Effect of grain size on the drawability of the niobium-stabilized ferritic stainless steel ASTM 430

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Tarcísio Reis de Oliveira 1

Abstract

Studies were carried out with samples of AISI 430 stainless steel stabilized with niobium produced by direct reduction (just one cold rolling process) to evaluate the effects of grain size on the material deep drawability. Recrystallized samples were heat treated in laboratory to promote the growth of the recrystallized grain. Microstructural characterizations were done by X-ray diffraction for evaluating crystallographic texture, optical microscopy and electron backscatter diffraction (EBSD). The drawability was evaluated by tensile tests for determination of the planar ($\Delta R$) and normal ($r$) anisotropy coefficients. An increase in the grain size promoted an increase in the intensity of the Gamma fiber (beneficial) and a reduction in the intensity of the Theta fiber (damaging) on the surface of the material’s thickness. At the centerline of the thickness, it was observed that the Gamma fiber’s intensity did not change significantly, and that the Theta fiber’s intensity was reduced. The change of the crystallographic texture, promoted by the increase of grain size, increased the coefficient of anisotropy ($r$) in all the analyzed directions and decreased the planar anisotropy coefficient.

Keywords: Anisotropy; Texture; Grain size; Deep drawability.

1 Introduction

Stainless steels have excellent corrosion resistance and elevated mechanical properties which allow them to be used in several applications in corrosive environments. In the stainless steel family, ferritic stainless steels are used in applications that require high corrosion resistance, good surface quality and good formability [1].

In the 90’s, Aperam South America developed a ferritic stainless steel ASTM 430 stabilized with Niobium, aiming to improve its formability [1]. The addition of Niobium inhibits the material recrystallization during the hot rolling process through the precipitation of Niobium carbonitride and the effect of Niobium in solid solution, resulting in a hot rolled coil with a more refined and homogeneous structure, improving the properties of the end product [2].

The stamping performance of ferritic steels is different in comparison to austenitic. During stamping, austenitic steels exhibit good stretching resistance and good strain distribution, allowing manufacturing of complex geometries. Ferritic stainless steels are commonly used for stamping pieces, where the deep drawing processes is predominant [3].

The ferritic stainless steel formability can be evaluated by the Lankford coefficient $r(\alpha)$, which is defined by the ratio of true strain in width and thickness of a specimen during a tensile test in relation to the rolling direction according to an angle $\alpha$. Materials with high average normal anisotropy value, $\bar{r}$, and low planar anisotropy value, $\Delta R$, have a low reduction in their thickness and low strain variation in their plane during the deep drawing process. Features that are advantageous for deep drawing applications [4,5].

In this paper, the influence of the recrystallized grain size of the 430Nb stainless steel with 0.6 mm thickness in the evolution of crystallographic texture, anisotropy and, consequently, their formability was evaluated.

2 Material and methods

The samples, labeled BF (identification for cold coil sample in portuguese), used in this study were taken out from a cold coil of a Niobium-stabilized ferritic stainless steel AISI 430 (430Nb) annealed in an industrial continuous furnace with a heating rate of 24 °C.s$^{-1}$ after a cold rolling process with 85% reduction in thickness. The chemical composition of the samples studied are shown in Table 1.

2.1 Heat treatment

The samples were annealed in an industrial furnace according to Table 2. After that, the samples identified as BF-02, BF-03 and BF-04 were submitted to an additional annealing process on a stationary furnace. This second annealing process had the objective to increase the sample recrystallized grain size.
2.2 Microstructure analysis

Samples were prepared through standard metallographic procedures (cutting, mounting, grinding and mechanical polishing) and analyzed through their longitudinal sections after etching with Villella reagent for 50s. The average grain size was measured using the Abrams Concentric Circles method according to ASTM E111-13 [6], considering the planar diameter. Six measurements fields along the thickness were made in each sample to measure the grain size and calculate the standard deviation.

2.3 Texture analysis

Crystallographic orientation intensity in the surface and center of the samples were analyzed using the X-ray diffraction method with the aid of X’Pert Philips® equipment and the software OIM Analysis®, which provided the orientation distribution functions ODFs (Euller angles according to Bunge’s convention) for $\beta_0 = 45^\circ$.

2.4 Mechanical properties

Tensile tests were carried out, at room temperature, using a traction test machine INSTRON 5583 according to ASTM E8/E8M standard [7]. Sample measurements were made in real time during the tensile test using a displacement point detector. A restricted uniform plastic deformation of 15% was used to determinate the ‘r’, ‘r45’ and ‘r90’. Finally, the average normal anisotropy value, $\bar{\Delta}R$, was computed using Equation 1.

$$ \frac{r_0 + 2r_{45} + r_{90}}{4} $$

(1)

where $r_0$, $r_{45}$ and $r_{90}$ correspond to $r(\alpha)$ in the $0^\circ$, $45^\circ$, $90^\circ$ directions in relation to the rolling direction of the sheet metal, respectively. The planar anisotropy value, $\Delta R$, was evaluated according to Equation 2 [8].

$$ \frac{r_0 - 2r_{45} + r_{90}}{2} $$

(2)

3 Results and discussion

Figure 1a shows the microstructure of a 430Nb 0.60 mm-thick coil annealed in an industrial continuous furnace. The samples microstructure with an additional heat treatment on a stationary furnace are shown in Figures 1b, 1c, and 1d. The additional heat treatment process promoted an increase in the average grain size through the migration of the previous grain boundaries in the recrystallized structure [9].

BF-01 and BF-02 samples presented a homogeneous recrystallized ferritic structure along their thickness, precipitates distributed throughout the matrix and 10μm and 12μm grain size, respectively. BF-03 and BF-04 samples presented heterogeneity on structure along their thickness, precipitates distributed throughout the ferritic matrix with 19μm and 28μm grain size, respectively.

The sheet formability can be evaluated through the average normal anisotropy value, $\bar{\Delta}R$, and through the low planar anisotropy value, $\Delta R$ [5,8]. The average normal anisotropy describes the sheet’s ability to resist thickness thinning when subjected to tensile or compressive stresses. While, planar anisotropy describes the strain variation in the sheet plane during the deformation.

During the stamping process, the sheet strain occurs in all directions (0° to 360°) on the sheet plan. Therefore, rupture occurs in the direction with less thinning resistance. Thus, the analysis of the formability considers r for all directions [1]. Materials with high values of average normal anisotropy and low values of planar anisotropy display better performance in deep drawing process [4].

Table 3 shows that an increase in the grain size from 10μm to 19μm promoted an increase of $\bar{\Delta}R$ in all directions ($r_0$, $r_{45}$ and $r_{90}$). However, the increase in the average grain size, from 19 μm 28 μm, promoted a reduction in $\Delta R$ in all directions. Regarding planar anisotropy, the increase in the average grain size, from 10μm to 28μm, promoted a reduction in the $\Delta R$ coefficient.

The increase in the $\Delta R$, i.e, the ability to withstand thinning, is strongly related to the increase in the Gamma fiber components intensity and the reduction of Theta fiber components intensity [4,8,10,11]. The increase in the Theta fiber’s length reduces the material formability due to the low $r$ of the crystallographic orientations present in this fiber, especially in $r_{45}$ [5].

Figure 2 shows the orientation distribution functions (ODFs) on Bunge convention used to plot the ODFs in figure 3. The coefficient “s”, shown in figure 3, indicates the measurements positions of the texture along the thickness. Figure 3 shows the crystallographic texture change along the grain growth in the samples. Where, s=0 indicates the

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Table 1. Chemical composition of the 430Nb samples

<table>
<thead>
<tr>
<th>Element</th>
<th>C</th>
<th>Mn</th>
<th>Si</th>
<th>Cr</th>
<th>N</th>
<th>Ni</th>
<th>Nb</th>
</tr>
</thead>
<tbody>
<tr>
<td>BF-01</td>
<td>0.018</td>
<td>0.202</td>
<td>0.36</td>
<td>16.2</td>
<td>0.032</td>
<td>0.31</td>
<td>0.24</td>
</tr>
<tr>
<td>BF-02</td>
<td>0.018</td>
<td>0.202</td>
<td>0.36</td>
<td>16.2</td>
<td>0.032</td>
<td>0.31</td>
<td>0.24</td>
</tr>
<tr>
<td>BF-03</td>
<td>0.018</td>
<td>0.202</td>
<td>0.36</td>
<td>16.2</td>
<td>0.032</td>
<td>0.31</td>
<td>0.24</td>
</tr>
<tr>
<td>BF-04</td>
<td>0.018</td>
<td>0.202</td>
<td>0.36</td>
<td>16.2</td>
<td>0.032</td>
<td>0.31</td>
<td>0.24</td>
</tr>
</tbody>
</table>

Units in % mass.
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In general, all samples (Figure 3) exhibited a recrystallization texture in the \((334)[4\bar{8}3]\) component, which is typical of a cold rolled ferritic stainless steel subjected to direct rolling process \([5,8]\). This texture can be explained by the preferential growth of the component \((334)[4\bar{8}3]\) around the rolled component \((112)[1\bar{1}0]\) of the \(\alpha\) fiber, due to the oriented growth mechanism caused by the selective drag of segregated particles in the grain boundaries \([8,10]\). The other higher intensity was found at \((111)[\bar{1}2\bar{1}]\). Both components, \((334)[4\bar{8}3]\) and \((111)[\bar{1}2\bar{1}]\) belong to the gamma fiber.

Table 4 and Table 5 show the evolution of the principal volumetric fraction texture of the samples for each grain size on surface \((s = 0)\) and center \((s = 0.5)\). Figure 3 and Figure 4 show the texture evolution throughout the ODFs.

It can be observed on the surface of the thickness (Table 4) that an increase in the average grain size, from 10\(\mu\)m to 19\(\mu\)m, promoted a reduce on volume fraction of Theta fiber \((100)[uvw]\) and an increase on volume fraction of fiber \(\gamma\) components \((111)[1\bar{2}\bar{1}]\) and \((111)[\bar{1}\bar{2}\bar{1}]\). Simultaneously, at the center of the thickness (Table 5), \(\gamma\) fiber remained stable and the Theta fiber fraction \((100)[uvw]\) was reduced with increased average grain size from 10\(\mu\)m to 28\(\mu\)m.

That the obtained values of \(\bar{r}\) (Table 3) showed good correlation with the surface Gamma/Theta ratio values (Table 4). By increasing the grain size from 10\(\mu\)m to 19\(\mu\)m, \(\bar{r}\) was increased from 1.44 to 1.98 and the Gamma/Theta ratio on surface was increased from 4.36 to 4.83. By further increasing

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**Table 3. Mechanical properties of samples**

<table>
<thead>
<tr>
<th>Samples</th>
<th>(r^{0})</th>
<th>(r^{45})</th>
<th>(r^{90})</th>
<th>(\bar{r})</th>
<th>(\Delta R)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BF01-10(\mu)m</td>
<td>1.59</td>
<td>1.10</td>
<td>1.96</td>
<td>1.44</td>
<td>0.68</td>
</tr>
<tr>
<td>BF02-12(\mu)m</td>
<td>1.64</td>
<td>1.33</td>
<td>2.31</td>
<td>1.65</td>
<td>0.65</td>
</tr>
<tr>
<td>BF03-19(\mu)m</td>
<td>1.80</td>
<td>1.70</td>
<td>2.73</td>
<td>1.98</td>
<td>0.57</td>
</tr>
<tr>
<td>BF04-28(\mu)m</td>
<td>1.42</td>
<td>1.66</td>
<td>2.23</td>
<td>1.74</td>
<td>0.16</td>
</tr>
</tbody>
</table>

\(\Delta R\) = Planar anisotropy; [%]; \(\bar{r}\) = Average normal anisotropy.

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Figure 1. Microstructure in 200 \(\mu\)m (red bar) and average grain size of 430Nb samples after the heat treatment. (a) No additional heat treatment; (b) with additional heat treatment of 915\(°\)C; (c) with additional heat treatment of 965\(°\)C; and (d) with additional heat treatment of 1015\(°\)C \((s = 0\) for surface layer and \(s = 0.5\) for center layer). SD It is an abbreviation for standard deviations.
Figure 2. Section for $\delta = 45^\circ$ showing the position of main orientations along with the RD, TD: transverse direction and ND. Euler angles on notations proposed by Bunge [12].

Figure 3. Recrystallization textures of the samples, where (a) $s = 0$ (BFXX-S) represents the surface layer and (b) $s = 0.5$ (BFXX-C) represents the center layer.
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Table 4. Volumetric fractions [%] and Gamma/Theta ratios of surface samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>Rotated Cube (001)[110]</th>
<th>Theta (100)[uvw]</th>
<th>Gamma (111)//ND</th>
<th>Gamma/Theta</th>
</tr>
</thead>
<tbody>
<tr>
<td>BF01-S-10μm</td>
<td>2.82</td>
<td>9.29</td>
<td>40.28</td>
<td>4.36</td>
</tr>
<tr>
<td>BF02-S-12μm</td>
<td>2.60</td>
<td>9.09</td>
<td>41.42</td>
<td>4.56</td>
</tr>
<tr>
<td>BF03-S-19μm</td>
<td>1.66</td>
<td>7.43</td>
<td>43.31</td>
<td>5.83</td>
</tr>
<tr>
<td>BF04-S-28μm</td>
<td>1.38</td>
<td>7.63</td>
<td>41.66</td>
<td>5.46</td>
</tr>
</tbody>
</table>

ND = Normal Direction.

Table 5. Volumetric fractions [%] and Gamma/Theta ratios of center samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>Rotated Cube (001)[110]</th>
<th>Theta (100)[uvw]</th>
<th>Gamma (111)//ND</th>
<th>Gamma/Theta</th>
</tr>
</thead>
<tbody>
<tr>
<td>BF01-C-10μm</td>
<td>2.83</td>
<td>9.56</td>
<td>40.11</td>
<td>4.20</td>
</tr>
<tr>
<td>BF02-C-12μm</td>
<td>2.90</td>
<td>10.09</td>
<td>39.40</td>
<td>4.90</td>
</tr>
<tr>
<td>BF03-C-19μm</td>
<td>2.10</td>
<td>8.76</td>
<td>40.42</td>
<td>4.62</td>
</tr>
<tr>
<td>BF04-C-28μm</td>
<td>1.40</td>
<td>8.27</td>
<td>39.68</td>
<td>4.81</td>
</tr>
</tbody>
</table>

ND = Normal Direction.

Figure 4. Texture change shown by subtraction of ODFs samples in function of grain growth. where (a) $s = 0$ represents the surface layer and (b) $s = 0.5$ represents the center layer.

the grain size from 19μm to 28μm, $r$ was reduced to 1.74 and the Gamma/Theta ratio on surface was reduced to 5.46. Regarding planar anisotropy, the $\Delta R$ was reduced from 0.68 to 0.16 due to the grain growth from 10μm to 28μm. This effect occurred due to the reduction of Rotated Cube (001)[110] and Theta fiber volumetric fractions improving the steel formability [5]. Figure 4 shows the texture change calculated by subtraction of ODFs samples, where each figure shows the texture differential between the grain sizes. Positive values indicate an increase in components intensity, while the negative values indicate a reduction on components intensity.

Figure 4a and 4b shows that, grain growth from 10μm to 19μm, the $\gamma$ fiber was intensified on surface and center
of thickness. On grain growth from 19 μm to 28 μm γ fiber intensity was decreased on surface and center of thickness. The grain growth from 10 μm to 19 μm was able to promote a formability accretion by γ fiber increase and, consequently, $r$ increase. Therefore, the formability reduction that occurred in grain growth from 19 μm to 28 μm was caused by selective drag effect by the segregated particles present in the grain boundaries [8,10]. This mechanism intensified the (334)[483] component around the (112)[110] rolled component of the α fiber and reduced the γ-fiber intensity.

4 Conclusions

Four 430Nb stainless steels samples with a thickness of 0.6 mm and from the same coil were submitted to an additional annealing process on a stationary furnace using different temperatures. The influence of the grain size on their formability was observed and the effect of the main texture components on the resulting $r$ values was analyzed. It was concluded that the formability can be improved through grain growth. Increasing the grain size from 10 μm to 19 μm, the $r$ was increased from 1.44 to 1.98. However, from 19 μm average grain size, the drag effect spins the γ fiber components to the (334)[483] reducing the $r$ from 1.98 to 1.74 and, consequently reducing the material formability.

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References


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