Effect of Parameters in the Physical Simulated Rough Rolling Stage on Microstructure Evaluation and Tensile Properties of a Bainitic Pipeline Steel

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Abstract. Microstructure evolution and tensile properties were studied in a bainitic pipeline steel grade by performing a number of physical simulations on samples machined out of an industrially produced transfer bar. In these simulations, the cooling interval between roughing and finishing stages ($t_V$) was varied from 5 s to 180 s. The austenite status after this cooling interval, regarding the prior austenite grain size and precipitates, simulates the condition of austenite before entering the finishing mill. The finishing parameters and the subsequent cooling strategy were kept unchanged throughout all the applied simulation processes. The gradual increase in $t_V$ resulted in a gradual increase of the granular bainite phase on the expense of the acicular ferrite. This resulted in an incremental decrease in ultimate tensile strength and yield strength with increasing $t_V$. However, this behavior approached a steady state condition after which the $t_V$ has limited/insignificant effect on the ultimate- and yield strength. This saturating value of $t_V$ is process parameter dependent.

Introduction

The demand of the pipeline industry for a more cost-effective pipeline design has pushed the standard pipeline steel grade requirements. Critical to the design of these steels is a low carbon equivalent for good field weldability [1, 2]. In these steels, the carbon is reduced to below 0.09 wt.%. The strength loss due to the low C content is compensated through alloy design philosophy based on the advanced use of cost effective micro-alloying elements, such as Nb, Ti and B in conjunction with moderate levels of other alloying elements, such as Mn, Si, Cr, Mo and Cu [3]. The use of aforementioned combinations of micro-alloying and alloying elements in conjunction with thermo-mechanical controlled processing (TMCP) lead to the development of API X80, X100 and X120 which exhibit yield strengths from 550 MPa up to 825 MPa [4]. In these steels, the desired balance of mechanical properties at a given steel composition are achieved through suitably designed thermo-mechanical processing schedules [5], which commonly involve controlled rolling, followed by controlled accelerated cooling. The controlled rolling compresses two stages, namely roughing and finishing rolling. Roughing starts after the austenitization process. During the rough rolling the austenite grain size is refined due to repeated cycles of work hardening and the recrystallization process. The finishing rolling starts subsequent to the roughing. During the finishing rolling the austenite is deformed in the non-recrystallization temperature regime, which brings significant refinement to the final microstructure. The accelerated cooling step aims to suppress the formation of polygonal ferrite and, instead, encourage non-equilibrium, non-equiaxed ferrite microstructures to be formed. The latter transformation products are known to contribute to increasing strength, through both small effective grain sizes and increased dislocation densities, while maintaining a reasonable level of toughness [1, 5, 6].

Objective. It was shown in [7] that decreasing the delay-time between the roughing and finishing rolling stages ($t_V$) from 180 s to 5 s resulted in pronounced improvement in both of ultimate tensile strength and proof stress. During the current work the same steel is processed under the same
thermo-mechanical (TM) processing parameters except that and intermediate values of $t_V$ between 5 s and 180 s are selected. The effect of the cooling time on the microstructure development and the mechanical properties is investigated.

Experimental Procedure

Material and Specimens Preparation. The current study is carried out on samples machined out of a transfer bar of API X80 pipeline steel. Salzgitter Flachstahl GmbH is acknowledged for providing the raw material. The chemical composition of the studied material is given in Table 1.

Table 1. Chemical analysis of the studied material (wt. %)

<table>
<thead>
<tr>
<th></th>
<th>C</th>
<th>Si</th>
<th>Mn</th>
<th>P</th>
<th>N</th>
<th>Cr</th>
<th>Mo</th>
<th>Ti</th>
<th>Nb</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.055</td>
<td>0.3</td>
<td>1.84</td>
<td>0.014</td>
<td>0.006</td>
<td>0.18</td>
<td>0.259</td>
<td>0.0256</td>
<td>0.101</td>
<td>0.0008</td>
</tr>
</tbody>
</table>

The provided steel slab has a thickness of 52 mm. All the specimens were taken with their longitudinal axes parallel to the rolling direction of the transfer bar and their thicknesses parallel to its thickness. The thickness-center region is excepted during machining of the specimens that is to avoid the zone of central segregation of the slab. The dimensions of the flat compression samples are shown in Fig. 1. The thickness of the specimen in the testing-zone is 6.4 mm. The specimen has 42 mm shoulders for clamping in the tensile testing machine after thermomechanical processing. The two Ø 6 mm holes are for reducing the heat dissipation from the testing-zone to the shoulders. All the simulation specimens are taken with their longitudinal axes parallel to the rolling direction and their thicknesses parallel to the thickness of the transfer bar.

![Fig. 1. Dimensions of the flat compressing sample](image)

TM simulator TTS 820 is used for carrying out the simulation process. For this purpose, a thermocouple is spot welded on the specimen, and then the specimen is placed on two ceramic rollers and fixed from the upper side by two ceramic rods. Two deformation stamps upset the specimen in its center. A detailed description of the flat compression setup of TTS820 is given in [7].

Thermo-mechanical Simulation. The samples with a geometry shown in Fig. 1 are subjected to the TM schedule sketched in Fig. 2. In this schedule, specimens were heated up to the austenitization temperature ($T_A$) and subjected to one deformation step with a true strain value of $\varphi_v$ at $T_V$. The austenite status at this stage - regarding the prior austenite grain size (PAGS) and precipitation - simulates the condition of austenite after the roughing process. The subsequent three deformation steps are to simulate the finishing rolling process, the time between roughing and finishing is designated in the figure by $t_V$. The studied parameters are varied according to the values listed in Table 2. The finishing rolling parameters and the subsequent cooling strategy were kept unchanged throughout all the applied simulation processes. The parameters in Table 2 are considered for varying the austenite status before entering the finishing mill. Two values of $t_V$ were
investigated in reference [7], namely 5 s and 180 s. The results of this project showed a strong
dependence of the ultimate and proof strength on tv. During the current study, additional
intermediate values for tv were investigated, written in italic font-style in Table 2.

![Schematic drawing of the applied thermo-mechanical schedule.](image)

**Table 2. Combination of parameters studied (Fig. 2)**

<table>
<thead>
<tr>
<th>TA (°C)</th>
<th>1285</th>
<th>1185</th>
</tr>
</thead>
<tbody>
<tr>
<td>TV (°C)</td>
<td>1000</td>
<td>1100</td>
</tr>
<tr>
<td>ϕv (-)</td>
<td>0.3</td>
<td>0.5</td>
</tr>
<tr>
<td>tv (s)</td>
<td>5</td>
<td>30</td>
</tr>
</tbody>
</table>

**Light Optical Microscopy.** Light optical microscopic (LOM) analysis of the as-received
samples as well as samples from various processing and conditions was performed by sectioning the
samples parallel to the deformation direction, and cold mounting. The samples were rough polished
using standard metallographic abrasive grinding papers ranging from coarse (180) to fine (1200).
The final polishing was done using 1.0 µm and 0.05 µm alumina, respectively. After polishing, the
samples were rinsed with ethyl alcohol and dried. The microstructure was developed by etching
with 2 % Nital.

**Tensile Testing.** The tensile tests were conducted in a computerized universal testing machine
(UTS) with a 250 kN load cell using a crosshead speed of 5 mm/min.

**Results and Discussion**

**Microstructure Evolution.** Metallographic investigations using LOM were conducted to
investigate the effect of tv on the microstructure development.

**TA = 1250 °C – TV = 1000 °C - ϕv = 0.3.** The effects of tv on the microstructure for the samples
austenitized at TA = 1250 °C and deformed at TV = 1000 °C with ϕv = 0.3 is shown in Fig. 3. The
micrographs of Fig. 3 show that the (Nb, Ti) (C, N) precipitates are well distributed in all
microstructures. The composition of these precipitates was investigated by using energy dispersive
X-ray spectroscopy (EDX) [7]. TiN precipitates form at higher temperature in the austenite region.
These precipitates serve as “cores” for the nucleation and epitaxial growth at lower temperatures of
shell of NbCN [8]. The obtained precipitates have an average size of about 132 nm. The addition of
Nb and Ti to pipeline steels effectively refines the austenite grain during the hot-rolling process
because the precipitates retard austenite recrystallization and, in turn, refine the final microstructure.
This microstructure refinement together with the existence of nano-size phase, which is the
precipitates themselves result in enhancing the mechanical properties of the pipeline steel. The microstructure is predominantly a mixture of acicular ferrite (AF) and granular bainite (GB). The microstructures for the samples having \( t_V = 180 \text{ s} \) (Fig. 3e) is dominated by the GB structure. For \( t_V = 5 \text{ s} \) (Fig. 3a), the microstructures show more AF and finer GB than that obtained for \( t_V = 180 \text{ s} \). The domination of the GB structure is also observed in the microstructures of the samples with \( t_V = 60 \text{ s} \) and 120 s. The sample with \( t_V = 30 \text{ s} \) shows more or less similar microstructural features to that for \( t_V = 5 \text{ s} \). The very tiny phase, e.g. the encircled phase in Fig. 3, is defined as a martensite/austenite (M/A) phase; this is confirmed by scanning electron microscopic investigations as shown in [7]. The occurrence of tiny martensite/austenite (M/A) phase is more pronounced for \( t_V = 5 \text{ s} \) and 30 s than for \( t_V = 60 \text{ s}, 120 \text{ s} \) and 180 s. The shorter cooling time between the roughing and finishing resulted in finer and/or pancaked prior austenite grains which motivated the formation of both, AF and fine M/A phases.

\[
T_A = 1150 \, ^\circ\text{C} - T_V = 1000 \, ^\circ\text{C} - \varphi_V = 0.3.
\]

A similar effect of \( t_V \) on the phase distribution of AF and GB in the samples with \( T_A = 1250 \, ^\circ\text{C} \) is observed in samples with \( T_A = 1150 \, ^\circ\text{C} \) (see Fig. 4). Explicitly, increasing \( t_V \) to 180 s resulted in the domination of the GB phase. Furthermore, for \( t_V = 60 \text{ s} \), the samples showed similar features compared to samples with \( t_V = 180 \text{ s} \) rather than to that with \( t_V = 5 \text{ s} \).
**Mechanical Properties.** Fig. 5 shows the effect of \( t_V \) on the stress-strain curves for different TM treatment conditions. The incremental increase in \( t_V \) generally results in an incremental decrease in ultimate tensile strength (\( R_m \)) and yield strength (\( R_p \)). e.g. 840 MPa was the highest recorded \( R_m \) value of the steel with \( T_A = 1250 \, ^\circ\text{C} - T_V = 1000 \, ^\circ\text{C} - \phi_V = 0.5 \) when \( t_V = 5 \) s. The lowest \( R_m \) value of 692 MPa was recorded for the same condition but with \( t_V = 180 \) s. However, it seems that this behavior has a saturation point after which \( t_V \) has a limited/insignificant effect on \( R_m \) and \( R_p \). The saturation point for large prior austenite grains (\( T_A = 1250 \, ^\circ\text{C} \)) deformed at a low temperature (\( T_V = 1000 \, ^\circ\text{C} \)) is not reached at \( t_V = 60 \) s (Fig. 5a and 5b) but rather at \( t_V = 120 \) s (Fig. 5a). On the other hand, a value of 60 s for \( t_V \) was enough to attain the saturation point for large prior austenite grains deformed at high temperature (Fig. 5b and 5c). For smaller PAG (\( T_A = 1150 \, ^\circ\text{C} \)), \( \phi_V \) is decisive for the saturating \( t_V \); for \( \phi_V = 0.3 \) a \( t_V \) of 60 s was enough for attaining saturation in \( R_m \) and \( R_p \) values (Fig. 5c). Higher \( t_V \) is required to attain this saturation for \( \phi_V = 0.5 \) (Fig. 5f).

Fig. 5 Stress-strain curves of samples processed under the prescribed TM treatment conditions.

**Summary**

Microstructure evolution and tensile properties were studied in a pipeline steel grade API-X80 by performing a number of physical simulations on samples machined out of an industrially produced transfer bar. In this physical simulation, specimens were heated up to the austenitization temperature (\( T_A \)) and subjected to one deformation step having a true strain value of \( \phi_V \) at \( T_V \). The cooling interval between roughing and finishing is designated by \( (t_V) \). The austenite status after this cooling interval, regarding the prior austenite grain size (PAGS) and precipitates simulates the condition of austenite before entering the finishing mill. The finishing parameters and the subsequent cooling strategy were kept unchanged throughout all the applied simulation processes. The results showed a strong dependence of the \( R_m \) and \( R_p \) on \( t_V \). The gradual increase in \( t_V \) results in a gradual increase of the granular bainite phase on the expense of the acicular ferrite. This results in an incremental decrease in \( R_m \) and \( R_p \) with increasing \( t_V \). However, it seems that this behavior has a saturation point after which the \( t_V \) has a limited/insignificant effect on the \( R_m \) and \( R_p \). This saturating value of \( t_V \) is process parameter dependent.
References


