Corrosion Resistant Cr\textsubscript{N\textsubscript{x}} Nanolayers Obtained by Low Temperature Ion Nitriding of Hard Chromium Coated AISI 1045 Steel

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Chromium nitrides Cr\textsubscript{x}N have attracted research interest due to their excellent wear and corrosion properties when are deposited on steels. Nevertheless, due to the considerable difference between Cr\textsubscript{x}N and steel expansion coefficients, microcracks and delamination are still persistent problems. In this regard, this research addressed the generation of Cr\textsubscript{N\textsubscript{x}} nanolayers, through an ion nitriding process, carried out on hard chromium coated AISI 1045 steel. The effect of nitriding temperature (500°C-550°C) and nitrogen content in plasma (25, 50, and 75%) on corrosion performance and layer characteristics were studied. Grazing Incidence X-ray Diffraction analysis revealed that modified surfaces are composed of nanolayers, constituted by a mixture of CrN, Cr\textsubscript{2}N\textsubscript{3}, Cr, and FeN. Rietveld quantification shows that the fraction of chromium nitrides in the region analyzed increased with increasing both temperature and nitrogen content in plasma, resulting CrN as the predominant phase for all evaluated conditions. The electrochemical behavior of the modified nanolayers was studied by potentiodynamic polarization technique, revealing an enhancement in corrosion performance of chromium coated 1045 steel by the nitriding treatment, showing a corrosion current density 10 times lower than the untreated sample and more positive corrosion potentials for the nitried samples concerning chromium coated 1045 steel.

Keywords: Ion nitriding, AISI 1045, Chromium nitrides, Hard chromium.

1. Introduction

Specific properties such as high corrosion resistance and excellent wear performance have been required through time in metallic components used in food, metalworking, and electronics industry. In the last decades the application of hard chromium coatings on the steel or alloys components has been a widely alternative to improve corrosion and wear resistance of components as engine parts or cutting tools\textsuperscript{1}. Although the development and extensive use of hard chromium coating processes, these coatings are still accompanied by important differences in thermal expansion coefficients, which generate microcracks or even delamination of the coating. For solving this problem, researchers have deposited a hard chromium coating on the surface of the steel since it has an intermediate thermal expansion coefficient between Cr\textsubscript{N} and the steel\textsuperscript{13,14}. Nevertheless, due to their extensive information related to the structure and phases of Cr\textsubscript{x}N (including effects on the corrosion resistance) applied by cathodic vapor deposition\textsuperscript{15,16}, there is still scarce current information involving the same phases synthesized by the ion nitriding process, a technique that offers high control of parameters, low temperature and relative short processing time\textsuperscript{11}.

Wierzchon et al.\textsuperscript{17} investigated the formation of chromium nitrides, oxynitrides and oxide by ion nitriding, in an Armco iron and AISI 1045 steel, which were electrochemically coated with hard chromium, reporting that the corrosion
resistance of these layers in the coating of hard chromium is elevated and can even be improved by modifying the process variables such as temperature and partial pressure of nitrogen. On the other hand, Menthe and Rietz reported a process of plasma nitriding on a steel with hard chromium coating at 560°C and 700°C, to different processing times (5 to 20 h), finding the formation of a chromium nitrides mixture (CrN, CrN and Cr) in the nitried region, showing a significant improvement in corrosion resistance. Sarraf et al. also reported the ion nitriding technique to repair microcracks in a hard chrome surface, revealing the formation of different CrN and CrN phases after the nitriding process. The results showed an improvement in corrosion resistance, in 3.5 wt% NaCl solution. Likewise, Aghajani et al. investigated the effect of plasma nitriding on H11 steel coated with hard chromium at ion nitriding temperature of 550°C for different process times, obtaining a mixture of CrN, CrN, Cr, FeN, and FeN phases in the modified zone. As well as Taktak et al. who studied the ion nitriding process in AISI 52100 and 8620 steels coated with hard chromium, using different gas mixtures, revealing an improvement in the wear properties due to the formation of CrN and CrN.

Therefore, this research is based on the formation of nanolayers composed mainly of corrosion-resistant CrN; it is also intended to show that the corrosion resistance of CrN can be efficient at a short time and low temperature at the nitriding process. These treatment conditions could result in a more efficient and profitable nitriding process with a high possibility for industrial applications (e.g. fuel cells bipolar plates or diluting magnetic semiconductors). On the other hand, the effect of nitrogen content in plasma on the formation of different types of CrN is studied, as well as its influence on corrosion resistance of coated steel.

2. Experimental Procedure

The CrN-type ceramic coatings were developed on commercial 10 x 10 x 5 mm³ coupons of AISI (American Iron and Steel Institute)-1045 steel, which were previously coated with a layer of electrolytic hard chromium with ~1.5 µm in thickness. The microstructure of base 1045 steel consists mainly of ferrite and perlite and its weight percentage chemical composition, determined by optical emission spectroscopy (PMI MASTER PRO, Oxford Instruments), was: C 0.49, Si 0.18, Mn 0.62, Cr 0.06, Mo 0.03, Ni 0.04, P 0.014, S 0.04, balance Fe.

The surface modification process was carried out in a 42 L laboratory chamber during 3 h, using a DC glow-discharge in a gas mixture of N₂/H₂ at 266 Pa, at potential of -580 V, at constant current density of 1.5 mA/cm². The temperature control was carried out through a closed circuit using a PID controller, coupled with a type K thermocouple in direct contact with sample. Mass flow controllers were used to modifying the N₂/H₂ rate in the plasma. After treatment, samples were cooled to room temperature inside the vacuum chamber with an argon inert atmosphere. Conditions of nitriding for different samples and their identification are shown in Table 1.

<table>
<thead>
<tr>
<th>Samples ID</th>
<th>Temperature (°C)</th>
<th>Nᵢ in plasma (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>500</td>
<td>25</td>
</tr>
<tr>
<td>S2</td>
<td>550</td>
<td>50</td>
</tr>
<tr>
<td>S3</td>
<td>500</td>
<td>75</td>
</tr>
<tr>
<td>S4</td>
<td>550</td>
<td>25</td>
</tr>
<tr>
<td>S5</td>
<td>500</td>
<td>50</td>
</tr>
<tr>
<td>S6</td>
<td>550</td>
<td>75</td>
</tr>
</tbody>
</table>

Surface and cross-section morphologies of the nitrided samples were analyzed by Scanning Electron Microscopy (SEM) (TESCAN MIRA 3). Nitrogen semi-quantitative profiles were determined by X-ray energy dispersion spectrometry (EDS) coupled to SEM. As corrosion resistance is related to the crystalline phases present on the surface, phases on nitrided samples were identified using Grazing Incidence X-ray Diffraction (GIXRD) with incidence angles set at 0.5 and 5°. This analysis was carried out on an Empyrean Panalytical X-ray diffractometer in a 2 theta range from 35 to 55°, with a scan speed of 175°/step with increments of 0.018°, using Cu kα radiation at 45 kV and 40 mA. In order to quantify the present phases in the modified surfaces, Rietveld refinement of XRD patterns (0.5 incidence angle) was performed for all nitrided samples.

The penetration depth of the X-ray during XRD analysis, was estimated by Equation 1, within the region of the most intense peaks in the patterns, assuming a location at 2θ angles of 43.65° for the main reflection of CrN and 44.58° for the main reflection of Cr₂ α,

$$d = \left[ \frac{1}{\mu \sin \alpha} + \frac{1}{\sin(2\theta - \alpha)} \right]^{-1}$$

where α represents the incidence angle and μ is the linear absorption coefficient. To calculate μ, the mass absorption coefficient (μ/ρ), which depends only of the composition, should be multiplied by the material density, ρ.

Mass absorption coefficients were consulted from the literature and the depth analysis was considered as the depth into the material measured along the surface normal where the intensity of X-ray falls to 1/e of its value at the surface. Likewise, the density of nitrided layers was estimated, considering the mixture of phases (CrN, CrN, FeN, and Cr) determined by Rietveld analysis for each different nitriding treatments.

The corrosion resistance was evaluated by the potentiodynamic method using a potentiostatic galvanic scatter Gamry 1000A interface equipment. An experimental arrangement of three electrodes was used with platinum as counter electrode, saturated calomel (SEC) as reference electrode, and nitrided sample as the working electrode. Nitrided samples were immersed in 0.5 M H₂SO₄ solution at room temperature, and the scanning potential was set in a range of -500 to 1200 mV, setting scanning rate at 1 mV/s and stabilizing time of 600 s.
3. Result and Discussion

3.1. Surface and cross morphology

SEM surface morphology of Cr-α coated AISI 1045 is shown in Figure 1. A smooth surface, with an extensive microcracks network throughout whole the surface of the sample can be observed. This morphology is characteristic of this kind of coatings, due to tensile stress generated during electroplating\(^\text{24}\).

Surface morphologies of nitrided samples are shown in Figure 2. Ion nitriding treatment at 500°C with different nitrogen contents in plasma (Figures 2 a, b, and c), does not considerably modify the surface morphologies, but it can be appreciated small precipitates on the surfaces and no presence of microcracks. The sealing of the microcracks is due to the unavoidable increase in volume that develops when the Cr\(_X\)N is formed, which provides a net volume increase of approximately 45%, causing an expansion of the chromium coating near the surface\(^\text{24}\).

As the ion nitriding temperature increases from 500 to 550°C, a higher density of precipitates is observed for all nitrogen contents (Figures 2 d, e, and f). The sizes of these irregular shape precipitates are in the range of 1µm; and it can be observed how the largest particles are formed by agglomeration mechanisms. In agreement with Shayan et al.\(^\text{20}\), who formed Cr\(_X\)N and Fe\(_X\)N on tools steel by traditional ion...
nitriding and active screen plasma nitriding, the development of this morphology can be attributed to the effect of cathode sputtering that commonly occurs in the ion nitriding processes.

Figure 3 presents a SEM cross-section micrograph of the sample nitrided at 550°C with 75% of nitrogen in plasma, the hard chromium coating and the nitride nanolayer can be observed. Nitrogen content is also shown for all nitrided samples until a depth of 900 nm, determined by point analysis using EDS. The depths analyzed in the nitrided layers were 0, 500 and 900 nm and an average of 5 microanalysis was recorded for each depth. EDS point analysis reveals a nitrogen profile in nitrided layers, showing a decrease in nitrogen content as depth increases for all treatments.

The samples treated at 550°C present a higher nitrogen content for the different depths analyzed in the nitrided layers, compared to the samples treated at 500°C, which indicates a greater nitrogen absorption at higher temperatures, suggesting an increase in the depth of chromium nitride. This behavior had also been observed by Ajikumar et al.\textsuperscript{25}, detecting a greater diffusion of nitrogen for temperatures above 700°C.

The Cr-N binary system provides information on the possible chromium nitride phases that can be obtained by manipulating the temperature and the nitrogen content. In this sense, it can be mentioned that the nitrided layers at 500°C can be shaped by a mixture of Cr and Cr\textsubscript{2}N phases in the outermost part of the layer, followed by an expanded structure of chromium with nitrogen. Meanwhile, for the samples treated to 550°C, the layers could be composed of a mixture of Cr\textsubscript{2}N and CrN phases up to a depth of 500 nm, according to the nitrogen content obtained in the point EDS analysis for the different treatments.

The formation of Cr\textsubscript{2}N on the hard chromium layer in ion and gaseous nitriding processes is controlled by the development of its own diffusion phenomena that depends on chemical kinetics. Buijnsters et al.\textsuperscript{26} reported that the formation of chromium nitride can be explained by the interstitial diffusion of atomic nitrogen in the Cr-α matrix until it reaches saturation of 33% of atomic nitrogen, then a layer of the hexagonal CrN phase begins growing parallelly to the direction of diffusion of nitrogen. As the concentration of nitrogen is higher in the outermost part of the layer due to constant nitrogen bombardment, the cubic CrN phase will begin growing on the surface of the hexagonal phase until a CrN monolayer is obtained.

### 3.2. Phases structure

Grazing incidence XRD analysis were carried in order to determine crystalline phases at different depths. Table 2 shows X-ray penetration depth (region from which...
the information is obtained), as well as the values of density and mass absorption coefficient used to determine them. Main reflections for Cr-α and CrN, located at 44.58° and 43.68° 20 angle respectively, were considered for penetration depth compute. In a general way, it can be appreciated that XRD information is obtained from a depth close to 50 nm for 0.5° incidence angle and about from 500 nm in depth for 5° incidence angle.

Figure 4 shows the XRD patterns obtained with an incidence angle of 0.5° for all treatments (information from approximately 50 nm in depth). It can be seen that after applying the ion nitriding treatment at 500°C and 550°C with different nitrogen contents in plasma, a mixture of phases is formed on the modified surface, which is mainly constituted by chromium nitrides (CrN and Cr₂N), in addition to the phases of Fe₂N and metallic Cr-α. These patterns are compared to JCPDS cards 04-003-5919, 04-014-1025, 00-064-0134, and 04-003-5597 as reference.

The treatment at 500°C with nitrogen content of 25% gives place to a mixture of CrN and Cr₂N phases with main reflection at 2θ positions of 37.5° and 42.7° respectively. The increasing of the plasma nitrogen concentration to 50% and 75% leaded the presence of a new peak at 43.6°, corresponding to the CrN phase. For a nitriding temperature of 550°C, an increase in the intensities of CrN and Cr₂N phases in the different orientations is observed, as well as a new peak at 40.2° related to CrN phase. The here reported phases are in accordance with the Cr-N phase diagram²⁷.

On the other hand, the reflection corresponding to metallic Cr-α is observed for all nitrided samples, which marks an incomplete transformation towards the different types of chromium nitrides²⁸. This effect could be related to the slow diffusion of atomic nitrogen in chromium nitrides. The nitrogen diffusion is affected by the high compression stresses developed during transformation of chromium nitrides since both phases generate a contribution of expansions, which are preferentially induced in Cr grains. Resulting in reduced kinetics for the transformations of Cr-α in chromium nitrides²⁵,²⁶.

The presence of Fe₂N reflections in 0.5° incidence angle GIXRD patterns for all nitriding samples could be explained by the sputtering phenomenon occurring on the fixtures of the nitriding system (cathode), which has been reported in thermochemical treatments assisted by plasma²⁰. The sputtering intensity is a function of nitriding temperature as well as the plasma composition (N₂/H₂ rate) so, variations in the intensity of Fe₂N reflections can be appreciated in 0.5° GIXRD patterns.

Figure 5 shows the GIXRD patterns obtained using an incidence angle of 5° (information from approximately 500 nm in depth). In this case, the reflections corresponding to CrN are absent for a nitriding temperature of 500°C regardless of nitrogen content in plasma. The phase Cr-α was identified for these conditions. At 550°C, CrN and Cr₂N are evidenced in positions 2θ of 37.3° and 43.7° for the CrN, and 40.2° and 42.6° for the Cr₂N, which can be an indicative that modified layers at this temperature have a thickness of at least 500 nm. Likewise, a slight shift of the Cr-α reflection towards lower 2θ angles can be observed for all nitriding conditions and both GIXRD evaluated conditions. This effect could be associated with the insertion of atomic nitrogen in interstitial position of the Cr-α crystalline structure, producing a slight expansion and an increase in lattice parameter²⁸,²⁹.

In Figure 5 it is also appreciated the disappearance of Fe₂N reflections, reinforcing the claim that this phase is only present on the top surface of the nitrided layers, which is attributed to cathodic sputtering phenomenon.

### 3.3. Phases quantification by Rietveld method

Table 3 shows the phases quantification, at 50 nm in depth, obtained by the Rietveld method for nitrided layers. Phase’s quantification reveals an increase in the total content of CrN as a result of the increase in the nitriding temperature. A greater nitriding potential (greater amount of N⁺ cations in the plasma) is expected at higher temperature as well as an increase in the nitrogen diffusion rate and consequently in the synthesized CrN volume. The real fraction of CrN in nitrided layers could be even higher than that presented in Table 3, based on the fact that presented quantification includes Fe₂N phase, which is located only on the most superficial part of the nitrided layer.
On the other hand, it is observed a greater fraction towards the CrN phase with the increase of the nitrogen flow and the nitriding temperature, these results are in good agreement with that reported by Taguchi and Kurihara.

They evidenced the formation of CrN and Cr2N at temperatures range of 700°C to 1100°C with a higher selectivity towards CrN, which increases at higher nitrogen concentration or processing time as follows:

\[2CrN + N_2 \rightarrow 2CrN\]

3.4. Electrochemical corrosion performance

Figure 6 shows the potentiodynamic polarization curves for nitrided samples at different conditions, as well as the curves for both; base material (1045-Cr) and nitrided 1045 steel.

For all evaluated samples, potentiodynamic polarization curves show a clear definition of the active-passive zone, which allows determining the corrosion rates for each treatment. The nitrided hard chromium layers present more positive values of corrosion potential than base metal and even than only nitrided 1045 steel, indicating a higher thermodynamic resistance to the initiation and evolution of corrosion phenomena. Likewise, a displacement to the left of the curves obtained for nitrided samples is also experienced in reference to the nitrided 1045 steel and the base material, which indicates a lower corrosion rate through a decrease of the corrosion current density. These results reveal that corrosion current densities for the nitrided layers are 10 times reduced regarding the base material.

Table 4 shows the values of corrosion current densities, which let to establish that the different nitrogen contents in plasma and the temperatures used during nitriding do not generate a significant change on them, since all systems have a composite nanolayer of CrN and Cr2N with a higher fraction of the CrN phase. These results are in good agreement with that reported by Taguchi and Kurihara, who studied the formation of a double structure with a phases mixture of CrN and Cr2N, which improved the corrosion resistance. They also found that a higher fraction of CrN improved the corrosion resistance. The protective efficacy (Pi) of the layers of CrN and Cr2N, determined using Equation 3, is also presented in Table 4,

\[Pi(\%) = \left(1 - \frac{i_{corr}}{i_{corri}}\right) \times 100\]

where \(i_{corri}\) and \(i_{corr}\) are the corrosion current densities of the base material and nitrided samples respectively. The protective efficiency of the nanolayers was in an approximated range from 78 to 90%. This result supports the claim that nanolayers composed of CrN and Cr2N represent an effective means of improving corrosion resistance. In general, transition metal nitrides are inert to chemical attack due to their relatively higher position in the electrochemical series. In agreement with Wierzchon et al., the corrosion resistance can be explained by the formation of a fine-grained homogeneous nanostructure of chromium nitrides (CrN and Cr2N), which acts as a barrier and prevents diffusion of the electrolyte to the substrate.

Figure 7 reveals the corroded surface morphologies (SEM) of base material, nitrided steel, and as reference, the samples nitrided at 500°C and 550°C with 75% of nitrogen. EDS analysis confirms that the layer hard chromium of base material was dissolved during the corrosion test.

![Figure 7](image-url)
Corrosion Resistant CrN\textsubscript{x} Nanolayers Obtained by Low Temperature Ion Nitriding of Hard Chromium Coated AISI 1045 Steel

(see Figure 7a), due to diffusion of the electrolyte through local defects, generating a galvanic couple between the hard chromium coating and the substrate. The corroded surface morphologies of the nitrided samples (Figures 7b and c), show a significant improvement because they do not present severe corrosion after the corrosion test, however small areas with localized corrosion can be observed. Pitting corrosion is also observed in the potentiodynamic polarization test in a potential range of 0.2 to 0.4 V. This performance could be associated with an incomplete sealing of the microcracks and the presence of new local defects in the ceramic nanolayers of Cr\textsubscript{x}N. It is known that Cr\textsubscript{x}N ceramic coatings present local defects such as microcracks and porosity, which act as channels to transport corrosive agents and generate localized corrosion\textsuperscript{32,33}. Nonetheless, the effect of these new defects is considerably less since the corrosion performance is significantly improved. The EDS analysis of the pitting confirms the diffusion of the electrolyte and the dissolution of the protective layer in the corroded area, leaving exposed the substrate. The nitrided steel 1045 presents generalized corrosion, according to Figure 7d) in which it is observed the presence of a large number of holes in the surface, indicative that the nitrided steel 1045 is susceptible to severe corrosion.

4. Conclusion

Nanolayers composed of a mixture of nitrides were developed through short time and relatively low temperatures (500 °C and 550 °C) plasma nitriding process on hard chromium coated AISI 1045 steel. The Rietveld refinement of the nitrided samples 0.5° GIXRD patterns confirms the fact that nitrided layers are composed of a mixture of CrN, Cr\textsubscript{2}N, Cr and Fe\textsubscript{4}N. It also shows that higher nitrogen flows and a higher nitriding temperature promote greater
selectivity towards the CrN phase. Increasing the temperature amplifies the potential for nitriding, developing nanolayers of chromium nitride of approximately 500 nm. The obtained Chromium nitride nanolayers showed better corrosion resistance compared to hard chromium coated 1045 steel and nitrided steel. Nitrided samples showed more positive values of corrosion potentials, as well as corrosion current densities 10 times lower than base material, with a protective efficiency value of up to 90% in an H\textsubscript{2}SO\textsubscript{4} electrolyte. The improvement in corrosion resistance could be attributed to the sealing of microcracks in the hard chromium layer and to the formation of a homogeneous nanolayer of fine-grained chromium nitrides within the area near the samples surface. These effects achieve without the need to use high temperatures and long periods nitriding times. These results confirm that the ceramic nanolayers of chromium nitrides developed at low temperatures and relatively short times by means of conventional ionic nitriding, could be potentialized for different technological applications, such as the development of bipolar plates or diluting magnetic semiconductors.

5. Acknowledgments

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6. References