels, with a chromium content that is not sufficiently high to form a Cr-rich oxide, instead these alloys rely on iron oxides as protection. Since the formed oxides Fe$_2$O$_3$ and Fe$_3$O$_4$ contains a high defect density, these oxides are insufficient as protection against rapid oxidation in corrosive environments or at higher temperatures and as a consequence, the temperature limit for these steels is 500 °C. In highly corrosive environments, this temperature has to be kept considerably lower. In more corrosive environments or at higher temperatures, iron/chromium oxide (Fe,Cr)$_2$O$_3$ or Cr$_2$O$_3$ formers are used instead. These can be divided into the classifications in Table 4. The Cr/Fe ratio in the alloy indicates the degree of protectiveness of the oxide. A higher ratio means that if the oxide is depleted in Cr, there is a reservoir left in the steel that can diffuse into the oxide, thus retaining the protective properties of the solid solution of Fe$_2$O$_3$/Cr$_2$O$_3$, as mentioned earlier. In very aggressive environments, a high Cr/Fe ratio in the alloy is required to withstand fast degradation. Therefore, all reactions that consumes Cr from the oxide are considered to be detrimental for the oxide’s protectiveness. During combustion ashes in the flue gas are deposited on top of the oxide (see Figure 4.4). The amount and composition of the deposits vary greatly with the fuel type. Fuels that have high concentrations of alkali and chlorine results in a corrosive deposit in contact with the oxide. As mentioned earlier, there are countermeasures available, including co-combustion and the introduction of additives.
Table 4. Steel classifications with compositions in weight %.

<table>
<thead>
<tr>
<th>Classification</th>
<th>Example</th>
<th>Cr %</th>
<th>Fe %</th>
<th>Ni %</th>
<th>Cr/Fe in alloy</th>
<th>Protective oxide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low alloyed steels</td>
<td>Fe-2.25Cr-1Mo</td>
<td>2</td>
<td>96</td>
<td>0</td>
<td>0.02</td>
<td>Fe₂O₃, Fe₃O₄</td>
</tr>
<tr>
<td>Ferritic stainless steels</td>
<td>X20</td>
<td>12</td>
<td>85</td>
<td>0.5</td>
<td>0.14</td>
<td>(Fe,Cr)₂O₃</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(Fe,Cr)₃O₄</td>
</tr>
<tr>
<td>Austenitic stainless steels</td>
<td>347H</td>
<td>17</td>
<td>68</td>
<td>11</td>
<td>0.25</td>
<td>(Fe,Cr)₂O₃</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(Fe,Cr)₃O₄</td>
</tr>
<tr>
<td>High alloyed austenitic steels</td>
<td>Sanicro 28</td>
<td>27</td>
<td>35</td>
<td>31</td>
<td>0.77</td>
<td>(Fe,Cr)₂O₃</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(Fe,Cr)₃O₄</td>
</tr>
<tr>
<td>Nickel based alloys</td>
<td>Inconel 625</td>
<td>22</td>
<td>5</td>
<td>58</td>
<td>4.4</td>
<td>Cr₂O₃</td>
</tr>
</tbody>
</table>

### 4.4 Boiler water wall corrosion

Compared to the corrosion issues of the superheater region, far less research has been conducted on waterwalls, however, researchers at Åbo Academy [39, 47] and KTH [40] are active in the field. Even though waterwalls consist of steel tubes similar to superheaters, there are differences. Instead of heating steam, waterwalls separate the pressurized water inside the tubes from the hot flue gases in the furnace. The temperature gradient between the flue gas and the water brings the water to a boil and the produced steam is collected in the vapor dome. With a water temperature of about 350 °C, the flue gas temperature is typically 750-1000 °C. A rule of thumb is that the water wall material temperature is up to 50 °C higher than the water temperature, i.e. up to 400 °C [40] (see Figure 4.5).
As a result of the lower material temperature at waterwalls than in superheaters, different alloys (low-alloyed steels) are normally used which in combination with the lower temperature, results in the altered corrosion mechanism that will be discussed in the following section.

### 4.5 Corrosion mechanism

The effect of HCl(g) or Cl\(_2\)(g) on the high temperature corrosion of pure metals [48-53] and alloys [48, 54-65] has been studied. Usually, the corrosive effect of chlorine-containing species, such as HCl(g), Cl\(_2\)(g), and alkali chlorides is explained by the proposed “active oxidation” mechanism [51, 52, 54, 56]. In this mechanism, Cl\(_2\) (formed from HCl through the Deacon process, reaction 4.1) is suggested to be transported through the protective oxide in gaseous form. When Cl\(_2\) reaches the metal/oxide interface, the oxygen partial pressure is low and the formation of transition metal chlorides, e.g. FeCl\(_2\), is hence thermodynamically favored. The transition metal chlorides formed are volatile at high temperatures and are thought to diffuse outward through the scale as gas molecules.

\[
2\text{HCl}(g) + 0.5\text{O}_2(g) \rightarrow \text{Cl}_2(g) + \text{H}_2\text{O}(g) \tag{4.1}
\]

\[
0.5x\text{Cl}_2(g) + \text{Me}(s) \rightarrow \text{MeCl}_x(s) \tag{4.2}
\]

\[
\text{MeCl}_x(s) \rightarrow \text{MeCl}_x(g) \tag{4.3}
\]
The following three reactions are an example of relevant chlorides that can be encountered in the corrosion process of steels. The equilibrium pressures gives an indication of the compound’s propensity to evaporate. At higher temperatures the evaporation increase.

\[
\text{FeCl}_2(\text{s}) \rightarrow \text{FeCl}_2(\text{g}), \quad p_{eq} = 5.13 \times 10^{-7} \text{ bar at 400 °C [66]} \text{ calculated with data from Barin.}
\]

\[
\text{NiCl}_2(\text{s}) \rightarrow \text{NiCl}_2(\text{g}), \quad p_{eq} = 1.32 \times 10^{-8} \text{ bar at 400 °C [66]} \text{ calculated with data from Landolt.}
\]

\[
\text{CrCl}_2(\text{s}) \rightarrow \text{CrCl}_2(\text{g}), \quad p_{eq} = 4.32 \times 10^{-12} \text{ bar at 400 °C [66]} \text{ calculated with data from Landolt.}
\]

The outward diffusion of metal chloride molecules has been suggested to be the rate-limiting step in the reaction mechanism [51, 56]. When the metal chloride molecules reach the oxide/gas interface, they decompose into non-protective porous metal oxide and Cl\(_2\) through reaction with O\(_2\).

\[
y\text{MeCl}_x(\text{g}) + 0.5z\text{O}_2(\text{g}) \rightarrow \text{Me}_y\text{O}_z(\text{s}) + (x*y)/2\text{Cl}_2(\text{g})
\]

(4.4)

The released chlorine gas may then penetrate the oxide again. This process is proposed to account for the porous and poorly adherent scale that is often formed in the presence of chlorine-containing species [51, 52].

An apparent difficulty [67] with the active oxidation mechanism is the transportation of Cl\(_2\)(g) through the oxide. At the metal/oxide interface, where oxygen activity is low, chlorine is assumed to react with the metal to form transition metal chlorides. This implies that the oxide is impermeable to the O\(_2\) molecule while, at the same time, the Cl\(_2\) molecule is transported through it. As illustrated in Figure 4.6, the Cl\(_2\) molecule (396 pm across) is substantially larger than the O\(_2\) molecule (264 pm across). It has not been explained why the oxide would be impermeable to the O\(_2\) molecule but permeable to the larger Cl\(_2\) molecule. Grabke [56] has suggested that chlorine creates fast diffusion paths in the oxide. However, regardless of how the pathways for Cl\(_2\) molecular diffusion are created, they must also be fast diffusion paths for oxygen molecules. If oxygen is able to use these fast diffusion paths, the thermodynamic conditions for metal chloride formation are no longer fulfilled. The outward diffusion of gaseous metal chlorides has similar difficulties. Metal chloride molecules are much larger than Cl\(_2\) and cannot penetrate an oxide that prevents the diffusion of gaseous O\(_2\). Another problem with the active oxidation model is its prediction that chlorides are confined to the metal/oxide.
interface where the oxygen partial pressure is low enough for them to form. Previous studies have reported that chlorine-rich areas are also present in the outer part of the scale [64]. This is peculiar, since according to the mechanism the metal chlorides is supposed to decompose at higher $p(O_2)$. Since studies have shown that metal chlorides are sometimes detected in the outer parts of the scale, the understanding of chlorine-induced high-temperature corrosion of metals is not satisfactory, and a better mechanism is needed.

![Diagram of Cl₂ and O₂ molecules](image.png)

**Figure 4.6:** Comparison of the size of the Cl₂ and O₂ molecules.

Metal chlorides are hygroscopic, and if they are present in the corrosion product layer on boiler alloys, it is likely that the chlorides will be hydrated by the absorption of water. This may have severe consequences for waterwalls and superheaters during the downtime of the boiler. The occurrence of such phases could make the alloy susceptible to aqueous corrosion and/or loss of oxide scale adhesion (see Figure 4.7). For this reason, the material degradation in boilers cannot entirely be explained by high temperature corrosion.
4.5.1 Chromate-forming corrosion

Stainless steels, relying on a chromium-rich corundum-type oxide for protection, have been shown to be susceptible to corrosion in the presence of K$_2$CO$_3$ and alkali chlorides, such as NaCl and KCl. This corrosion attack has been suggested to be initiated by a reaction with the chromium in the oxide, forming chromates [28-33].

\[
xK_2CO_3(s) + (Fe_{1-x}Cr_x)_{2}O_3(s) + 1.5 \times O_2(g) \leftrightarrow xK_2CrO_4(s) + (1-x)Fe_2O_3(s) + xCO_2(4.5)
\]

\[
4xKCl(s) + (Fe_{1-x}Cr_x)_{2}O_3(s) + 1.5xO_2(g) + 2xH_2O(g) \leftrightarrow 2xK_2CrO_4(s) + (1-x)Fe_2O_3(s) + 4xHCl(g) \quad (4.6)
\]

These reactions deplete the oxide of chromium, rendering it iron-rich and non-protective. After these reactions, the material is more susceptible to chlorine attack. The destructive effect of KCl and K$_2$CO$_3$ is illustrated in Figures 4.8-4.10. Figure 4.8 shows the initial stages of exposure before the salt has reacted with the oxide. It can be seen, however, that the salt has begun to diffuse over the surface. In Figure 4.9, the salt has reacted with the protective oxide according to reactions (4.5) and (4.6) resulting in an oxide depleted of chromium through alkali chromate formation. At this point the oxide has reached a chromium level at which the protective properties are lost and the steel suffers from breakaway corrosion (see Figure 4.10). At the point of breakaway corrosion, a thick and non-protective oxide starts to form. The oxide is divided into an outward growing hematite (Fe$_2$O$_3$) and an inward growing spinel-type oxide.
(Fe,Cr,Ni)$_3$O$_4$. Oxygen and chloride ions can readily penetrate the scales, thus, resulting in further oxide growth and the formation of subscale metal chlorides. At this stage, the salt particle is consumed and its original position is marked by an irregular formed oxide.

Figure 4.8: Illustration of the sample at the start of exposure with a protective chromium rich oxide (Cr,Fe)$_2$O$_3$. [68]

Figure 4.9: Illustration of the initiation of breakaway corrosion. The oxide is depleted of chromium due to chromate formation. [68]

Figure 4.10: Illustration of the propagation of breakaway corrosion. [68]
4.6 Waterwall corrosion mechanisms

Molten species in the deposits of waste and biomass-fired boilers have been suggested to cause accelerated corrosion [13, 18, 37, 69-71]. Kofstad [18] has suggested two main reasons for the corrosiveness of liquid phases. First, the liquid itself provides fast transport of ions. Second, a liquid phase provides an electrolyte that can act as a pathway for an ionic charge transfer for an electrochemical attack. The melt can dissolve the otherwise protective oxide, which exposes the metal to the surrounding environment. Both potassium and sodium chloride form eutectic compositions with iron chloride, with melting points between 200 – 600 °C see Table 4 [37]. These melts have been suggested to cause severe corrosion of steels exposed above the melting temperature. In waste-fired boilers the presence of zinc and lead compounds has also been suggested to account for severe corrosion by lowering the melting point of the deposits [70].

Table 5. Melting points of pure salts and mixtures.

<table>
<thead>
<tr>
<th>Salt and mixtures</th>
<th>Melting point [°C]</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZnCl₂</td>
<td>318</td>
</tr>
<tr>
<td>PbCl₂</td>
<td>489</td>
</tr>
<tr>
<td>KCl</td>
<td>772</td>
</tr>
<tr>
<td>NaCl</td>
<td>801</td>
</tr>
<tr>
<td>FeCl₂</td>
<td>673</td>
</tr>
<tr>
<td>KCl-ZnCl₂</td>
<td>230</td>
</tr>
<tr>
<td>KCl-PbCl₂</td>
<td>412</td>
</tr>
<tr>
<td>KCl-FeCl₂</td>
<td>355</td>
</tr>
<tr>
<td>NaCl-ZnCl₂</td>
<td>262</td>
</tr>
<tr>
<td>NaCl-PbCl₂</td>
<td>415</td>
</tr>
<tr>
<td>NaCl-FeCl₂</td>
<td>375</td>
</tr>
</tbody>
</table>

Ruh, et al have shown with thermodynamic calculations that FeCl₂ is soluble in molten KCl–ZnCl₂ near the eutectic compositions, while the solubility of CrCl₃ and NiCl₂ are very limited [67]. In this manner, the outward diffusion of FeCl₂ is favored and diffuses easily to outer part of the salt melt layer. As FeCl₂ reaches the outer part of salt melt, FeCl₂ will be oxidized due to the high p(O₂). The authors have concluded that the solubility of metal chlorides in molten KCl–ZnCl₂ leads to a higher diffusion rate that results in a higher oxidation rate. Therefore, the solubility of metal chlorides in the molten salts influences the extent of corrosion of each metal. A proposed mechanism for the chloride melt-induced dissolution of the steel can be seen in Figure 4.11 [67].
Figure 4.11: Model of chloride melt induced high temperature corrosion, displaying the transport of gas species and reactions that results in iron oxide and iron chloride [67].

Bankiewicz et al have in several papers studied the corrosive effect of ZnCl₂ and PbCl₂. The results indicate that PbCl₂ is comparatively more corrosive compared to ZnCl₂, with formation of an uneven oxide and signs of internal attack, already below the melting point of the lead chloride [47]. This observation is in agreement with the findings in this thesis, (even though the amount of PbCl₂ was far less) in which a localized attack with an uneven oxide was observed on T22 in the presence of PbCl₂ at 400 °C. Furthermore, the findings in this study is in line with [47], in which PbCl₂ seems to be more corrosive than ZnCl₂. Studies with salt mixtures containing ZnCl₂ and PbCl₂ have also been conducted by Bankiewicz et al. Those authors found a correlation between increased amounts of ZnCl₂ in the mixture and accelerated corrosion when T22 was exposed to a mixture of ZnCl₂ – K₂SO₄ [72]. When T22 was exposed in the presence of PbCl₂ containing salt mixtures (PbCl₂ with KCl or K₂SO₄), a severe corrosion attack was observed already below the melting point of the mixture [73]. This indicates that a melt is not a prerequisite for elevated corrosion rates. In comparison in the presence of a PbCl₂ - K₂SO₄ mixture at 400 °C, the formed corrosion layer was twice as thick as the one formed in the presence of a ZnCl₂ - K₂SO₄ mixture [73].

Schaal et al [12] have studied the influence of ZnCl₂ and PbCl₂ at 450 °C on the syntetic ashes that are representative for waste combustion. The tested materials were 16Mo3 and Inconel 625. Contrary to Bankiewicz et al, they detected a higher mass loss in the presence of ZnCl₂ than in the presence of PbCl₂. This was explained by a higher percentage of molten phase in the ZnCl₂ containing ashes. However, both studies, detected the highest corrosion rates in ashes in which ZnCl₂ and PbCl₂ were present simultaneously.
Lu W.M et al have investigated the corrosion of five commercial steels with a deposit of ZnCl$_2$–KCl under reducing conditions at 400 and 500 °C (material temperature) [74]. For all the investigated materials, the corrosion rates were significantly enhanced in the presence of ZnCl$_2$–KCl compared to salt free exposures. Consequently, the scales grown beneath the ZnCl$_2$–KCl deposit differed considerably from those grown in the absence of salts. The corrosion attack in the former case was characterized by fast growth and the scales were porous and poorly adherent. Accelerated corrosion beneath the ZnCl$_2$–KCl deposit was observed for all materials during the initial stage. This was proposed to be an effect of the presence of chlorine induced by the ZnCl$_2$–KCl deposit. Even though the presence of the ZnCl$_2$–KCl deposits caused an accelerated corrosion of all materials throughout the corrosion tests, the corrosion rate decreased after about 3 hours. This was suggested to be due to the continuous volatilization of ZnCl$_2$, and thereby, the increase in the melting temperature of the deposit. The melting point of the ZnCl$_2$–KCl salt mixture prepared for the tests was about 250 °C, well below the exposure temperatures of 400 and 500 °C. Consequently, the presence of the melt makes the establishment of a protective scale more difficult. The authors suggest that this is mainly caused by the presence of the Cl$^-$ anions in the eutectic ZnCl$_2$–KCl system. The molten salt partly dissolves the iron oxide scale and diffuses through it, degrading its protective properties. Thus, the corrosion rate is expected to be rather fast during the initial stage, in good agreement with the experimental kinetic curves. In oxidizing environments, as corrosion proceeds, ZnCl$_2$ is consumed continuously due to its volatilization as well as to its participation in the following reactions (4.7-4.9):

- \[ 2\text{ZnCl}_2 + 2\text{Fe}_2\text{O}_3 + 2\text{H}_2\text{O}(g) \leftrightarrow 2\text{ZnFe}_2\text{O}_4 + 4\text{HCl}(g) \]  
\[ \Delta G = 27 \text{ kJ}, \ p_{eq}(\text{HCl}) = 0.06 \text{ bar} \] [66] calculated with data from Knacke.

- \[ \text{ZnCl}_2 + \text{H}_2\text{O}(g) \leftrightarrow \text{ZnO} + 2\text{HCl}(g) \]  
\[ \Delta G = 46 \text{ kJ}, \ p_{eq}(\text{HCl}) = 9.7\times10^{-3} \text{ bar} \] [66] calculated with data from Landolt.

- \[ \text{ZnO} + \text{Fe}_2\text{O}_3 \leftrightarrow \text{ZnFe}_2\text{O}_4 \]  
\[ \Delta G = -20 \text{ kJ} \] [66] calculated with data from Knacke.
According to these reactions, KCl is enriched in the molten salt mixture, producing a rise in the melting point of the KCl–ZnCl₂ mixture, according to the phase-stability diagram of the KCl–ZnCl₂ system (see Figure 4.12). Within the work of this thesis, ZnCl₂ was exposed on gold. It was noted that after 15 min of exposure, all ZnCl₂ had either evaporated or reacted into ZnO, which under oxidizing conditions will deplete the mixture in ZnCl₂ and shift the composition towards KCl. This fast evaporation of ZnCl₂ has also been reported by Lehmosto et.al [75], in which 30% of the salt had evaporated after 3 hours at 400 °C. Bankiewicz et.al [47] have performed thermodynamic calculations regarding saturation pressure of ZnCl₂ over a range of temperatures. It was noted that, above 350 °C a rapid increase in the saturation pressure commence, which theoretically can explain the fast evaporation.

![Phase diagram of the KCl-ZnCl₂ system](image)

Figure 4.12 Phase diagram of the KCl-ZnCl₂ system calculated with Factsage [76].

Under this condition, the salt mixture will gradually become solid, unless it is supplied with new ZnCl₂ and the corrosion rates decrease during the later stages.

Folkeson et al have reported that the acceleration of the corrosion of Fe-2.25Cr-1Mo steel caused by KCl is very strong at 400 °C [77]. The resulting thick scale consists of several oxide layers, with iron chloride accumulating at the scale/metal interface. The bottom oxide layer in
that study was formed by the inward diffusion of oxygen and consisted of iron chromium spinel oxide. The upper part of the scale was chromium free and made up of a magnetite layer in the middle and a hematite layer on top.

In contradiction to the active oxidation model, a new mechanism is proposed to explain the effect of KCl. Jonsson et al have shown with ESEM in-situ oxidation that the initial corrosion of a Fe-2.25Cr-1Mo steel deposited with KCl starts at approximately 355 °C [78], which is the eutectic temperature for the KCl-FeCl₂ system (see Figure 4.13). At this temperature, it is argued that a continuous liquid surface film is formed, which causes the KCl from the large particles to spread over the surface due to a concentration gradient. Simultaneously, FeCl₂ diffuses towards the KCl particles. It is proposed that KCl reacts with O₂ and water on the scale surface forming potassium hydroxide. The chloride ions released in this way diffuse towards the metal via the oxide grain boundaries, forming iron chloride. The increase in oxidation rate caused by KCl is attributed to the presence of iron chloride at the oxide grain boundaries, the iron chloride facilitating the grain boundary transport of both iron ions and oxygen ions through the scale. (Compare this finding with the active oxidation mechanism, in which gas is assumed to diffuse through the scale.) Additionally, the buckling of the scale and the poor scale adhesion are also attributed to the presence of iron chloride.

![Phase diagram of the KCl-FeCl₂ system calculated with Factsage](image)

Figure 4.13 Phase diagram of the KCl-FeCl₂ system calculated with Factsage [76].
4.6.1 Corrosion mitigating techniques

Since the shift from fossil fuels has resulted in very high corrosion rates, there have been attempts to find means to mitigate this material wastage. The more successful techniques have been the methods, which in different ways, raise the levels of SO$_2$ and/or SO$_3$ in the flue gas. The sulfur oxides can then react with chlorides in the flue gas and consequently, convert these into less corrosive sulfates. This can be achieved with various methods. The first method is to co-combust fuels with a high chlorine content with a different fuel stream that is known to contain relatively high amounts of sulfur. Examples of such fuels are sewage sludge or peat, both of which have been shown to lower the levels of alkali chlorides in the flue gas with lower corrosion rates as a result [79, 80]. A different method that has shown promising results, has been developed by Vattenfall, the so called ChlorOut$^\text{®}$ technique. With this method ammonium sulfate ((NH$_4$)$_2$SO$_4$) is sprayed into the flue gas, thereby converting the chlorides into sulfates [81, 82]. The method that is included in this thesis is a technique in which the sulfur in the fuel, which is released during combustion, is extracted from the flue gas in a wet scrubber. The scrubber contains hydrogen peroxide (H$_2$O$_2$) which reacts with the SO$_2$, resulting in sulfuric acid (H$_2$SO$_4$) which is recirculated back into the furnace. In the furnace, the sulfuric acid decomposes into SO$_2$, thus, closing the loop. The advantage of this technique is that no externally source of sulfur is needed, instead, the method utilizes existing sulfur in the fuel. This method is further explained in paper V.

The most commonly used method to mitigate corrosion at the waterwalls is to weld or coat the original wall tubes with a nickel based alloy. These materials are, however, very expensive compared to ferritic steels, which is why new less expensive preventative techniques are needed.
5 Experimental techniques

5.1 Laboratory exposures

5.1.1 Sample preparation

The alloy studied was the Fe-2.25Cr-1Mo steel T22 (see Table 2.1 for composition). The sample dimensions were 1.5*1.5*0.2 cm. A hole (1.5 mm in diameter) was drilled for handling. The surface area was 5.56 cm². The corresponding surface area of the TG-exposed samples were 2.27 cm². Prior to exposure, the samples were ground to a 320 mesh in water and polished with 9, 3 and 1µm diamond solutions to a mirror-like appearance. The samples were degreased in acetone and ethanol using ultrasonic agitation. Gravimetric measurements were made using a Sartorius balance with microgram resolution.

5.1.2 Experimental set-up

Isothermal exposures were performed in a horizontal silica tube furnace (see Figure 5.1). The atmosphere consisted of N₂ with 5 % O₂ with 40 % or 20 % H₂O. The linear flow rate was 3 cm/s. All parts of the system were kept above 100 °C to prevent the condensation of water. The samples were mounted three at a time on an alumina sample holder. The samples were positioned parallel to the gas flow. The temperature was 400 and 600 ±1 °C. The samples were exposed for 24 hours.

Figure 5.1: Schematic of the tube furnace setup.
Isothermal exposures were also performed in-situ in a Setaram Setsys Evolution® system. The sample was suspended vertically from the scale as shown in Figure 5.2. This arrangement makes it possible to record the mass change of the sample in real time. A counter flow of inert N₂ was introduced from above the sample to protect the scale from corrosive gases. At the bottom of the system, the gases that came in contact with the sample were introduced. This setup allows the system to be run with varying levels of H₂O(g), HCl(g), as well as SO₂(g). In order to check that the desired concentration of HCl(g) was present during exposure, the outlet from the furnace was bubbled in 1 L of water for 1 hour. The water was then analyzed with Ion chromatography (IC) and the resulting amount of Cl⁻ matched the calculated theoretical value that is expected in 1 L after 1 hour.

Figure 5.2: Schematic of the in-situ thermo gravimetric setup.

5.2 Field exposures

Field exposures were carried out in Renovas waste-fired grate boiler in Sävenäs Gothenburg, Sweden, a schematic of the plant is shown in Figure 5.4. Corrosion probes were exposed to the flue gas in the superheater region. Three different materials were studied, the low-alloyed 16Mo3, the high-alloyed Sanicro 28 and the nickel based Inconel 625 that was welded onto a 16Mo3 tube (for compositions see Table 1). The material temperature was regulated with
compressed air. The material temperature was logged for each sample. One of the sample temperatures was chosen to control the cooling. The thermocouples were directed towards the gas stream, referred to as windward (see Figure 5.3). The samples were exposed for 24 hours and 1000 hours. Before exposure, the samples were cleaned and degreased in ethanol and acetone using ultrasonic agitation. After exposure, the samples were stored in desiccators over P₂O₅. The samples exposed for 1000 hours were analyzed to determine material loss (as a measurement of corrosion).

Figure 5.3: Corrosion probe used in the Sulfur recirculation exposures.

Figure 5.4: Schematic overview of the Sulfur recirculation system and boiler.
6 Analytical techniques

A wide range of analytical techniques can be used to evaluate the exposed samples. All available techniques have advantages and drawbacks and a single technique is rarely able to provide the complete story of a corrosion attack. Therefore it is important to couple the results from different analytical tools in order to obtain a more complete picture. The analytical techniques used for high temperature corrosion can be divided into three categories: imaging, elemental analyzing and phase detection techniques. Imaging techniques includes different types of microscopes such as Scanning Electron Microscopes (SEM), Transmission Electron Microscopy (TEM) and Optical Microscopy (OM). Both SEM and TEM are often used together with a technique from the second type namely Energy Dispersive X-rays (EDX), which is used to determine the elemental composition on the sample. Another technique is IC (Ion Chromatography), used to quantify the amount of water soluble ions on a sample. These techniques gives information about the elemental distribution, however knowing only the distribution is not sufficient to explain a corrosion attack. The aim is to describe oxide growth, which requires knowledge about the crystalline phases. These are detected with diffraction techniques, such as X-ray Diffraction (XRD) and electron diffraction in the TEM.

6.1 Imaging techniques and elemental analysis

6.1.1 Electron microscopy

Electron microscopy was used to investigate the microstructure of a corrosion attack in detail. This method offers high magnification imaging at high resolution by scanning a focused electron beam over the sample surface. It also offers information about the chemical composition of the imaged area. Thus, electron microscopy is an essential tool for the study of high-temperature corrosion.
Figure 6.1: Some of the different processes that occur when atoms are bombarded with electrons [83].

As the atoms in the sample are hit by the electrons, several different interactions take place that generate various types of signals (see Figure 6.1). All these processes occur throughout the entire volume of interaction but, because the specific energy of the emitted electrons or photons differs, the depth from which they can escape also differs. This is illustrated in Figure 6.2.

Figure 6.2: Illustration of the interaction volume (the size corresponds to an acceleration voltage of 20 kV). A signal with more energy can escape from larger depths [83].
Secondary electrons are generated when the primary electron beam causes ionization of inner core electrons in the surface atoms. Because of their low energy, secondary electrons may only escape from a small volume, 5-10 nm from the surface [83]. They are, therefore, well suited for high resolution imaging.

Backscatter electrons are generated by the scattering of the primary electron beam by the sample atoms. Consequently, the yield of backscatter signal depends on the atomic number of the scattering atoms, heavy elements appearing brighter. Backscatter electrons offer both a topographical and a compositional contrast. However, because of their higher energy (>50 eV) [83], they escape from a larger volume than secondary electrons, and as a result, the resolution is worse (compare images a) and b) in Figure 6.3).

Figure 6.3: An example of the difference between a) back scattered electron image and b) secondary electron image of the same area

Chemical information about the surface atoms is provided by characteristic X-rays and Auger electrons. These types of signals are generated in similar ways, starting with the ionization of an inner core electron. The resulting ion is, thus, left in an excited state. An electron in a higher orbital can relax to the vacancy left by the ionization. The energy of the system can be lowered by the emission of either an X-ray photon or a loosely bound outer shell Auger electron. The specific energies of both X-ray photons and Auger electrons are highly dependent on the electronic structure of the element from which they are emitted and thus, offer chemical
information. However, characteristic X-rays have higher energy and escape from the entire volume of interaction (as illustrated in Figure 6.2).

6.1.2 Scanning electron microscopy, SEM

The instrument used in this study was a FEI Quanta ESEM 200. It was equipped with a Field Emission Gun, providing excellent spatial resolution. The instrument was also equipped with an Oxford Inca EDX (Energy Dispersive X-ray) system. The EDX system enables chemical analysis through the detection and quantification of the characteristic X-rays emitted by a sample. The accuracy of the EDX analysis was tested by performing a point analysis in the steel below the oxide scale. By comparing the result with the theoretical composition of the steel the precision of the analysis is confirmed. The instrument was operated at high vacuum, with an acceleration voltage between 10 and 25 kV.

6.1.3 Broad ion beam, BIB

When a sample is covered with thick porous corrosion products, manually polished cross sections generally are challenging to perform. This is due to the fact that polishing may pull out parts of the corrosion layer and redeposit it somewhere else, or if water sensitive corrosion products are present (like metal chlorides). Instead, ion-etched cross sections can be used, which is considerably less destructive. To obtain wide ion etched cross sections Broad ion beam (BIB) milling can be used. The BIB uses a broad argon ion beam for milling, which makes cross sections in the range 500-1000µm possible. The BIB cross sections in this study were produced using a LeicaTIC3X BIB instrument. The three ion guns were operated at 6.5 kV for a duration of 7 hours. Prior to milling, a 0.5mm silicon wafer was glued on top of the sample with Loctite® 415. After hardening, the sample was cut using a low-speed saw and a consumable protective screen was positioned on the sample with silver glue to render a smooth cross section (see Figure 6.4).
Figure 6.4: SEM image overview of a BIB cross section with high magnification to the right.

6.1.4 Focused Ion Beam, FIB/SEM workstation

The BIB is a very coarse milling technique in the sense that it generates wide cross sections, but there is limited possibilities to pinpoint features. When a certain area on a sample is interesting, a technique called Focused Ion Beam (FIB) is used instead. A 10-50µm wide cross section is then prepared by sputtering Ga ions through the oxide scale (see Figure 6.5).

A FEI Versa was used to study cross sections of selected features. The cross sections in the study were prepared with a geometry that enabled SEM/EDX analysis. Since the ion beam has an incidence angle of 52° relative to the sample surface, the samples were tilted 7° in order to achieve a 45° cross section relative to the sample surface. The sample was then transferred to the FEI Quanta ESEM 200 where a 45° pre-tilted sample holder was used to achieve a 90° angle between the electron beam and the cross section (see the right image in Figure 6.5).
Transmission Electron Microscopy (TEM)

The transmission electron microscopy technique enables the investigation of microstructural features through high resolution imaging. Contrary to an SEM in which the electrons are scattered back by the sample, in a TEM, most of the electrons pass the sample that must be very thin (20-200 nm) and the accelerating voltage high, typically 200-300 kV. Lift-out samples (from exposed samples) for TEM analysis has been performed in the FIB/SEM workstation. All sample preparation and TEM analysis has been performed by Dr M. Sattari and A. Persdotter.

In this study the TEM was operated in Scanning Transmission Electron Microscope (STEM) mode, which was used for High Annular Dark Field (HAADF) and Bright Field (BF) imaging. In the HAADF mode, it is possible to detect the variation in atomic number over a sample cross section (the Z- contrast), where heavier elements appear brighter and comparatively lighter elements attain a darker contrast. In the BF mode the weakening of the transmitted beam (by its interaction with the sample) is imaged. Thick areas and areas with heavier elements appear with a darker contrast. On the contrary, voids, due to less interaction with the beam, appear bright.
6.2 Phase detection techniques

6.2.1 X-Ray diffraction, XRD

The analytical techniques mentioned above give the elemental composition of a sample surface, but they cannot determine the form in which the elements are associated, i.e. which crystalline compounds are present. Crystalline chemical compounds can be identified by means of X-ray diffraction. In this study Cu Kα radiation (λ = 1.5418 Å) was used to be diffracted in the crystalline phases in the sample. Diffraction occurs when the X-rays scattered by the atoms in the crystal interfere constructively. Bragg’s law (illustrated in Figure 6.6) links the angles, θ, at which the scattered X-rays interfere constructively with the spacing, dhkl, between the hkl crystal planes for a given wavelength, λ (n is an integer):

\[2d_{hkl} \sin \theta = n\lambda \text{ (Bragg’s law)} \quad (5.1)\]

Figure 6.6: Illustration of Bragg’s law for diffraction with crystal planes.

Several different set-ups with different detector angles and X-ray source angles are used in diffraction. The most useful set-up for studying the corrosion of metal surfaces is the Grazing Incidence technique (see Figure 6.7). In this set-up the X-ray source is fixed at a low angle of incidence to minimize the depth of interaction, and thus also the bulk contribution. The detector scans the θ range to map diffraction intensity.
Figure 6.7: Set-up for grazing incidence X-ray diffraction.

The diffraction data from the samples were compared to a database of known crystalline phases to identify the compounds present on the sample. The requirements of the sample for identification are that the corrosion products are crystalline and that they are included in the database. As the data for one compound may overlap the data of another, it is crucial to know which elements are present in the corrosion products. In this manner, the different analytical techniques complement each other.

In this study a Siemens D5000 powder diffractometer, equipped with a Göbel mirror was used. Analyses were performed in grazing incidence mode, with an angle of incidence of 0.3 – 1 ° for sample coupons from the lab exposures and 5 – 10 ° for samples from field exposures. The measuring range was 10 ° < θ < 70 °.

6.3 Elemental analyzing of water soluble ions

6.3.1 Ion chromatography, IC

Ion chromatography is a technique that is used to determine the concentration of water soluble ions in a solution. The solution is separated in an analytical column in which ions are separated according to size and charge. The concentration of each ion is determined after the column where conductivity measurements are made and compared with standards of known ion concentrations (see Figure 6.8). To determine the amount of water soluble chlorides and sulfates after exposure the samples were analyzed with a Dionex ICS-90 with an IonPac AS4A-SC analytic column. The samples were leached in MilliQ water under ultrasonic agitation for 10 + 10 minutes. The flowrate was 2 mL/min and 1.8 mM Na₂CO₃/ 1.7 mM NaHCO₃ was used as eluent fluid.
6.4 Material loss measurement

Samples exposed for 1000 hours in the field study were analyzed to determine material loss. Material loss is a measure of the thickness of the sample prior to and after exposure, i.e. it is a quantification of the corrosion rate. The thickness of the sample rings were measured prior to exposure at 8 evenly distributed positions around the ring with a micrometer screw. After exposure, the samples were cut in two with a precision saw to obtain a cross section of each sample ring. With a high resolution camera each sample ring was photographed. The images were then digitalized using the Engauge™ software. The thickness of the sample rings were measured prior to exposure on 8 evenly distributed positions around the ring with a micrometer screw. After exposure, the samples were cut in two with a precision saw to obtain a cross section of each sample ring. In order to measure the material losses, each sample ring was photographed with a high-resolution camera. The images were then digitalized using the Engauge™ software. The sample thickness was measured at approximately 400 individual positions (see Figure 6.9). Thus, enabling a more realistic evaluation of the material loss with superior accuracy compared to the previously used method in which 8 positions around the sample ring was measured with a micrometer screw.
Figure 6.9: A section of the digitized photo of the exposed corrosion probe sample ring cross section.
7 Results and Discussion

7.1 Laboratory exposures simulating waterwall conditions

In this part of the thesis, the laboratory results will be discussed with the aim of increasing the understanding of waterwall corrosion. The first section will discuss the corrosive effect of pure chloride salts on Fe-2.25Cr-1Mo (T22) (composition is given in Table 1) as well as compare the effect of alkali chlorides with the effect of heavy metal chlorides. This is of importance since both varieties are commonly detected in waterwall deposits, and it is not completely understood which of these species that are the main contributor to the corrosion.

7.1.1 Exposures in presence of pure chlorides

In order to determine the corrosive effect of chloride-containing salts towards T22 at 400 °C, it is first important to investigate the T22 alloy in the absence of salt. As reference, samples exposed without salt were used for comparison. The mass gain of the reference can be seen in Figure 7.1.

![Figure 7.1: Mass change of Fe-2.25Cr-1Mo in the presence of chlorides at 400 °C with 20 % H₂O.](image)
The plan view and cross section are displayed in Figure 7.2. The surface morphology is homogeneous where an even smooth oxide with whiskers covers the surface. The XRD analysis of the surface in Table 6 detected $\text{M}_3\text{O}_4$, $\text{Fe}_3\text{O}_4$ and Fe. The detection of the substrate (as Fe), indicates that the oxide is relatively thin.

Table 6. Crystalline phases detected with XRD analysis of Fe-2.25Cr-1Mo exposed 24 hours with chloride salts with 20 % $\text{H}_2\text{O}$ at 400 °C.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\text{M}_3\text{O}_4$</th>
<th>$\text{Fe}_2\text{O}_3$</th>
<th>ZnO</th>
<th>ZnCl₂</th>
<th>NaCl</th>
<th>KCl</th>
<th>PbCl₂</th>
<th>PbO</th>
<th>Fe</th>
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<tr>
<td>Reference</td>
<td>X</td>
<td>X</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>X</td>
</tr>
<tr>
<td>KCl</td>
<td>X</td>
<td>X</td>
<td>-</td>
<td>-</td>
<td>X</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>NaCl</td>
<td>X</td>
<td>X</td>
<td>-</td>
<td>X</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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</tr>
<tr>
<td>PbCl₂</td>
<td>X</td>
<td>X</td>
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<td>X</td>
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<tr>
<td>ZnCl₂</td>
<td>X</td>
<td>X</td>
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<td>-</td>
<td>X</td>
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</tbody>
</table>

The cross section in Figure 7.2 and the TEM analysis in Figure 7.19 shows that the scale is thin (approximately 1 µm), well adherent to the steel and can be divided into three parts as follows: from the bottom up ($\text{Fe},\text{Cr})_3\text{O}_4$, $\text{Fe}_3\text{O}_4$ with an outer $\text{Fe}_2\text{O}_3$. Additional growth of the oxide occurs by the formation of $\text{Fe}^{2+}$ at the steel/oxide interface see reaction (7.1), simultaneously oxygen is reduced at the oxide/gas interface see reaction (7.2). The outward growing part of the oxide is formed by the outwards diffusion of $\text{Fe}^{2+}$ and the inward part by the diffusion of $\text{O}^{2-}$.

$$\text{Fe}(s) \rightarrow \text{Fe}^{2+} + 2e^- \quad (7.1)$$

$$\frac{1}{2}\text{O}_2(g) + 2e^- \rightarrow \text{O}^{2-} \quad (7.2)$$
Seemingly this steel is able to perform well in this environment during 24 hours at 400 °C without the presence of chlorides. However, what happens with the performance of the steel when chloride-containing species are introduced?

In all exposure cases with chlorine containing species present, the mass gain of T22 increased greatly compared to the reference. The mass gain of the T22 in the presence of 0.1 mg/cm² alkali and 0.185 mg/cm² heavy metal chlorides is plotted in Figure 7.1. 0.185 mg/cm² PbCl₂ was chosen to obtain equal moles of chlorine in both exposure cases. The highest mass gain was observed in the presence of KCl. PbCl₂ and NaCl resulted in similar mass gains and ZnCl₂ in the lowest. A comparison of the surface morphologies can be seen in Figure 7.3.
Compared to the reference sample the morphology is considerably rougher in the presence of KCl with some sporadically spread areas of smooth oxide (see Figure 7.3 a)). KCl particles overgrown with oxide are visible in the image (an image of an unexposed sample with applied KCl can be seen in Figure 4 in paper II). The reason for the high oxidation rate of the KCl coated T22 has been reported by Folkeson et al. [78] to be due to the formation of a liquid thin film of the system KCl/FeCl$_2$ at 355°C, (see Figure 4.13), which increases the diffusion of ions over the surface. It has also been proposed that FeCl$_2$ is located at oxide grain boundaries thereby facilitating the transport of ions through the scale suggested by Jonsson et al. [77]. The reaction path where FeCl$_2$ is formed is suggested to be the following. KCl is assumed to decompose according to reaction (7.3).
2KCl(s) + H₂O(g) + ½O₂(g) + 2e⁻ → 2KOH(ads) + 2Cl⁻(ads)  \hspace{1cm} (7.3)

Simultaneously, iron is oxidized at the oxide/steel interface according to reaction (7.4).

\[ \text{Fe(s)} \rightarrow \text{Fe}^{2+} + 2\text{e}^- \hspace{1cm} (7.4) \]

The sum reaction thus becomes:

\[ 2\text{KCl(s)} + \text{H}_2\text{O(g)} + ½\text{O}_2\text{(g)} + \text{Fe(s)} \rightarrow 2\text{KOH(ads)} + \text{FeCl}_2\text{(s)} \hspace{1cm} (7.5) \]

Since FeCl₂ is thermodynamically unstable at the oxide scale/atmosphere interface it readily decomposes according to reaction (7.6). This can result in the formation of porous poorly protective Fe₂O₃ on the surface.

\[ 2\text{FeCl}_2\text{(s)} + ½\text{O}_2\text{(g)} + 2\text{H}_2\text{O(g)} \rightarrow \text{Fe}_2\text{O}_3\text{(s)} + 4\text{HCl(g)} \hspace{1cm} (7.6) \]

\[ \Delta G= -209 \text{kJ at 400 °C} \text{ [66]} \] calculated with data from Landolt.

It could be argued that reaction (7.7) competes with reaction (7.6) in the formation of Fe₂O₃(s), however, Cl⁻ has a lower charge than O²⁻ which is suggested to increase the mobility of Cl⁻ inwards along the oxide grain boundaries.

\[ 2\text{Fe(s)} + 3½\text{O}_2\text{(g)} \rightarrow \text{Fe}_2\text{O}_3\text{(s)} \hspace{1cm} (7.7) \]

\[ \Delta G= -641.4 \text{kJ at 400 °C} \text{ [66]} \] calculated with data from Barin.

The severe corrosion attack which was observed in the presence of KCl, may be attributed to the presence of iron chloride. Either at the steel/oxide interface, where the scale adhesion can be affected in a detrimental way, or in the oxide grain boundaries, where the presence of chlorine is expected to increase the diffusion rate of ions. Consequently, resulting in a thick scale and material wastage.

The attack in the presence of NaCl is similar to the presence of KCl on Fe-2.25Cr-1Mo, which results in rough oxides and chloride particles partially overgrown with oxide, as is illustrated in
Figure 7.3. There is also a eutectic composition in the NaCl/FeCl\(_2\) system with the formation of a liquid film at 375 °C. This causes NaCl to spread away from the chloride particles and induces a widespread attack all over the surface. Compared to KCl, more unreacted salt particles remain on the surface in the case of NaCl, which can be seen as dark grey structures in Figure 7.3. The comparatively greater stability of NaCl is also indicated by the IC analysis in Figure 7.4. As shown, more Cl\(^-\) remains on the surface in the case of NaCl compared to KCl (vapor pressures of KCl(s) and NaCl(s) has been calculated in paper I). Furthermore, the HCl equilibrium vapor pressure is higher in reaction (7.8) than in reaction (7.9), which is part of the reason why the NaCl particles is relatively unaffected after exposure compared to the KCl particles. Another factor that may explain the higher stability of NaCl than KCl, is the difference in eutectic temperature for the systems KCl/FeCl\(_2\) and NaCl/FeCl\(_2\) which is 355 °C and 375 °C respectively. Both of these factors may explain the comparatively higher stability of NaCl.

\[
\text{KCl(s)} + \text{H}_2\text{O(g)} \leftrightarrow \text{KOH(l)} + \text{HCl(g)} \quad (7.8) \\
\Delta G = +118 \text{ kJ, } p(\text{HCl}) = 2.8\times10^{-10} \text{ bar at 400°C [66]} \text{ calculated with data from Barin.}
\]

\[
\text{NaCl(s)} + \text{H}_2\text{O(g)} \leftrightarrow \text{NaOH(l)} + \text{HCl(g)} \quad (7.9) \\
\Delta G = +138 \text{ kJ, } p(\text{HCl}) = 8.6\times10^{-12} \text{ bar at 400°C [66]} \text{ calculated with data from Barin.}
\]

Figure 7.4: IC results of remaining Cl\(^-\) on Fe-2.25Cr-1Mo after 24 hours in the presence of KCl, NaCl or ZnCl\(_2\) at 400 °C (PbCl\(_2\) not analyzed).
The corrosion attack in the presence of ZnCl\(_2\) and PbCl\(_2\) is similar in the sense that no widespread attack all over the surface is detected as in the presence of KCl and NaCl. Instead, the thickest part of the oxide scale is located underneath or in the vicinity of the original position of the chlorides prior to exposure. This is clearly seen in Figure 7.3 c) where an interface between the bright Zn-rich area and the darker Zn deficient is visible. The reason for the local corrosion can be attributed to the fast reaction of ZnCl\(_2\) into ZnO and ZnFe\(_2\)O\(_4\) according to reaction 7.10 and 7.11.

\[
\text{ZnCl}_2(s) + \text{H}_2\text{O(g)} \leftrightarrow \text{ZnO(s)} + 2\text{HCl(g)} \quad (7.10)
\]
\[\Delta G^\circ = +47 \text{ kJ}, \ p(\text{HCl}) = 9.7*10^{-3} \text{ bar at } 400 \ ^\circ\text{C} \ [66] \text{ calculated with data from Landolt.}
\]

\[
\text{ZnO(s)} + \text{Fe}_2\text{O}_3(s) \leftrightarrow \text{ZnFe}_2\text{O}_4(s) \quad (7.11)
\]
\[\Delta G^\circ = -20 \text{ kJ}, \text{ at } 400^\circ\text{C}[66] \text{ calculated with data from Landolt.}
\]

In addition to the reactions above, the ZnCl\(_2\) has a high vapor pressure at 400 °C (see reaction 7.12). Because of these reactions, the time of duration on the steel surface for ZnCl\(_2\) compared to the alkali chlorides is substantially shorter. Accordingly, ZnCl\(_2\) is not detected in the XRD analysis in Table 6 and low amounts of Cl\(^-\) are detected with IC in Figure 7.4. This might be the reason that the lowest mass gain is detected in the presence of ZnCl\(_2\). Large amounts of chlorine leaves the sample during the early stages of exposure and thus do not influence the corrosion rate. Furthermore, ZnCl\(_2\) is the only chloride that has a melting temperature below the exposure temperature. However, no signs of molten phases were detected after exposure, possibly due to the fast transformation of ZnCl\(_2\) into ZnO. With larger amounts of salt, melt formation might be more prevalent, as reported by [47].

\[
\text{ZnCl}_2(s) \rightarrow \text{ZnCl}_2(g) \quad (7.12)
\]
\[p_{eq}(\text{ZnCl}_2) = 9.78*10^{-2} \text{ bar}, \text{ at } 400 \ ^\circ\text{C} \ [76]
\]

Similarly the PbCl\(_2\) in Figure 7.3 d) has reacted into PbO, which is seen as the bright structures in the image. No diffraction peaks from PbCl\(_2\) can be detected with XRD in Table 6. The decomposition and evaporation occurs according to reactions 7.13 and 7.14. The vapor pressure is lower compared to ZnCl\(_2\) which means that more chlorine is present for a longer duration.
\[ \text{PbCl}_2(s) \rightarrow \text{PbCl}_2(g) \quad (7.13) \]

\[ p(\text{PbCl}_2) = 5.7 \times 10^{-6} \text{ bar} \quad [76] \]

The \( \Delta G \) values of the decomposition reactions of KCl, NaCl and PbCl\(_2\) are in the same range, however, the partial pressure of HCl is orders of magnitude higher in reaction 7.14. This means that the reaction into PbO is more likely, especially in a system with gas flow. The formed HCl is transported away from the sample, thus the reaction is shifted to the right, which could be the reason that KCl and NaCl are detected with XRD after exposure and PbCl\(_2\) is not.

\[ \text{PbCl}_2(s) + \text{H}_2\text{O}(g) \leftrightarrow \text{PbO}(s) + 2\text{HCl}(g) \quad (7.14) \]

\[ \Delta G = +120 \text{ kJ} \quad [66] \text{ calculated with data from Barin,} \]

\[ p_{eq}(\text{HCl}) = 1.5 \times 10^{-5} \text{ bar at } 400 \text{ °C} \quad [76] \]

As mentioned earlier, there exists an eutectic composition in the KCl/FeCl\(_2\) and NaCl/FeCl\(_2\) system with melting points below the exposure temperature. In the PbCl\(_2\)/FeCl\(_2\) system, the eutectic composition has a melting point above 400 °C, which could explain why widespread attacks are detected in the presence of alkali chlorides, while the PbCl\(_2\) remains in its original position, where it induces local corrosion. This is clearly seen in Figure 7.5, where large amounts of FeCl\(_2\) is located at the steel/scale interface.

By comparing the surface morphology of a sample, it is possible to get an indication of the corrosion attack. However, by observing cross sections of the exposed samples, not only can the oxide scale thickness be measured, but also the porosity and scale adhesion is seen.

The widespread nature of the attack in the presence of KCl is clearly shown in Figure 7.5 a). In the figure, a thick layered oxide with poor adhesion has formed throughout the KCl exposed sample. In contrast to the widespread attack in the presence of KCl, the localized attack of PbCl\(_2\) can be seen in image c) where a layered oxide structure is located in correlation with the original position of the PbCl\(_2\). The dark spots, visible at the interface between the steel and the oxide, are metal chlorides. EDX analysis of this area is shown in Figure 7.6, where it is evident that the presence of metal chloride has caused poor adhesion, which may have resulted in the thick layered oxide structure.
Figure 7.5: Cross sections of Fe-2.25Cr-1Mo exposed for 24 hours with 20 % H₂O at 400 °C a) KCl, b) NaCl, c) PbCl₂ and d) ZnCl₂.
Figure 7.6: Cross section SEM/EDX maps of Fe-2.25Cr-1Mo exposed for 24 hours at 400 °C, 20 % H_2O(g) in the presence of PbCl_2.

In the presence of ZnCl_2, the lowest mass gain was detected, which is in agreement with the cross section in Figure 7.5 d), where the thinnest scale is observed. As mentioned above, this can be attributed to the fast decomposition and evaporation of the chloride, which might be the reason for the relative mild attack.

To summarize, it can be concluded that the most severe corrosion attack was observed in the presence of KCl and not in the presence of ZnCl_2, which is supposed to be molten. This indicates that the presence of a bulk melt is not a requirement for high corrosion rates. However ZnCl_2 readily reacts into ZnO, which means that a molten state may perhaps only be present during the initial stages of exposure. The next section will investigate the influence on corrosion in the presence of a low melting mixture of NaCl/ZnCl_2 or KCl/ZnCl_2 with melt temperatures far below the exposure temperature. Will these mixtures induce a more severe attack than KCl? It should be noted that all the exposures presented below were performed in 40 % H_2O(g).
7.1.2 Exposure in the presence of mixed chlorides

Figure 7.7: Mass change of Fe-2.25Cr-1Mo in the presence of pure chlorides and mixtures at 400 °C with 40 % H₂O.

A mixture of the salts (KCl/ZnCl₂ and NaCl/ZnCl₂) substantially lowers the melting temperature of the salt mixture compared to the melting point of each individual salts (250 °C and 260 °C respectively). If the melting temperature is directly related to the corrosion rate, the samples coated with the salt mixtures should exhibit the highest corrosion rates. However, this is not the case. The sample exposed in the presence of NaCl/ZnCl₂ resulted in a mass gain, which is similar to pure ZnCl₂ (see Figure 7.7). The mass gain with KCl/ZnCl₂ is slightly higher, almost equal to NaCl and the greatest increase was again observed in the exposure with pure KCl. The exposure temperature is above the melt temperature for both KCl/ZnCl₂ and NaCl/ZnCl₂ as well as ZnCl₂. Despite this fact there is no profound difference between the mixtures and the individual salts regarding mass gain. The melt temperature of the salts/mixtures seems to be less detrimental to the oxidation rate than the characteristics of the individual alkali chlorides.

The T22 in the presence of NaCl/ZnCl₂ and KCl/ZnCl₂ behaves similar, in that the morphology is altered compared to the samples coated with pure salts. With pure NaCl, the crystals remain relatively intact on the surface of the sample, whereas, when mixed with ZnCl₂ no intact NaCl crystals are left intact after exposure, instead, Zn, Na, O and Cl containing nodules have formed on the surface (see Figure 7.8 a)). The plan view in the presence of KCl/ZnCl₂ is visible in Figure 7.8 b). Similar to the NaCl/ZnCl₂ case, nodules are scattered over the surface. The bright regions are Zn- rich with K and Cl in the range of 7-10 atomic %, which could indicate the
presence of an area where a melt was present during exposure. However, there is no signs of thick oxide in the vicinity of this area.

Figure 7.8 Plan view BSE SEM image of Fe-2.25Cr-1Mo exposed 24 hours at 400 °C and 20 % H$_2$O(g) in the presence of a) NaCl/ZnCl$_2$ b) KCl/ZnCl$_2$.

The oxide on the pure KCl-coated sample was slightly thicker than the oxide formed on the NaCl-coated sample. The same trend is observed for the salt mixtures. If melting temperature is the main factor regarding oxide growth, there should be a significant difference in oxide thickness between the individual pure chlorides and the mixtures. However, one has to take into consideration that sufficient amounts of ZnCl$_2$ have evaporated and/or have been converted into ZnO, thus raising the melting temperature. If that is true, then melt formation may only influence oxidation to a minor extent. Cross sections of T22 samples coated with NaCl/ZnCl$_2$ and KCl/ZnCl$_2$ are shown in Figure 7.9 where no signs of molten phases are present. Instead, clear signs of poor adhesion can be seen in both images, which indicates that metal chlorides were present at the interface between the inward/ outward growing parts of the oxide scale during exposure.
Figure 7.9: Cross section SEM/EDX maps of T22 exposed for 24 hours at 400 °C and 40 % H₂O(g) in the presence of a) NaCl/ZnCl₂ and b) KCl/ZnCl₂.
Figure 7.10: Cross section SEM of T22 exposed for 24 hours at 400 °C and 40 % H₂O(g) in the presence of a) NaCl/ZnCl₂ and b) KCl/ZnCl₂.

Both salt mixtures investigated severe spallation (more clearly seen in Figure 7.10), which explains why less Cl⁻ is detected in the IC analysis than pure alkali chlorides (see Table 7). Cl⁻ is also lost due to the decomposition and evaporation of ZnCl₂ see reactions 7.10 and 7.12.
Table 7. Remaining Cl\(^{-}\) on T22 after 24 hours in the presence of the different salts at 400 °C and 40 % H\(_2\)O(g) analyzed with IC.

<table>
<thead>
<tr>
<th>Salt</th>
<th>Remaining Cl(^{-}) after exposure as [mole %]</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaCl</td>
<td>65</td>
</tr>
<tr>
<td>KCl</td>
<td>57</td>
</tr>
<tr>
<td>ZnCl(_2)</td>
<td>23</td>
</tr>
<tr>
<td>NaCl/ZnCl(_2)</td>
<td>25 (spallation)</td>
</tr>
<tr>
<td>KCl/ZnCl(_2)</td>
<td>20 (spallation)</td>
</tr>
</tbody>
</table>

7.1.3 Exposures in the presence of HCl(g) and KCl-HCl(g)

In order to more closely investigate the role of chlorine in the corrosion attack, the simultaneous presence of KCl and HCl(g) were investigated.

The T22 was exposed to KCl, HCl(g) and KCl-HCl(g), both in a tube furnace and in a TG. As can be seen in the gravimetric analysis in Figure 7.11, the introduction of HCl(g) resulted in a doubling of the mass gain compared to the reference.

![Mass change of T22 in the presence of 500vppm HCl and KCl at 400°C with 20 % H\(_2\)O.](image)

Figure 7.11: Mass change of T22 in the presence of 500vppm HCl and KCl at 400°C with 20 % H\(_2\)O.

In comparison with the exposed reference samples, there is no profound difference in surface morphology (compare Figure 7.2 and the left-hand image in Figure 7.12). In images with higher
magnification of both the reference sample and in the presence of HCl(g), oxide blades/whiskers appear all across the surface. This would indicate a top layer in the oxide scale consisting of Fe$_2$O$_3$, since this oxide is known to form these structures. On some samples in the presence of HCl(g) however, two different areas can be distinguished. Where one area has a similar oxide thickness as the reference sample. Whereas the other area is roughly 3 times thicker, which may indicate the presence of an incubation time.

Figure 7.12 Plan view and cross section BSE SEM images of Fe-2.25Cr-1Mo exposed 24 hours at 400 °C in the presence of 500 vppm HCl(g) and 20 % H$_2$O(g).

With the addition of HCl(g), the oxide is similar to the reference sample consisting of an outward and inward growing part, however the thicker areas, are thicker by a factor of 3. Interestingly, the scale is more or less devoid of chlorine, which can be seen in the Cl map in the lower right-hand corner in Figure 7.12. Although no Cl was detected with EDX in the marked area, it might be present at some other position. Moreover, the minor amounts that were present during exposure may have either evaporated or may have been under the detection limit of the EDX. Since the introduction of HCl(g) resulted in a doubling of the mass gain, Cl has influenced the oxidation kinetics in some way. If the entire sample is leached and analyzed, minor Cl concentrations can be detected with IC (see Figure 7.13).
In order to obtain more information about the oxidation kinetics, exposures in a thermo gravimetric in-situ system were performed. Mass gain recorded in-situ provides more detailed information about the reaction kinetics during exposure and not just one data point as is the case with samples exposed in a tube furnace. As can be seen in Figure 7.14, both the reference and the HCl(g) exposures exhibit low mass gains with parabolic behavior. Oxidation behavior in the presence of KCl is rather different from the reference and HCl(g) exposures. The mass gain curve exhibits several bumps during the exposure. This indicates cracking of the oxide, thus revealing the underlying steel which would result in a short period of fast oxidation until a diffusion-controlled regime is once again established. Resulting once again in an outward and inward growing part of the scale.
When KCl and HCl(g) are present simultaneously, the oxidation is altered dramatically as is shown in Figure 7.14. During the first 2 hours, the mass gain was extremely rapid, but after this point it levelled off.

![Figure 7.14: Mass change of T22 for the different exposure cases (as denoted in the image) at 400 °C with 20 % H₂O.](image)

The difference between the KCl and KCl-HCl(g) exposed samples in the early stages of exposure is shown in Figure 7.15, where the exposures were terminated after 1 hour. In the KCl exposure, oxidation is initiated between the KCl particles and around the edges of the particles (see Figure 7.15 a)). On the contrary, in the case of KCl-HCl(g) exposure, (see Figure 7.15 b)) the oxidation was much more localized around KCl particles. The plan view comparison does not explain the difference in mass gain during the early stages, however, by comparing the cross sections in Figure 7.15 it becomes more evident. Figure 7.15 c) shows the KCl cross section with a relatively thin uniform oxide. Figure 7.15 d) shows the cross section of KCl-HCl(g), where the square highlights a region in which a thick oxide has formed around a KCl particle. In the Cl EDX map, Cl is observed as a continuous layer underneath the oxide scale.
Figure 7.15: Plan view and cross section SEM/EDX maps for comparison of T22 exposed for 1 hour at 400 °C and 20 % H₂O(g) in the presence of a) KCl plan view and b) KCl- HCl(g) plan view, c) KCl cross section and d) KCl- HCl(g) cross section.

An image with higher magnification of the marked region in Figure 7.15 d) is shown in Figure 7.16. Almost no inward growing part of the oxide scale is detected. Instead it appears that the inward transportation of Cl was faster than that of oxygen, furthermore the outward diffusion of iron was extremely rapid. Similar areas with thick oxide were formed all over the sample surface, which can explain the comparatively much higher mass gain during the early stages than in the KCl exposure in the absence of HCl(g).
After 24 hours of exposure, a notable difference between KCl samples and KCl- HCl(g) samples can be seen in Figure 7.17 a) and b). It appears that the corrosion attack is widespread all over the surface in the presence of KCl. The attack on the KCl- HCl(g) sample is much more localized to the vicinity of the KCl particles. Another difference is that the KCl particles in the presence of HCl(g) appear to be less consumed after exposure than KCl in the absence of HCl(g).
A BIB cross section of the KCl- HCl(g) exposed sample with EDX maps is shown in Figure 7.18. It can be clearly seen that the thickest scale is located close to the KCl particles. Cl is detected in high concentrations at the steel/scale interface underneath the KCl and accordingly, higher amounts of Cl\(^-\) are detected with IC (see Figure 7.13). Between the KCl particles, the oxide scale is much thinner and Cl is almost absent at the interface. However, the scale is thicker than in the case with only HCl(g) see Figure 7.19. As shown in Figure 7.14 the rate of mass gain is extremely rapid during the initial stages of exposure. The cross section analysis indicates that most of the oxide growth takes place in the vicinity of the KCl, where fast oxidation kinetics reign during the early phases of exposure. After some time, the oxide has reached a point where the transport of ions is greatly reduced and slow oxidation kinetics reign during the remaining period of exposure.
Figure 7.18: Cross section SEM/EDX maps of Fe-2.25Cr-1Mo exposed for 24 hours at 400°C in the presence of KCl - HCl(g) and 20% H₂O(g).

7.1.4 TEM microstructure analysis of T22

To further examine the microstructure of the formed oxides, TEM analysis has been performed in collaboration with Dr M. Sattari and A Persdotter. In order to enable comparison between samples with and without KCl, the area selected for analysis is located at a distance from KCl particles. In this way it is possible to compare the influence of Cl regarding the microstructure of the base oxide, denoted by region 2 in paper II. Both the reference and the HCl(g) samples in Figure 7.19 a) and b) has formed a considerably thicker outward growing part of the scale compared to the inward growing, which indicates that the outward diffusion of iron is faster than the inward diffusion of oxygen. There is a large void density in the Fe₃O₄ layer on both samples, however somewhat higher in the case with HCl(g). This might be the reason for the higher oxidation rate, where the voids is an indicator of fast outward diffusion of Fe. In Figure 7.19 c) the poor scale adhesion and layered structure is seen on the sample in the presence of KCl. Together with the gravimetric analysis in the TG, this is strong evidence of crack formation and subsequent rehealing with a new scale. The TEM analysis showed that the “rehealed” layers similarly consist of a three layered structure with hematite whiskers on top as is marked in the image. The case with KCl – HCl(g) in Figure 7.19 d) shows that the ratio between the outward/inward growing part is decreased. Thus, the simultaneous presence of KCl and HCl(g) has resulted in a faster inward diffusion of oxygen compared to the reference and the HCl(g) exposures. Similar to the HCl(g) case there are numerous voids in the Fe₃O₄ layer.
To further investigate the role of chlorine in the corrosion process it is important to perform more time resolved exposures and detailed microstructural analysis. TEM can be useful in order to map out where chlorine is present during the initiation of the corrosion process where rapid mass gains are initiated. Also if chlorine induces a change of the oxide microstructure (thereby increasing the grain boundary diffusion), but without clear signs of chlorine in the scale. As was seen in the presence of HCl(g).

Figure 7.19: Cross section TEM images of Fe-2.25Cr-1Mo exposed for 24 hours at 400°C and 20% H₂O(g) in the presence of a) Reference, b) HCl(g), c) KCl, d) KCl – HCl(g).
7.1.5 Concluding discussion of laboratory exposures

In all studied exposures in which chlorine-containing compounds were present, compared to the reference, accelerated corrosion rates were observed. The increased corrosion rates may be attributed to the formation of transition metal chlorides. These metal chlorides are located at the steel/oxide scale interface. This will lead to a decrease in scale adhesion. Combined with the fact that the oxide is growing, stresses are introduced which can explain the buckling and cracking of the oxide that was observed in the presence of NaCl/ZnCl$_2$ and KCl/ZnCl$_2$ for example. Transition metal chlorides can also increase the diffusion of ions along oxide grain boundaries.

In the presence of KCl, a widespread corrosion attack was observed all over the sample surface. This is attributed to the presence of a very thin liquid film of KCl/FeCl$_2$, were KCl diffuses out from the particles, which can explain the widespread nature of the attack [78]. The cross section analysis indicated the presence of FeCl$_2$ during exposure. It is believed that the reaction between FeCl$_2$ and adsorbed KOH in the outer parts of the scale have caused the poor scale adhesion and the occurrence of large voids. A similar mechanism is thought to be at play in the presence of NaCl.

In the presence of PbCl$_2$, a more localized attack was observed. A thick and layered scale was observed below the area where a PbCl$_2$ particle was positioned prior to exposure. The localized attack can be explained by the absence of a liquid film of PbCl$_2$/FeCl$_2$. Since the eutectic temperature was above the exposure temperature, less diffusion of corrosive species occurred. After 24 hours of exposure no PbCl$_2$ was detected due to the conversion into PbO, which may explain the high amounts of Cl that were present at the steel/oxide interface. In the reaction in which PbO is formed, Cl is released. This Cl can (as the results indicate) diffuse through the oxide to form metal chlorides. Since the nature of the attack was localized, increased amounts of PbCl$_2$ would probably result in a much more severe attack, than was observed under the studied conditions.

The ZnCl$_2$ should be in a molten state under the studied conditions, which is expected to initiate accelerated corrosion. However, no signs of corrosion induced by a melt were observed. This was probably due to the fact that large amounts of ZnCl$_2$ were lost by evaporation and by
reactions in which ZnFe$_2$O$_4$ and ZnO were formed. Similar to in the case with PbCl$_2$, higher amounts would most likely result in higher corrosion rates, as has been reported by other researchers [12, 47, 72].

The mixtures KCl/ZnCl$_2$ and NaCl/ZnCl$_2$, with a low melting point, did not show any clear signs of bulk melt induced corrosion. Instead, out of all the Fe-2.25Cr-1Mo samples, exposed in this study, the presence of low melting mixtures resulted in the most severe spallation. However ZnCl$_2$ is readily converted into ZnO, why the presence of a molten phase may only influence the corrosion during the early stages of exposure.

The hypothesis before the exposures with KCl and HCl(g), was that the simultaneous presence of these species would cause a more severe attack than if only KCl was present. On the contrary, the results showed a lower mass gain and a denser and a better adhered oxide scale. The cross sections showed that metal chlorides were located in a band underneath KCl particles. The fast oxidation that was observed in the TG during the early stages of exposure could be explained by the fast scale growth above could be connected to the presence of this band. Furthermore the KCl particles was less consumed and the nature of the corrosion attack was much more localized in the presence of HCl(g). These results has led to the proposition of a new mechanism. Where differences in the chemistry at the surface can be coupled to the severity of the corrosion attack.

In the case with KCl basic conditions (KOH adsorbed on Fe$_2$O$_3$) reigns at the top oxide surface and as was observed with both the TEM and the TG analysis, repeated breakdowns (cracking) of the scale has occurred during the exposure. It is suggested that the cracking is initiated by the reaction of FeCl$_2$ at the bottom of the scale with excess alkali in the upper parts according to: FeCl$_2$(s) + 2KOH(ads@Fe$_2$O$_3$)→Fe(OH)$_2$(s) + 2KCl(s). It is suggested that both KOH and FeCl$_2$ is transported along oxide grain boundaries, to provide the contact between the two species. The decomposition will disrupt the scale adhesion and leads to the formation of voids. The cracked, void dense scale gives the surrounding atmosphere access to the unprotected metal, which would explain the occurrence of the “bumps” in the mass gain curve and the appearance of the TEM cross section.

Whereas in the case in which HCl(g) is present the top surface becomes acidic. Therefore, compounds that are stable under alkaline conditions decompose according to: KOH(ads@Fe$_2$O$_3$) + HCl(g)→KCl(s) + H$_2$O(g).
The large accumulation of FeCl$_2$ beneath the scale in the KCl-HCl(g) case compared to the KCl(only) exposure, is attributed to the HCl(g) which eliminates the alkaline chemistry at the scale surface. Thus, the formation of cracks does not initiate fast decomposition of the sub-scale FeCl$_2$ as in the KCl(only) exposure.

The rapid deceleration of the mass gain in the KCl-HCl(g) exposure is attributed to the electronically insulating properties of the FeCl$_2$ layer. Thus, disturbing the O$_2$ reduction at the surface, which results in slower oxidation kinetics.

### 7.2 Superheater exposures

This section of the thesis investigates the corrosiveness of alkali chlorides towards superheater materials, which operate at a higher material temperature and lower flue gas temperatures than the waterwalls.

By switching the fuel from coal to biomass and waste, the result is a considerably more corrosive environment inside the boiler. The flue gas from these fuels is relatively rich in reactive alkali chlorides and typically, the sulfur dioxide concentration is low. As a consequence, the deposit formed on superheater tubes tends to have substantial amounts of corrosive alkali chlorides. Numerous studies have shown the corrosiveness of KCl and NaCl towards superheater materials (mainly stainless steels) predominantly due to alkali chromate formation. One way to mitigate this is by raising the SO$_2$(g) concentration in the flue gas. SO$_2$(g) reacts with the alkali chlorides, which are converted into the corresponding and at the reigning temperature range less corrosive, alkali sulfates. The availability of sulfur in the boiler can be increased in several ways, for example by adding sewage sludge which contains sulfur in the form of sulfates. Other ways can be to add sulfur in the form of granules, or by spraying ammonium sulfate into the flue gas [81]. In the field study of the thesis, a novel technique was used in which existing sulfur in the fuel was recirculated back into the furnace. The laboratory part of this section investigates the corrosive effect of 304L in the presence of relatively large amounts of KCl (1mg/cm$^2$) and if the presence of SO$_2$(g) can mitigate the corrosiveness of large amounts of KCl.
7.2.1 **In-situ sulfation of KCl in laboratory exposures**

Previous studies on 304L with 1.35 \( \mu \text{mol/cm}^2 \) KCl (referred to as small amounts of KCl) have shown that the corrosion attack is initiated by chromate formation according to reaction 7.15.

\[
\text{Cr}_2\text{O}_3(s) + 4\text{KCl}(s) + 2\text{H}_2\text{O}(g) + \frac{3}{2} \text{O}_2(g) \rightleftharpoons 2 \text{K}_2\text{CrO}_4(s) + 4\text{HCl}(g)
\]  
\[\Delta G = 147 \text{ kJ/mol, } p_{eq}(\text{HCl}) = 1303 \times 10^{-6} \text{ bar at 600°C [66] calculated with data from Barin.}\]

The chromate formation leads to a chromium depletion that results in a breakdown of the Cr-rich protective oxide that initially forms on stainless steel. As a result, a fast growing, iron-rich oxide scale forms, (see Figure 7.19 a)).

![Figure 7.19: BSE image of samples exposed 24 hours at 600 °C in 5 % O\(_2\)(g) + 40 % H\(_2\)O(g) + 1.35 μmol/cm\(^2\) KCl for a) without SO\(_2\)(g) and b) with SO\(_2\)(g).](image)

The small amount of KCl applied onto the sample was consumed very fast, after 1 hour almost 40 % of the KCl had reacted and was more or less consumed after 24 hours of exposures. Chromate formation releases chlorine as HCl(g), which is believed to cause accelerated corrosion. However, there were no signs of chlorine in the cross-section, which indicates that HCl(g) leaves the surface after formation in the chromate reaction.

Using sulfur-rich additives or co-combustion with sulfur-rich fuels are strategies to decrease the corrosion rate in biomass and waste-fired boilers. The aim is to transform corrosive alkali
chlorides to less corrosive alkali sulfates. In contrast to KCl, potassium sulfate does not induce an accelerated corrosion attack on 304L at 600 °C. Consequently, the reaction of K$_2$SO$_4$ with the protective scale to form K$_2$CrO$_4$ is not thermodynamically favored under the present experimental conditions. The corrosion rate of 304L in the presence KCl+SO$_2$(g) is considerably lower than the corresponding exposure in the absence of SO$_2$, which is illustrated in Figure 7.19 (compare images a) and b)).

By increasing the amount of KCl to 13.5 µmol/cm$^2$ in the absence of SO$_2$(g), the corrosion morphology of the surface did not change significantly compared to samples exposed with small amounts of salt. In both cases the general corrosion attack can be explained by the formation of chromate, which destroyed the protective properties of the Cr-rich oxide. However, samples exposed to large amounts KCl showed more than twice the amount of chromate formation than samples exposed to small amounts. Consequently, the general corrosion attack was more severe on samples with large amounts of KCl.

The presence of large amounts of KCl also accelerates the corrosion attack along steel grain boundaries (see Figure 7.20 a)). This attack may be the result of increased chromate formation, leading to more Cr-depleted steel grain boundaries. Another explanation could be that chlorine, released as HCl(g) during the formation of chromate, induces the attack.
When 304L was exposed to large amounts of KCl + SO$_2$(g), the general corrosion attack was similar to the corresponding attack in the presence of small amounts of KCl + SO$_2$(g). However, the attack along the steel grain boundaries became much more severe (see Figure 7.20 b)). The steel grain boundary attack could clearly be related to the presence of sulfides and chlorides. Chlorination is expected to be the result of the formation of both K$_2$CrO$_4$ and K$_2$SO$_4$, resulting in (locally) high levels of HCl(g). Since the gas flow is expected to be low beneath the porous KCl particles, the HCl(g) concentrations should reach higher levels in these areas. This implies that the steel grain boundary attack should be more severe in these areas, as indicated by the results. Furthermore, since a large number of the attacked steel grain boundaries contain sulfides, it is suggested that sulfidation of steel grain boundaries facilitates the steel grain boundary attack. For this reason, both the chlorination and sulfidation of steel grain boundaries are expected to accelerate this attack.

Figure 7.20: BSE image of samples exposed 24 hours at 600°C in 5 % O$_2$ + 40 % H$_2$O + 13.5 μmol/cm$^2$ KCl for a) without SO$_2$ and b) with SO$_2$. 
7.2.2 **Concluding discussion of in-situ sulfation**

The corrosiveness of different amounts of KCl on 304L in both the absence and presence of SO$_2$(g) at 600 °C was investigated. With small amounts of KCl (1.35 µm/cm$^2$) the general corrosion attack was initiated by the formation of chromate. This resulted in chromium depletion of the protective oxide that initially forms on this stainless steel, and a fast growing iron-rich oxide was formed. No steel grain boundary attack was observed.

With larger amounts of salt (13.5 µm/cm$^2$ KCl), the formation of chromate increased and an attack along the steel boundary was introduced. It is suggested that this attack is accelerated by the increased presence of chlorine, released from the chromate formation.

The presence of 13.5 µm/cm$^2$ KCl on 304L in a SO$_2$(g) atmosphere resulted in the simultaneous formation of K$_2$SO$_4$ and K$_2$CrO$_4$. However, the resulting general attack was considerably slower. In contrast to the general corrosion attack, the presence of SO$_2$ accelerated the steel grain boundary attack.

Larger amounts of KCl in the presence of SO$_2$(g) did not increase the general corrosion attack. However, the corrosion attack along the steel grain boundaries was greatly increased. This attack is suggested to be due to the formation of metal chlorides and metal sulfides in the steel grain boundaries.

7.2.3 **The effect of sulfur recirculation (field study)**

To investigate the effect of sulfur recirculation, both 24 hour and 1000 hour exposures were performed in order to study both the initial and the long-term corrosion. Reference exposures without recirculation were also performed and the material temperature for both exposures was set to 525°C. In the reference exposures (denoted REF), municipal and industrial waste was used as fuel. The flue gas concentration of HCl and SO$_2$ were 300 ppm and 90 ppm, respectively. In the exposures in which sulfur recirculation (denoted S REC) was deployed, both the HCl(g) and SO$_2$(g) concentrations increased to 400 ppm and 270 ppm, respectively. In both exposure conditions (REF and S REC), the three materials Sanicro 28, Inconel 625 (over-
welded on a 16Mo3 tube) and 16Mo3 (for composition see Table 8) were exposed simultaneously.

Table 8. Chemical composition of the tested materials in weight%

<table>
<thead>
<tr>
<th>Alloy</th>
<th>% C</th>
<th>% Fe</th>
<th>% Cr</th>
<th>% Ni</th>
<th>% Mo</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>16Mo3</td>
<td>0.12</td>
<td>99</td>
<td>-</td>
<td>-</td>
<td>0.3</td>
<td>0.6 % Mn</td>
</tr>
<tr>
<td>Inconel 625</td>
<td>&lt; 0.10</td>
<td>5</td>
<td>22</td>
<td>58</td>
<td>9</td>
<td>0.5 % Mn. 3.15-4.15 Nb+Ta</td>
</tr>
<tr>
<td>Sanicro 28</td>
<td>&lt; 0.02</td>
<td>35</td>
<td>27</td>
<td>31</td>
<td>3.5</td>
<td>1 % Cu</td>
</tr>
</tbody>
</table>

Figure 7.21 shows two optical cross sections images of 16Mo3 sample rings exposed 1000 hours, to the left REF and to the right S REC. The REF sample is covered with a grey/brown deposit, underneath is the corrosion product layer located on top of the sample ring. Also the S REC sample is covered with deposit, however not as thick as on the REF sample. The corrosion product layer is located between the deposit and the steel ring, it can clearly be seen that this layer is considerably thinner on the S REC sample than on the REF sample.

Figure 7.21: Optical microscopy images of 16Mo3 cross sections exposed 1000 hours for the REF and S REC samples.

In addition to optical microscopy, SEM/EDX, XRD, IC and material loss measurements were performed. In all exposures, two samples of each material were exposed on the corrosion probe. One of the samples was analyzed with XRD to identify crystalline corrosion products. After the XRD analysis the deposit and corrosion product layers were leached in MilliQ water for the quantification of water soluble ions with IC. The other sample was mounted in epoxy and cut
in two in order to obtain cross sections. One of the sample cross sections was used to analyze the deposit, corrosion product layer and metal with SEM/EDX. The other cross section was used to determine material loss.

7.2.3.1 24 hours exposure at 525 °C

The IC analysis showed that the deposit was dominated by sulfates on both the REF and the SREC samples. However, a larger fraction of Cl\(^{-}\) was detected on the REF samples than on the SREC. It is, thus, evident that sulfur recirculation results in deposits with less Cl\(^{-}\) (see Figure 7.22). Low concentrations of Fluoride (F\(^{-}\)) were also detected, but did not seem to be involved in the corrosion attack.

![Figure 7.22: Ion chromatography analysis of deposit on 24 hour samples, presented as mole %.

On the 24 hour samples the XRD results coincides fairly well with the IC analysis above. For the samples exposed with SREC no alkali chlorides were detected with XRD, whereas alkali chlorides showed weak to medium intensity peaks in the REF exposure (see Table 9). Consequently, the results clearly indicate that sulfur recirculation suppresses the amount of alkali chlorides in the deposit.
Table 9. Crystalline phases detected by XRD analysis of samples exposed 24 hours.

<table>
<thead>
<tr>
<th></th>
<th>CaSO₄</th>
<th>Na₂SO₄</th>
<th>K₂Ca₂(SO₄)₃</th>
<th>KCl</th>
<th>(K,Na)Cl</th>
<th>NiO</th>
<th>Fe₂O₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sanicro28</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reference</td>
<td>S</td>
<td>-</td>
<td>-</td>
<td>M</td>
<td>-</td>
<td>M</td>
<td>W</td>
</tr>
<tr>
<td>S-Recirculation</td>
<td>S</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>W</td>
</tr>
<tr>
<td>Inconel625</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reference</td>
<td>S</td>
<td>-</td>
<td>-</td>
<td>W</td>
<td>-</td>
<td>M</td>
<td>W</td>
</tr>
<tr>
<td>S-Recirculation</td>
<td>S</td>
<td>-</td>
<td>W</td>
<td>-</td>
<td>-</td>
<td>W</td>
<td>W</td>
</tr>
<tr>
<td>16Mo3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reference</td>
<td>S</td>
<td>-</td>
<td>-</td>
<td>M</td>
<td>W</td>
<td>-</td>
<td>M</td>
</tr>
<tr>
<td>S-Recirculation</td>
<td>S</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>W</td>
</tr>
</tbody>
</table>

W = Weak, M = Medium and S = Strong

Figure 7.23 shows a BSE image of the REF 24 hour Sanicro 28 ring together with EDX maps of that area. The figure shows that the deposit mainly consists of Na, K, Ca, O and S, most likely as sulfates in various compositions, which is in agreement with the IC and XRD analyses. The marked area in the Cl map marks where the highest concentration is located, this location corresponds to an outward migration of Cr in the Cr map. Fe, Cr and Ni can be seen in scattered locations above the steel surface and not as a coherent scale, which indicates the inability to form a protective oxide scale during the exposure.

![Figure 7.23: BSE image and EDX maps of the REF 24 hours Sanicro 28 cross section.](image)

Figure 7.23: BSE image and EDX maps of the REF 24 hours Sanicro 28 cross section.

Figure 7.24 displays a BSE image of the S REC 24 hour Sanicro 28 ring together with EDX maps of that area. With recirculated sulfur, it can be seen that the deposit has shifted from mixed Na, K, and Ca sulfates, which were detected on the REF sample, towards relatively pure CaSO₄.

![Figure 7.24: BSE image and EDX maps of the REC 24 hours Sanicro 28 cross section.](image)
Whereas the REF sample did not form a coherent protective oxide scale, the Cr map in Figure 7.24 indicates that Cr is located in a thin continuous scale together with Fe, Ni and O. This behavior and appearance of an oxide scale is what usually constitutes protective properties against corrosion. These findings support the theory which states that alkali chlorides can be transformed into less corrosive alkali sulfates. IC, XRD and SEM/EDX analyses shows that sulfur recirculation results in a less corrosive environment. The initial property of an oxide is vital to the long term protection of an alloy. If a thick and porous oxide forms during the initial stages of exposure, the oxide will fail to provide protection during later stages. If, instead, a thin oxide with protective properties forms initially, its protective properties seems to last for longer durations of exposure.

Figure 7.24: BSE image and EDX maps of the S REC 24 hours Sanicro 28 cross section.

7.2.3.2 1000 hours exposure at 525°C

The XRD results for the 1000 hour samples do not coincide well with the IC results (see Figure 7.25 and Table 10). The IC results indicate that the amount of Cl⁻ decreases by roughly 50 % with sulfur recirculation. However, the XRD analysis detected medium intensity peaks for alkali chlorides in both the REF and S REC samples. Unfortunately, the Sanicro 28 REF sample was damaged during sample preparation and no IC analysis was performed. It is, however, relatively safe to assume that the deposit on the Sanicro 28 REF sample is similar in
composition to the deposit on the 16Mo3 and Inconel 625 samples due to their close positioning on the sample probe.

Figure 7.25: Ion chromatography analysis of deposit on 1000 hour samples, presented as mole %.

Table 10. Crystalline phases detected by XRD analysis of samples exposed 1000 hours.

<table>
<thead>
<tr>
<th></th>
<th>CaSO₄</th>
<th>Na₂SO₄</th>
<th>K₂Ca₂[SO₄]₃</th>
<th>KCl</th>
<th>(K,Na)Cl</th>
<th>NiO</th>
<th>Fe₂O₃</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sanicro28</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reference</td>
<td>S</td>
<td>W</td>
<td>W</td>
<td>M</td>
<td>W</td>
<td>M</td>
<td>-</td>
</tr>
<tr>
<td>S-Recirc.</td>
<td>S</td>
<td>-</td>
<td>-</td>
<td>M</td>
<td>M</td>
<td>-</td>
<td>W</td>
</tr>
<tr>
<td><strong>Inconel625</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reference</td>
<td>S</td>
<td>W</td>
<td>W</td>
<td>M</td>
<td>W</td>
<td>S</td>
<td>-</td>
</tr>
<tr>
<td>S-Recirc.</td>
<td>S</td>
<td>-</td>
<td>-</td>
<td>M</td>
<td>M</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td><strong>16Mo3</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td>M</td>
<td>W</td>
<td>W</td>
<td>-</td>
<td>-</td>
<td>W</td>
</tr>
<tr>
<td>S-Recirc.</td>
<td>S</td>
<td>-</td>
<td>-</td>
<td>M</td>
<td>M</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

W = Weak, M = Medium and S = Strong

Figure 7.26 shows a BSE image of the REF 1000 hour Sanicro 28 ring together with EDX maps of that area. The image shows the corrosion product layer formed on the Sanicro 28 REF sample, which is relatively thick and full of pores and cracks, indicating non protective behavior. Fe, Cr and Ni constitute the oxide with minor inclusions of Zn, Na, K and S. In the Cr map, two bright bands are visible. The first is located close to the metal surface and the second in the outer parts of the layer. Cl is spread throughout the layer, mainly correlating to
the Na map, indicating NaCl. Closer to the metal surface, the Cl map overlaps with the metals originating from the steel, indicating the presence of metal chlorides, which could explain the porous morphology and the thick corrosion product layer.

![Figure 7.26: BSE image and EDX maps of the 525 °C REF 1000 hours Sanicro 28 cross section.](image)

In Figure 7.27 a BSE image of the S REC 1000 hour Sanicro 28 ring together with EDX maps of that area is shown. In the image, it is evident that the oxide thickness has decreased dramatically with S REC from ~600 µm with REF to ~30 µm with S REC. In the EDX maps, it is clear that the deposit consists of sulfates, mainly potassium sulfate. The Cl map is totally dark, which indicates that most alkali chlorides have been transformed into alkali sulfates, which is in agreement with the IC analysis in Figure 7.25, in that figure the amount of Cl-decreased by approximately 50 % on the S REC sample. In the Cr map, a bright band is visible above the steel surface, indicating that a Cr rich oxide has formed. This can again be coupled to the initial properties of the Sanicro 28 S REC sample exposed for 24 hours, on which a thin continuous oxide scale was present. The protective property of the oxide scale seems to be sufficient as protection also after 1000 hours of exposure. The arrows mark a correlation between S and Ni, indicating the occurrence of Ni$_2$S$_3$ on top of the oxide scale. The formation of sulfide instead of an oxide or sulfide incorporated into an oxide on top of the steel surface decreases the oxide’s ability to protect the steel from further oxidation. Sulfide formation can also cause poor adhesion of the oxide scale to the steel. In other words, the presence of sulfides is an indication that the SO$_2$(g) levels were too high during the exposure.
The material loss measurement in Figure 7.28 shows the beneficial effect of sulphur recirculation on corrosion. The Sanicro 28 sample exposed in the S REC exposure exhibited a material loss approximately 85% lower than the sample in the REF exposure.

Figure 7.28: BSE image and EDX maps of the S REC 525 °C 1000 hour Sanicro 28 cross section.

Figure 7.28: Material loss measurements of 1000 hour sample.
7.2.4 Concluding discussion of superheater corrosion (laboratory and field)

The laboratory study showed that the corrosion attack on the 304L alloy at 600 °C is similar in the presence of 0.1 mg/cm$^2$ and 1 mg/cm$^2$ KCl. By introducing SO$_2$(g), it was possible to satisfactorily mitigate the corrosion attack of 0.1 mg/cm$^2$ KCl by the formation of less corrosive K$_2$SO$_4$. The introduction of SO$_2$(g) in the presence of 1 mg/cm$^2$ KCl resulted in a reduction of the general corrosion, but instead chlorination and sulfidation of the steel grain boundaries were observed.

When the waste fired boiler in the study was operated with sulfur recirculation, the Sanicro28, Inconel 625 and 16Mo3 all formed relatively thin and dense oxides after 24 and 1000 hours compared with Ref samples. This was in agreement with both SEM and material loss measurements. The Ref samples were covered with a thick deposit rich in corrosive alkali chlorides. Sulfur recirculation substantially decreased the amount of alkali chlorides in the deposit, which was seen with IC. A novel imaging technique was developed and used for evaluating material loss on the samples exposed for 1000 hours. It proved to be a fast technique that resulted in a more accurate estimation of the material wastage compared to the previously used method, in which a micrometer screw was used.
8 Acknowledgements

First of all I would like to thank my supervisors Professor Lars-Gunnar Johansson, Professor Jan-Erik Svensson, Jesper Liske Ph.D. and Torbjörn Jonsson Ph.D., for all their support and interesting conversations.

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