

Demonstration of Elemental Partitioning During Austenite Formation in Low-Carbon Aluminium Alloyed Steel

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This work investigates the influence of aluminium, in solid solution, on austenite formation in a low-carbon aluminium alloyed (0.48 wt. %) steel during continuous heating. A thin section across an untransformed ferrite and austenite interface was prepared for transmission electron microscopy by focused ion beam milling. Microstructural characterization using imaging and elemental analysis demonstrates that aluminium partitions from austenite to ferrite during very slow heating conditions, stabilizing this latter phase and shifting the final transformation temperature for austenite formation (A_{c3}).

Keywords: Low-carbon steel, Austenitization, FIB-TEM.

1. Introduction

Austenite formation is an important and much studied process in steel research, being intensively investigated in low and medium carbon steels.. In ferritic-pearlitic steels this process can be split into two: nucleation of austenite in pearlitic colonies with rapid growth of austenite consuming pearlite followed by the slower growth of austenite consuming ferrite [1-6]. Studies of the effect of substitutional elements such as manganese and chromium on austenite formation have shown that they slow down the transformation when partitioning of the elements occurs during austenitization [2, 3]. There has been little research concerning the effect of aluminium as a substitutional element in steel in amounts higher than 0.1 wt. % despite a general awareness that Al is a ferrite stabilizer and tends to expand the ferrite + austenite phase field of the equilibrium phase diagram. This could be due to technological concerns over the production of high Al-content steels. Commercial interest has emerged recently with studies on Al additions, of the order of 0.5–2 wt.%, to low carbon high strength strip steels to produce a multi-phase

microstructure containing retained austenite for cold forming applications [7-10]. There is still limited knowledge regarding the role of Al as a solute in such steels; work is required to understand its influence on phase transformations and microstructures. This study investigates the influence of Al in solid solution on austenite formation during continuous heating.

2. Experimental procedure

Steels with a base composition of: 0.02 wt.% C, 1.41 wt.% Mn, 0.001 wt.% N and different concentrations of aluminium: 0.02 wt.% (Steel E) and 0.48 wt.% (Steel F) were produced. The initial microstructure of the steel is ferritic with some small pearlite islands. For more details regarding the processing, microstructure and sample preparation readers are referred to other publications [11-14]. A high resolution dilatometer (Adamel Lhomargy DT1000) was employed to determine the critical transformation temperatures for the end of the austenite formation process (A_{c3}); samples of 12 mm in length and 2 mm in diameter were continuously heated at 0.05 °C/s and 7 °C/s.

3. Results and discussion

Table 1 displays the A_{c3} temperature determined from dilatometry plots, for the two heating conditions. The addition of 0.48 wt % Al increases A_{c3} from ~910 to 987-1025 °C, consistent with Al acting as a ferrite stabilizer and expanding the ferrite + austenite phase field. The Al-alloyed steel (F), has an A_{c3} that is very sensitive to the applied heating rate, in contrast to steel E. For steel F, the A_{c3} increases (from 987 to 1026 °C) with a slower heating rate. Previous work on the effect of heating rate on A_{c3} in low and medium steels without Al show the opposite, a faster heating rate gives a higher A_{c3} since a faster heating rate takes the steel from equilibrium conditions [1, 15]. The most likely interpretation for the observation here is the diffusion or partitioning of Al to the untransformed ferrite grains, delaying their transformation to higher temperatures. Thus, at temperatures close to A_{c3} a greater concentration of Al is expected in the untransformed ferrite grains of an Al-rich steel (steel F); this would be most prominent at a slow heating rate (such as 0.05 °C/s) since this allows Al time to diffuse and partition. We have investigated Al levels (by energy dispersive X-ray spectroscopy [EDX] in TEM) in an untransformed ferrite grain and a former austenite grain of the partially transformed steel F (details below). It is well known that during austenitization heat treatment, partitioning of alloying elements may occur, that the extent is dependant on the relative solubilities in austenite and ferrite and that this phenomenon can affect the transformation kinetics in steel [2, 3]. It is well documented that annealing in the ferrite-austenite phase region can produce Mn partitioning [16, 17] however, there is little literature concerning partitioning of Al during the austenitization process. Koo et. al. [18] have shown that Al partitions under certain heating conditions during intercritical annealing (ferrite/austenite region) of steels alloyed with 1 wt.% Al and

Amirthalingam et.al.[19] have shown higher concentrations of Al in ferrite grains in welded, high silicon and high aluminium TRIP steels.

To study the austenite/ferrite interface in the Al added steel (F), a specimen (2 cm cube) was heated continuously at 0.05 °C/s up to 970 °C and water quenched to room temperature. Standard metallographic procedures were carried out to reveal the microstructure (Fig. 1a). To ensure a TEM thin foil sample containing an appropriate untransformed ferrite and former austenite boundary was produced site-specific focused ion beam (FIB) preparation was utilized. A TEM thin foil (thickness ~60 nm, determined by the ratio of inelastic to elastically scattered electrons in the electron energy loss spectrum) was prepared using a FEI NOVA200 dual beam FIB/SEM (Fig. 1b). A 30 keV Ga ion beam was used, and for final thinning the beam current was varied from 500 to 50 pA. An FEI CM200 TEM, fitted with a Gatan imaging filter and an Oxford Instruments EDX spectrometer was used to examine the FIB section.. EDX spectra were taken in TEM mode with a spot size of ~6 nm diameter (we estimate this increases to 10 nm after beam broadening by the specimen) and were quantified using Oxford Instruments' ISIS software (including an appropriate absorption).

The heat treated steel contains a mixture of large ferrite grains (approximately 10 % of the bulk) and relatively fine ferrite grains (approximately 90 % of the bulk; Fig. 1a and b). Due to the low carbon and high Al content of the steel, the hardenability of austenite at temperatures close to A_{c3} is extremely low, resulting in the unavoidable formation of ferrite, even at cooling rates greater than 300 °C/s (confirmed through dilatometry experiments [14]). Consequently, for this water-quenched specimen, the fine grained ferrite is presumed to be formed on quenching austenite grains formed at temperature and the large ferrite is presumed to be untransformed ferrite consistent with the volume fraction of ferrite and austenite at 970 °C based on dilatometry data. We have assumed that the interface between coarse and fine ferrite grains represents the interface between an untransformed ferrite and austenite grain at high temperatures and that the rapid quench has prevented any Al re-diffusion. This suggests that the rhomboidal indentation shown in Fig. 1b is located at the interface between an untransformed ferritic region (Grain A in Figs 1(b-d) and 2(a, b)) and a transformed, former austenitic region (Grain B). A thin foil of this identified boundary region was prepared for TEM by dual beam FIB (Fig. 1c). Electron diffraction confirms that the two grains in the FIB section are ferrite (Fig. 1d and 2a). Atomic lattice imaging of the section revealed a sharp grain boundary interface (Fig. 2b). The spatial distribution of Al and Mn respectively across the interface was determined by spot EDX (Figs 3a and b). The areal concentration of Al is significantly higher in grain A (mean 0.94 +/- 0.06 wt % Al) than grain B (mean 0.53 +/- 0.03 wt % Al). The variation in the areal concentrations of Al and also Mn could be either due to the fact that the Al and Mn distribution is inhomogenous within ferrite or, following the assumptions above, that we are examining

the dynamic at-temperature transformation interface. If the latter, such a distribution would be expected if partitioning/diffusion of Al into the untransformed ferrite phase is taking place at high temperature. MTDData [20] calculations indicate that the equilibrium Al concentration in both grains should be lower (grain A 0.7 wt % and grain B 0.48 wt %) and may be the case if observed at greater distances from the interface (i.e. in the bulk). The corresponding concentration profile for Mn is inverse to that for Al (i.e. it increases from grain A to B) as, in contrast to Al, Mn is an austenite stabilizer and so partitions to austenite (Fig. 3b). EDX elemental analysis was also performed across what is presumed to be a former austenite/austenite interface (in the region indicated with an arrow in Fig. 1a) and Fig. 3c shows the areal concentration profile of Al across this interface. As expected, the results do not show any significant difference in the level of Al between these two former austenite grains and is similar to the base composition of the steel.

4. Conclusion

This investigation shows an increase in A_{c3} temperature in the Al alloyed steel during slow (0.05 °C/s) heating as compared with fast (7 °C/s) heating. TEM-EDX analysis of an untransformed ferrite and former austenite interface shows that the rise in A_{c3} is due to partitioning of Al atoms from austenite to the untransformed ferrite, so further stabilizing this phase. In addition, the results show that partitioning of Mn to austenite takes place during slow heating, consistent with previous studies [2, 3, 16, 17] and is expected because Mn is a known austenite stabilizer.

Reference:

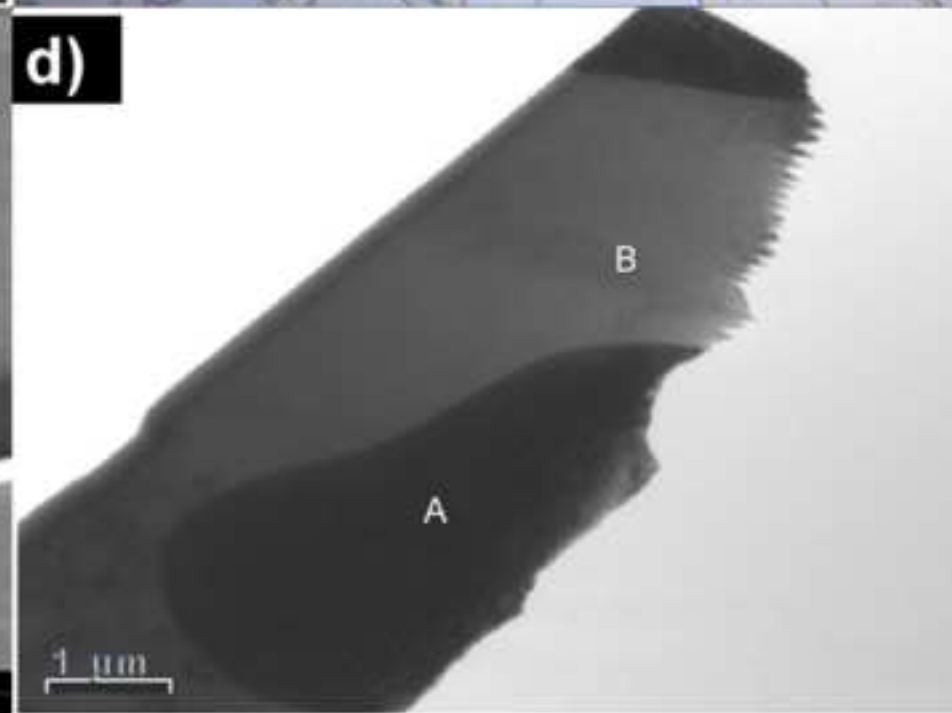
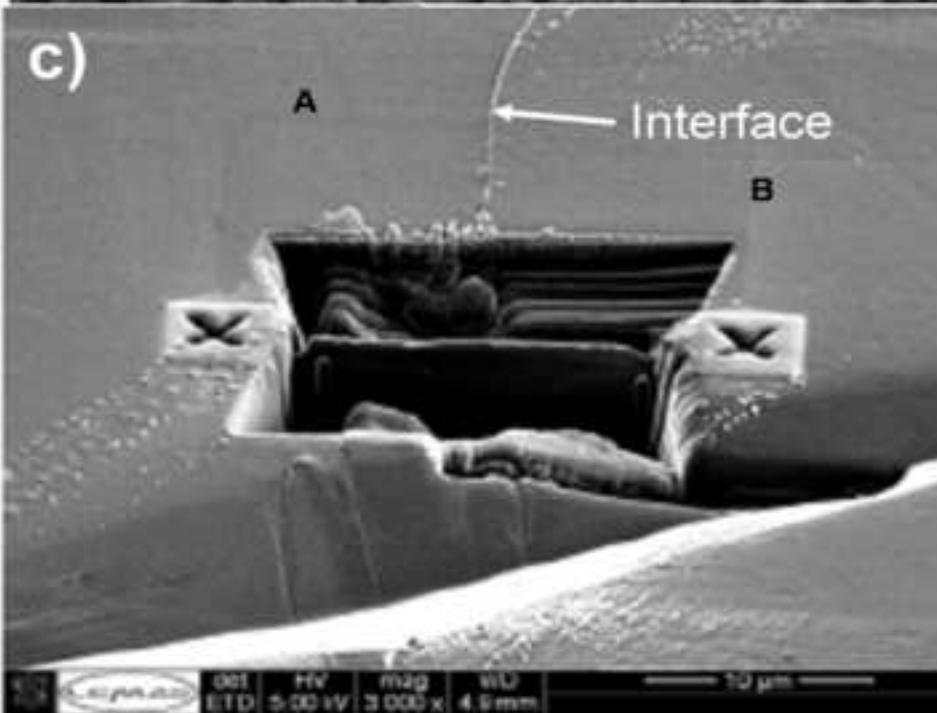
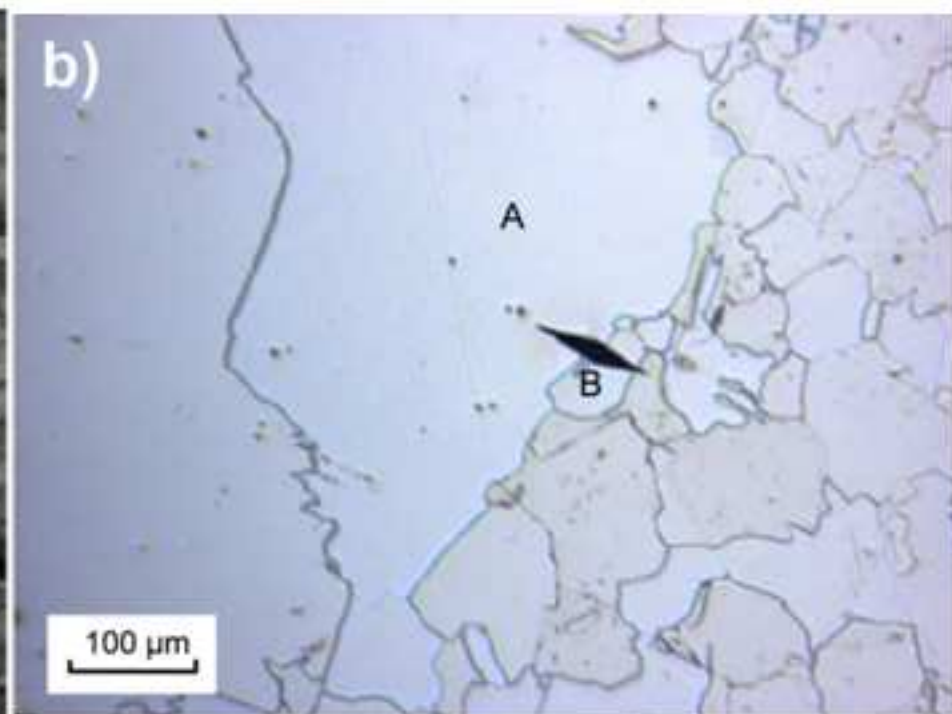
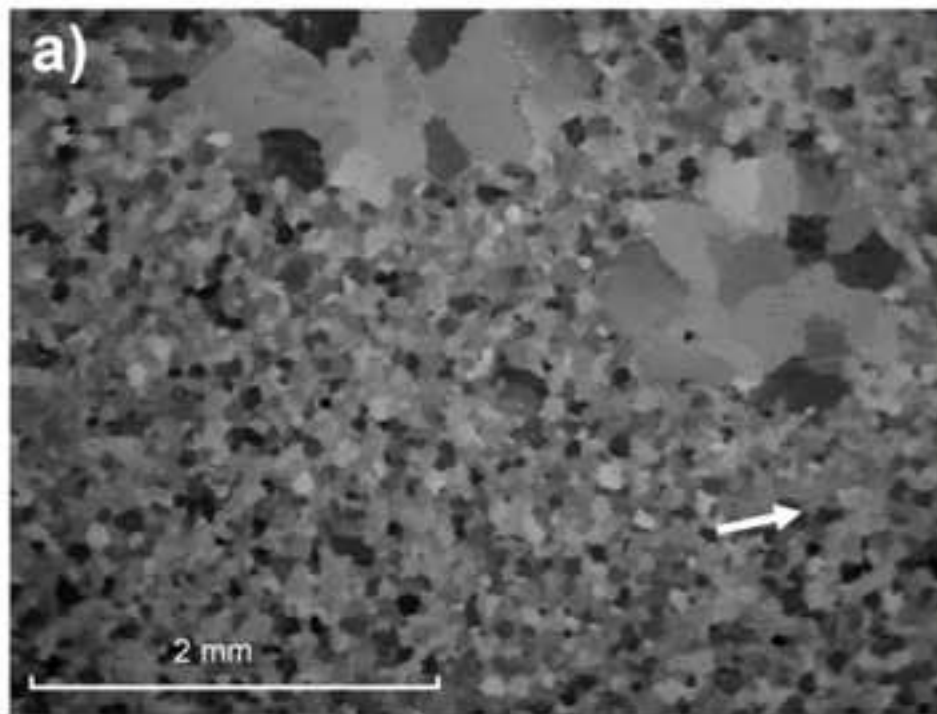
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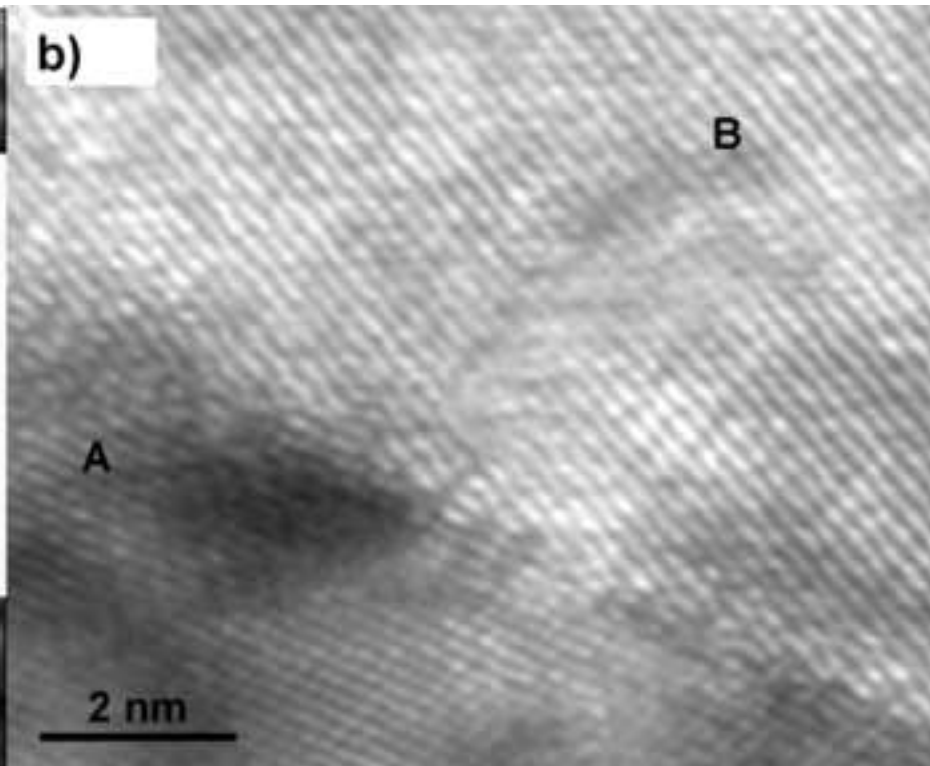
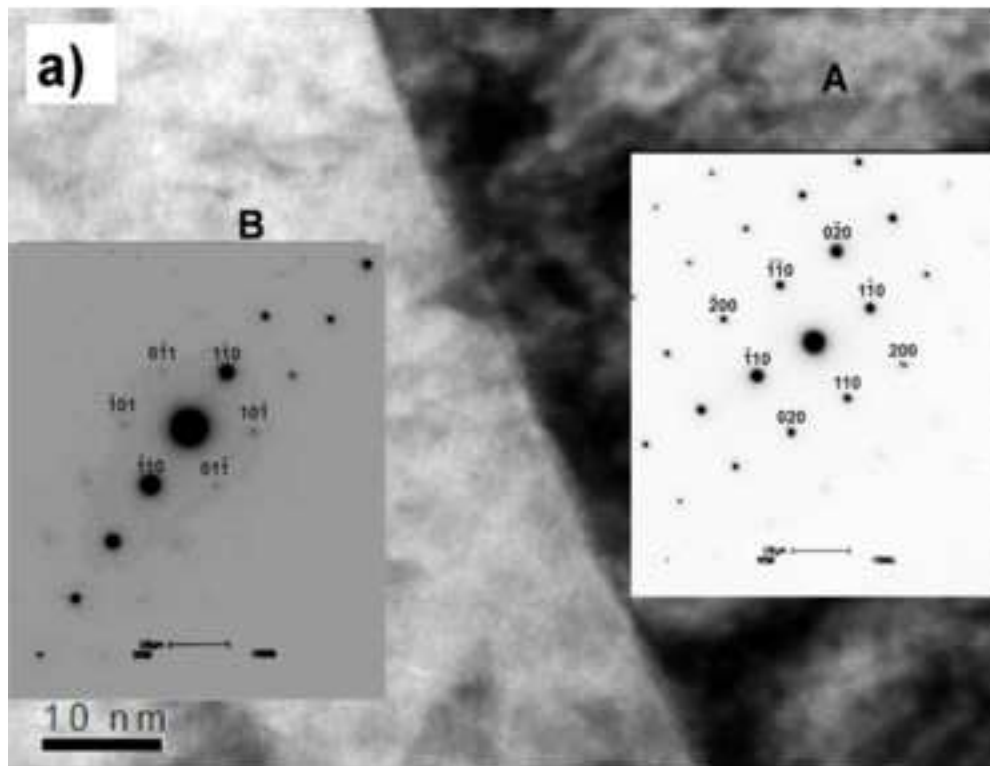
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Fig. 1 Micrographs of the microstructure of the heat treated Steel F: a) Back scattered electron image; b) optical metallography image of an interface between a ferritic region and a former austenitic region marked A and B respectively; c) secondary electron image recorded during FIB sectioning of the selected area marked by the rhomboid in b; d) FIB section imaged in the TEM at low magnification, grains A and B are marked.

Fig. 2 a) Bright field TEM image of the untransformed ferrite and former austenite interface identified in Fig. 1, inset are the selected area electron diffraction patterns from the two grains, both index to ferrite in [111] orientation in grain A and [001] orientation in grain B b) TEM lattice image showing that the interface is atomically abrupt

Fig. 3 Elemental concentrations (EDX) of Al (a) and Mn (b) across the former austenite/ untransformed ferrite interface and Al (c) across a former austenite/austenite interface.





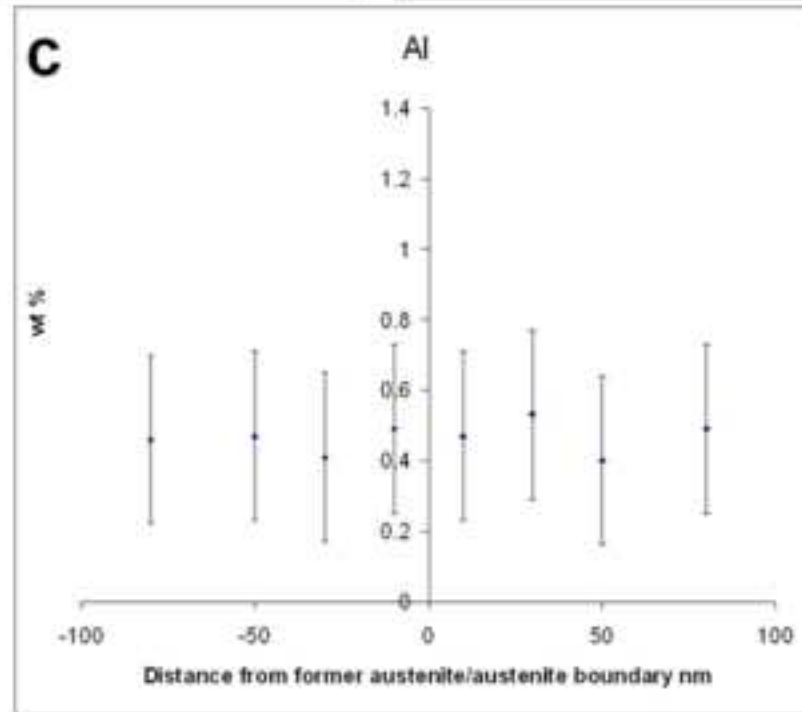
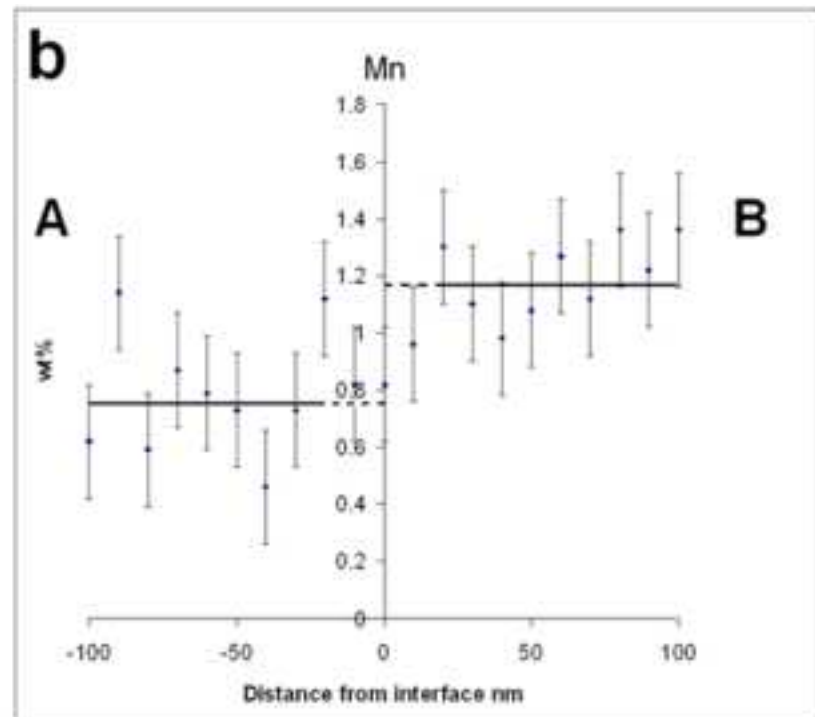
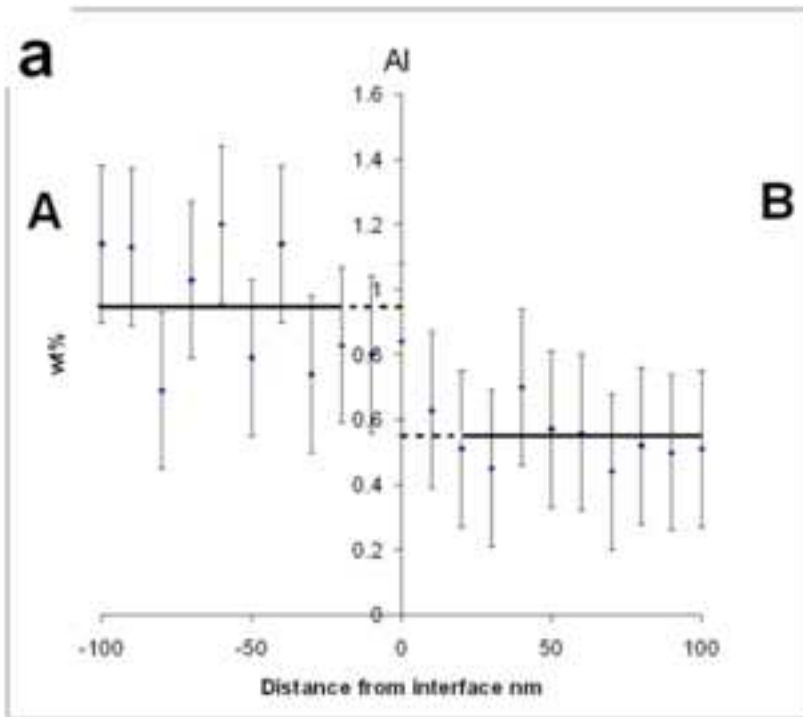


Table 1 Influence of heating rate on temperature A_{c3} for steels E (0.02 wt% Al) and F (0.48 wt% Al)

	Heating rate	
	0.05 °C/s	7 °C/s
Steel E: A_{c3}	910±7 °C	904±4 °C
Steel F: A_{c3}	1025±3 °C	987±2 °C