Nitriding Process Characterization of Cold Worked AISI 304 and 316 Austenitic Stainless Steels

Waldemar Alfredo Monteiro, Silvio Andre Lima Pereira, and Jan Vatavuk

1 Nuclear and Energy Research Institute, IPEN, Avenida Lineu Prestes 2242, 05508-000 São Paulo, SP, Brazil
2 MAHLE Metal Leve S/A, Rod. Anhanguera, s/n, Tijuca Preto, 13205-700 Jundiaí, SP, Brazil
3 School of Engineering, Presbyterian Mackenzie University, Rua da Consolação 930, 01302-901 São Paulo, SP, Brazil

Correspondence should be addressed to Waldemar Alfredo Monteiro; wamontei@ipen.br

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1. Introduction

The austenitic stainless steels are plenty utilized in chemical processes equipment in pharmaceutical, foodstuff, textile, petroleum, and cellulose industries, where these components are exposed at aggressive ambient conditions and low temperatures. These steels are also employed in orthopedic implant due to its biocompatibility [1].

The austenitic stainless steels, despite the elevated corrosion resistance, have low hardness that can only be partially improved by cold deformation reducing its application in components submitted to severe wear conditions [2–5].

To surpass this problem there are alternatives in the surface engineering to improve surface properties, without hindering all the other properties like the corrosion resistance. One alternative is the thermochemical heat treatment, the well-known nitriding process that basically can run by one of the three methods: gaseous, salt bath, or plasma. The nitride process, independent of the method, increases the superficial hardness of austenitic stainless steels and promotes higher wear resistance.

The literature has shown the possibility of austenitic stainless steels nitride processing, which is the occurrence of nitrogen in solid solution, increasing the hardness of the surface and the wear resistance without loss of corrosion resistance. FCC phase expanded by nitrogen atoms is obtained. This identification is relating to the obtained X-rays diffraction reflections that are dislocated with low Bragg angles relating to the positions in a normal austenite phase [6–13].

Gontijo et al. [8] mention that many literatures [7–13] define this phase formed at low temperature as a nitrogen supersaturate phase, with a distorted crystalline structure (tetragonal structure). The plasma nitriding permits the introduction of nitrogen in steels with temperatures as low as 450°C, producing a harder structure and metastable phases resistant to the corrosion.

The austenitic stainless steels response to the conventional nitriding processes is surprising (500°C ≤ T ≤ 600°C);
Table 1: Chemical composition (weight%) of the studied steels (NBR 5601:1981).

<table>
<thead>
<tr>
<th>AISI</th>
<th>% C</th>
<th>% Si</th>
<th>% Mn</th>
<th>% P</th>
<th>% S</th>
<th>% Ni</th>
<th>% Cr</th>
<th>% Mo</th>
</tr>
</thead>
<tbody>
<tr>
<td>304</td>
<td>0.08</td>
<td>1.00</td>
<td>2.00</td>
<td>0.045</td>
<td>0.03</td>
<td>8.0/10.0</td>
<td>18.0/20.0</td>
<td>—</td>
</tr>
<tr>
<td>316</td>
<td>0.08</td>
<td>1.00</td>
<td>2.00</td>
<td>0.045</td>
<td>0.03</td>
<td>10.0/14.0</td>
<td>16.0/18.0</td>
<td>2.0/3.0</td>
</tr>
</tbody>
</table>

Figure 1: Nondeformed and deformed samples: (a) AISI 304 and (b) AISI 316.

Figure 2: Schema of distribution of each divided sample.

2. Materials and Methods

The austenitic stainless steels have substantial work hardening after cold work processing. In the present study cold worked and heat treated steels sheets were utilized (AISI 304 with 1.0 mm thickness and AISI 316 with 1.5 mm thickness). The chemical composition is presented in Table 1.

2.1. Samples Deformation Process. Tensile test samples were prepared to induce cold plastic deformation by tensile test using a universal machine tensile test MTS810, by the ASTM E8MM-98 test procedure. The employed deformation degree in heat treated steels was 0%, 10%, 20%, 30%, and 40% (Figure 1).

2.2. Nitriding Processes. The three possible nitriding processes were explored. Each tested sample was divided in four identical samples (Figure 2):

(i) Samples 01: for characterization of initial condition.
(ii) Samples 02: gaseous nitriding at 570°C during 3.5 h.
(iii) Samples 03: plasma nitriding at 430°C during 10 h.
(iv) Samples 04: liquid nitriding at 570°C during 2 h.

2.3. Metallographic Preparation for Optical Microscopy. Metallographic samples were obtained by abrasive cutting. Samples (squares of 5 mm) with and without the nitriding process were transversally sectioned with adequate "cut-off." The metallographic specimen mounting is done in a hot thermosetting powder mounting material (bakelite) and after this it was utilized the traditional steels metallographic
The final polished samples were etched with Marble’s reagent for Leica optical microscope observation with image program A4iDocucapture, and the nitriding layer of the steels samples was measured and also documented.

For SEM observations, the samples were completely repolished, cleaned, and covered with a fine conductive Al layer. Through a WDS MICROSPAC analyzer in a SEM PHILIPS Quanta 600 the N and C elements were quantified.

The composition curves were determined for N and C elements from surface for the core direction with points spaced out to 5 𝜇m. The nitrogen element was quantified automatically by the SEM, but the automatic method was not well succeeded for carbon.

The carbon calibration method was very effective with the SEM parameters (Table 2) and through five carbon steel patterns (0.060% C; 0.163% C; 0.354% C; 0.838% C; 0.926% C) the X-ray peak curve of the carbon element was obtained (Figure 3).

Utilizing the expression $y = 41.40x + 26.02$ (see Figure 3) it is possible to quantify the carbon element.

### Table 2: SEM parameters for the N and C analysis.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Work distance</td>
<td>12.25 mm</td>
</tr>
<tr>
<td>“Spot size” of the beam</td>
<td>7.5</td>
</tr>
<tr>
<td>kV</td>
<td>12</td>
</tr>
<tr>
<td>Current density</td>
<td>50 nA</td>
</tr>
<tr>
<td>Exposition time</td>
<td>20 s</td>
</tr>
</tbody>
</table>

### Figure 3: Calibration curve of carbon element.

**3. Results and Discussion**

#### 3.1. Mechanical Behavior

The tensile test curves of the samples are shown in Figure 4. Due to the low elastic (deformation) behavior of the samples, their results are not considered in this study.

Austenitic stainless steels are susceptible to induced martensite by cold plastic deformation that are strongly related to the alloy chemical composition, mainly the carbon content, due to its effect of the martensite transformation temperature lowering, reducing the possibility of the martensite induced by plastic cold deformation.

#### 3.2. Optical Microscopy Characterization

Two sublayers inside the nitride area are observed in all samples (Figures 5 and 6).

There is a solid tendency to the occurrence of martensite during the plastic cold deformation, mainly in low carbon steels as 304L [18]. In conditions where the steels were acquired, an enough defect concentration to evaluate this influence in the kinetics study of the nitrogen diffusion during the nitriding processes is expected.

Moreover, the thickness variations were random (Table 3) and do not suggest any tendency relating to the influence in thickness due to employed cold plastic deformation degree.

Concerning Figure 6, an example of optical micrographs referring to AISI 316 and AISI 304 nitriding process is shown. In all studied conditions the thicknesses of nitride layer was measured.

#### 3.3. Chemical Analysis by WDS (Scanning Electron Microscopy)

The nitrogen and carbon concentration, illustrated in Figure 7, obtained by wave dispersive X-rays spectrometry
(WDS) using SEM can be observed, where the first layer has a nitrogen concentration of 5.5% and the second layer shows a 1% carbon concentration peak. Both steels can have up to 0.08 weight% of carbon and, with the exception of the liquid nitriding process, the gaseous and plasma processes are free of carbon. Then it can be concluded that in this region a decarbonization with nitrogen diffusion possibly occurs and these carbon atoms are diffusing to regions just below the real nitride layer, forming a second layer with a superior carbon concentration. Consequently two layers originate, the first a matrix of austenite expanded by nitrogen atoms, followed by another layer formed by matrix of austenite expanded by carbon atoms.

3.4. Microhardness Analysis. Figure 8 shows the hardness value of the samples with different degree of cold work before nitriding processes. There is a considerable hardness increase, more with the AISI 304 steel, possibly due to the strong tendency for the occurrence of martensite induced by plastic cold work.
Table 3: Average thickness of the nitride layer (µm) relating to Figure 5.

<table>
<thead>
<tr>
<th>AISI</th>
<th>Deformations</th>
<th>Gaseous A layer</th>
<th>Liquid A layer</th>
<th>Gaseous B layer</th>
<th>Liquid B layer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0%</td>
<td>71</td>
<td>33</td>
<td>34</td>
<td>10</td>
</tr>
<tr>
<td>304</td>
<td>10%</td>
<td>67</td>
<td>29</td>
<td>33</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>20%</td>
<td>65</td>
<td>34</td>
<td>33</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>30%</td>
<td>66</td>
<td>26</td>
<td>32</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td>40%</td>
<td>71</td>
<td>26</td>
<td>33</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>0%</td>
<td>51</td>
<td>29</td>
<td>28</td>
<td>12</td>
</tr>
<tr>
<td>316</td>
<td>10%</td>
<td>48</td>
<td>20</td>
<td>23</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>20%</td>
<td>50</td>
<td>27</td>
<td>22</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>30%</td>
<td>47</td>
<td>28</td>
<td>24</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>40%</td>
<td>51</td>
<td>20</td>
<td>26</td>
<td>14</td>
</tr>
</tbody>
</table>

Figure 7: Nitrogen and carbon distribution (weight%) regarding the layer depth in (a) AISI 304 steel and (b) AISI 316 steel.

Figure 8: Hardness values of the cold work samples of AISI 304 and 306 steels.
Figure 9: Hardness curves in samples after diverse nitriding processes.

Figure 9 shows the hardness curves of the samples after nitriding process; it can be observed, with the data, that the applied precold work shows a little influence in all the utilized diffusion processes.

Concerning that both AISI 304 and 316 were nitrated in same conditions, more high hardness values in AISI 304 samples depth are observed. Besides, higher values were obtained in samples with plasma nitriding processes,
probably due to the applied process temperature (plasma, \( T = 430^\circ C \); gaseous and liquid, \( T = 570^\circ C \)).

4. Conclusions

The initial cold work (hardening) in both AISI 304 and 306 austenitic stainless steels does not show special influence in all applied nitriding kinetics. In both studied materials two distinguished layers are observed in the microstructure, one formed by the austenite expanded due to the presence of nitrogen atoms followed by another layer formed by austenite expanded due to the presence of carbon atoms. The gaseous and liquid nitriding processes showed hardness values in the layers of 1100 HV and the plasma nitriding process 1200 HV. Better diffusion kinetics occurrence in AISI 304 steel is also observed.

Competing Interests

The authors declare that they have no competing interests.

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