Grain Boundary and Microstructural Characterization of Heat-Treated As-Rolled 2205 Duplex Stainless Steel

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1. INTRODUCTION

Metal such as duplex stainless steel 2205 possesses an equivalent amount of ferrite and austenitic phases. The phases formed the basis for improved strength, including ductility [1-3]. Previous research has demonstrated that 2205 DSS has exceptional intrinsic and functional attributes. The outstanding characteristics of 2205 duplex stainless steel have attracted and continue to attract a vast number of users for different applications such as antimicrobials [4], industrial-marine environment [5], and hemocompatibility [6], to mention a few. Its extensive uses result from its alloying elements, such as molybdenum and chromium, which are resistant to corrosion and oxidation processes [7]. Due to these excellent properties, the DSS 2205 is broadly applied in the petrochemical, oil and gas, chemical, and desalination sectors, including maritime environments [7-9].

However, the intrinsic mechanical properties and functionality could improve or depreciate after heat treatment or hot deformation [10, 11]. The improvement or depreciation is caused by the change in the bulk materials’ microstructure, morphology and phase orientations. Although, the hot-rolling, heat treatment or hot-deformation process is known to cause changes in the characteristics of materials. For example, when magnesium ZEK100 alloy was subjected to hot rolling at temperatures from 350°C to 450°C, there was grain refinement as a result of thermo-mechanical processing, and it improved both the ultimate tensile strength and yield strength [12]. Elongated grains of the primary Alpha (α) phase at 790°C and Beta (β) at 870°C rolling temperatures were seen in high-strength beta titanium alloy [13]. β-stabilised Y-TiAl based alloy hot-rolled at temperatures; 1200°C, 1250°C and 1350°C showed that the grains became coarsened and elongated along the rolling direction [14]. It is also worth knowing that 10 Mn steel warm-rolled from 250°C to 600°C also caused changes in the microstructure and eventually caused improvement in the mechanical properties [15]. Heat treatment of laser melted 316L steel caused a lower fatigue life of the material [16]. A reduced hardening rate is observed at elevated temperatures for both A6061 and A6063 aluminum alloys [17]. These changes in properties are based on the interactions of the grains in the microstructures. The precipitation of "detrimental phases in the microstructure has been reported to either impact or have no effect on the yielding properties [18-20].

The interior structure of crystals is said to contain a network of interfaces known as grain boundaries. Its presence and design have an effect on a material’s mechanical characteristics and overall functionality [21]. The three forms of boundaries associated with materials are low angle grain boundaries (LAGB), high angle grain boundaries (HAGB), and special high angle grain boundaries (SHAGB). Tilt and twist angle grain boundaries are divisions of LAGB. HAGB are created when the dislocation cores of low-angle tilt boundaries cross at an angle greater than 15° [22]. These limits have a haphazard, or more accurately, uneven structure [23]. Boundaries with a high orientation and a high density of coincidence locations relationship are referred to as SAGB [24]. The rate at which grain boundaries move tells us what propels them in that direction [18]. This movement leads to the microstructure of polycrystalline materials. Even though the
physics of grain boundary migration is unclear, it thus depends on atoms passing through grain boundaries to gain energy. At low-angle grain boundaries (LAGB), migration is caused by cross- and slide-dislocations [25]. However, dynamic recovery happens due to grain boundary displacement because dynamic recrystallization does not proceed [26]. In low stacking fault metals, grain boundary migration results in dynamic recrystallization due to an increase in dislocation density, which restricts dynamic recovery [26]. Coherent, semi-coherent, and incoherent interfaces are the interfaces that can exist in metal. These groups of interfaces are visible in the equiaxed and Widmanstätten austenite morphologies of 2205 duplex stainless steel: the austenite-ferrite (A/F), ferrite-ferrite (F/F), and austenite-austenite (A/A) interfaces. Heterophase interfaces called A/F interfaces mimic metal interfaces, while solid-solid interfaces are associated with the A/A or F/F interfaces [27]. Furthermore, the resulting grain of A/F exhibits a Kurdjumov-Sachs (K-S) orientation relationship when phase transition occurs. This is indicated by a rotation of the angle of the crystallographic axis [28]. The orientation connection boosts its ductility by altering the deformation behaviour and hot workability of 2205 DSS [29].

There is still a dearth of investigation on the effects of heat treatment on the abolition of precipitation of hazardous phases on the DSS 2205 equiaxed and Widmanstätten austenite morphologies. Previous studies conducted did not critically unravel what happens at the interfaces and subgranular levels. They rather focused on the macroscopic behaviour of heat-treated hot rolled 2205 DSS in applications such as corrosion resistance [30, 31], strength, wear resistance [32] and crack resistance if inclusions are avoided [33]. Some researchers investigated grains evolution and interface relationship but did not examine changes in phase after heat treatment [34, 35]. Therefore, the 2205 DSS grains will be characterized to determine the boundaries, interfaces and phases that may exist after heat treatment which could further influence its properties.

Here, the microstructure of the heat-treated as-rolled 2205 DSS is characterized using three different image analyses, including electron backscatter diffraction (EBSD) technology which establishes a stereological-based analysis. It does measure all parameters and statistics associated with microstructure and micrographs, including mono- and polycrystalline materials [36]. The characterization would be supplemented by x-ray diffraction. To properly investigate the characteristics (grains, boundaries, interfaces and the microstructure) of 2205 DSS, at least two morphologies are required for adequate comparison and relative. These methods are used to demonstrate that the heat treatment eliminates detrimental phases and establishes the formation of both intragranular and intergranular grains. Also, additional phases, such as cementite and martensites, depend on the cooling after heat treatment.

The study will assist engineers and scientists in understanding the behaviour of this material when subjected to heat treatment and enable the development of an efficient microstructure and properties for a particular application.

2. MATERIALS AND METHOD

2.1 Material

Metal Centre in Gauteng, South Africa, supplied the 2205 duplex stainless steel sample. The as-received was in slab and then formed into a 10mm diameter cylindrical bar.

2.2 Heat treatment

The as-rolled material was solution heat-treated to form a DSS microstructure with equiaxed and Widmanstätten austenite morphologies. The treatment was carried out in a controlled environment using a tube furnace to avoid oxidation during heating. The material was heated at the rate of 10°C/s from 25°C to a region of single-phase delta ferritic with a temperature of 1340°C. After reaching 1340°C, there were 40 minutes of holding to ensure delta ferritic microstructure was attained. The specimens were then cooled in the furnace to 970°C at the rate of 0.002°C/s. The cooling was achieved in 48 hours, and then the samples were subjected to rapid cooling in water to achieve equiaxed morphology. The goal of rapid cooling from 970°C was to avoid the creation of harmful phases within 800 - 400°C. Figure 1 depicts a pictorial representation of the process. In the case of Widmanstätten morphology, the sample was rapidly cooled after solution treatment at 1340°C.

![Figure 1. Solution heat treatments of 2205 DSS to obtain equiaxed and Widmanstätten austenite morphologies](image)

2.3 Characterizations

For accurate characterization, the samples were prepared according to standards ASTM E3-01, ASTM E7-03, and ASTM E407-99. A ZEISS VisionSE64 optical microscope was utilized for phase identification. For the microstructure and grain boundary analysis, Merlin VP Compact scanning electron microscope equipped with HKL EBSD technology operated at 20kV, 7nA electron beam size and electron beam detector at 0.00° tilt was used. The comprehensive orientation maps’ dimensions were determined at 1.05mm in width and 1.4mm in height to investigate the substructure properties. The process had a dwell time of 19 μs. X-ray diffraction investigation was carried out on the heat-treated 2205 DSS in order to identify phase compounds. Using Rigaku Ultima IV, the patterns were acquired by scanning from 5° to 90° equal 2-theta angle at 0.5°/min with an XRD diffraction. The X-ray source’s voltage, current, and wavelength were set to 40kV, 30mA, and 1.54, respectively. The PDXL software was used to analyze the data.
3. RESULTS AND DISCUSSION

3.1 Microstructure

3.1.1 Scanning electron microscope

For the investigation of the grain distribution, an optical microscope was used at low magnifications, Figure 2. The images were taken at low magnification to ascertain the phases in the microstructure. Figure 2a represents the microstructure of the as-rolled 2205 DSS, while Figure 2b displays the Widmanstätten morphology and Figure 2c as the equiaxed microstructure. Both Figures 2a and 2b are attained after the systemic cooling down process.

The as-rolled 2205 duplex stainless steel grains in Figure 2a appear to be significantly pancaked in both ferrite and austenite grains. The ferrite grains, which are dark grey, are surrounded by light grey austenite grains that are plastically elongated in the direction of rolling. Due to compression during rolling, the austenitic grain showed mixed continuous and discontinuous lamellae structures. The exterior and core of the as-rolled showed to be compact and continuous, while in-between (core) morphology showed to be discontinuous though the grains were elongated. The occurrence of this kind of structure is a result of the existence of low stack energy between the austenite-ferrite when compared with ferrite-ferrite or austenite-austenite grain [37]. The Widmanstätten morphology in Figure 2b shows ferrite and austenite of needle-like structure as well as dendritic in nature as a result of the accelerated cooling after heat treatment. The Widmanstätten austenite nucleates discretely on the austenite allotriomorphs grain boundaries and also grows on the ferrite matrix by a diffusional mechanism with a close Kurdjumov–Sachs orientation relationship [38]. Kurdjumov-Sachs orientation is ideal because the lattice planes between the 100-planar of ferrite and the 111-planar of the austenite correspond. From Figure 2b, it is evident that both the intragranular and intergranular austenite are formed. The austenite grows in this region, and the defect-free phase barrier results in a good transition between the phases [36]. Figure 2c illustrates the ferrite and austenite grains produced when the sample was heated to 1350°C, allowed to homogenize for 40 minutes, and then quickly cooled to 1000°C to produce an equiaxed structure. After heat treatment, the equiaxed microstructure recovered from elongated structures back to rigid and discrete phases. Therefore, the equiaxed structure underwent recrystallization. The secondary austenite was generated as a result of the hot-rolling process and dissolved into ferrite in both morphologies. This process is enabled by the reduced Kurdjumov-Sachs fractions produced as a result of slow cooling associated with the microstructure. Therefore, it means that after heat-treating hot-rolled 2205 DSS, the secondary austenite-ferrite transition occurred prior to the primary austenite-ferrite transition, thereby making all the secondary austenite transmogrified to ferrite before the completion of heat-treatment.

3.1.2 Scanning electron microscope

The chemical composition was determined by energy-dispersive spectrometry (EDS) attached to the scanning electron microscope (SEM) as a first step toward identifying the phases in equiaxed and Widmanstätten structures [39]. The outcomes of EDS studies of the phases are shown in Figure 3 and detailed in Table 1. According to the EDS investigations, the primary balance phases of ferrite and austenite were correctly detected. According to previous studies, chromium, nickel, and molybdenum increase steel's ability to resist corrosion, including pitting and crack corrosion [9]. Chromium and molybdenum are ferrite formers, while manganese, nickel, and nitrogen are austenite stabilizers. The EDS study of the microstructure of the hot-rolled 2205 DSS demonstrates the partitioning of elements between the ferrite and the austenite, and it is consistent with the findings of ferrite transforming into austenite. In Table 1, the ferrite formers are distributed into the old ferrite phase according to their chemical compositions of 23.10 and 01.75%, respectively. In contrast, the austenite stabilizers are transformed into coarsened austenite, which is demonstrated by their greater chemical compositions of 05.10 and 01.82% in Table 1.

Figure 2. Microstructure of 2205 DSS taken at 100 µm of a) as-rolled, b) Widmanstätten and c) equiaxed
Table 1. Elements of normalized analyzed by SEM/EDS

<table>
<thead>
<tr>
<th>Element</th>
<th>Equiaxed Ferrite (%)</th>
<th>Austenite (%)</th>
<th>Widmanstätten Ferrite (%)</th>
<th>Austenite (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>0.00</td>
<td>0.00</td>
<td>03.84</td>
<td>04.28</td>
</tr>
<tr>
<td>Si</td>
<td>0.31</td>
<td>0.00</td>
<td>0.32</td>
<td>0.33</td>
</tr>
<tr>
<td>Cr</td>
<td>21.15</td>
<td>22.66</td>
<td>23.10</td>
<td>21.21</td>
</tr>
<tr>
<td>Mn</td>
<td>01.38</td>
<td>21.63</td>
<td>01.75</td>
<td>01.82</td>
</tr>
<tr>
<td>Fe</td>
<td>61.70</td>
<td>02.41</td>
<td>63.32</td>
<td>62.76</td>
</tr>
<tr>
<td>Ni</td>
<td>03.82</td>
<td>0.23</td>
<td>0.00</td>
<td>05.10</td>
</tr>
<tr>
<td>Mo</td>
<td>00.00</td>
<td>0.00</td>
<td>00.00</td>
<td>02.64</td>
</tr>
</tbody>
</table>

The ferrite formers are partitioned to the old ferrite phase as well, whereas nickel and manganese, the austenite stabilizers, are converted to the coarsened austenite. In Figures 3a and c, iron (Fe) and chromium (Cr) displayed the largest peaks. The situation exists due to the size of the precipitate at the submicron scale and interaction with the ferrite matrix. The amounts of chromium and nickel are listed for the ferrite and austenite phases in Table 1. The lack of balance when ferrite changed into austenite may have been primarily caused by the low nickel concentrations, which could contribute to its poor hot workability. As previously indicated, the transition from ferrite to coarsened austenite was caused by the weaker interatomic bonds brought about by elevated temperatures. The matching lattice planes between the ferrite and austenite (100 and 111) enable this change. In the ensuing microstructure, the coarse austenite formed more sporadic contact with the A/A, F/F, and A/F interfaces [40].

3.1.3 Electron back-scattered diffraction
EBSD phase mapping was done on a 1.47 mm² area and presented in Figure 4. The three constituents visible on the phase map are the Fe-BCC, Fe-FCC, and minor percentages of TiC lining around the coarse BCC grain boundaries. Table 2 displays the mapping statistics for the area of interest. A higher percentage of iron FCC, 67.72%, is present and is in accordance with the microstructure. According to the microstructure in 4b, a higher fraction of iron FCC-67.72% is present, while the BCC fraction was discovered to be 28.96%. The high iron FCC fractions suggest that austenite is more prevalent than ferrite. It was established that low stacking fault energy causes the reduction in mobility of dislocations at grain boundaries associated with austenite. Due to the significant proportions of austenite, it is therefore expected that the corresponding grain boundaries are stationary [41]. The locations at which the electron beam of the SEM with EBSD was unable to detect fractions are represented by the zero solution in Table 2.

Figure 3. Represents the SEM morphology (i) and the corresponding EDS (ii) analysis of 2205 duplex stainless steel for a) equiaxed ferrite phase, b) equiaxed austenite phase c) Widmanstätten austenite phase, and d) Widmanstätten ferrite phase

Table 2. Full-mapping statistics of the heat-treated 2205 DSS

<table>
<thead>
<tr>
<th>Microstructure</th>
<th>Phase Type</th>
<th>Phase Fraction (%)</th>
<th>Phase count</th>
<th>Mean MAD</th>
<th>Standard Deviation MAD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Equiaxed</td>
<td>Fe-BCC (old)</td>
<td>28.96</td>
<td>4128</td>
<td>0.77</td>
<td>0.12</td>
</tr>
<tr>
<td></td>
<td>Fe-FCC</td>
<td>67.72</td>
<td>9654</td>
<td>0.80</td>
<td>0.12</td>
</tr>
<tr>
<td></td>
<td>TiC</td>
<td>0.33</td>
<td>47</td>
<td>0.84</td>
<td>0.09</td>
</tr>
<tr>
<td></td>
<td>Zero Solutions</td>
<td>3.00</td>
<td>427</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Widmanstätten</td>
<td>Fe-BCC (old)</td>
<td>60.79</td>
<td>6128</td>
<td>0.64</td>
<td>0.13</td>
</tr>
<tr>
<td></td>
<td>Fe-FCC</td>
<td>24.24</td>
<td>2443</td>
<td>0.72</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>TiC</td>
<td>0.04</td>
<td>4</td>
<td>1.36</td>
<td>0.78</td>
</tr>
<tr>
<td></td>
<td>Zero Solutions</td>
<td>14.93</td>
<td>1505</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

*MAD= mean angular deviation

An inverse pole figure is a two-dimensional graphical depiction of the orientation of a selected plane that is relatively normal to the sample reference frame. The mapping obtained from the analysis results in the microstructural evolution in the X, Y, and Z directions, as shown in Figure 5 (a, b, and c) for as-rolled, equiaxed and Widmanstätten. The as-rolled grains, Figure 5a, demonstrated a deficient stage orientation of A/F grain distribution along 111 and 110 planar with respect to XYZ coordinates. It is a testament to the fact that the interface is incoherent due to the random orientation of the nucleus...
matrix. The orientation is induced by the deformation the material suffered after hot-rolling. In Figure 5b, the iron BCC dominates the matrix, and the iron FCC makes up the coarsened grains. With reference to Figure 5, the IPF orientation triangle is utilized to depict the direction in which these grains orient. Because the green grains in this example transpose to the blue grains, the grains are orientated in the [111] direction. The purple grains also transpose to the blue grains in Figure 5b; the grains are oriented towards the [100] direction.

Furthermore, the grains also aligned in the [110] direction. In a polycrystalline ferrite matrix, plate-like monocrystal austenites are distributed almost randomly in space. They are orientated in accordance with the Kurdjumov-Sachs relationship when the interface possesses semi-coherent interphase boundaries, according to Zhang [34]. The K-S relationships between (011) Ferrite/(111) austenite planar and (111) Ferrite/(110) austenite planar can also be used to establish the Nishiyama-Wassermann (N-W) relationship by rotating (011) Ferrite by 5.26° [42]. Due to the semi-coherent interfaces that Widmanstätten have, these semi-coherent interfaces go from semi-coherent to incoherent following the rolling process.

Figure 4. The EBSD mapping of the yielded microstructure of a) as-rolled, b) equiaxed, and c) Widmanstätten

Figure 5. Inverse pole figures in the x, y, and z directions of the a) as rolled, b) the equiaxed, and c) the Widmanstätten
Figure 6. Grain orientation of the as-rolled, equiaxed and Widmanstätten after heat-treatment of the as-rolled

Figure 6 shows a frequency distribution plotted against the orientation angle for the as-rolled, equiaxed austenite and Widmanstätten morphologies. The as-rolled showed mixed and alternate low and high grain boundaries having orientation angles from 0° to 55°, with orientation within 5° being predominant. This orientation within 5° results from grain breakage emanating from the rolling process. According to the literature, high angle grain boundaries are visible when $\theta > 15^\circ$ and low angle grain boundaries are visible when $\theta < 15^\circ$ [43]. The equiaxed indicates that the LAGB predominate in the microstructure since the largest peak is seen when the angle is less than 15°. Dislocations are placed sporadically along the grain borders. Additionally, these grain boundaries are located in regions where crystals fit together perfectly. As a result, the low-angle grain boundaries have a cohesive interface. The interfaces that result from the exact alignment of two crystals at the interface planes are known as coherent interfaces [44]. These interfaces will have low interfacial energy, which is insufficient to support dislocation climbing, resulting in static grain boundaries.

However, the Widmanstätten structure displayed almost purely high grain angles as the angle is greater than 15°. The Widmanstätten histogram is dispersed to the right having predominant grains within 25° – 30°. With a high percentage of high-angle grain boundaries, Widmanstätten austenite is likely to crack since these borders will have high interfacial energy, which will facilitate the climbing and cross-slipping of the grain boundaries. It will be least workable at 2205 DSS during hot workability. It also carries an incoherent nature that does not prevent grain boundaries from sliding. High grain boundaries also pose to have excessive hardness and internal strength at the expense of ductility and toughness, thereby subjecting the grain to the point of crack initiation [44]. It is more likely that the steel will crack because of the incoherent interfaces’ K-S orientation connection that prevents the sliding of grain boundaries. The lack of workability of the steel is further explained by the incoherent nature and K-S orientation connection of austenite-ferrite (A/F) interfaces, which suggests that a higher percentage of A/F interfaces have these characteristics.

3.2 X-ray diffraction

Figure 7 shows the as-received 2205 DSS, normalized (equiaxed), and quenched (Widmanstätten) sample’s diffraction pattern. The purpose of the XRD was to identify the phases present in each sample after the heat treatment.

In the as-rolled sample, three phases were identified: alpha ferrite, gamma austenite, and martensite. This contradicts the results obtained from the optical micrograph. Falodun [31] asserts that during the hot rolling of 2205 DSS, strain-deformation causes the austenite phase to change into the martensite phase. Based on Bain's theory, heat treatment can
turn martensite back into the austenite phase because the 2205 DSS only needs the ferrite and austenite phases. Hence, applying heat converts the martensite phase to the austenite phase [45]. In order to determine whether any intermetallic phases in the Widmanstätten structure might impair the functionality or use of the 2205 DSS, an XRD was conducted on the heat-treated sample. Two phases were found; martensite is profound at 76.10%, and the volume fraction of the austenite is around 23.89%. During rapid cooling, the presence of C and N led to the precipitation of the martensite phase, which tends to harden the 2205 DSS. Therefore, rapid cooling causes a rise in martensite formation [46]. The high-volume martensite component will increase the hardness of 2205 DSS, making the material brittle and prone to cracking when subjected to stress.

Martensite is formed when austenite cools quickly and traps carbon atoms that can't exit the crystal structure. The parent phase- austenite, becomes mechanically unstable when it reaches the martensite temperature [47]. Generally, martensite is formed during annealing at temperatures lower than 400°C. The austenite volume proportion in the equiaxed sample is large since the amount of ferrite stabilizer components in the chemical composition has decreased (Ni and Cr). The three phases were detected with the equiaxed structure: ferrite, austenite, and a small quantity of cementite. The volume fraction of the ferrite phase was found to be 58.7%, the austenite phase, 40.3%, and the cementite phase, 9.22%. The presence of cementite could be a result of cooling to room temperature. The carbon that did not retain in the solid solution during the ferrite phase formed cementite, preventing the carbon from evaporating as free carbon [48]. The modest volume proportion of cementite will only slightly increase the hardness of 2205 duplex stainless steel.

From the observation, both samples have no detrimental phases detected by the XRD, indicating that neither structure creates detrimental phases such as the sigma phase, chi phase, or alpha prime. Furthermore, there were no intermetallic phases observed. The equiaxed sample will not quickly fail due to its greater ductility potential during the application, while the Widmanstätten sample, which has a higher volume fraction of martensite, could result in brittle failure during application.

4. CONCLUSIONS

2205 DSS is associated with poor hot workability. Hence, the effect of the cooling method on the microstructural evolution is successfully characterized. The goal also characterizes the grain boundaries that exist in the microstructures which define its functionality. After cooling, the equiaxed and Widmanstätten austenite morphologies are yielded. The rapid cooling associated with water quenching completely transformed the austenite phase into martensite. This resulted in a volume fraction of 76.10%, which can increase the hardness of the 2205 DSS. In contrast, the normalized quenching process only partially transformed the austenite phase into cementite; and culminated in a volume fraction of 0.92%, which could reduce the hardness of the 2205 DSS. The precipitation of the detrimental phases was not observed in both morphologies. Furthermore, from the microstructural observations, adequate and accurate heat treatment eliminates austenite protrusions usually associated with low-angle sub-boundaries. The cooling method demonstrated its effect on the yielded Widmanstätten morphology as there are large numbers of partially transformed austenite in the matrix observed due to the dissolution of the austenite. The EBSD data indicated that both morphologies oriented more toward 111-planar to establish its austenite intervariant boundaries plane distribution. The XRD analysis results showed no initiation of the detrimental phases, such as chi, sigma, and alpha prime, in both microstructures. This means that the 2205 duplex stainless steel will not be susceptible to cracking because of detrimental phases. Due to the different cooling methods, phases such as martensite and cementite are present in Widmanstätten and equiaxed morphologies, respectively. The formation of martensite in the Widmanstätten morphology could result from low carbon content in the chemical composition of 2205 DSS. The equiaxed structure will enhance sliding and minimize cracks associated with hot rolled 2205 DSS while Widmanstätten obeys an ideal K-S relationship. It is clear that heat treatment also causes the recrystallization of grains in structures. Heat treatment of hot-rolled 2205 DSS will eliminate poor hot working conditions related to hot-rolled 2205 DSS.

Therefore, the equiaxed structure will function better in hot-working conditions compared to the morphology associated with Widmanstätten.

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REFERENCES


NOMENCLATURE

A/A Austenite/Austenite
A/F Austenite/Ferrite
bcc Body-centred cubic
Cr Chromium
DSS Duplex stainless steel
EDS Energy-dispersive spectrometry
F/F Ferrite/Ferrite
fcc Face-centred cubic
Fe Iron
HABG High-angle grain boundaries
K-S Kurdjumov-Sachs
LAGB Low angle grain boundaries
Mn Manganese
Mo Molybdenum
N-W Nishiyama-Wassermann
SAGB Special angle grain boundaries
SEM Scanning electron microscope

Greek symbols

\( \alpha \) Alpha, Ferrite
\( \alpha' \) Martensite
\( \beta \) Beta
\( \gamma \) Gamma, Austenite