1. Introduction

Deoxidation of steel using aluminum (Al) is an essential step in secondary steelmaking for the production of clean steel. The dissolution behavior of Al in molten steel and the liquid steel flow pattern determine the location where Al is released and, therefore, the Al deoxidation yield. Due to the large temperature difference and the heat transfer between Fe and Al, solid Al in contact with Fe heats up and partly melts. At the same time, liquid Fe solidifies at the interface, forming a shell. This solid shell grows to a maximum thickness controlled by a heat transfer balance. As the melting temperature of Al is much lower than that of steel, Al should melt completely inside the shell. During the shell period, interactions between solid steel and liquid Al may lead to the formation of Fe–Al intermetallic compounds, as predicted by the phase diagram of the system, i.e., αFe(Al), FeAl, FeAl₂, Fe₂Al₅, and FeAl₃. Based on the microstructure, the concentration profiles, diffusion processes in solid and liquid phases, and the behavior of Al₂O₃ inclusions in the reaction zone, it was demonstrated that the reaction zone was completely liquid and that the solid Fe shell melted for the samples with holding time of 30 and 60 s.

KEY WORDS: Fe–Al intermetallic compounds; deoxidation; Fe–Al–O ternary system; diffusion.

The phenomena taking place at the early stage of Al deoxidation were investigated with specific attention on the formation of Al₂O₃ inclusions. In a quartz tube, Al was exposed to liquid Fe containing different amounts of dissolved oxygen (O₂) for 1, 5, 30 and 60 s. In this paper, quenched microstructures of the diffusion couple were examined to identify the interactions and the nature of the phases at the experimental temperature. They revealed that the reaction zone was composed of successive layers of Fe–Al intermetallic compounds, as predicted by the phase diagram of the system.

Based on the microstructure, the concentration profiles, diffusion processes in solid and liquid phases, and the behavior of Al₂O₃ inclusions in the reaction zone, it was demonstrated that the reaction zone was completely liquid and that the solid Fe shell melted for the samples with holding time of 30 and 60 s.

KEY WORDS: Fe–Al intermetallic compounds; deoxidation; Fe–Al–O ternary system; diffusion.

The phase diagram of the binary Fe–Al system is shown in Fig. 1. The left hand side of the diagram shows that the solubility of Al in fcc γ-Fe is limited to a few percents. In disordered bcc α-Fe, up to 28 wt% Al can be dissolved depending on temperature. The corresponding phase is noted αFe(Al) in this text. Intermetallic phase formation can occur from 12 wt% Al on. Six non-stoichiometric intermetallic compounds are listed in the Fe–Al binary system. Depending on the Al content and temperature, Fe₃Al (D₀₃-ordered), FeAl (ordered B2), FeAl₂, Fe₃Al₂ (ε), Fe₅Al₃, and FeAl₃ are formed. Information on the physical, thermodynamic properties and phase reactions in the Fe–Al system are provided by Shahverdi. On the right hand side of the phase diagram, the solubility of Fe in solid Al is close to zero.

The microstructural analysis of interface layers that form when molten Al comes into contact with solid Fe or steel is mainly documented at temperatures ranging between 700 to 900°C. The interfacial reactions in this binary system have implications on many processes including aluminizing, mold casting, bimetals fabrication, joining and welding, etc. The types of interfacial phases formed, their morphology as well as their kinetics of growth were examined.
The major intermetallic phase identified is Fe₃Al₃ close to the steel substrate. Minor phases were also reported, principally the formation of Fe₅Al₈ close to the solidified Al. These studies highlight that the interfacial phases observed are not those given by the equilibrium phase diagram of the system. Some phases predicted by thermodynamic equilibrium are not found, indicating that the formation mechanisms and the diffusion conditions in the phases influence greatly the growth process and the overall interfacial structure at a given temperature.

This paper is the first part of an experimental study of the phenomena involved in the early steps of the deoxidation process, when Al rich melts are brought into contact with oxidized Fe for a very short time. It focuses on the comprehension of the interactions between Al and liquid Fe containing various dissolved oxygen contents of 160 to 1 800 ppm during the initial dissolution stage. These interactions define thermal history, mass transfer, concentration profiles and supersaturation degree, which determine the nucleation and growth conditions of the inclusions. The second part of the study discusses in more detail the formation and the morphology of the inclusions formed at the onset of deoxidation.

2. Experimental

One hundred grams of electrolytic Fe (99.97% Fe) was filled in an alumina crucible (30 mm ID, 35 mm OD, 50 mm H) and placed inside a high temperature vertical tube furnace (GERO HTRV 100-250/18, MoSi₂ heating elements) and melted at 1 600°C under purified Ar atmosphere. The oxygen content in the off-gas was measured with a solid state ceramic oxygen sensor (Rapidox 2100, Cambridge Sensotec Ltd.). A typical value of the oxygen concentration in purified Ar is about 10⁻¹⁸ ppm. The temperature profile between Fe and Al was slowly increased, approximately 4 cm in length, in which the temperature varies by less than ±1 K. The oxygen content in the melt was adjusted by the addition of reagent grade iron oxide powder (Fe₂O₃), followed by stirring with an Al₂O₃ rod. Since the impurity level of electrolytic Al is low, the dissolved oxygen (O) is assumed equal to the total oxygen content. The latter was determined by combustion analysis on the bulk Fe of the samples.

A piece of Al (99% Al) was cut in a cubic shape (~5 mm side), ground, pickled in 5% NaOH solution for 1 min, followed by washing in demineralised water, rinsing in acetone and drying. The Al piece was placed inside a quartz tube (8 mm inside diameter, 10 mm outside diameter). The last 3 cm of the quartz tube were narrower in order to fix the Al piece inside (6 mm inside diameter, 8 mm outside diameter). After 30 min at 1 600°C to stabilize the temperature of the melt, the quartz tube containing the Al piece was quickly introduced in the furnace and lowered into the liquid Fe. The Al piece was not preheated to avoid melting before the interaction. A small volume of molten Fe was sucked in the tube and brought into contact with Al (Fig. 2(a)). Except for sample 4, the contact between Fe and Al took place at the lower facet of the cubic piece of Al. In sample 4, the contact was made at a corner of the Al piece. During the interaction between Fe and Al, the quartz tube was maintained at that position in the liquid Fe, so that the material inside the tube was subjected to the furnace heat (Fig. 2(b)). After the desired reaction time, the quartz tube was rapidly withdrawn from the metal bath, removed from the furnace and quenched in water. Maximum 4 s elapsed during the sample withdrawal procedure. The contact time between liquid Fe and Al in the furnace hot zone varied from 1 to 60 s. The conditions for each test are listed in Table 1.

The upper part of the samples was cut along the longitudinal axis and prepared for Scanning Electron Microscopy investigation (SEM XL30 FEG). The SEM is equipped with an EDS spectrometer from EDAX with a Super Ultra thin window allowing the analysis of light elements from B, with which concentration distributions were measured.

3. Results

3.1. Macroscopic Observations

Macroscopic examination of the samples after the interaction shows that mixing between Fe and Al was not complete, even after 1 min. The Al piece has completely changed from the initial cube to a more rounded shape (Fig. 3). After 1 s interaction, the height of the remaining Al piece on top of Fe has not much changed (Fig. 3(a)). As a result of Al dissolution in Fe, it has considerably decreased after 60 s of interaction (Fig. 3(b)).

3.2. Evolution of the Reaction Zone Microstructure

During the experiments, a diffusion couple is initiated between Fe and Al (Fig. 4). Interdiffusion of Fe and Al re-
results in the formation of a reaction zone (Fe–Al), bounded by two diffusion fronts indicated by the dashed lines in Fig. 4. Diffusion front 1 is located at the interface between the reaction zone and Fe. Diffusion front 2 is located at the reaction zone/Al interface. We are mainly interested in the interactions occurring at diffusion front 1. Therefore in this text, diffusion front refers to the Fe/Fe–Al interface.

The diffusion couple involves O, which is present in the liquid Fe. Due to their high affinity, Al and O will react to form solid Al2O3 inclusions at the reaction front, which is indicated by the dotted line in Fig. 4. Since both Al and O must be available for the reaction to take place, the reaction front is located in the reaction zone. To simplify the microstructure description and the discussion, it is assumed that the Al2O3 formation reaction is instantaneous, that equilibrium is reached and that Al and O equilibrium compositions are negligibly small. With these assumptions, the position of the reaction front corresponds to that of the diffusion front, as will be shown during the investigation of the sample reaction zone.

The reaction zone consists of several product layers composed of intermetallic compounds in a sequence dictated by the phase diagram. The various product layers between the Al-rich region and bulk Fe were identified by SEM and EDS analysis. Based on the occurrence, the morphology of the product layers, differences were indicated by the dark phase in Fig. 5(b), presumably richer in Fe. This secondary phase would have a higher melting point than the FeAl phase due to the presence of intermetallic compounds FeAl and FeAl2 (eutectoid reaction). Higher resolution images of this intermetallic layer show that a fine eutectoid structure has formed (Fig. 6(b)), probably during quenching at the end of the experiment. Based on its composition ranging between 26 and 35 wt% Al, the ordered-B2 FeAl phase layer was identified next to the two-phase FeAl1 + FeAl2 layer towards the Fe-rich side. In the direction of the Al-rich area, the BSE contrast reveals that the FeAl grains in sample 1 are surrounded by a brighter phase (Fig. 7(b)), presumably richer in Fe. This secondary phase would have a higher melting point than the FeAl phase due to the higher Fe content, indicating that this structure is formed during quenching of the liquid. Finally, α(FeAl) with a composition ranging from 0 to 26 wt% Al and thickness listed in Table 1 was identified between the diffusion front and the FeAl product layer (Figs. 6(a) and 7(a)).

The Al-rich intermetallic layer sequence (above 60 wt% Al) in the samples with high initial O content (samples 4 to 6, 1800 ppm O) is similar to that observed in samples 1 to
3 (Figs. 8(a), 9(c) and 10(a), and Table 1). The main difference compared to the samples 1 to 3 is the presence of, towards the Fe-rich areas, a two-phase layer consisting of an Al-rich eutectic (FeAl₃–Fe₂Al₅, at approximately 58 wt% Al, 1 160°C), with a very fine eutectic structure formed during quenching (Figs. 8(b), 9(b) and 9(c)). Between the αFe(Al) and the two-phase layer (Fig. 8(b)), two layers can be identified by the BSE contrast. The lower layer consists of FeAl and the upper layer has a composition lying in the domains FeAl₁₋₃FeAl₂ and FeAl₂. The thickness of the layers in samples 4 to 6 is provided in Table 1. Neither microsegregation nor a eutectoid structure was detected in these layers (Figs. 8(b), 9(a) and 9(b)). As regards sample 4, contact between Fe and the Al piece was made at a corner of the Al piece,
resulting in Fe penetrating deeper in the Al, as seen in Fig. 10(a). Though the sequence and the morphology of the solidified intermetallic layers are identical to those in samples 5 and 6, the layers are much thicker compared to other samples with similar holding time, as seen in Table 1. Owing to the high initial $O$ content in the Fe, numerous $Al_2O_3$ inclusions are found distributed through the solidified intermetallic layers (Figs. 9 and 10(b)). Near the diffusion front, the inclusions are mostly individual (Figs. 9(a) and 10(b)). In sample 5, nm sized $Al_2O_3$ inclusions were found (see Sec. 3.3, Fig. 12(a)). Those in samples 4 and 6 have an average size of, respectively, 0.7 and 2 µm (Fig. 12(b) and 12(c)). Towards the Al-rich area, $Al_2O_3$ inclusions tend to form aggregates (Figs. 9(b), 9(c), and 10(b)).

### 3.3. Concentration Gradient

Figure 11 shows the Al concentration profile measured on the samples with various interaction time and initial $O$ contents as a function of the distance from the diffusion front. The concentration profile was measured perpendicular to the diffusion front. With a contact time of 1 or 5 s (samples 2 and 5, Fig. 11(a)), the Al profiles, which are shown only for the first 100 µm, are very steep until the first 20 µm. The concentration is then slowly increasing. The effect of the initial $O$ content is hardly noticeable. The less steep Al concentration profile near the diffusion front measured in sample 4 is due to a different position of the Al inside the quartz tube, creating a two-dimensional and thereby faster diffusion process compared to the other samples. The Al profile in sample 3 is not shown due to lack of measurement data.

Figure 11(b) shows part of the Al concentration profiles in samples 1 and 6, with a longer interaction time. The Al profile extended over a larger area due to longer diffusion time compared to samples 2 to 5. The concentration profile for sample 1 (30 s, 160 ppm $O$) is steep over approximately 100 µm near the diffusion front, though less steep compared to samples 2 to 5. At a distance between 100 and 170 µm from the diffusion front, it remains almost constant at 10 wt% Al. Beyond that point, the Al concentration increases significantly again. The fluctuations in the concentration profiles are due to local inhomogeneities, especially with the presence of two-phase layers. The shape of the Al profile of sample 6 (60 s, 1 800 ppm) is similar to that of sample 1, yet different in magnitude. The Al profile is steep within approximately 120 µm, remains nearly constant at approximately 21 wt% Al for another 10 µm and increases again, almost linearly. Near the interface, the Al profile of sample 6, with longer interaction time, is steeper than that of sample 1, with shorter holding time. With an identical diffusion process, the profile at the interface is expected to become less steep with holding time. It is believed that the difference comes from the $O$ content in the bulk Fe which is significantly higher in sample 6.

Because the $O$ level in the Fe matrix was below the detection limit of the EDS system, the $O$ concentration profile near the interface could not be quantified. However, for samples 4, 5 and 6 in which the initial $O$ content of the liquid Fe was high, the presence of an $O$ gradient near the interface is suggested by the formation of a “precipitate-free” zone separating the Fe–O from the Fe–Al domain (Fig. 12). In the Fe–O domain, numerous FeO inclusions are found, due to high $O$ content in liquid Fe and the decreasing $O$ solubility in Fe with temperature. The absence of precipitates means therefore a lower $O$ content at high temperature. The extent of liquid Fe with low $O$ content is determined by the
thickness of the precipitate-free zone. As most of the mass transport under the experimental conditions is assumed to be driven mainly by diffusion processes, a thin precipitate-free zone corresponds to a steep O content profile near the diffusion front. In the Fe–Al layer next to the interface, only Al2O3 inclusions are found, resulting from the diffusion of Fe, Al and O and the deoxidation reaction at the diffusion front when Al and O meet. As seen in Fig. 12, the thickness of the precipitate-free zone is clearly varying between the samples. For sample 5, this zone is 1.5 to 2.5 μm thick (Fig. 12(a)), whereas it reaches approximately 17 μm for sample 4 (Fig. 12(b)) and 63 μm for sample 6 (Fig. 12(c)). After comparing these values with the Al concentration profiles given in Fig. 11, the precipitate-free zones tend to be thinner when the Al gradient is steeper near the interface. The latter is due to the deoxidation reaction taking place at the interface, which determines the relation between the O and the Al gradients at both sides of the diffusion front in order to satisfy the mass balance at the diffusion front and the reaction stoichiometry.

4. Discussion

4.1. Mass Transport of Fe in the Al-rich Region

Bouché et al.12,13 also observed a thin needle-shaped FeAl layer uniformly dispersed in the Al matrix. The formation of these crystals is attributed to a eutectic reaction during solidification of liquid Al containing some dissolved Fe. Rapid solidification conditions in hypereutectic Al alloys containing Fe can lead to the formation of Al solid solution as primary phase to solidify, with intermetallics forming in the intercellular region.17 This transition is due to kinetic effects.

The presence of Fe in the Al solid solution is an indication that diffusion of Fe in Al occurred even when the immersion time was very short. The diffusion of Fe atoms in the Al-rich area seems to be much faster than the conventional diffusion process. According to Shahverdi et al.,10 Fe is brought to the Al-rich area through the breakage, flotation, re-melting and dissolution of FeAl3 into molten Al. Under the present conditions, the flotation of Al-rich intermetallics in liquid Al seems not possible owing to their higher densities compared to liquid Al.15 Instead, turbulent flow might be initiated by the large temperature difference between liquid Fe, initially at 1 600°C, and solid Al, initially at 25°C, combined with the furnace heat surrounding the quartz tube (Fig. 2(b)) and reheating the charge. In the case of turbulent flow, mixing is significantly accelerated and Fe can be transported faster in the liquid phase. This mechanism is likely responsible for the Fe transport to the Al-rich region, indicating that the latter was in the liquid state during the interaction.

Si was not measured (or detected) in Fe nor in Al. SiO2 from the quartz tube could not be reduced by Al due to the short contact time and the limited contact surface between Al and the quartz tube.

4.2. Influence of O on the Diffusion Process and the Microstructure

The influence of O on the Al concentration profiles was noticeable for samples 1 and 6 with longer holding times. As shown in Fig. 11(b), an increase in the O content seems to slow down the Al penetration, resulting in a steeper Al concentration profile compared to the case with low O content. Since Al cannot diffuse in oxidized Fe, O, which is transported faster than Al towards the diffusion front by concentration gradient and surface tension effects, must be consumed by the reaction before Al can diffuse further. The process of Al penetration is therefore hindered with higher O content, resulting in a steeper Al profile near the diffusion front.

The two-phase layer FeAl+FeAl3 with a eutectoid structure in samples 1 to 3 (Fig. 6(b)) was not observed in samples 4 to 6. Instead, a single homogeneous phase was found in that composition range and an Al-rich eutectic formed the Al-richer area. The reasons for this are not clear. The main differences between these samples are the O content and the numerous Al2O3 inclusions distributed in most of the solidified layers. The presence of Al2O3 inclusions brings an additional element in the system, O. The ternary system Fe–Al–O should be considered instead of the binary system Fe–Al. In some alloys, such as Al–Ti alloys,19 O influences the solidification behavior by shifting phase transitions and extending stability domains compared to binary systems. Further work is needed to clarify the influence of O on the solidification behavior in Fe–Al alloys.

4.3. Identification of the Solid and Liquid Phases at Experiment Temperature

4.3.1. Reaction Zone

It was shown in Sec. 4.1 that the upper Al-rich part of the sample should be in the liquid state to enable fast transport of Fe in liquid Al. The observation of the microstructures revealed a typical quenched-liquid structure in all samples, especially in the eutectic and eutectoid composition range. Considering that the quenched eutectoid structure forms during solidification of the ρ phase (FeAl3), the local temperature during the experiment should have been at least 1 102°C, according to the Fe–Al phase diagram (Fig. 1). Moreover, assuming that the region of the sample with a composition lying in the FeAl3 domain was liquid, the temperature in that area should have been above 1 160°C according to the Fe+Al phase diagram (Fig. 1). No evidence of a liquid, solid or mushy zone was found for the αFe(Al) and FeAl layers.

The growth rate of the intermetallic layers during interaction between solid Fe and liquid Al is documented in the temperature range of 700 to 1 050°C.10–13,20–25 After 1 min hot-dip aluminizing treatment at 800°C, the maximum reported layer thickness of FeAl3 is about 200 μm.12 It attains about 700 μm after 45 min at 800°C.11 After a diffusion treatment of the aluminized specimen during 15 min at 1 000°C, αFe(Al) and FeAl layers were formed and attained a thickness of, respectively, 32 and 16 μm.20,21 The growth kinetics of the intermetallic layers is reported to be diffusion-controlled, following a general parabolic equation11–13,20,25 from which activation energies for the growth of FeAl3, FeAl and αFe(Al) were evaluated. The thickness of αFe(Al) and FeAl heated at, respectively, 1 811 K and 1 583 K, which are the respective melting point of these compounds, were calculated for diffusion times of 35 and 65 s (samples 1 and 6), using the reported growth
law relations.\textsuperscript{29} Five seconds were added in the calculations to consider the time during the sample withdrawal procedure. These temperatures and durations are employed in the calculations to obtain an estimate of the maximum possible layer thickness. As seen in Table 2, the calculated thickness is much smaller than the observed one, indicating that the reaction zone could not be formed entirely by growth of solid intermetallic compounds. Most of the reaction zone results from the solidification of a liquid phase, which corroborates the microstructure observations.

In binary systems, only single-phase intermetallic layers, which are separated by straight interfaces, can occur in the reaction zone under isothermal conditions.\textsuperscript{16} In the present observations, some areas in the reaction zone consist of more than one single-phase. Also the Al concentration profile across the reaction zone is very gradual (Fig. 11), which is not the case for a binary diffusion couple with the reaction zone in the solid state. Both arguments suggest that the reaction zone consists of a liquid phase. Moreover, the presence of solid intermetallic layers at the diffusion front would have limited the diffusion of Fe to the liquid phase, which is in disagreement with the large mass transport of Fe observed in the samples.

Numerous Al\textsubscript{2}O\textsubscript{3} inclusions were observed in the samples with high initial O\textsubscript{Fe} content. They were formed immediately behind the diffusion front. In the first tens of \textmu m from the diffusion front, mainly individual inclusions were found (Figs. 9, 10 and 12). Beyond that point, the inclusions tend to form aggregates through a collision-coagulation mechanism (Figs. 9 and 10). A collision-coagulation mechanism implies that the inclusions could move, which indicate that the metal matrix was in the liquid state. At the position where the aggregates are observed, the Al content reaches 33 wt\% for sample 4 and 20 wt\% for sample 1. The local temperature was therefore not less than, respectively, 1300°C and 1420°C, according to the Fe–Al phase diagram (Fig. 1). Moreover, the average size of the individual inclusions near the diffusion front attained 0.7 and 2 \textmu m in, respectively, samples 4 and 6 (Figs. 12(b) and 12(c)), which cannot be attributed to solid state precipitation.\textsuperscript{26} This also corroborates that this area is most likely liquid.

To summarize the discussion on the reaction zone, it was shown that most of it was in the liquid state during the experiment. In the case of samples 1 and 6 with longer holding times and of sample 4, it was concluded that the reaction zone was completely liquid before quenching. In case of samples 2, 3 and 5, it is unsure whether the thin layers of FeAl and \textalpha Fe(Al) next to the shell interface were liquid or solid.

### 4.3.2. Solid Fe Shell

The main consequence of the presence of a solid shell at the interface is the appearance of a significant discontinuity in the Al profile, owing to the large difference in mass transport between the liquid and the solid phase. The shell provides a barrier for the transport of Al, creating a step in the concentration profile. The height of the step gradually decreases as Fe dissolution progresses, yet the discontinuity remains as long as the shell exists. A steep Al concentration profile was measured at the interface with samples 2 and 5 (Fig. 11(a)), which may be interpreted as a step, indicating the presence of the solid shell at the other side of the interface. The measured Al concentration profiles showed a different behavior for samples 1, 4 and 6. The step was not present and the profiles exhibited a gradual increase in Al concentration. Al seemed to have diffused in Fe. This suggests that the solid Fe shell was not present anymore in samples 1 and 6. Under the present conditions, the shell melting time was less than 30 s.

In Sec. 3.3, it was shown that the presence of a precipitate-free zone suggests the formation of an O\textsubscript{Fe} gradient at the Fe side of the interface. The development of the O\textsubscript{Fe} concentration gradient in pure Fe implies that the O\textsubscript{Fe} should be able to be transported to some extent. The range of the diffusivity of O\textsubscript{Fe} in solid Fe\textsuperscript{26} is in general comprised between \(7 \cdot 10^{-5}\) and \(7 \cdot 10^{-7}\) cm\textsuperscript{2} · s\textsuperscript{-1} in the temperature interval 1538–600°C. In liquid Fe, \(D_0\) is reported to be \(1.4 \cdot 10^{-7}\) cm\textsuperscript{2} · s\textsuperscript{-1} at 1600°C,\textsuperscript{28} which is higher than in solid Fe. The O\textsubscript{Fe} content is very low in the cold Fe matrix (low solubility of O\textsubscript{Fe} in solid Fe, most of the O\textsubscript{Fe} is present as FeO\textsubscript{Fe} inclusions) and its diffusion in the solid phase is very limited. For that reason, one would expect the absence of the precipitate-free zone and the occurrence of FeO\textsubscript{Fe} inclusions at the diffusion front if Fe was in solid state next to the Fe/Fe–Al diffusion front. From the experimental observations on the samples with high initial O\textsubscript{Fe} content (samples 4 to 6), it is clear that this precipitate-free zone exists and that O\textsubscript{Fe} diffusion took place from the O\textsubscript{Fe} rich area to the Fe/Fe–Al diffusion front. Considering the short interaction time, Fe in the precipitate free zone at the Fe/Fe–Al diffusion front was most likely in liquid state to allow O\textsubscript{Fe} to be transported at a sufficiently high rate.

### 5. Conclusions

Inside a quartz tube, a piece of Al was brought in contact for a short time (1, 5, 30 and 60 s) with liquid Fe containing different O\textsubscript{Fe} levels at 1600°C. The purpose was to investigate the interactions between Fe and Al shortly after the deoxidation stage, as these interactions control the nucleation and growth conditions of the inclusions. The present work examined the quenched microstructures of the diffusion couple to identify the interactions and the nature of the phases at the experimental temperature. The reaction zone consisted of several layers of Al-rich and Fe-rich intermetallic compounds of the Fe–Al system.

Based on the microstructure, the concentration profiles, diffusion processes in solid and liquid phases, estimations of the intermetallic layer growth rates, and on the behavior of Al\textsubscript{2}O\textsubscript{3} inclusions in the reaction zone, it was shown that the latter was completely liquid and that the solid Fe shell melted for the samples with holding time of 30 and 60 s. For holding times of 1 and 5 s, the liquid or solid nature of

---

**Table 2.** Comparison of the calculated and measured thickness of \(\alpha\)Fe(Al) and FeAl for samples 1 and 6. The calculations were performed using growth law relations from Sasaki and Yakou\textsuperscript{20} at \(T = 1811\) K for \(\alpha\)Fe(Al) and at \(T = 1583\) K for FeAl.

| Sample | Calcd (\textmu m) | Measured (\textmu m) |
|--------|-------------------|----------------------|
| Sample 1 | 35                | 99                    |
| Sample 6 | 65                | 135                   |

© 2010 ISIJ
the thin layers of FeAl and αFe(Al), which are adjacent to the shell/reaction zone interface, could not be established.

Acknowledgements

The authors would like to thank ArcelorMittal Research and Development and the Belgian National Fund for Scientific Research (F.R.I.A.-F.N.R.S.) for their financial support.

REFERENCES

1) R. I. L. Guthrie, R. Clift and H. Henein: Metall. Trans. B, 6 (1975), 321.
2) S. A. Argyropoulos and R. I. L. Guthrie: Metall. Trans. B, 15 (1984), 47.
3) S. Sanyal, J. K. Saha, S. Chandra and C. Bhanu: ISIJ Int., 46 (2006), 779.
4) L. Pandelaers, F. Verhaeghe, B. Blanpain, P. Wollants and P. Gardin: Metall. Mater. Trans. B, 40 (2009), 676.
5) P. Gardin, J. F. Domgin, M. Simonnet and J. Lehmann: Rev. Metall. Cah. Inf. Tech., 105 (2008), 84.
6) K. Beskov, N. N. Viswanathan, L. Jonsson and Du Sichen: Metall. Mater. Trans. B, 32 (2001), 319.
7) M. Palm: Intermetallics, 13 (2005), 1286.
8) M. Hansen: Constitution of Binary Alloys, MacGraw-Hill, New York, (1958), 90.
9) U. R. Kettner: Binary Alloy Phase Diagrams, ASM International, Materials Park, OH, (1990), 147.
10) H. R. Shahverdi, M. R. Ghomashchi, S. Shabestari and J. Hejazi: J. Mater. Process. Technol., 124 (2002), 345.
11) A. Bouayad, C. Gerometta, A. Belkebir and A. Ambari: Mater. Sci. Eng. A, 363 (2003), 53.
12) K. Bouché, F. Barbier and A. Coulot: Mater. Sci. Eng. A, 249 (1998), 167.
13) H. R. Shahverdi, M. R. Ghomashchi, S. Shabestari and J. Hejazi: J Mater. Sci., 37 (2002), 1061.
14) V. I. Dybkov: J. Mater. Sci., 25 (1990), 3615.
15) M. A. Van Ende, M. Guo, J. Proost, B. Blanpain and P. Wollants: ISIJ Int., 51 (2011), No. 1, in press.
16) A. A. Kodentsov, G. F. Bastin and F. J. J. van Loo: J. Alloys Comp., 320 (2001), 207.
17) D. M. Stefanescu: Science and Engineering of Casting Solidification, Springer, New York, (2009), 273.
18) Y. Plevachuk, I. Egly, J. Brillo, D. Holland-Moritz and I. Kaban: Int. J Mat. Res., 98 (2007), 107.
19) J. Zollinger, J. Lapin, D. Daloz and H. Combeau: Intermetallics, 15 (2007), 1343.
20) T. Sasaki and T. Yakou: ISIJ Int., 47 (2007), 1016.
21) T. Sasaki and T. Yakou: Surf. Coat. Technol., 201 (2006), 2131.
22) T. Sasaki and T. Yakou: J Mater. Process. Technol., 197 (2008), 89.
23) Y. J. Li, J. Wang and X. Holly: Mater. Sci. Technol., 19 (2003), 657.
24) Y. J. Li, Y. L. Zhang and Y. X. Liou: J. Mater. Sci., 30 (1995), 2635.
25) S. Kobayashi and T. Yakou: Mater. Sci. Eng. A, 338 (2002), 44.
26) E. Schroer, S. Hopfe, P. Werner, U. Gösle, G. Duscher, M. Rühle and T. Y. Tan: Appl. Phys. Lett., 70 (1997), 327.
27) E. T. Turkdogan: Fundamentals of Steelmaking, Cambridge University Press, London, (1996), 134.
28) T. Isida and R. I. L. Guthrie: The Physical Properties of Liquid Metals, Clarendon Press, Oxford, (1993), 221.