High-resolution topochemical analysis and thermochemical simulations of oxides and nitrides at grain boundaries and within the grains of a low alloy Mn-Cr hot-rolled steel sheet

Vera G. Praig, Michael Stöger-Pollach, Arno Schintlmeister, Michael Auinger, Herbert Danninger

1. Introduction

Advanced high strength steels are very important for the automotive and construction materials sector. Due to their superior mechanical performance such as high tensile and yield strength combined with excellent ductility, they are often used for the safety structure in cars as well as for crash-sensitive parts. Various steel grades are available such as dual phase steels (DP), complex phase steels (CP) or TRIP/TWIP-steels which are each iron-based alloys containing 2 wt% manganese and moderate amounts of either chromium, aluminium or silicon. After (continuous) casting, these alloys are often hot rolled to sheet material with an optional hot forming stage to obtain the final shape of the workpiece. Especially during hot rolling and coil cooling, the material suffers considerably from oxidation, both externally and internally. Whereas the external oxides (termed "scale layer") can be removed rather easily, the formation of internal oxides both inside the grains (termed "bulk oxides" or "inner oxides") and along grain boundaries (termed "grain boundary oxides") lead to problems during subsequent pickling, cold forming and coating processes. Before strategies for mitigating the effect of internal oxides can be applied, the formation and composition of internal oxides needs to be understood further into detail, which has been the main scope of this study. Since these oxides are formed underneath the external scale layer, the local oxygen partial pressure is below the onset of wüstite formation but sufficiently high for selective oxidation of the added alloy elements. This oxidation not only lowers the concentrations of dissolved alloy elements and thus changes the stability of phases, it also causes problems with crevice corrosion along grain boundaries in subsequent pickling and/or weakens cohesion between individual grains during subsequent forming processes. The removal of the oxide-affected zone thus requires additional post-treatment of hot formed steel grades, which increases both cost and loss of material. Besides the wealth of information that can be found on the (grain boundary) oxide formation in such industrial steel grades, the present work aims at more detailed investigation of
the fine structure of formed grain boundary precipitates as well as the role of nitrides on the oxide formation and microstructure of grain boundary oxides and inner oxides. Accordingly, the present study focuses on a low/medium manganese dual phase (DP) steel grade from industrial production, as opposed to synthetic model alloys from lab exposures. This will account for the multitude of elements that ultimately define the precipitate microstructure. Our findings on the internal oxide and nitride formation, obtained from industrial steel grades under typical industrial hot rolling and coil cooling conditions, are compared to numerical simulations of element transport and thermodynamic stability of the formed phases. Consequently, the drawn conclusions can be applied more generally on steels and high temperature alloys. Furthermore, theoretical results can also be helpful to understand the formation and microstructure of high temperature corrosion products inside the metallic grains and along grain boundaries in boiler steels [17,18], or for turbine blades [19,20] and how the formation of precipitates can be affected by small amounts of nitride inclusions in the alloys.

2. Experimental procedures

In order to obtain results representative for industrial production, an industrial steel grade (composition see Table 1) was sampled directly after hot-rolling under production conditions and used for the lab-scale heat exposures. The topmost zone, which contained the formed scale layer as well as possible segregation of elements near the alloy/oxide interface, was removed mechanically by grinding and the samples were then polished with SiC grinding paper down to 2500 grit size. Obtained samples were then subjected to a heat treatment at 700 °C for 1 h in dry air and then slowly cooled down to 550 °C over a period of 3.5 h, which is termed in this work as a “complete coil cooling cycle”. After cooling down to room temperature, the heat-treated samples were cut into square shaped pieces of 15 mm × 15 mm × 2 mm in size using a diamond saw.

Where necessary, the samples were mounted in epoxy resin (PolyFast, Struers) for further preparation. Normal cross sections (90°) for standard analysis were prepared as well as polished sections with a 25° tilt angle relative to the specimen surface in order to enlarge the depth of the oxidised zone by a factor of 2.37. All sections were mechanically ground by using grinding papers from 400 down to 2500 grit size and polished with 1 μm diamond paste. Prior to analysis, all samples were ultrasonically cleaned in ethanol and dried in a cold stream of dry air. On the samples intended for nanometer-scale secondary ion mass spectrometry (NanoSIMS), thin AuPd films were sputter-deposited to prevent electrical charging during the NanoSIMS measurement process. The nominal coating thicknesses were 50 nm on the front side and 180 nm on the back side of the sample. NanoSIMS measurements were performed on a NanoSIMS from Cameca (Genevilliers, France). Prior to data acquisition, analysis areas were pre-sputtered by scanning of a high-intensity, slightly defocused Cs+ ion beam (spot size ca. 2 μm at 400 pA beam current) to remove the AuPd coating within the field of view and to establish the steady state signal intensity regime. In order to avoid crater edge effects, scanning during pre-sputtering was conducted over square-sized areas with an edge length exceeding the frame size of the subsequently recorded images by at least 15 μm. Imaging was conducted by sequential scanning of a finely focused Cs+ primary ion beam (2 pA beam current) over areas ranging from 19 μm × 19 μm to 65 μm × 65 μm at a physical resolution of 70 nm (probe size) and an image resolution of 512 × 512 pixels. Data were acquired as multilayer image stacks with a per-pixel dwell time of 10 ms per cycle. The total per-pixel dwell times ranged from 150 msec to 270 msec, corresponding to a total acquisition time of 11–20 h per measurement. 12C, 16O, 14N, 28Si, 34S, 31P and 52Cr as well as secondary electrons were simultaneously detected and the mass spectrometer was tuned for achieving a mass resolving power of > 8000 (according to Cameca’s definition). Evaluation of the image data was carried out, using the WinImage software package (version 2.0.8) provided by Cameca. Prior to stack accumulation, the individual images were aligned to compensate for positional variations arising from primary ion beam and/or sample stage drift. Secondary ion signal intensities were corrected for detector dead-time on a per-pixel basis. Electron microscopy analysis comprised scanning electron microscopy (SEM, FEI Quanta 200 k, The Netherlands) and transmission electron microscopy / electron energy loss spectroscopy (TEM/EELS, FEI TECNAI 20, The Netherlands) at 20 nm lateral resolution [21]. SEM was applied on all sample cross sections, TEM/EELS was conducted on thin lamellae obtained by focused ion beam (FIB) sectioning of polished cross sections using a FIB FEI Quanta 300 2D dual beam equipped with a Ga liquid metal ion source (5–30 kV).

3. Mathematical modelling

Numerical simulations of the internal oxidation process and local phase distributions were carried out by the subsequent two-step algorithm "ASTRID" [22], consisting of element migration and local thermodynamic equilibrium calculations. Transport of the atomic species (oxygen, nitrogen, iron and the alloying elements) has been derived from the system of partial differential equations. Diffusion coefficients are listed in Table 2 and were chosen from literature. The considered thermodynamic phases are provided in Table 3.

The temperature program comprised of an isothermal holding at 700 °C for 1 h and subsequent slow, linear cooling down to 550 °C over a period of 3.5 h. As stated above, in the present work this is termed as a “complete coil cooling cycle”. Since the industrial steel grade appears to be mostly ferritic at process temperatures (see Fig. B.8 in the appendix), the ferrite phase specific diffusion properties were considered. The alloy content of the samples in this study was either too low to stabilise the austenite phase or did not stabilise the austenite phase at all. Hence, depletion of dissolved alloy element additions (C, Al, Cr, Mn, Si) due to oxide formation does not lead to a phase change in the experiments. The oxygen concentration at the surface (upper boundary) was set to a constant value, according to the maximum solubility of oxygen for the iron/â"ıtësitë equilibrium at the applied temperatures (0.45 ppm [O] at 700 °C, see [25]) which corresponds to a scenario of maximum internal oxidation underneath the external scale layer. This corresponds to the constant source model firstly proposed by Wagner [26], which applies to the actual conditions at the iron/â"ıtësitë interface with sufficient accuracy. Accordingly, this assumption is most commonly used in theoretical simulation studies on internal oxidation [9,14,10,27-29]. Since the external scales of our industrial samples contained a large number of cracks, maximum nitrogen uptake was assumed as well [24,30]. Hence the nitrogen concentration at the surface (upper boundary) was set to a constant value which corresponds to the solubility of nitrogen at 1 bar (25.0 ppm [N] at 700 °C, see [31]). The calculation of elemental transport via diffusion has been carried out for small time intervals (≤ 1 min). The results after each calculation step were used to derive the local concentrations of

| C (wt%) | Al (wt%) | Cr (wt%) | Mn (wt%) | Si (wt%) | N (ppm wt.) | S (ppm wt.) | P (ppm wt.) | Fe |
|--------|---------|---------|---------|---------|------------|------------|------------|----|
| 0.1    | 0.1     | 0.8     | 1.8     | 0.1     | <50         | <10        | <200       | bal. |

Table 1 Composition of an industrial Cr-Mn steel, provided by voestalpine Stahl GmbH. Concentrations were determined by atomic absorption spectrometry (AAS).
Table 2
Diffusion coefficients of alloy elements in ferrite.

| Element     | D<sub>o</sub> / m<sup>2</sup>s<sup>-1</sup> | E<sub>a</sub> / kJ mol<sup>-1</sup> | D<sub>0</sub>×10<sup>-18</sup> | Refs. |
|-------------|---------------------------------|-------------------------------|-----------------------------|-------|
| iron (Fe)   | 1.21 × 10<sup>-2</sup>          | 281.6                         | 9.27 × 10<sup>-18</sup>    | [23.9]|
| oxygen (O)  | 3.72 × 10<sup>-6</sup>          | 42.40                         | 1.97 × 10<sup>-8</sup>     | [23.9]|
| nitrogen (N)| 7.80 × 10<sup>-7</sup>          | 79.04                         | 4.46 × 10<sup>-11</sup>    | [23.24]|
| carbon (C)  | 3.94 × 10<sup>-7</sup>          | 80.22                         | 1.95 × 10<sup>-11</sup>    | [23.14]|
| aluminium (Al)| 1.8 × 10<sup>-4</sup>      | 228.2                         | 1.01 × 10<sup>-16</sup>    | [23.9]|
| chromium (Cr)| 8.52 × 10<sup>-4</sup>        | 250.8                         | 2.94 × 10<sup>-17</sup>    | [23.9]|
| manganese (Mn)| 7.60 × 10<sup>-5</sup>  | 224.6                         | 6.68 × 10<sup>-17</sup>    | [23.9]|
| silicon (Si) | 1.7 × 10<sup>-4</sup>          | 229.1                         | 8.57 × 10<sup>-17</sup>    | [23.9]|

4. Results and discussion

Industrial steel grades, as opposed to perfectly homogeneous binary and ternary model alloys, contain a multitude of elements and are prone to small variations in local composition. Thus, results from a single measurement may not be representative for the behaviour of the entire alloy which is why several replicate analyses were conducted and the discussion focuses on the most common observations. Numerical simulations of the spatial distribution of stable phases reveal three distinct regions of thermodynamically stable precipitates (oxides - nitrides - carbides), as shown in Fig. 1. After the heat treatment, a 5–6 μm wide zone of internal oxides has emerged beneath the surface. According to the chemical composition, the most dominant precipitates are thermodynamically stable spinel-type oxides (M<sup>III</sup>N<sub>2</sub>O<sub>4</sub>), followed by other mixed and pure alloy element oxides. This domain is followed by a 40 μm wide zone containing nitride precipitates. During the high temperature treatment, the zone of nitrides gradually moves inwards as the oxidation proceeds. This so-called “push-effect” [33] on the macroscopic scale in fact relies on a substitution of nitrogen atoms in the existing precipitates by oxygen, since oxides are thermodynamically more stable than the corresponding nitride phases (see Fig. B.8). A narrow region of silicon-oxynitrides is visible between the oxide and nitride zones, which is consistent with the thermodynamic predictions. It should be noted that the local nitrogen activity in-between the oxide/nitride zones reaches values close to the partial pressure of nitrogen in the gas atmosphere (≈ 1 bar) since there are no stable nitrides within the oxide zone but only diffusion of dissolved nitrogen. In this context, it should further be noted that transformation of aluminium nitrides into oxides at p(N₂) = 1 bar necessitates an oxygen partial pressure of 10⁻³³ bar. Such an oxygen partial pressure is remarkably close to the onset of oxide formation for manganese and, to some extent, also for chromium (Fig. B.8). This is also indicated by the slight overlap in the computed oxide and nitride phase distributions in Fig. 1. Very little evidence was found on the miscibility of nitrides in literature [34–36]. Hence, the formed nitrides will indeed separate the alloy elements whereas oxides can thermodynamically form mixed spinel phases - if not already spatially separated by previous nitride formation. The bulk of the alloy contains carbides, which are stable at more than 45 μm beneath the alloy surface. Again, a “push-effect”, mimicking migration of carbides by nitrides, is indicated. However, in contrast to the above described transformation of nitrides into oxides, the released carbon is enriched underneath the nitride zone which manifests itself in an increase of the carbide content directly underneath the nitride zone. This behaviour is well known and very common during nitride hardening processes [33,37]. In extreme cases, the local activity of carbon reaches values by which a small peak of dissolved carbon in the iron matrix can be observed (see the region of dissolved carbon in Fig. 1). This fraction of mobile carbon diffuses towards the bulk of the sample and forms the sudden hump in the carbide profile at around 48 μm below the surface.

The simulation results for the depth of the bulk oxide zone (5–6 μm) agree fairly well with experimental results from Fig. 2 (1–3 μm). A pronounced oxidation along grain boundaries can be observed, which matches with the formation of silicon-rich oxides. The distributions of Mn, Cr and Al were not measured due to limited NanoSIMS measurement time and the interested reader is referred to another publication that discusses the oxidation behaviour in ternary model alloys [9]. Although it has been shown in previous studies that the oxidation along grain boundaries follows tabulated diffusion values, it should be noted that diffusion coefficients along grain boundaries are 100–1000 times larger than bulk diffusion coefficients which have been used for the simulations in Fig. 1.

In the same work [9] it is also mentioned that for internal oxidation, the bulk diffusion of oxygen is slowed down, which explains the lower bulk oxidation depth of ≈ 2 μm in Fig. 2 as compared to the 5–6 μm.
wide zone in the simulations of Fig. 1. A clear difference between simulations and experiment can be seen for the existence of the nitride zone, which is absent in the experimental data. This discrepancy is caused by the formation of an external scale during the industrial processing. This (oxide) scale layer is known to inhibit transport of nitrogen from the gas phase into the material [17]. In contrast to this, a maximum uptake of nitrogen from the gas atmosphere was assumed in the simulations, depending on the nitrogen partial pressure. Albeit only indicative, it should be noted that the concentration of nitrogen in the zone of oxides appears to be slightly lower than deeper inside the material. Here, the nitrides seem to have mixed with carbides that emerge below the oxidised zone. Larger precipitates can be observed in this area, and the lamellar structure of the carbon distribution within individual grains indicate the presence of pearlite [38] in the alloy. A strong segregation of sulphur and phosphorus at grain boundaries has also become evident from the NanoSIMS measurements (Fig. 2).

Detailed scans of the oxide-affected zone are shown in Fig. 3, which reveal the difference between inner oxides and grain boundary oxides. The concentration of sulphur and phosphorus along grain boundaries is lower in close proximity to the surface, which indicates that both elements also suffer from a “push-effect” due to oxide formation (e.g. transformation of MnS to MnO [18]) because the concentration of sulphur and phosphorus clearly reaches a maximum near the internal oxidation front. This observation qualitatively underpins the assumption from literature that sulphur and phosphorus are pushed inwards as the front of oxides/nitrides and carbides grows, since the concentration of sulphur and phosphorus suddenly drops towards the center of the alloy.

TEM images of heat-treated samples (Fig. 4) reveal a fine dispersion of 50–100 nm sized precipitates near the internal corrosion front. As can be seen from the corresponding elemental scans, both nitrides and oxides have been formed. Here, the distribution of oxides and nitrides is not homogeneous but instead a distinct peak of nitrides is exhibited in the center of the grain boundary domain (i). According to the higher thermodynamic stability of oxides relative to nitrides (see Fig. A.7 in the appendix), this nitride peak could be explained by the partial substitution of nitrogen by oxygen during oxidation. Consequently, bound nitrogen is released into the material [24,30] or pushed to the center of the grain boundary precipitate where the nitrogen activity exceeds the critical limit to stabilise the nitride phase. According to the elemental scans of manganese and chromium in Fig. 4, it is likely that the center of this particular grain boundary is composed of manganese/iron-rich nitrides. Since the concentration of aluminium and silicon was below the limit of detection in this measurement, their presence in minor amounts, however, cannot be safely excluded in these nitrides as well. In contrast to many literature sources [10,6,5,39,40], which assume the presence of the thermodynamically most stable spinel (Fe1-x-yMnxCry)O4 for internal oxidation of manganese-chromium steels, the distribution of chromium and manganese in Fig. 4 shows a small overlap of both elements, indicating the presence of binary Mn and Cr oxides, respectively. It is unlikely that this separation was caused by a decomposition of the spinel phase during cooling from
Fig. 2. NanoSIMS secondary ion signal intensity distribution images of selected elements in an industrial grade Fe-1.8Mn-0.8Cr steel after a complete coil cooling cycle of 4.5 h. The scale/metal interface of the steel sample is located close to the top edge of the images. Polished cross sections were prepared with a 25° tilt angle, to enlarge the vertical view of the oxidised zone by a factor of 2.37. Signal intensities, given in counts per second, are displayed on a logarithmic false-color scale, ranging from dark blue to red for low to high intensities, respectively.

Fig. 3. Detailed NanoSIMS secondary ion signal intensity distribution images of selected elements in the near-surface region of an industrial grade Fe-1.8Mn-0.8Cr steel after a complete coil cooling cycle of 4.5 h. The scale/metal interface of the steel sample is located close to the left edge of the images. Cross sections were prepared with a regular 90° angle. Signal intensities, given in counts per second, are displayed on a logarithmic false-color scale, ranging from dark blue to red for low to high intensities, respectively.
700 °C down to room temperature. The resulting Cr₂O₃ and (Fe₁₋₅Mnₓ)O would still be more evenly distributed throughout the internal oxides [18,9,39].

Two-dimensional elemental maps of another grain boundary close to the internal corrosion front (see Fig. 5) confirm the separation of manganese and chromium. Here, the center of the grain boundary is composed of chromium-rich oxides which again show a small overlap with the surrounding manganese oxides. Small amounts of nitrides were detected in the center of the grain boundary and at the outer periphery of the manganese-rich oxides. The higher thermodynamic stability of oxides (see Fig. A.7) suggests that oxygen diffused along grain boundaries and substituted nitrogen in the formed precipitates. Arguably, the continuous band of chromium oxides along the original grain boundary might indicate that this phase was formed first. However, since the equilibrium conditions of manganese and chromium oxide/nitride formation are very similar in this alloy, the initial formation of chromium may have simply been caused by lattice misfits that affect nucleation and growth of formed oxidation products [10,41–43].

Remarkably, the results from Figs. 4 & 5 give rise to the hypothesis that transport properties along oxidised grain boundaries are varying. Whereas the vast majority of available literature assumes the properties of grain boundaries to be homogeneous (due to lack of detailed data), the observed spatial oxide/nitride distributions clearly indicate that element transport attains another level of complexity that is beyond the reported values and texture dependence of grain boundary diffusion coefficients for high temperature corrosion simulations [10,23]. Oxygen transport along grain boundaries may thus not occur through the entire grain boundary, as proposed by the classical diffusion models of Wagner [26] and Fisher [44,45] but preferentially takes place at the oxide/metal interface. Interfaces and grain boundaries are often the origin for fast transport in ionic crystals [46,16,47]. As a consequence, the diffusion model by Wagner and Fisher may need to be refined such that the grain boundary domain is further separated into "interface" and "bulk oxide" regions. Although this would theoretically explain why nitrogen has been found in the center of grain boundary precipitates, it should be noted that a definite conclusion on the oxygen transport mechanism along grain boundaries cannot be drawn purely from the obtained results. The separation of manganese, aluminium and silicon in internal oxides follows similar trends as observed for grain boundaries (Fig. 6). Due to the smaller size of the inner oxides (20 nm) and hence shorter diffusion paths, the separation of the alloy elements is smaller than within grain boundary oxides (90 nm).

Fig. 4. TEM-image (left) and EELS profile (right) of a grain boundary oxide (i) and internal oxides (ii-iv) in a hot-rolled Fe-1.8Mn-0.8Cr industrial-grade steel sheet after the completed coil cooling programme. The direction of the grain boundary is indicated by the dashed line in the left TEM-image and located close to the internal corrosion front.

Fig. 5. TEM-EELS elemental maps of (left) manganese (green), chromium (red) and (right) oxygen (blue), nitrogen (green) near the grain boundary of a hot rolled Fe-1.8Mn-0.8Cr industrial grade steel sheet after the completed cooling programme. Note the marginal overlap of the elements in both images. The dashed yellow line in both graphs indicates a string of chromium-rich oxides and the direction of the original grain boundary close to the internal corrosion front. These chromium-rich oxides are thermodynamically stable but may originate from pre-existing nitride and/or carbide precipitates that formed first during heat exposure.
The center of the grain boundary in Fig. 6 contains a high amount of aluminium, which has a high affinity to both nitrogen and oxygen (Fig. A.7). It is thus assumed that this phase was formed first. Again, according to predominance diagrams (Fig. B.8), an oxygen partial pressure \( \sqrt{p(O_2)} = 10^{-33} \) bar is required for transformation of AlN into \( Al_2O_3 \) at 700 °C and a nitrogen partial pressure of 1 bar. This value matches almost exactly with the oxygen partial pressure of the Mn/MnO and Cr/Cr_2O_3 equilibria. For the upper limit of the oxygen partial pressure \( (\sqrt{p(O_2)} = 10^{-22} \) bar) defined by the iron/wustite equilibrium at the alloy/oxide scale interface, the calculated nitrogen partial pressure to stabilise aluminium nitrides is much higher than \( p(N_2) = 10^{-15} \) bar. This means that based on bulk thermodynamic principles, no nitride phases are expected to be stable near the surface because the thermodynamic nitrogen partial pressure required would be far too high. However, closer to the internal oxidation front, the presence and/or absence of sufficient amounts of manganese and chromium will indeed affect the transformation of nitrides into oxides and hence the structure of the formed grain boundary precipitates.

5. Conclusions

High temperature oxidation studies and numerical simulations of phase stabilities for an industrial low-alloy Mn-Cr steel have been carried out. Numerical simulations of spatial phase distributions, in which more than 40 thermodynamic phases were considered, show three clearly separated zones from the surface to the core containing oxide, nitride and carbide precipitates in the alloy. The values of the oxidation depth agree reasonably well between theoretical predictions and experimental observations but deviate for the amount of formed nitrides. This deviation originates from the presence of an external wüstite scale at process temperatures, which serves as an oxygen reservoir for selective oxidation but blocks further nitrogen uptake from the reaction gas. High resolution images of internal precipitates clearly show heterogeneities in their chemical composition instead of a homogeneous spinel-type oxide. Grain boundary oxides often seem to be composed of a layered structure with chromium, aluminium and nitrogen in the center, surrounded by manganese-rich oxides. This layered structure has not been reported before and is likely caused by distinct thermodynamic stabilities of the oxides which show similar oxygen activity values for manganese and chromium as well as for aluminium and silicon. However, the thermodynamically most stable spinel phases which are often speculated to be present at grain boundaries, are only observed in minor amounts. This is potentially due to the initial formation of immiscible alloy element nitrides which are eventually transformed into oxides. This fine-structure of oxides along grain boundaries may be an indication that oxygen transport is not uniform across the entire grain boundary domain as proposed by Fisher. Whereas the same spatial separation between alloy elements exists for inner oxides as well, it appears to be much weaker compared to oxidation products formed along grain boundaries. In addition to the detailed analyses of oxygen, nitrogen, carbon and the alloy elements, it could also be shown that a strong segregation of sulphur and phosphorus exists along the grain boundaries near the internal oxidation front. Finally, it should be noted that the experimental investigations in this study were carried out on an industrial-grade steel. In contrast to perfectly homogeneous binary and ternary model alloys, industrial alloys contain a multitude of elements and are prone to small variations in local composition. Accordingly, several replicate analyses have been carried out and discussions have focussed on the most common observations. Future efforts are suggested to consider and characterize the microstructure of formed precipitates in model alloys more into detail.

CRediT authorship contribution statement

Vera G. Praig: sample preparation, experiments, discussions. Michael Stöger-Pollach: TEM/EELS measurements. Arno Schintlmeister: NanoSIMS measurements, discussions, publication writing, review. Michael Auning: simulations, discussions, publication writing, review. Herbert Danning: discussions, publication writing, review.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

The authors thank Susanne Strobl and Kurt Piplits (both Technische Universität Wien, Austria) for their support with sample preparations and NanoSIMS measurements and Michael Wagner, head of the Large-Instrument Facility for Environmental and Isotope Mass Spectrometry at the University of Vienna (Austria) for kindly providing access to the NanoSIMS instrument. Scientific discussions and industrial inputs from Bernhard Linder and Dieter Paesold (voestalpine Stahl GmbH, Austria) are gratefully acknowledged.
Appendix A. Phase stability diagrams

Fig. A.7. Ellingham-Richardson diagrams of oxide (left) and nitride stability (right) of pure elements. The diagrams have been calculated with the FactSage steels database (FSstel [32,48]). Process temperature ranges for the heat exposure of industrial alloys in this study are highlighted in yellow.

Appendix B. Predominance diagrams

Fig. B.8. Predominance diagrams of oxide vs. nitride stability in binary iron-based alloys containing Al, Cr, Mn and Si at 700 °C. The diagrams have been calculated with the FactSage steels database (FSstel [32,48]). Thermodynamic conditions where the pure metallic alloys without any oxides/nitrides are stable, have been highlighted in yellow.
