SA508 grade 3 steel is widely used in the manufacture of large-scale forged components for nuclear reactor applications. Numerical models have already been established to simulate industrial forging process of grade 3 steel; however, limited information is available on the microstructural evolution of this steel during hot forging operation. This work focuses on the flow behavior and related microstructural evolution in grade 3 steel with detailed analysis on the interfacial friction, texture and hardness evolution. Uniaxial hot compression tests were conducted over a range of test temperatures (880-1130 °C) and true strain rates (0.001-1/s), representative of the industrial hot forging conditions. Two different deformation mechanisms, MDRX at the lowest forging temperature and DRV along with DRX at the highest forging temperature, were observed showing marked impact on the final microstructure and hardness. A random fiber-type weak deformation texture was observed irrespective of the test temperatures and strain rates used. The microstructural changes from the as-received to the various deformed conditions were quantified. The quantitative data are the key to obtain accurate parameters for DRV and DRX processes that affect the accuracy of the mathematical models.

1. Introduction

ASME SA508 is a widely used steel grade in nuclear reactor pressure vessels (RPV), steam generators (SG) and pressurizers. Among the SA508 steel variants, the grade 3 steel is a more preferable choice over other grades (grade 1 and grade 2) due to its improved strength, excellent toughness, high weldability and superior resistance to irradiation embrittlement. Bowen et al. (Ref 1) reported that the large rod-shaped and spherical cementites formed at grain/lath boundary reduced the toughness and increased the ductile–brittle transition temperature (DBTT) of the heat-treated SA508 grade 3 steel. Im et al. (Ref 2) showed that the nano-sized needle-shaped M2C carbides significantly improved the yield and tensile strength of this material and simultaneously lowered the DBTT, thereby increased the cleavage fracture stress. The needle-shaped M2C carbides, densely formed within the ferritic matrix, attracted interest due to their significantly higher resistance to irradiation embrittlement and improved strength. Pous-Romero et al. (Ref 3) summarized the influence of multi-step heat-treatments on this steel grade in achieving optimum austenite grain size and modeled the austenite grain growth in RPVs. The optimum grain size is important in order to achieve desired mechanical properties and therefore longer service life of grade 3 steel. These studies were primarily focused on establishing a basic understanding of the structural, mechanical and service behavior of this steel grade. The outcomes from these studies were used to modify the microstructure for meeting the demanding requirements of hostile service environment involving high temperature, high pressure and exposure to high-energy neutrons. On the other hand, limited studies have been conducted on the hot deformation behavior of the SA508 grade 3 steel focussing mainly on the effects of process parameters and prediction of the material behavior during hot forging by establishing different numerical models. Sun et al. (Ref 4) reviewed the previous studies on SA508 grade 3 steel and identified lack of information about the mechanical response at different temperatures, strains and strain rates in plastic deformation region, which was important for designing heavy forging processes of large components. They modeled mechanical behavior of this steel during hot deformation (in the range of 700-1200 °C with strain rates of 0.001-10/s) by incorporating the work hardening (WH), dynamic recovery (DRV) and dynamic recrystallization (DRX) using Bergstrom dislocation-based model and Avrami-type equation. As the temperature reached just above Ac3 (temperature at which transformation from ferrite to austenite occurs), they measured the prior austenite grain size (PAGS) by quenching the sample and then using an empirical formula, they calculated the initial PAGS before hot deformation. This PAGS value is an important input parameter for calculating both DRX and DRV parameters and final mixed grain size of the forged product. The effectiveness of this mathematical model depends on the accuracy of these parameters. Although Sun et al. (Ref 4) did not provide any insights into the microstructural evolution, but they proved the effectiveness of their FEM model for a nuclear conical shell forging process. Sui et al. (Ref 5) carried out similar type of simulation work focused on inhomogeneous hot deformation (IHD) process of SA508 grade 3 steel. They used average grain
size calculated from optical micrographs as an input to the FEM model, but did not provide details on the microstructural evolution. The model showed that the temperature had more influence over the DRX than the strain and strain rates involved during IHD. Typically, the forging of large components involves multi-passes, where the material is often subjected to dynamic recovery (DRV), dynamic recrystallization (DRX), static recrystallization (SRX) and metadynamic recrystallization (MDRX) leading to subsequent changes in the microstructural and mechanical characteristics. As the strain in the first pass exceeds the critical strain of DRX, the MDRX takes place by continuous growth of the nuclei formed by DRX during deformation at high temperature. The MDRX significantly reduces stress in the subsequent passes and therefore refines the grain size during hot forging process. Recently, Dong et al. (Ref 6) investigated the MDRX behavior of SA508 grade 3 steel during isothermal double-hit hot compression tests performed at a wide range of forming temperatures (950-1250 °C), strain rates (0.001-0.1/s) and inter-stage delay times. They observed significant effects of temperature and strain rate on MDRX behavior of grade 3 steel as compared to the effects of both pre-strain and initial austenite grain size. Based on the experimental results obtained, they also established the kinetic equations and the grain size model for predicting the softening fractions during MDRX. They used it as an input in the commercial FE software for modeling and optimizing the hot forging process of grade 3 steel. In summary, these studies simulated the hot forging operation of grade 3 steel within a wide range of forming temperatures (950-1250 °C), strain rates (0.005-10/s) and true strain of unity. The standard specific range of test temperatures (880-1130 °C) and true strain range of 0.001-1/s with a true strain of unity. The strain measurements were deduced from the displacement of the machine crosshead and adjusted to take account of the machine compliance. The measured flow curves were then corrected for both the interfacial friction and the adiabatic heating. The values of barreling and ovality coefficients calculated from the deformed samples were within the acceptable limits as per the NPL good practice guide, which details the experimental practice and measurement of the flow stress during isothermal uniaxial axisymmetric compression tests (Ref 10). The strain measurements were deduced from the displacement of the machine crosshead and adjusted to take account of the machine compliance. The measured flow curves were then corrected for both the interfacial friction and the adiabatic heating. The values of barreling and ovality coefficients calculated from the deformed samples were within the acceptable limits as per the NPL good practice guide, and thus, the validity of each compression test was ensured (Ref 10). After deformation, the deformed samples were quenched within 3-5 s from end of the tests to freeze the respective deformed microstructure. The samples were then sectioned and mounted in conductive resin molds. The mounted samples were ground with SiC abrasive grit papers and finish-polished using UltraPol 9 colloidal silica suspension. Then, the samples were etched using nital (solution of 2% HNO3 in ethanol) for PAGS

2. Experimental Details

Table 1 shows the elemental composition of the SA508 grade 3 steel blocks (12 blocks, each having a dimension of 63 mm length × 19 mm width × 15 mm height) as supplied by Rolls Royce Plc. Figure 1(a) shows the orientation of machined cylindrical compression test samples (12 mm diameter × 18 mm height) from these blocks, and Fig. 1(b) shows the sample co-ordinate system consisting of compression direction (CD) and radial direction (RD) as used in this study. Before the compression tests, the machined samples were coated with glass lubricant (Prince Grey Glass Lubricant 87.2.237) in order to avoid any oxidation and sticking issues with the platen of the testing machine at high temperature. Zwick Z250 and HA250 mechanical test machines were used for conducting isothermal hot compression tests at atmospheric condition in the temperature range of 880-1130 °C and the true strain rate range of 0.001-1/s with a true strain of unity. Figure 2 shows the schematic diagram of the hot deformation process, and Table 2 details the compression test matrix. The samples were soaked at the required deformation temperatures for 30 min to achieve a uniform temperature distribution, and it was maintained within ±3 °C throughout the duration of the test. The compression tests were performed as per the National Physical Laboratory (NPL) good practice guide, which details the experimental practice and measurement of the flow stress during isothermal uniaxial axisymmetric compression tests (Ref 10). The strain measurements were deduced from the displacement of the machine crosshead and adjusted to take account of the machine compliance. The measured flow curves were then corrected for both the interfacial friction and the adiabatic heating. The values of barreling and ovality coefficients calculated from the deformed samples were within the acceptable limits as per the NPL good practice guide, and thus, the validity of each compression test was ensured (Ref 10). After deformation, the deformed samples were quenched within 3-5 s from end of the tests to freeze the respective deformed microstructure.

The samples were then sectioned and mounted in conductive resin molds. The mounted samples were ground with SiC abrasive grit papers and finish-polished using UltraPol 9 μm, Trident 3 μm, MasterTex 1 μm and Microcloth with 0.02 μm colloidal silica suspension. Then, the samples were etched using nital (solution of 2% HNO3 in ethanol) for PAGS

Table 1 Elemental composition of as-received SA508 grade 3 machined steel blocks

| Element | C  | Mn  | Ni  | Cr  | Mo  |
|---------|----|-----|-----|-----|-----|
| Wt%     | 0.206 | 1.280 | 0.644 | 0.140 | 0.493 |

| Element | Si  | S   | P   | Fe  |
|---------|-----|-----|-----|-----|
| Wt%     | 0.321 | 0.014 | 0.002 | 97.093 |
measurement. A Leica DM1200M optical microscope was used to capture the micrographs and conduct related grain size analysis using the Leica LAS-X image analysis software coupled with the microscope. Figure 3 shows an example of PAGS analysis using the image analysis software where the automatic grain detection technique is used to identify the prior austenite grains (PAGs) in the as-received sample. The software detects the different PAGs with different colors and provides a grain size distribution. The quantitative grain size analysis carried out in this study is as per the ASTM standard E112 (Ref11). The as-received sample and the four deformed samples covering the maximum and minimum parametric range of the compression test matrix, i.e., 880 °C-0.001/s, 880 °C-1/s, 1130 °C-0.001/s and 1130 °C-1/s, were further vibro-polished for EBSD analysis. A FEI Quanta FEG 250 SEM was used to collect the EBSD data, from the longitudinal section, along the sample axis at a distance of h/3 from one end of the sample, where h was the deformed sample height. Evans and Scharring (Ref 12) reported that the portion of deformed specimen at h/3 distance from each end of the cylindrical sample was representative of the bulk strain experienced during hot axisymmetric compression tests under constant strain rate. Hence, it is very important to capture microstructure from this region for quantitative analysis to avoid areas within the deformed specimens that have inhomogeneous strain distribution, i.e., dead zones. The orientation mapping was performed using a rectangular grid with a step size of 0.4 μm and 0.3 μm, respectively, for as-received and deformed samples with an indexing rate of 98-99%. Both the bainitic and martensitic packets, i.e., effective grains, were identified by the high-angle grain boundaries (HAGB) with a misorientation angle of $\theta > 15^\circ$, and the minimum size of accepted effective grains
was taken as 5 pixels in this study (Ref 13). The EBSD data processing was carried out using HKL Channel-5 post-processing software. A Struers hardness tester was used to measure the average hardness of these five samples according to ASTM standard E384-11 (Ref 14). The indents (> 100 indents) were made in a rectangular matrix using a Vickers indenter with a fixed load of 5 kgf, and the distance between any two indents was kept as 1 mm along both X and Y directions ensuring more than 3 diagonal widths spacing between any two indents as recommended for the steel samples (Ref 15).

### 3. Evolution of Mechanical Properties during Hot Deformation

#### 3.1 Analysis of Flow Behavior

Figure 4 summarizes the flow curves of SA508 grade 3 steel samples at different true strain rates (0.001-1/s) with a variation in the test temperature (880-1130 °C). These flow stress curves have been corrected for frictional and adiabatic heating losses as outlined in slab analysis method (Ref 16) using an in-house

| Temperature (°C) | Zwick Z250 (for low strain rates) | Zwick HA250 (for high strain rate) |
|------------------|-----------------------------------|-----------------------------------|
| 880              | 0.001                             | 0.1                               | 1 |
| 963              | 0.001                             | 0.1                               | 1 |
| 1046             | 0.001                             | 0.1                               | 1 |
| 1130             | 0.001                             | 0.1                               | 1 |

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Fig. 3 Representative example of PAGS analysis using Leica LAS-X software—(a) as-received microstructure, (b) PAGS identified using software (different colors indicate different grain sizes) and (c) PAGS distribution in as-received sample
developed Matlab code. Figure 5 shows combined flow stress curves before and after implementing the necessary corrections for one of the test samples, i.e., 1130 °C-1/s. The blue line, named as ‘true stress without friction correction,’ shows the true stress curve as a direct output of the compression test, i.e., raw flow stress data, without taking into account any influence of interfacial friction and adiabatic heating during the test. The red line, named as ‘true stress with friction correction,’ shows the flow stress curve considering corrected for interfacial friction loss. The green line, named as ‘true stress with friction and adiabatic heating correction,’ shows the corrected flow stress curve considering both frictional and adiabatic heating losses.

Equation 1 and Equation 2 summarizes the steps for correcting those losses according to the slab analysis method. In this method, the energy under the flow stress curve is calculated subtracting the conduction heat losses from the ends and then the remainder is converted to a temperature rise. A linear temperature sensitivity is assumed during the temperature correction to this flow curve, and the value is obtained from the flow stress data. These corrections on the flow stress data are performed by an in-house developed Mathcad code, and Table 3 lists the input data used in that Mathcad code. Wan et al. (Ref 17) have described and performed similar analysis during hot deformation of a nickel-based U720Li alloy.

The corrected (for both frictional and adiabatic heating losses) flow stress curves in Fig. 4 show a typical stress–strain behavior influenced by both work hardening and thermally activated softening mechanisms. DRV and DRX are two important softening mechanisms that significantly influence the flow stress behavior during high-temperature deformation of metals. Typically, DRX is dominant in face-centered cubic (FCC) materials owing to their low stacking fault energy (SFE), whereas DRV has been found dominant in body-centered cubic (BCC) materials due to their high SFE. However, under certain combination of deformation temperature and true strain rate values during hot deformation, the flow behavior of BCC materials is influenced by either the presence of both DRX and DRV or a change in softening mechanism from DRX to DRV. For example, Zhang et al. (Ref 18) showed the presence of both DRX and DRV during hot deformation behavior of BCC Ti-15-3 metastable beta titanium alloy. The DRX was dominant at a particular combination of deformation temperature and strain rate, i.e., 1050 °C-0.01/s, and DRV was dominant under either lower or higher temperature and strain rate combinations (850 °C-0.01/s and 1100 °C-1/s, respectively). DRX results in grain refinement and annihilation of deformation defects, thereby significantly improving the mechanical properties and hence has advantages over DRV. On the other hand, Davenport et al. (Ref 19) mentioned a change in flow softening mechanism from, DRX to DRV, for BCC plain C-Mn steel (max. 0.25% C and 1.5% Mn). The DRX was dominant at higher temperature and lower strain rate combination (1100 °C-1/s), whereas the softening mechanism completely changed from DRX to DRV at lower temperature and significantly higher strain rate combinations (900 °C-10/s and 1000 °C-50/s, respectively). These variations are typically characterized using Zener–Holloman parameter Z, where low Z value indicates occurrence of rapid DRX and high Z value shows the presence of either DRV or a combination of both WH and DRV in the BCC materials. The flow instability is observed at very high Z values, which varies depending on the material. In the current study, Fig. 4 shows a linear increase in peak stress values with decrease in deformation temperature (at any fixed strain rate) and with increase in strain rate (at any fixed temperature). Particularly, a significant decrease in the flow stress value is observed when the deformation temperature is 1046 °C and above. Feng et al. (Ref 20) reported similar behavior during high-temperature deformation of a high-nitrogen martensitic 30Cr15Mo1N stainless steel. They reported that the low strain rate provided enough time for energy accumulation, whereas the high deformation temperature promoted dissolution of precipitates, dislocation movements and grain boundary migration for nucleation and growth of DRX grains eventually leading to a decrease in the flow stress level. Beside this, another
notable fact in the current study is the presence of hump in the flow stress curves for all three low strain rate tests (0.1-0.001/s) irrespective of the deformation temperatures. The initial rise of the flow stress is attributed to the WH due to increase in dislocation density. With increase in the strain value, the effect of WH is balanced by both DRV and DRX, which leads to formation of a hump in the flow curves. A steady-state condition is achieved when the balance is reached to an equilibrium state. Specifically, for 1/s strain rate tests, the hump becomes wider and the steady-state condition is achieved faster as compared to other low strain rate tests. Thus, 880°C-1/s and 1130°C-0.001/s test conditions exhibit maximum and minimum effect of WH, respectively, along with significant effect of flow softening for the later. This is also evident from Fig. 6 showing relationship between WH rate ($\dot{h}$) and true stress as derived from the experimental data. Similar observations were also reported by Sun et al. (Ref 4). After the steady-state condition, all the flow curves in Fig. 4 show a downward trend irrespective of the strain rates and deformation temperatures used. This is attributed to the occurrence of DRX followed by DRV. Feng et al. (Ref 20) showed that the flow stress curves having a single peak followed by a steady-state condition typically indicated the appearance of DRX during hot deformation. The waviness observed at the end of the flow curves was attributed to DRV due to high migration of dislocations and with further deformation, and the dislocation density continued to increase leading to a second peak in the flow stress curve. Besides this, Fig. 7 shows the strain rate sensitivity ($m$) of the deformed samples at different test temperatures. The highest and lowest peak stresses were observed for 880°C-1/s and 1130°C-0.001/s test conditions, respectively. The $m$ value was observed as 0.131, 0.148, 0.167 and 0.182 for test temperatures 880°C, 963°C, 1046°C and 1130°C, respectively. This slight difference in the $m$ values indicates the stability of the forging process, if performed within the specified parametric range as observed from the corrected stress–strain curves in Fig. 4.

**Equation 1: Correction of flow stress for frictional loss**

(a) Defining the effect of deformation on sample:

$$Shape_t = \mu \cdot \sqrt{\frac{\dot{h}^2 \cdot e^{\gamma \cdot \frac{x}{h}}} {h \cdot e^{\gamma \cdot \frac{y}{h}}}$$

(b) Corrected flow stress after frictional loss:

$$\sigma_{fric_t} = \frac{2 \sigma_t} {\left( \frac{1}{Shape_t} \right)^2 \cdot (e^{2Shape_t} - 2 \cdot Shape_t - 1)}$$

**Table 3 List of input data used in the Mathcad code for correction of flow curves**

| Input factors                      | 0.001/s | 0.01/s | 0.1/s | 1/s  |
|------------------------------------|---------|--------|-------|------|
| Coefficient of friction            | 0.05    |        |       |      |
| Density, kg/m³                     | 8050    |        |       |      |
| Heat transfer coefficient, W/m² K⁻¹ | 50      |        |       |      |
| Sample diameter                    | Variable (12 mm ± x* mm) |       |       |      |
| Sample length                      | Variable (18 mm ± y* mm) |       |       |      |
| Test temperatures                  | Variable (880°C, 963°C, 1046°C and 1130°C) |       |       |      |
| Stress-temp coefficient, MPa/K     | 0.107   | 0.101  | 0.105 | 0.134|
| *where x and y are different for each sample |
where \( \mu \) = friction coefficient of the hot deformation process, \( r \) and \( h \) = initial radius and height of the compressed test sample, respectively, and \( \varepsilon \) = measured strain.

Equation 2: Correction of flow stress for loss due to adiabatic heating

(a) Defining the adiabatic temperature rise:
\[
\Delta T = \sum_{i=0}^{n-1} \left( \frac{\pi r^2 \cdot h \cdot \sigma_{\text{fric}_i} \cdot 10^6 \cdot (\varepsilon_i - \varepsilon_{i-1})}{C_p \cdot \rho \cdot \pi r^2 \cdot h} \right)
\]
where \( \Delta T \) = maximum temperature rise assuming no heat loss, \( C_p \) = specific heat of the material, \( \rho \) = density of the material and \( n \) = data length

(b) Defining the thermal losses due to conduction:
\[
\text{loss}_{n+1} = dq_n \cdot 2\pi r^2 h \cdot \Delta T_n = \left( h \cdot \frac{\varepsilon_i - \varepsilon_{i-1}}{\dot{\varepsilon}} \right) \cdot 2\pi r^2 h \cdot \Delta T_n
\]
where \( dq_n \) = heat flow per increment, \( 2\pi r^2 h \) = die contact area and \( \dot{\varepsilon} \) = required strain rate

(c) Defining the corrected test temperature:
\[
\text{testtemp}_n = T + \Delta T
\]
Corrected flow stress after incorporating temperature rise and energy loss:
\[
\sigma_{\text{adia}} = \sigma_{\text{fric}} + \text{tempcoeff} \cdot (\text{testtemp}_n - \text{testtemp}_1)
\]
where \( \mu \) = friction coefficient of the hot deformation process, \( r \) and \( h \) = initial radius and height of the compressed test sample, respectively, and \( \varepsilon \) = measured strain.
3.2 Analysis of Interfacial Friction

Interfacial friction is a key factor that influences the hot deformation behavior of all metallic materials. Li et al. (Ref 21) determined the average Tresca friction coefficient (μ) of a high-temperature barrel compression test by measuring the degree of barreling as shown in Equation 3 using upper-bound theory. This friction coefficient was dependent on the surface quality of the material, deformation temperature as well as strain values; thus, it covered the entire range of friction that could be observed during hot deformation processes starting from the perfect sliding (μ = 0) to the sticking friction (μ = 1). The same approach has been used in this study, and Fig. 8 shows the variation in friction coefficient at different true strain rates (0.001-1/s) and test temperatures (880-1130 °C). Mostly high sticking friction (μ = 0.78-1.04) is observed for all true strain rates within the temperature range of 880-1046 °C, whereas a minor reduction in friction coefficient (μ = 0.63-0.75) is observed at 1130 °C irrespective of the strain rates used.

This reduction in friction could be attributed to the formation of thick oxide scale containing lubricious and low shear strength iron oxides, such as FeO and Fe3O4, at high temperature (Ref 22). In the current work, formation of such oxide layers is clearly evident from the SEM images and the related EDX analysis confirms the chemical composition as shown in Fig. 9. For example, both 1130 °C-0.001/s and 1130 °C-1/s samples show formation of thick oxide scales (400-760 μm) with porosity and large cracks (Fig. 9a and b, respectively). The sharp peaks of both iron and oxygen are clearly observed from the EDX spectra when collected from the interlayers (i.e., the thin layer between steel sample and oxide layer) and the oxide layers at the top.

Equation 3: coefficient of friction calculation using upper-bound theory

\[ \mu = \frac{(r/h) \times b}{4/\sqrt{3} - 2h/(3\sqrt{3})} \]

where \( \mu \) = friction coefficient of the hot deformation process, \( r, h \) = average radius and height of the compressed test sample, respectively, and \( b \) = barreling coefficient.

4. Evolution of Microstructural Properties during Hot Deformation

4.1 Microstructure Analysis

Figure 10 shows the microstructures of as-received material and undeformed samples soaked at different test temperatures (880-1130 °C) for 30 min followed by water quenching. The as-received sample shows characteristic lower bainitic microstructure, i.e., thin plates of ferrite containing very fine laths of cementite, which is transformed into austenite at elevated temperatures. At the end of the soaking period, the austenitic microstructure is transformed into fine acicular martensitic lath structure due to water quenching. The quenched undeformed samples, soaked at 880 °C and 963 °C, contain a microstructure exhibiting thicker martensitic laths, whereas samples soaked at 1046 °C or higher show the presence of finer martensitic laths. Furthermore, the as-received microstructure contains coarse carbide particles which dissolve completely when heated at those high temperatures.

Figure 11 shows microstructure of the compressed test samples deformed at the highest strain rate of 1/s. The martensitic laths observed in this case are finer in the samples deformed at 1046 °C and 1130 °C when compared to others. Similarly, Fig. 12 shows the microstructure of the compressed test samples deformed at the lowest strain rate of 0.001/s. In this case, martensitic laths appear coarser as compared to the 1/s strain rate cases due to longer exposures at the test temperatures. It should be noted here that the contrast of these optical images might lead to the confusion in reader’s mind that the prior austenite grain boundaries (PAGB) are not clearly distinguishable from the lath boundaries; however, this is not the case. The PAGBs were identified using the LAS-X software at a significantly high magnification in order to perform the measurement of PAGS as already demonstrated in Fig. 3.

Figure 13 summarizes the PAGS analysis for as-received material and all water-quenched undeformed and deformed samples. The as-received material has an average PAGS of \( \sim 32 \) μm, which shows no significant changes when soaked up to 963 °C before deformation. The undeformed samples show a drastic rise in the average PAGS, i.e., \( \sim 70 \) μm.
and \(~ 240 \mu m\), when soaked at 1046 °C and 1130 °C, respectively. In brief, almost no grain growth is observed in the undeformed samples up to 963 °C, over which a rapid grain growth takes place. Slightly different behavior is observed for the deformed samples. The samples deformed up to 1046 °C show an average PAGS of \(~ 49 \) to \(~ 60 \mu m\) indicating insignificant grain growth irrespective of the different strain rates. When deformation temperature is increased to 1130 °C, a drastic difference is observed in the average PAGS of samples deformed at two different strain rates. The sample deformed at 1/s shows minor increase in average PAGS, i.e., \(~ 70 \mu m\), whereas the sample deformed at 0.001/s shows a rapid increase in average PAGS, i.e., \(~ 150 \mu m\). This significant grain growth observed particularly in the undeformed 1130 °C sample and deformed 1130 °C-0.001/s sample could be attributed to three different phenomena—(1) reduction in Zener drag at that high temperature, (2) accumulation of enough energy for such grain growth at a very low strain rate and (3) the absence of carbide particles at the high temperature, which in turn accelerates the austenitic grain growth. Pous-Romero et al. (Ref3) reported similar behavior for SA508 grade 3 steel under a series of different austenitization heat-treatments. They observed an insignificant austenite grain growth up to 965 °C for 30 min soaking time. With increase in soaking time as well as the test temperature, a rapid austenite grain growth was observed over \(~ 940 \) °C due to reduction in Zener drag. Besides this,
Papaefthymiou et al. (Ref 23) reported that the presence of undissolved carbides in a medium-carbon steel prevented the austenite grain growth at 950 °C resulting in a fine-grained microstructure. In this study, the specific temperature over which the rapid austenitic grain growth occurred is at 963 °C in case of undeformed sample and at 1046 °C for the deformed sample.

The as-received material and four deformed samples, i.e., 880 °C-0.001/s, 880 °C-1/s, 1130 °C-0.001/s and 1130 °C-1/s, were further subjected to EBSD analysis for better understanding of the microstructure and texture evolution during hot deformation of the SA508 grade 3 steel. The strength and toughness of the martensitic steels are strongly dependent on the packet size, and hence, the packets are considered as 'effective grains' in a typical lath martensitic microstructure (Ref 24, 25). The bainitic microstructure shows similar microstructural features, but the difference lies in their mechanical properties. Morphologies are identified visually or using EBSD analysis where the thickness of the lath martensite is found to be finer than that of the bainite (Ref 26). Figure 14 shows the band contrast (BC) images and phase maps of the as-received and the deformed samples. The green dots in BC images indicate the areas of no indexing. The as-received sample exhibits bainitic microstructure with larger effective grains, whereas deformed samples contain martensitic structure with significantly smaller effective grains as a result of water quenching. The phase maps show the BCC and FCC phases in yellow and blue colors, respectively, clearly demonstrating the

![Fig. 10 Optical images showing the microstructure of as-received material and undeformed samples soaked for 30 min at different deformation temperatures (880-1130 °C) before compression test](image-url)
difference between sizes and shapes of the bainitic and martensitic effective grains. The effective grain size is found to be smaller in the samples deformed at 880 °C and slightly larger for the samples deformed at 1130 °C. Depending on the concentration of carbon, either BCC or BCT martensitic microstructure is formed in the quenched steel samples. At carbon concentration typically < 0.6 wt.%, the microstructure is mostly BCC (c/a ratio is 1), whereas it changes to BCT (c/a ratio > 1) with further increase in carbon concentration (0.6-1.75 wt.%) (Ref 27). Brust et al. (Ref 28) pointed out that the EBSD technique was inadequate in accurately measuring the lattice dimension of c axis in high-carbon steels, and therefore, the BCT martensitic microstructure was often indexed as BCC martensitic microstructure due to pseudo-symmetry between BCT and BCC phases. The SA508 grade 3 steel used in this study is a low-carbon steel with only ~ 0.2 wt.% carbon, which upon water quenching forms BCC martensitic microstructure. Thus, yellow color indicates the formation of BCC bainitic microstructure in as-received material, but BCC martensitic microstructure in all the four deformed samples was analyzed in this work. Only 1130 °C-1/s sample shows the presence of negligible amount of FCC phase (in blue) along with BCC martensitic microstructure.

Figure 15 shows the effective grain size maps and respective inverse pole figure (IPF) maps for effective grains in the as-received and the four deformed samples. The effective grain size maps show very fine and large grains in blue and red colors, respectively, and the scale bar below every map indicates the equivalent circle diameter of grains (in micron scale) present in that particular scan profile. The IPF maps, taken in the CD–RD plane, show the orientation of effective grains for each scan profiles. The as-received material mostly contains equiaxed effective grains with diameter range of 2-20 μm and a few large grains in the range of 50-75 μm. A drastic decrease in the effective grains size is observed when the samples are deformed at 880 °C. Irrespective of the strain rate, both 880 °C-0.001/s and 880 °C-1/s samples show similar equiaxed microstructure, where most of the effective grains are below 6 μm with a few grains in the diameter range of 10-16 μm. The IPF maps also show random grain orientation for the as-received material, which remains same following deformation at 880 °C. A significant difference in both the effective grain size and shape results when the samples are deformed at 1130 °C. Both 1130 °C-0.001/s and 1130 °C-1/s samples contain a significant amount of elongated effective grains (aspect ratio in the range of 3-18). 1130 °C-0.001/s sample contains majority of the effective grains below 25 μm diameter with a few grains in the range of 55-70 μm. In case of the 1130 °C-1/s sample, almost all the effective grains are below 10 μm along with very few grains within the 28-34 μm diameter range. The IPF maps show that the majority of the effective grains are oriented in between 001 and 111 directions for 1130 °C-0.001/s sample, and in between 101 and 111 directions for 1130 °C-1/s sample.

A summary of the average effective grain size (in terms of equivalent circle diameter) is shown in Fig. 16. The average effective grain size of the as-received sample (~ 10.5 μm) is decreased to almost 1/3rd for the samples deformed at 880 °C (~ 3 to 3.7 μm) and almost half for the samples deformed at 1130 °C (~ 4 to 6 μm). Irrespective of the deformation
temperature, the samples experiencing the highest strain rate of 1/s show more grain refinement as compared to those deformed at the lowest strain rate of 0.001/s. Sun et al. (Ref 4) reported the occurrence of DRX in SA508 grade 3 steel in the temperature range of 800-900 °C and observed finer grains within that range. Dai and Yang (Ref 9) reported occurrence of DRX in SA508 grade 4 steel from 950 °C onwards and showed that lower strain rate deformation favored the growth behavior of DRX grains more. In this study, the finer effective grains observed at 880 °C indicate the occurrence of DRX and further grain refinement, particularly during 1/s test, is attributed to the occurrence of MDRX as evident from Fig. 15. A noticeable growth of these DRX grains is observed when the deformation temperature reached 1130 °C. It is expected that the DRX

![Fig. 12](image12.jpg)

Optical images showing the microstructure of samples deformed at true strain rate of 0.001/s and at different test temperatures (880-1130 °C)

![Fig. 13](image13.jpg)

Summary of PAGS analysis for as-received, undeformed and deformed SA508 grade 3 steel samples
Fig. 14  Band contrast image (BC) and phase map (BCC in yellow and FCC in blue) showing microstructural evolution of the deformed SA508 grade 3 steel samples.
Fig. 15  Effective grain size maps and respective IPF maps as observed in as-received and deformed SA508 grade 3 steel samples
grains would experience some recovery (as evidence of DRV during deformation was clearly seen from the nature of the flow curves shown in Fig. 4) at that high temperature leading to grain growth particularly at a strain rate of 0.001/s. However, it is difficult to quantify the DRX and the DRV separately due to austenitic to martensitic phase transformation during the water quenching stage after compression test.

4.2 Texture Analysis

The evolution of crystallographic texture from the as-received to the various deformed conditions of SA508 grade 3 steel samples is presented in this section. Hu (Ref 29) showed that when uniaxial compression test was performed on BCC materials at room temperature, a preferred texture developed along a specific direction. During deformation, the active slip planes seem to have rotated toward the compression axis and as a result, the crystal orientation changed which led to development of compression texture. The compression texture was a mixture of 111 and 100 fibers, in which 111 was the major component. Figure 17 shows the pole figures of the as-received and deformed SA508 grade 3 steel samples for {111}, {001} and {101} planes. In order to understand the effect of deformation temperature and strain rate on the texture strength, all the pole figures are plotted in the same scale, i.e., with a maximum density of 10. The as-received material shows random fiber-type texture with a maximum density of 4.58. As expected, the texture of the as-received material weakens when deformed at 880 °C and a decrease in texture strength is observed with increase in strain rate from 0.001 to 1/s. Initially, a significantly strong peak-type texture with a maximum density of 22.14 was observed in 1130 °C-0.001/s sample. However, this was the case when the scanning was done within a small 175 μm × 150 μm area. The strong texture was attributed to the same orientation of the acicular martensitic laths within this area. As observed in Fig. 13, the average PAGS of this samples is ~ 151 μm; therefore, the small scanning area was likely to have covered almost same orientation of the acicular martensitic laths within a single large parent austenite grain (observed from the IPF map in Fig. 15). This led to a very strong peak-type texture, which was not representative of the forging operation at that condition, rather it mostly presented the texture of a random large parent austenite grain. In order to avoid this misrepresentation of texture, a large scanning area of 1 mm × 1 mm is further chosen with 10 μm step size for the EBSD scan. The obtained pole figure shows random fiber-type texture with a maximum density of 3.66, which is representative of forging operation at that condition. With an increase in strain rate to 1/s, the density of the random fiber-type texture increases to 5.84. This random fiber-type weak deformation texture is expected from any forging operation, and it can be attributed to the exposure of a very high temperature leading to greater softening and coarsening of microstructure with higher degree of deformation. Hence, the EBSD analysis has been useful in quantifying the complex microstructure and texture, which is not possible by other means such as optical micrographs. The texture data, however, reveal no significant difference despite the changes of bainitic microstructure present in the as-received sample to the martensitic microstructure of the deformed samples.

4.3 Hardness Analysis

Figure 18 summarizes the average hardness of as-received and deformed steel samples. The hardness values of as-received material measured at different locations are averaged at ~ 194 HV. The samples deformed at 880 °C exhibit an average hardness of ~ 427 HV and ~ 431 HV, in case of 0.001/s and 1/s strain rates, respectively. The average hardness of 1130 °C-0.001/s and 1130 °C-1/s samples is ~ 402 HV and ~ 425 HV, respectively, slightly lower than that observed at 880 °C. It should be noted that the samples experiencing most work hardening and softening behavior, i.e., 880 °C-1/s and 1130 °C-0.001/s, exhibit the highest and lowest average...
Fig. 17  Texture representation of as-received and deformed SA508 grade 3 steel samples
hardness, respectively. The hardness characteristics can be directly linked with the respective microstructure and the average effective grain size. The as-received material contains bainitic microstructure with an average grain size of $\sim 10.5\ \mu m$, which exhibits the lowest average hardness value ($\sim 194\ HV$) among all the samples. However, all the deformed samples show martensitic microstructure with substantially low average effective grain sizes ($\sim 3$ to $6\ \mu m$) resulting in significantly higher average hardness values ($\sim 402$ to $431\ HV$) when compared to the as-received sample. Among the deformed samples, $880\ ^\circ C$-1/s and $1130\ ^\circ C$-0.001/s samples show the smallest and largest average effective grain sizes, i.e., $\sim 3\ \mu m$ and $\sim 6\ \mu m$, respectively, thus resulting in the highest and lowest average hardness values, respectively.

5. Discussion

Table 4 summarizes the key findings from the corrected flow stress behavior and the microstructural evolution of the SA508 grade 3 steel during hot deformation. The as-received material has bainitic microstructure with average effective grain size of $\sim 10.5\ \mu m$ and an average hardness of $\sim 194\ HV$. The combined effects of deformation temperature and true strain rate on this steel are apparent in case of two samples—(1) sample experiencing the highest work hardening, i.e., $880\ ^\circ C$-1/s and (2) sample showing most softening behavior, i.e., $1130\ ^\circ C$-0.001/s. The corrected flow stress curves (Fig. 4) show the highest peak stress (210.56 MPa) and a friction coefficient of 0.82 for $880\ ^\circ C$-1/s sample, whereas totally opposite behavior is observed for $1130\ ^\circ C$-0.001/s sample (peak stress of 19.7 MPa and the lowest friction coefficient of 0.63). Both deformed samples exhibit similar austenitic to martensitic phase transformation during water quenching stage after hot deformation. The average PAGS for $880\ ^\circ C$-1/s and $1130\ ^\circ C$-0.001/s samples is observed as $\sim 49\ \mu m$ and $\sim 151\ \mu m$, respectively (Fig. 13). The high deformation temperature eventually reduces the Zener drag and eliminates the carbide particles from the initial microstructure, whereas low strain rate provides sufficient energy for such grain growth particularly in $1130\ ^\circ C$-0.001/s condition as already explained in Sect. 4.1. The martensitic microstructure of the deformed samples is quantified by calculating average effective grain size. The $880\ ^\circ C$-1/s sample exhibits equiaxed effective grains (i.e., finer laths) with random grain orientation, whereas the $1130\ ^\circ C$-0.001/s sample shows elongated effective grains (i.e., coarser laths) with a specific grain orientation between 001 and 111 directions (Fig. 15). MDRX appears to be the dominant deformation mechanism in $880\ ^\circ C$-1/s case, but no significant grain growth is noticed due to little deformation time before water quenching. As a result, finer laths are observed for $880\ ^\circ C$-1/s sample with an average effective grain size of $\sim 3\ \mu m$. On the other hand, the deformation time is more for the $1130\ ^\circ C$-0.001/s sample due to the combination of highest test temperature and lowest strain rate. In this case, the recrystallization occurs simultaneously with the deformation, which eventually leads to elongation of the already recrystallized grains, i.e., formation of coarser laths. The deformation is stopped due to water quenching before a subsequent cycle of recrystallization takes place; therefore, the average effective grain size remains highest (i.e., $\sim 6\ \mu m$) for $1130\ ^\circ C$-0.001/s sample when compared to others (Fig. 16). The DRV is normally dominant in BCC materials, but DRX can also occur under certain conditions, which is observed true in this study for samples deformed at $1130\ ^\circ C$. This also implies that if the samples deformed at $1130\ ^\circ C$ are furnace-cooled instead of water-quenched, then the effective grains would be initially equiaxed and finer due to having sufficient time for subsequent cycles of recrystallization. Then, these recrystallized grains would grow depending on the slow cooling rate of the furnace. In this study, the quantified microstructural data, i.e., average PAGS and average effective grain size calculated for different deformation conditions, can be directly correlated with the
average hardness of the respective samples according to Hall–Petch relationship. For example, 880 °C-1/s and 1130 °C-0.001/s samples have smallest and largest average effective grain sizes (≈ 3 μm and ≈ 6 μm, respectively); thus, they exhibit highest and lowest average hardness values (≈ 431 HV and ≈ 402 HV, respectively, Fig. 18). Beside this, another important factor is the strain rate sensitivity (m), which is observed as 0.131 and 0.182 at 880 °C and 1130 °C, respectively, indicating the stability of the hot forging operation performed within this temperature range (Fig. 7). This is further supported by the texture analysis, where a random fiber-type weak deformation texture is appeared in all deformed samples indicating the stability of the hot forging operation within the specified parametric range (Fig. 17). The significance of these findings is the quantification of the key mechanical and microstructural data for the test conditions that closely represent the industrial open die forging of SA508 grade 3 steel. The data produced in this work provide insight into the softening mechanisms occurring in this steel during hot forging. The quantified average PAGS and average effective grain size, the corrected flow stress behavior and the hardness data altogether provide vital inputs to the FE process models making the prediction of mechanical and microstructure properties in the end products more accurate. Thus, the effective forging conditions should always be identified based on the requirement of microstructural and mechanical properties for the end application.

### 6. Conclusions

Following conclusions are drawn from the current work:

- Highest work hardening in SA508 grade 3 steel is observed at the lowest test temperature and at the highest strain rate condition, i.e., 880 °C-1/s, exhibiting highest peak stress. MDRX is the dominant deformation mechanism for this sample resulting in significant grain refinement (finer laths having average effective grain size of ≈ 3 μm) and therefore highest average hardness (≈ 431 HV) as compared to rest of the conditions. In this case, no stable iron oxide film at the interface between the sample and the platens is formed, which leads to highest interfacial friction coefficient (0.82) during hot deformation.

- Flow softening is observed dominant at the highest test temperature and at the lowest strain rate condition, i.e., 1130 °C-0.001/s, showing lowest peak stress. DRV is found as dominant deformation mechanism as opposed to DRX. In this case, the recrystallization occurs simultaneously with the deformation due to prolonged deformation time. As a result, the recrystallized grains are elongated during deformation leading to coarser laths with an average effective grain size of ≈ 6 μm. The highest average effective grain size leads to lowest average hardness (≈ 402 HV) as compared to others maintaining Hall–Petch relationship. Beside this, the prolonged deformation time is beneficial in order to form a stable iron oxide film at the interface between the sample and the platens. This lubricious oxide scale leads to a substantial decrease in the friction coefficient (0.63) as compared to rest of the conditions.

- Within the temperature range of 880-1130 °C, the strain rate sensitivity (m) is observed in between 0.131 and 0.182 indicating stability of the hot forging process within the specified parametric range.

- The hardness of deformed samples is ranged from ≈ 402 to ≈ 431 HV, which is more than twice the amount of hardness exhibited by the as-received material of ≈ 194
Despite the high-temperature deformation and the softening processes, it is the characteristic transformation from bainitic to martensitic microstructure, resulting from water quenching at the end of the hot deformation that leads to this significant hardness increase.

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Data Availability

The raw/processed data required to reproduce these findings cannot be shared at this time due to legal or ethical reasons. Currently, the authors do not have permission to share this data.

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