Effect of B and N Content and Austenitization Temperature on the Tensile and Impact Properties of Modified 9Cr-1Mo Steels

Ravindran Rejeesh 1,*, Rakesh Kumar Barik 1, Rahul Mitra 1, Andrii Kostryzhev 2, Chitta R. Das 3,4, Shaju K. Albert 3,4 and Debalay Chakrabarti 1

Abstract: The present study investigates the relative effect of B and N concentrations and the austenitization temperature on the microstructure and mechanical properties (tensile and Charpy impact) of modified 9Cr-1Mo (P91) steels. Initially, a B-free P91 steel (with 500 ppm N) and four different B-containing steels (25–100 ppm) with varying N concentrations (20–108 ppm) were hot-rolled, normalized from different austenitization temperatures (1000–1100 °C/1 h) and finally tempered at 760 °C for 1 h. A Charpy impact test shows that the ductile–brittle transition temperature (DBTT) of all the B-added steels decreases with an increase in the austenitization temperature, where the 100 ppm B steel offers the lowest DBTT (−85 °C). Similarly, the strength increases with the increase in the austenitization temperature (1100 °C), with a slight drop in ductility. The influence of precipitates on the microstructure and mechanical properties is explained considering the B enrichment at the precipitates and the thermodynamic stability of the precipitates. The 100 ppm B steel (containing the maximum B and minimum N), normalized from 1100 °C austenitization, shows the best combination of tensile and Charpy impact properties, owing to the effective dissolution of coarse M23C6 and MX precipitates during the normalization treatment and the formation of fine B-rich (Fe,Cr)23(B,C)6 precipitates during the subsequent tempering.

Keywords: modified P91 steels; B and N contents; austenitization temperature; auger electron spectroscopy; precipitate stability; microalloyed MX precipitates; tensile testing; impact properties

1. Introduction

Boron-added-modified 9Cr-1Mo (P91) steel is suitable for use as a structural component of steam generators and boilers considering its excellent oxidation resistance and superior creep rupture strength at elevated temperatures, along with its good tensile strength [1,2]. Although these steels are subjected to irradiation, impact toughness remains an important requirement for ensuring structural integrity. Particularly, it is necessary to understand whether the addition of B and N has any adverse effect on the impact toughness. Ensuring sufficient toughness and avoiding hydrogen-assisted cracking (HAC) are among the significant technological obstacles encountered during the welding process of 9Cr-1Mo steel [3,4].

Some recent studies on the effect of the composition have led to the development of the P92 grade steels, where some fraction of Mo in the P91 steel is replaced by tungsten (W) containing tungsten (W) with a lower fraction of Mo compared to P91 steel developed to achieve better mechanical properties and a higher creep rupture strength [5]. These benefits make the 9% Cr steel a dependable option for demanding applications in sectors with
high temperatures and high levels of stress. The higher chances of laves phase nucleation and rapid growth during long-term aging also play a key role in the variation in the static mechanical properties of P92 steels [6]. Apart from the steel grades containing 9 wt.% Cr, some research has also been devoted to studying the influence of zirconium (Zr) and hafnium (Hf) addition in refining the precipitates (mostly the oxide particles) in 14 wt.% Cr and 16 wt.% Cr steels, respectively, to improve both the strength and ductility [7,8]. The improved strength–toughness combination was mainly attributed to the stronger bonding force at the interface between the fine particles and matrix, as well as the increased hindrance provided by the smaller particles against dislocation movements.

Besides the steel composition, the heat treatment schedule, particularly the austenitization temperature, also plays an important role in regulating the impact toughness of high-Cr steels. Chatterjee et al. [9,10] studied the effect of the austenitization temperature (950–1100 °C) on the ductile-to-brittle transition behavior of B-free P91 steel by performing Charpy impact testing. It was found that the austenitization treatment at an intermediate temperature of 1025 °C improved the impact properties due to a smaller ‘effective grain size’, a higher fraction of high-angle boundaries, the presence of a beneficial γ-fiber texture and the dissolution of the pre-existing coarse precipitates. Pandey et al. [11] performed a similar study on B-free P91 grade steel and found that an increase in the normalizing temperature (from 950 °C to 1150 °C) resulted in a decrease in toughness (particularly above 1050 °C), primarily due to the coarsening of M23C6 precipitates, which promotes the cracking of secondary particles and de-cohesion at the precipitate/matrix interface. Such studies are rare in the same grade of steel with different B and N concentrations, which are expected to influence the precipitate coarsening and dissolution during the austenitization treatment.

In general, B is added in high-strength, low-C bainitic/martensitic steels for increasing the hardenability by segregating at the prior austenite grain (PAG) boundaries and thereby suppressing the nucleation of ferrite and pearlite at those sites [12,13]. Such a phenomenon is certainly beneficial in 9Cr-1Mo steel with a tempered martensitic microstructure, as B segregation can strengthen the PAG boundaries, preventing the local softening at those locations during high-temperature exposure. B significantly reduces the coarsening rate of M23C6 precipitates by forming fine and stable boro-carbides by partially replacing C with B, which provides a stronger pinning effect of grain boundaries and sub-grain boundaries [14–23]. In addition to M23C6-type precipitates, MX precipitates (M = V and/or Nb, X = C and/or N) also play a vital role in strengthening. Moreover, the presence of a high amount of N in P91 steel can easily form enough fine and stable MX nitrides which contribute to creep resistance [24–26]. However, excess N can also form brittle BN particles, which act as crack initiation sites [27,28]. Therefore, an optimum concentration of B and N is necessary for improving the creep strength of P91 steel. Our recent work [19] showed that, in base steels containing high B (70–100 ppm), a considerably higher N content (90–110 ppm) is beneficial for creep resistance (contributed by fine and stable MX nitrides) compared to a low N content (20 ppm).

Limited studies have been carried out on the impact toughness of B-containing high-Cr steels. Usually, B is found to be detrimental to the impact toughness of these steels by promoting a temper embrittlement phenomenon due to the stabilization and coagulation of M23(B,C)6 precipitates along the PAG boundaries or lath boundaries [29,30]. In addition, Mishnev et al. [30] observed a deterioration in the impact toughness of 10% Cr-containing steel with an increase in the B content (from 0 to 80 ppm) and a decrease in the N content (from 500 ppm to 30 ppm) due to the formation of chains of closely spaced M23C6 carbides along the lath boundaries. The nucleation of micro-voids from those closely spaced precipitates leads to early void coalescence and cracking. However, the exact effect of N on influencing the toughness is not deliberated in this study.

The present study aims to investigate the synergetic effect of B and N concentrations and the austenitization temperature on the microstructure and tensile and impact properties of modified 9Cr-1Mo steel.
2. Materials and Methods

2.1. Steel Composition

Five different steel grades of modified 9Cr-1Mo steel (B-free and B-added P91 steels) were prepared by an Indian manufacturer following identical vacuum induction melting and casting practices, each grade weighing between 10 and 15 kg (also detailed in our previous work [19]). The chemical compositions of all the prepared steels were determined using optical emission spectroscopy (OES), which are provided in Table 1. The B-free steel is a standard modified 9Cr-1Mo steel that contains a high amount of N (500 ppm), prepared according to the ASTM A213 (or ASME SA213) specifications [31]. On the other hand, in the remaining B-containing steels, different combinations of low-B medium-N (25 ppm B and 90 ppm N), medium-B high-N (70 ppm B and 108 ppm N), high-B medium-N (90 ppm B and 90 ppm N) and high-B low-N (100 ppm B and 20 ppm N) were used for a better understanding of the relative effect of both of the alloying elements. Steels are named after the amount of B addition, e.g., B-free, 25 ppm B, 70 ppm B, 90 ppm B and 100 ppm B (Table 1). The B and N concentrations are chosen in such a way that a controlled compositional variation can be achieved by avoiding the harmful BN formation, which is considered as a potential crack initiator that can deteriorate the steel toughness. To prevent the BN formation, we adhered to the BN solubility curve [19] derived from Equation (1), following [27]. Additionally, all of the other alloying elements in the steel are kept within the standard specification range.

\[
\log[\text{wt.}\%\text{B}] = 2.45 \log[\text{wt.}\%\text{N}] - 6.81 \tag{1}
\]

Table 1. Composition of investigated steels (wt.%).

<table>
<thead>
<tr>
<th>Elements (wt.%)</th>
<th>C</th>
<th>Cr</th>
<th>Mo</th>
<th>Mn</th>
<th>Si</th>
<th>V</th>
<th>Nb</th>
<th>S</th>
<th>P</th>
<th>Ni</th>
<th>Al</th>
<th>Ti</th>
<th>N</th>
<th>B</th>
</tr>
</thead>
<tbody>
<tr>
<td>B-free</td>
<td>0.10</td>
<td>8.93</td>
<td>0.86</td>
<td>0.41</td>
<td>0.21</td>
<td>0.20</td>
<td>0.08</td>
<td>0.005</td>
<td>0.020</td>
<td>0.20</td>
<td>0.010</td>
<td>-</td>
<td>0.050</td>
<td>-</td>
</tr>
<tr>
<td>25 ppm B</td>
<td>0.13</td>
<td>8.65</td>
<td>1.06</td>
<td>0.22</td>
<td>0.11</td>
<td>0.24</td>
<td>0.11</td>
<td>0.006</td>
<td>0.02</td>
<td>0.08</td>
<td>0.003</td>
<td>-</td>
<td>0.009</td>
<td>0.0026</td>
</tr>
<tr>
<td>70 ppm B</td>
<td>0.15</td>
<td>9.35</td>
<td>0.98</td>
<td>0.37</td>
<td>0.33</td>
<td>0.11</td>
<td>0.10</td>
<td>0.004</td>
<td>0.002</td>
<td>0.05</td>
<td>0.004</td>
<td>0.002</td>
<td>0.011</td>
<td>0.0070</td>
</tr>
<tr>
<td>90 ppm B</td>
<td>0.11</td>
<td>8.70</td>
<td>0.96</td>
<td>0.25</td>
<td>0.10</td>
<td>0.25</td>
<td>0.08</td>
<td>0.007</td>
<td>0.02</td>
<td>0.11</td>
<td>0.002</td>
<td>-</td>
<td>0.009</td>
<td>0.0090</td>
</tr>
<tr>
<td>100 ppm B</td>
<td>0.10</td>
<td>8.50</td>
<td>1.00</td>
<td>0.30</td>
<td>0.40</td>
<td>0.23</td>
<td>0.09</td>
<td>0.002</td>
<td>0.005</td>
<td>0.02</td>
<td>0.030</td>
<td>-</td>
<td>0.002</td>
<td>0.0100</td>
</tr>
</tbody>
</table>

2.2. Material Processing and Characterization

The as-cast ingots were homogenized at 1250 °C, hot-forged and hot-rolled (finish-rolled at ~1050 °C) to 10 mm thick plates. The normalization (N) treatment comprised austenitization at 1000/1050/1100 °C for 1 h and air-cooling followed by tempering (T) at 760 °C for 1 h. Details of the heat treatment schedule can be found in [19].

Detailed microstructural characterization was performed on the RD-ND plane of the heat-treated plates (shown in Figure 1 and also reported in our previous article [19]), which includes optical microscopy, scanning electron microscopy (SEM), transmission electron microscopy (TEM) and auger electron spectroscopy (AES) techniques. The details of all the characterization techniques employed are provided in our previous article [19].

2.3. Tensile and Charpy Impact Testing

Tensile specimens were fabricated from the heat-treated plates as per the ASTM E8 standard [32] and were tested at room temperature with a strain rate of $3.3 \times 10^{-4}$ s$^{-1}$ using an Instron servo hydraulic test system with a 100 kN load capacity. Standard Charpy V-notch specimens were fabricated and tested in a 400 J capacity pendulum-type machine following ASTM E23 [33] at different temperatures over the range of +100 °C to −196 °C. The dimensions and orientations of the tensile and Charpy impact specimens, extracted from the rolled and heat-treated plates, are shown in Figure 1. As a note, for both the tensile and Charpy tests, three specimens were evaluated under each testing
condition, and the reported properties represent the average results obtained for each heat-treatment condition.

Figure 1. Schematic representation of the samples used for the different microstructural analysis and mechanical testing with respect to rolled and heat-treated plate.

3. Results

3.1. Microstructural Characterization

The microstructural characterization performed on the investigated steels is reported in detail in our previous article [19]. Some of the important microstructural observations are summarized below.

- The investigated specimens display a tempered lath martensitic structure, with finer PAG and packet sizes in B-free steel compared to B-added steels, indicating boron’s role in stabilizing the microstructure;
- With an increase in the austenitization temperature (AT), the PAG size increases in all the steels. It was found that 70 ppm B steel shows the minimum variation in the PAG size, packet size and lath width with a change in AT, followed by 100 ppm B steel. However, the 25 ppm B and 90 ppm B steels show a significant increase in the PAG size (>50 μm) and packet size (>30 μm) with an increase in AT. Optical micrographs showing the change in the prior austenite grain size (PAGS) of 100 ppm B steel with the increase in AT are presented in Figure 2a–c;
- The low-angle boundaries (LABs) were sub-grain boundaries (particularly in B-free steel) and lath boundaries (in B-containing steels). The fraction of LABs (2–15° misorientation angle), estimated from the EBSD analysis, was found to be at the maximum in the 1000 °C-austenitized B-free specimen and decreased with the increase in AT. The same trend was also followed by the 100 ppm B steel. In contrast, the 70 ppm B steel displayed the minimum existence of LABs for 1000 °C AT, whilst it reached the maximum value at 1100 °C AT;
- High-angle grain boundaries (HABs) comprised PAG boundaries, a packet and block boundaries. The HABs of B-free steel were consistently high for all the ATs (due to the refined grain structure). It was found that 70–100 ppm B steels revealed a consistently low occurrence of HABs that remained stable with AT. The density of HABs decreased linearly with the increase in AT for the 25 ppm B steel. At 1100 °C, both the 25 ppm B steel and the 90 ppm B steel reached the minimum intensity of HABs;
- Undissolved precipitates were more prevalent in samples austenitized at lower temperatures, mainly due to the decreased likelihood of the pre-existing precipitates dissolving at lower temperatures. After austenitizing at a higher temperature (1100 °C), M$_{23}$C$_6$-type precipitates completely dissolved, while only partially or undissolved MX 
precipitates were retained. During the subsequent tempering, the dissolved solute atoms formed fresh and fine precipitates. SEM (M_{23}C_{6} precipitates) and TEM (M_{23}C_{6} and MX precipitates) images, showing precipitates formed after tempering, are exhibited in Figure 2d–i. For a lower AT, the undissolved precipitates tend to coarsen during tempering:

- As the AT increases, there is a reduction in the average precipitate size (>50 nm; mostly M_{23}C_{6}) and an increase in the volume fraction after the tempering treatment;
- Among the B-added steels, 100 ppm B steel possesses the smallest precipitate size, whereas the 70 ppm B steel exhibits larger precipitates and a higher volume fraction, regardless of the AT, compared to other B-added steels;
- The AES study confirms the segregation of B exactly at the precipitates in all the steels nearby PAG boundaries and sub-grain boundaries closer to the PAG boundaries.

All of the above findings reported need to be correlated with tensile and impact testing results to understand the influence of alloying elements, precipitates and the microstructure.

Figure 3 presents the AES results for B segregation in 100 ppm B steel under different heat treatment conditions. In the as-normalized condition (N), B segregated preferentially at PAG boundaries and nearby lath boundaries. B segregation was higher in the specimen austenitized at 1100 °C (as shown in Figure 3a,b) compared to the specimen austenitized at a lower temperature of 1000 °C (Figure 3c,d). As a note, B segregation near PAG boundaries usually promotes the formation of B-rich precipitates (M_{23}(B,C)_{6}) during subsequent tempering [34–37]. The enrichment of B in the (Fe,Cr)_{23}C_{6} precipitates after the tempering treatment is evidenced in Figure 3e–h in 100 ppm B and 70 ppm B steels (also elucidated in [19]). The B enrichment in the M_{23}(B,C)_{6} precipitates reduces the coarsening rate of those precipitates.
Table 2. Tensile properties of the investigated steels after normalization followed by tempering (1000/1100 °C N+T).

<table>
<thead>
<tr>
<th>Tensile Properties</th>
<th>B-free</th>
<th>25 ppm B</th>
<th>70 ppm B</th>
<th>90 ppm B</th>
<th>100 ppm B</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1000 °C N+T</td>
<td>1100 °C N+T</td>
<td>1000 °C N+T</td>
<td>1100 °C N+T</td>
<td>1000 °C N+T</td>
</tr>
<tr>
<td>Yield strength (YS)</td>
<td>458.1 ± 5</td>
<td>467.9 ± 6</td>
<td>490.6 ± 6</td>
<td>511.2 ± 7</td>
<td>478.1 ± 6</td>
</tr>
<tr>
<td>UTS</td>
<td>656.1 ± 6</td>
<td>669.4 ± 4</td>
<td>682.6 ± 5</td>
<td>705.0 ± 5</td>
<td>660.2 ± 4</td>
</tr>
<tr>
<td>%EL</td>
<td>30.5 ± 1.3</td>
<td>28.9 ± 0.9</td>
<td>25.6 ± 0.9</td>
<td>22.7 ± 0.7</td>
<td>25.9 ± 1.2</td>
</tr>
</tbody>
</table>

* Tests were repeated, and the deviations in measurements were within 5%.

The important observations are as follows.
• All the steels austenitized at 1100 °C possess superior tensile strength as compared to those austenitized at 1000 °C;
• B-containing steels possess higher YS and UTS compared to B-free steel irrespective of the austenitizing conditions;
• The B-free steel that was austenitized at 1000 °C exhibited the lowest YS (458 MPa) and UTS (656 MPa), whereas the highest YS (543 MPa) and UTS (711 MPa) were observed in the 90 ppm B steel after austenitizing at 1100 °C. The strength of the steel is as follows: 90 ppm B > 25 ppm B > 100 ppm B > 70 ppm B > B-free;
• Compared to other B-containing steels, the 100 ppm B steel displays good tensile strength and significant ductility. However, at 1100 °C, the 25 ppm B and 90 ppm B steels exhibit the highest tensile strength, but they have the least ductility among all the steel types and heat treatment conditions, with only 23% and 24% total elongation, respectively. Among the steels heat-treated at 1000 °C, both these steels exhibit the lowest ductility (26% each). B-free has the highest percentage elongation, measuring 31% and 29% at 1000 °C and 1100 °C, respectively, followed by the 100 ppm B steel, with 29% total elongation at 1000 °C and 28% at 1100 °C.

3.3. Charpy Impact Properties

The Charpy impact transition curves of the B-free and B-added steels for different N+T conditions are shown in Figure 5. All the transition curves were fitted with the ‘tanh’ function in the equation below using the data measured via the Charpy impact testing,

\[ C_v = A + B \times \tanh \left( \frac{(T - T_0)}{C} \right) \]  \hspace{1cm} (2)

where A, B and C are constants, \( T \) = testing temperature and \( T_0 \) = ductile–brittle transition temperature (DBTT) estimated from the mean of the corresponding impact energy observed in the upper shelf region (USE) and lower shelf region (LSE) [38]. The major findings observed from the curves are listed below:

• The impact properties of 100 ppm B steel appear to be superior although more sensitive to heat treatment conditions compared to those of the other steels. In the case of 100 ppm B steel, the USE and DBTT vary over a wider range with the increase in AT (175 J to 235 J USE with –55 °C to –85 °C DBTT);
• Among the austenitization temperatures, all the B-added steels show the best impact properties at 1100 °C AT, in which 100 ppm B steel possesses the lowest DBTT (–85 °C), followed by 70 ppm B (–65 °C), 90 ppm B (–51 °C) and 25 ppm B (–36 °C);
• At a lower AT (1000 °C N+T), the 70 ppm B steel possesses the lowest DBTT (–60 °C). Moreover, the least change in DBTT (–60 °C to –65 °C) and USE (147 J to 175 J) with increasing AT was observed in 70 ppm B steel among the investigated steels;
• The USE values of 25 ppm B and 90 ppm B steels vary from 153 J to 184 J and from 147 J to 178 J, respectively, with increasing AT;
• For B-free steel, the best impact properties were obtained at 1050 °C AT, with lower USE (197 J) and considerably lower DBTT (–59 °C) compared to those of the 1100 °C AT sample (216 J USE, –32 °C DBTT).

Figure 6 displays the fracture surfaces of the Charpy impact-tested 100 ppm B steels, which depict a transition from the purely ductile mode of fracture at higher temperatures to the mixed mode at intermediate temperatures and the purely brittle mode of fracture at very low temperatures. Different arrows are shown in each regime. The red arrows indicate the presence of cleavage facets, which indicates the brittle mode of fracture, especially for the micrographs from the lower shelf region containing numerous facets along with de-cohesion along the PAG boundaries. The presence of cleavage facets and dimples (green arrows) represents the transition zone or the brittle-to-ductile transition of the fracture surface, and it becomes completely ductile (only green arrows) once it reaches the upper shelf region of the transition curve.
The USE values of 25 ppm B and 90 ppm B steels vary from 153 J to 184 J and from 147 J to 178 J, respectively, with increasing AT; for B-free steel, the best impact properties were obtained at 1050 °C AT, with lower USE (197 J) and considerably lower DBTT (−59 °C) compared to those of the 1100 °C AT sample (216 J, −32 °C DBTT).

Figure 5. Ductile–brittle transition curves (Charpy impact energy vs. temperature) of the investigated steels normalized at 1000/1050/1100 °C for 1 h followed by tempering at 760 °C for 1 h of (a) B-free steel, (b) 25 ppm B steel, (c) 70 ppm B steel, (d) 90 ppm B steel and (e) 100 ppm B steel. (f) Table showing the upper shelf energy and DBTT of all the investigated steels.

Figure 6. (a–i) Fractographs of Charpy impact-tested specimens from different zones of the ductile-to-brittle transition curve generated for 100 ppm B steels. Blue, Red and Green arrows represent decohesion along PAG boundaries, cleavage facets and ductile dimples, respectively. The heat-treatment condition, Charpy impact testing temperature and steel codes are marked in each micrograph.

4. Discussion
Superior tensile properties at higher austenitization temperatures can be attributed to the proper dissolution of pre-existing precipitates and the formation of finer precipitates during tempering. Figure 7 depicts the detailed characterization of both the M23C6 and MX undissolved precipitates observed in the TEM study, confirmed using energy dispersive spectroscopy (EDS) and the selected area diffraction pattern (SADP) on the steels after normalizing (i.e., before tempering). As the AT decreases, there is an increase in the number of undissolved precipitates that tend to coarsen during tempering, leading to a deterioration in strength. Additionally, highly stable MX nitrides/carbides containing mainly V and Nb also significantly contributed to precipitation strengthening. MX nitrides possess a higher stability than MX carbides, and 70 ppm B steel contains the highest fraction of N among the B-containing steels.
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![Figure 7. TEM bright field and dark field images of (a–d) B-free, (e–h) 70 ppm B and (i–l) 100 ppm B steels after austenitization at 1000/1100 °C (N). Different undissolved precipitates are marked based on their corresponding EDS data (shown as table insets) and SAD patterns.](image)

In 100 ppm B steel, even though the C content is less (0.10 wt.%), the finest ($\text{Fe,Cr})_{23}(\text{B,C})_6$ precipitates (with the highest fraction) along with a relatively low fraction of coarser $\text{Cr}_{23}\text{C}_6$ contributed to the good strength for both austenitization conditions, particularly at 1100 °C AT. Despite the highest C content (0.15 wt.%) and sufficient N content (108 ppm) present in the 70 ppm B steel, a moderate tensile strength and ductility were observed, possibly due to the largest average $\text{M}_{23}\text{C}_6$ precipitate size among the B-containing steels. The higher N content in 25 ppm B and 90 ppm B steels (both 90 ppm N) compared to that in 100 ppm B steel (20 ppm N) can potentially form a higher fraction of MX nitrides along with B-containing carbides, resulting in a superior tensile strength. At the same time, a comparatively higher C content (0.11 and 0.13 wt.%) might possibly reduce the ductility. Additionally, the 70 ppm B steel showed the least change in the $\text{YS}$, $\text{UTS}$ and percentage elongation with AT, which can be attributed to the consistent microstructural features. In B-free steel, the average $\text{M}_{23}\text{C}_6$ size was measured to be the highest irrespective of the austenitization condition, which possibly reduced its tensile strength.
The presence of elements such as C and Cr, which dissolved during austenitization, promotes solid solution strengthening in steels and can lower the impact toughness. It was found that 70 ppm B steel contains a higher fraction of both C and Cr compared to other steels, and this may have reduced the impact energy absorbed in the upper shelf region of the transition curve. The 100 ppm B steel has the least amount of C and Cr; hence, the effect of solid solution strengthening is reduced. Consequently, it exhibits the highest upper shelf energy and lowest DBTT among all the steels investigated. The addition of Ni is beneficial for impact properties [3], and 70 ppm B steel has the least amount of Ni, which may have not contributed enough for better impact properties. Similarly, B-free steel also possesses good impact properties owing to its low C and Cr content as well as a higher Ni content. The 25 ppm B and 90 ppm B steels contain a lower amount of Cr and a higher amount of Ni than the 70 ppm B steel, which resulted in a slight improvement in the impact energy absorbed.

Auger electron spectroscopy (in Figure 3) confirmed the segregation of B and the formation of B-enriched precipitates, which are more stable and fine than the B-free M
\(_{23}\)C
\(_{6}\) precipitates [19]. The formation of (Fe,Cr)\(_{23}\)(B,C)\(_{6}\) reduces the Cr\(_{23}\)C\(_{6}\) fraction, and these precipitates along with a small fraction of fine MX precipitates restrict the prior austenite grain refinement during the austenitization of B-added steels. The stabilization of fine precipitates along the martensite lath (and block) boundaries facilitates the growth of austenite as fine films leading to the reconstruction of PAG [34,35,39]. During the progress of austenite reversion, the austenite films grow rapidly (possibly following a shear transformation mechanism) and consume the martensite block, within which the laths are situated. Thus, austenite films grow ‘block-by-block’, finally coalescing with each other to reconstruct the entire PAG. The retention of fine and stable precipitates along the lath boundaries further reduces the Cr\(_{23}\)C\(_{6}\) fraction, and these precipitates along with a small fraction of fine MX precipitates restrict the prior austenite grain refinement during the austenitization of B-added steels. The stabilization of fine precipitates along the martensite lath (and block) boundaries facilitates the growth of austenite as fine films leading to the reconstruction of PAG [34,35,39]. During the progress of austenite reversion, the austenite films grow rapidly (possibly following a shear transformation mechanism) and consume the martensite block, within which the laths are situated. Thus, austenite films grow ‘block-by-block’, finally coalescing with each other to reconstruct the entire PAG. The retention of fine and stable precipitates along the lath boundaries further ensures the transformation of austenite into a lath martensite structure upon cooling instead of a sub-grain structure. Thus, stable (Fe,Cr)\(_{23}\)(B,C)\(_{6}\) precipitates can lead to a coarser PAG and packet size in B-containing steels, which can reduce the fraction of HABs. Now, the HABs are known to retard and divert the cleavage crack propagation, particularly at low test temperatures, and a finer PAG structure can be beneficial in this respect [9]. However, in B-free steel, although the PAG size is fine, mostly, a sub-grain structure formed (with a high density of low-angle sub-boundaries), as the lath martensite structure could not be retained due to the lower precipitate stability. The LABs are ineffective in retarding the cleavage crack propagation. The B-free steel also contained coarser precipitates, which lowered the precipitation strengthening contribution and, consequently, the overall strength. A lower strength is expected to promote plastic deformation and improve the impact toughness, particularly at the upper shelf region. As a result, B-free steel showed relatively higher USE and an intermediate DBTT.

The best impact properties (high USE and low DBTT) of 100 ppm B steel can be attributed to (i) the fine precipitate size (due to the B stabilization effect) and (ii) the low fraction of MX precipitates (due to the low N content). The fractions of M\(_{23}\)C\(_{6}\) and MX precipitates are predicted by Thermo-Calc® software for the investigated steel compositions and are reported in our previous study [19]). Fine precipitates nucleate small voids, and the scope for the sufficient growth of those voids contributes to the impact toughness. Fine precipitates are also less prone to initiating cleavage crack than coarse precipitates. The lower precipitation strengthening can also be beneficial for toughness. The best impact properties of 100 ppm B steel (with just 20 ppm N) indicate the dominant role of the precipitate size and fraction on the impact toughness compared to the microstructural refinement in the B-added P91 steels. In spite of the high precipitate and microstructural stability, 70 ppm B steel contained relatively coarser M\(_{23}\)C\(_{6}\) precipitates and a higher fraction of MX precipitates than 100 ppm B steel. Therefore, the impact properties of 70 ppm B steel were not as good as those of 100 ppm B steel. The combination of the precipitate size and fraction (and the extent of microstructural refinement) also resulted in inferior impact properties of 25 ppm and 90 ppm B steels as compared to those of 100 ppm
B steel. Thus, the impact properties obtained in the present study contradict the findings of Mishnev et al. [30].

Besides the level of DBTT, its variation with AT is different between B-free steel (lowest DBTT at an intermediate AT of 1050 °C) and the B-containing steels (decrease in DBTT with the increase in AT). In B-free P91 steel, Chatterjee et al. [9] also obtained the lowest DBTT at the intermediate AT (1025 °C). Such a finding can be explained by the optimum precipitate size and grain size (‘effective grain size’ in this case) at an intermediate AT, since the precipitates become coarse at a low AT, whilst the grains become coarse at a high AT. In B-containing steels, precipitates appear to govern the impact properties, and, hence, DBTT decreases with the increase in AT. The presence of undissolved precipitates at a lower AT and the subsequent coarsening of those precipitates during tempering were detrimental to the impact toughness. At higher testing temperatures, where the ductile fracture dominates, the coarse precipitates nucleate large voids which coalesce into cracks without sufficient growth, thereby lowering the impact toughness. On the other side, at lower test temperatures, coarse particles can act as the initiation sites for cleavage cracks, again affecting the toughness.

5. Conclusions

After performing different microstructural characterization and mechanical testing, the main points are summarized below.

- The B-free steel possesses the best ductility (%EL) among all the investigated steels (including the austenitization temperature, AT and variation) but poor YS and UTS. The poor strength of B-free steel is mainly attributed to the presence of large, undissolved $M_{23}C_6$ and MX precipitates. The lower C and Cr contents as well as the higher Ni content in B-free steel greatly contributed to its high ductility as well as good impact properties (mainly the USE). Moreover, the high fraction of HAB (due to refined PAGs) in the B-free steel greatly improves the low temperature impact toughness, which thereby results in a lower DBTT;

- In the B-containing steels, both the strength as well as the impact properties are mainly controlled by the precipitate characteristics rather than the PAG size. Increasing the AT dissolves the coarse $M_{23}C_6$ precipitates, while increasing the B content helps in stabilizing the formation of fine B-rich ($\text{(Fe,Cr)}_{23}(\text{B,C})_6$) precipitates during the subsequent tempering. Increasing the N content results in larger and more abundant MX precipitates that are detrimental for toughness. As a whole, the presence of a high fraction of fine ($\text{(Fe,Cr)}_{23}(\text{B,C})_6$) precipitates and low fractions of coarse $M_{23}C_6$ and MX precipitates leads to an enhanced strength–impact toughness combination, which can be achieved by increasing the AT and B content and decreasing the N content of the B-added P91 steels;

- The 100 ppm B steel (containing the maximum B and minimum N), normalized at the maximum AT of 1100 °C, shows the best combination of tensile properties (YS: 513 MPa, UTS: 676 MPa and %EL: 28 %) and Charpy impact properties (USE: 235 J and DBTT: −85 °C), owing to the abundance of fine B-rich ($\text{(Fe,Cr)}_{23}(\text{B,C})_6$) precipitates and the deficiency of coarse $M_{23}C_6$ and MX precipitates.


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