Influence of Plasma Nitriding with a Nitrogen Rich Gas Composition on the Reciprocating Sliding Wear of a DIN 18MnCrSiMo6-4 Steel

Rafael Luciano Dalcin* a, Alexandre da Silva Rocha a, Victor Velho de Castro a, Victor Velho de Castro b, Leonardo Fonseca Oliveira a, Julio Cesar Klein das Neves c, Carlos Henrique da Silva c, Célia de Fraga Malfatti b

a Universidade Federal do Rio Grande do Sul, Laboratório de Transformação Mecânica, Porto Alegre, RS, Brasil
b Universidade Federal do Rio Grande do Sul, Laboratório de Pesquisa em Corrosão, Porto Alegre, RS, Brasil
c Universidade Tecnológica Federal do Paraná, Laboratório de Superfícies e Contato, Curitiba, PR, Brasil

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In this study, the sliding wear of a DIN 18MnCrSiMo6-4 continuous cooling bainitic steel plasma nitrided with a nitrogen rich gas composition was investigated. To evaluate the influence of processing time and temperature on mechanical and microstructural characteristics of nitrided layer, the samples were nitrided at 400, 450, 500 and 550 °C for 3, 6 and 9 h. The produced nitrided layers were characterized concerning the microstructure, phase composition, microhardness and surface roughness. The samples were tested by ball-on-flat reciprocating dry sliding for friction coefficient and wear analysis. The tests were stopped after a given damage criteria involving the rapid growth of the friction coefficients and wear. The correlation of the different treatment parameters and resulting case depths and surface hardness with sliding distance at the time of microcracks formation or delamination of the surface layer was evaluated statistically by the analysis of variance (ANOVA). The plasma nitrided samples at 550 °C showed better wear performances in the ball-on-flat tests than the other groups investigated, since these samples have a thicker compound layer and diffusion zone higher than the other conditions investigated. In general, the beginning wear is slower because of the hardest region of the compound layer.

Keywords: Plasma Nitriding, Continuous Cooling Bainitic Steel, Wear Resistance, Reciprocating Dry Sliding.

1. Introduction

Continuous cooling bainitic steel has an increasing use in industrial processes owing to its excellent outstanding combination of yield strength and toughness1-3, the possibility to reduce the process chain and reduced energy consumption4-7. Although their surface properties are acceptable for many purposes8-11, surface hardness and wear resistance are insufficient for some highly loaded automotive components as gears for example. Therefore, the improvement of surface properties is essential. Several surface treatments have been considered for advanced high strength steels for automotive applications12-14.

The plasma nitriding process can be used to develop outstanding surface properties while carrying treatments using relatively low treatment temperatures15,16, thus representing a reliable alternative to conventional thermal and thermochemical treatments. Since long nitriding times can have a detrimental effect on the mechanical properties of steel, the treatment temperature should be carefully selected to avoid overheating of the steel itself and to preserve the bainitic microstructure of these steels, as well as softening of the plasma nitrided layer due to the excess aging of nitrides15,17. The microstructure of the plasma nitrided layer must be well controlled to achieve the desired increase in surface hardness and provide significant improvements in wear resistance18,19.

This study is a sequence of a previous work about plasma nitriding of continuous cooling bainitic steel15. The previous investigations showed excellent results with respect to case depth, surface hardness and compressive residual stresses in the diffusion zone. Despite the variety and particularities of investigations concerning the wear resistance of plasma nitrided steels20-23, the influence of plasma nitriding on the reciprocating dry sliding wear of continuous cooling bainitic steels is relatively scarce. The main objective of this work is to improve the surface properties of DIN 18MnCrSiMo6-4 continuous cooling bainitic steel by plasma nitriding, aiming to increase surface hardness and optimize its wear resistance in reciprocating sliding. In order to study the influence of
plasma nitriding with a nitrogen rich gas composition on the reciprocating sliding wear of a DIN 18MnCrSiMo6-4 continuous cooling bainitic steel, a systematic variation of temperature and time parameters has been carried out. The major contribution of the work is to attempt, through statistical methods, to establish correlations between performance parameters (wear, layer thickness, hardness) and temperature and time employed to produce the modified surfaces.

2. Materials and Methods

The steel to be plasma nitrided was a DIN 18MnCrSiMo6-4 steel (0.19% C, 1.16% Si, 1.35% Mn, 1.14% Cr, 0.26% Mo and 0.06% Ni) which is a continuous cooling bainitic steel. The material’s microstructure is composed of pro-eutectoid ferrite and granular bainite after hot rolling. In the case of continuous cooling bainitic steels there is no annealing (tempering) treatment applied as the microstructure was obtained directly after hot rolling and controlled cooling. Prior to the surface treatments, samples flat surfaces were ground with silicon carbide grinding paper in a sequence with increasing grit sizes (#100, #220, #320, #400, #600, #1200) and then polished with 3 μm diamond paste, in order to obtain low roughness, and almost no plastic deformation.

The surface treatment was carried out in a plasma nitriding furnace equipped with a DC power supply which was developed by the Technology Center and Metallurgy Department of Federal University of Rio Grande do Sul. The samples were degreased and cleaned with acetone in an ultrasonic bath before being placed into the plasma nitriding furnace. The sputtering was performed for 15 min, using pure hydrogen with a flow of 140 sccm, and in the heating step until reaching the treatment temperature, a gas mixture composed of argon and hydrogen was used, with 150 sccm of argon and 140 sccm of hydrogen (H₂). For the plasma nitriding process, a gas mixture composed of 76 vol.% nitrogen (N₂) and 24 vol.% H₂ was used. The treatment parameters are shown in Table 1.

In this work it was studied the influence of processing time (3, 6 and 9 h) and temperature (400, 450, 500 and 550 °C) on mechanical and microstructural characteristics of plasma nitrided layer. The temperature range based on common plasma nitriding temperatures for low alloyed steels, from 400 to 550 °C, to evaluate the maximum temperature that could be used to accelerate the layer growth without causing a decrease in core hardness. The nitrogen rich gas composition (76 vol.% N₂) is commonly used in gas nitriding, as it provides formation of higher layer depths. Therefore, in this work we firstly choose to employ this fixed composition and vary temperature and time. The current density was calculated by dividing the measured current by the total area covered by the plasma sheath. The current density cannot be directly controlled, as we did not use an auxiliary heating system. The current density is dependent on the desired treatment temperature and depends on the gas mixture being used, and on the heat exchange in the furnace.

For the microstructural analysis of the plasma nitrided layers, samples were carefully cut perpendicular to the upper nitrided face in a precision diamond blade cutting machine. After hot pressing mounting in bakelite, the cross-sections were ground with silicon carbide grinding papers in a sequence with increasing grit sizes (#100, #220, #320, #400, #600, #1200) and polished with a diamond paste of 3 μm particle size. A Nital 3% solution was used to etch the samples revealing the microstructure of the nitrided layers. The cross-sectional microstructural images were obtained by an Olympus metallurgical microscope (BX51M model). The compound layer thickness was measured using the Image-J software.

X-ray phase analysis was carried out using a XRD M – Research Edition diffractometer (GE Seifert Charon model) equipped with a Meteor 1D fast line position sensitive detector. Phase analysis was performed on samples surface in the Bragg–Brentano geometry (θ–2θ) with Cr-Kα (λ = 2.2897 Å) radiation. Diffraction lines were recorded in the range of 20 from 55° to 80°, with a step size of 0.01° and a scan time of 200 s per step. The phases present in the non-nitrided and in the plasma nitrided material were determined by comparative analysis between the standards contained in the crystallographic information files (CIFs) from the Crystallography Open Database and the Inorganic Crystal Structures Database by analysis using the Profex-BGMN software.

Vickers microhardness tests were applied to the plasma nitrided samples for determination of the surface hardness and Vickers microhardness profiles of the cross-sections.

| Temperature (°C) | Nitriding time (h) | Gas mixtures (vol. % N₂ in H₂) | Current density (mA/cm²) | Pressure (Pa) |
|-----------------|--------------------|--------------------------------|--------------------------|--------------|
| 400 ± 2         | 6                  | 76 vol.% N₂ in 24 vol.% H₂    | 2.0 ± 0.06               | 300 ± 3      |
|                 | 9                  |                                |                          |              |
| 450 ± 2         | 6                  | 76 vol.% N₂ in 24 vol.% H₂    | 2.2 ± 0.06               | 300 ± 3      |
|                 | 9                  |                                |                          |              |
| 500 ± 2         | 6                  | 76 vol.% N₂ in 24 vol.% H₂    | 2.4 ± 0.10               | 300 ± 3      |
|                 | 9                  |                                |                          |              |
| 550 ± 2         | 6                  | 76 vol.% N₂ in 24 vol.% H₂    | 2.9 ± 0.06               | 300 ± 3      |
|                 | 9                  |                                |                          |              |
In the tests, an Insize micro hardness tester (ISH-TDV 1000 model) was used. Five microhardness profiles per sample were constructed, using a load of 100 gf with a dwell time of 10 s\(^2\). The case depth (compound layer + diffusion zone) was conventionally determined by the distance from the surface where the core hardness is exceeded in 50 HV. The determination of the case depth was carried out based on the microhardness profiles and following the recommendations established in DIN 50 190\(^2\).

The roughness parameters (R\(_z\) and R\(_q\)) were measured with a Mitutoyo contact profilometer (SJ-210 model) equipped with an 8 µm tip radius probe, based on ISO 4287\(^2\). Due to the topographic characteristics of the samples, measurements were made using the aperiodic roughness profile configuration\(^5\). In this case, a cutoff value of 0.8 mm and a measurement length of 4.0 mm was set on the profilometer.

Wear tests were conducted using a CETR UMT (Universal Materials Tester) tribometer with a reciprocating ball-on-flat configuration, according to the standard ASTM G133\(^3\). Tests were performed in a room with 50 to 55% of relative humidity and the temperature at 23 °C. The treated surfaces were placed in contact against a 4.76 mm zirconia ball (1150 HV\(_{0.1}\) ± 28). The choice of a ball with a close hardness to the nitrided surface is due to the need to simulate the contact conditions with a close hardness, such as for example, the contact between the flanks of the gears, to be tested in future works. The load of 6 N was applied (maximum contact pressure nearly 1351 MPa), and the tests were conducted until the microcracks formation or delamination of the surface layer. The applied load was selected aiming to analyze the wear resistance of the whole case (compound layer + diffusion zone), and not the specific wear resistance of the compound layer. The maximum contact pressure (p\(_c\)) of 1351 MPa was estimated for wear tests through the equation \[ p_c = \frac{3W}{2\pi a^2}, \] being (W) is the normal load and (a) is the radius of the contact area\(^2\). For this reason, it was chosen to use a high contact pressure. Sliding proceeded in the reciprocating mode over a 4 mm track at a frequency of 4 Hz. The surface topography and track profiles were acquired in a Bruker interferometer (Contour Elite model). The cross-sectional area was measured, using the Image-J software, in three regions of each worn track.

Finally, to identify wear mechanisms, some specific analysis of the balls and wear tracks were carried out in a Zeiss scanning electron microscope (EVO MA 15 model). Interaction plots for case depth, surface hardness, and sliding distance at the time of microcracks formation or delamination of the surface layer, with nitriding temperature and time were generated. The data was then submitted to an analysis of variance, in order to evaluate the statistical significance of temperature variation and treatment time on wear, following the Tukey’s-b post hoc test\(^1\) by using the statistical package program (Minitab 16). Three repetitions and a confidence level of 95% (significance level \(\alpha = 0.05\)) were used.

3. Results and Discussion

3.1. Microstructure, phase analysis and microhardness

As a representative result of treatments using the gas mixture of 76 vol.% N\(_2\) and 24 vol.% H\(_2\), cross-sectional microstructures of a plasma nitrided sample at 550 °C and 400 °C are shown in Figure 1a, b, respectively. The white portion on top distinguishes the compound layer, as seen at Figure 1a. For plasma nitrided samples at the lowest temperature, Figure 1b, the case depth is predominantly composed of the diffusion zone with a very thin compound layer (see Table 2). Under conditions where samples were plasma nitrided at temperatures higher than 400 °C, effective compound layer appears in the metallography. All different nitriding conditions in the present work, besides the diffusion zone, led to the formation of compound layer. Table 2 shows the average case depth of the plasma nitrided samples for different treatment temperatures and times. In general, the use of higher temperatures and longer treatment times favored the expansion of case depth.

Previous work\(^15\) showed that all the plasma nitrided samples exhibited diffraction peaks that indicated the formation of a compound layer. Since the penetration depth of radiation decreases with 20 angle\(^24\), the peaks of the \(\varepsilon\)-Fe\(_{23}(C)N\) phase, contained in the compound layer, are easily identified with smaller 20 angles. Therefore Figure 2a-c present the diffractograms for the (20) angular interval from 55° to 80°.

![Figure 1. Optical microstructure of cross-section of plasma nitrided samples at: (a) 550 °C for 6 h (compound layer predominant); (b) 400 °C for 3 h (diffusion zone predominant).](image-url)
Table 2. Compilation of results from case depth, compound layer thickness, surface hardness and sliding distance at the time of microcracks formation or delamination of the surface layer.

| Nitriding parameters | Case depth (µm) | Compound layer thickness (µm) | Surface hardness (HV<sub>0.1</sub>) | Sliding distance* (m) |
|----------------------|----------------|-----------------------------|-----------------------------------|---------------------|
| 400 °C - 3 h         | 67 ± 20        | 0.6 ± 0.2                   | 1060 ± 38                         | 11 ± 2              |
| 400 °C - 6 h         | 98 ± 12        | 1.5 ± 0.2                   | 1051 ± 28                         | 13 ± 1              |
| 400 °C - 9 h         | 125 ± 21       | 2.7 ± 0.3                   | 1095 ± 95                         | 18 ± 1              |
| 450 °C - 3 h         | 97 ± 20        | 2.6 ± 0.3                   | 1253 ± 56                         | 19 ± 3              |
| 450 °C - 6 h         | 131 ± 24       | 3.9 ± 0.2                   | 1105 ± 90                         | 18 ± 2              |
| 450 °C - 9 h         | 169 ± 40       | 4.8 ± 0.4                   | 1101 ± 56                         | 37 ± 15             |
| 500 °C - 3 h         | 139 ± 30       | 4.3 ± 0.2                   | 1246 ± 95                         | 21 ± 6              |
| 500 °C - 6 h         | 192 ± 25       | 6.7 ± 0.3                   | 1295 ± 108                        | 29 ± 7              |
| 500 °C - 9 h         | 245 ± 40       | 8.2 ± 0.2                   | 1079 ± 87                         | 30 ± 8              |
| 550 °C - 3 h         | 253 ± 23       | 7.9 ± 0.3                   | 1195 ± 45                         | 50 ± 3              |
| 550 °C - 6 h         | 308 ± 24       | 9.6 ± 0.3                   | 1089 ± 46                         | 58 ± 0**            |
| 550 °C - 9 h         | 335 ± 15       | 11.5 ± 0.2                  | 1029 ± 50                         | 58 ± 0**            |

*Sliding distance at the time of microcracks formation or delamination of the surface layer. **Without COF’s sudden growth.

Diffractograms of non-nitrided samples shows the α-Iron peaks, Figure 2a-c, from the ferritic phases. The intensity of the peaks of the α-Iron decreases and the retained austenite completely disappeared with the increase of the compound layer thickness, as the opposite takes place with ε-Fe<sub>2</sub>(C) N and γ'-Fe<sub>N</sub> phase peaks, because to the absorption of the radiation by the most superficial portions of the material. The α phase peaks of the nitrided samples are slightly shifted to the region of lower 20 angle with respect to the non-nitrided samples. This is an information coming from the diffusion layer, as the compound layer is composed by (carbo)nitrides only<sup>15</sup>. In general, the peaks intensity related to the γ'-Fe<sub>N</sub> and ε-Fe<sub>2</sub>(C)N nitride phases increase with the temperature and treatment time, since compound layer thickness (Table 2) follow the increase of these parameters.

The surface hardness decreases, especially for longer treatments of 6 and 9 h (Table 2). The core hardness is also reduced in samples treated at 550 °C, which is not observed for other treatment temperatures<sup>15</sup>. Higher plasma nitriding temperature leads to the intensification of defects formed in the compound layer, such as pores and cracks, and therefore the measure hardness value decreases. Our previous work shows that at high temperatures, there is also a competition between the hardening effect of nitriding and the hardness decrease due to the overheating of the steel matrix, so that the maximum hardening potential is not reached.

Figure 3a-c show microhardness profiles for the different plasma nitriding conditions. From the microhardness profiles, a case depth was estimated. As expected, the nitriding depth follows the behavior of a diffusion controlled process<sup>15,16</sup>. Plasma nitriding under the conditions implemented in this work could lead to hardness levels from 1029 to 1295 HV<sub>0.1</sub>, and case depths up to 300 µm, see Table 2. Plasma nitriding leads to an increase in hardness on the surface (Table 2), for the reason it also ends up creating a less ductile region more prone to brittle fracture<sup>15</sup>. The mechanical properties of the diffusion zone influence the fracture properties of the compound layer, since the diffusion zone provides support for the surface compound layer<sup>15,16,36</sup>. Previous work<sup>15</sup> showed that the higher nitriding temperature and time promoted an increase in the fracture toughness of the compound layer. This would also support the proposition that a harder substrate results in an increase in fracture toughness through improved mechanical support (load bearing capacity) of the compound layer by the diffusion zone.

3.2. Roughness

To investigate the influence of temperature and nitriding time on the surface topography, the roughness values (parameters R<sub>a</sub> and R<sub>s</sub>) measured before and after plasma nitriding are shown in Figure 4a, b. In the cases shown in Figure 4a, b, the exposure time to ion bombardment ends up changing the final a lot, confirming the results reported by<sup>37,38</sup>. It can be seen in Figure 4a, b that after the plasma nitriding treatment the roughness (R<sub>a</sub> and R<sub>s</sub> parameter) increases when compared to the polished surface prior to nitriding, and that the increased temperature and the nitriding time caused an increase in the roughness in the two parameters evaluated.

From a statistical point of view, the ANOVA reveals that the roughness was influenced by the nitriding temperature (P<sub>Value</sub> = 0.00 < α = 0.05) and that there was a synergistic effect between the temperature and the nitriding time (P<sub>Value</sub> = 0.02 < α = 0.05) on the increase in roughness, but it was not possible to determine the effect of treatment time (P<sub>Value</sub> = 0.25 > α = 0.05), due the standard deviation. Tukey’s-b post hoc test shows that the greatest roughness was found in the plasma nitrided samples for 9 h at 550 °C, Figure 4a, b.

The three-dimensional topographic measurements revealed that the plasma nitrided surface morphology is modified by the nitriding parameters, confirming the previously presented roughness values. Figure 5a, b show the 3D surface representative images of plasma nitrided samples with gas mixture of 76 vol.% N<sub>2</sub> and 24 vol.% H<sub>2</sub>. In general, there is an intense change in topography after nitriding. This generalized increase in roughness after nitriding can be related to the ion bombardment during plasma treatment and the formation mechanism of the nitrided layer<sup>37,38</sup>. Nevertheless, this increase in topography roughness may have contributed to the increased wear of the zirconia ball.
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3.3. Reciprocating dry sliding wear tests

In this work, the friction coefficients (COF) approach obtained on the wear tests will be carried out in two regions: at the beginning on the wear tests, Figure 6a-d, and at the time of microcracks formation or delamination of the surface layer, Figure 7a-d. The compound layer of the group of plasma nitrided samples at temperatures of 400, 450 and 500 °C is similar (from point of view of the COF), with less roughness and more importantly the characteristics of the compound layer, as specifying the images shown in Figure 6a-d. For the group of plasma nitrided samples at 550 °C, the COF is very low since the beginning of the tests, regardless of the treatment time. Regardless of the nitriding temperature, at the beginning of the reciprocal sliding wear test, the behavior of the nitrided samples in 9 h is similar at all temperatures. This is justified due to the fact that the hardness of the compound layer is similar among the group of plasma nitrided samples in 9 h, Table 2. Even with the
Figure 4. Roughness measurements before and after plasma nitriding at different temperatures and times: (a) $R_a$; (b) $R_z$.

Figure 5. Surface representative images of plasma nitrided samples for 9 h at: (a) 400 °C; (b) 500 °C.

Figure 6. Friction coefficient obtained at the beginning on the reciprocating wear tests on non-nitrided samples and plasma nitrided samples for 3, 6 and 9 h at (a) 400 °C, (b) 450 °C, (c) 500 °C and (d) 550 °C.
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roughness being lower in the non-nitrided samples and in the nitrided samples in 3 h, the COF is higher, and this is related to an adhesion effect as being preponderant for this tribological system.

The average COF of the non-nitrided samples was 0.56 ± 0.05 (admitting that the steady-state occurs from 30 m), Figure 7a-d. The non-nitrided samples stabilizes the friction values around 0.56 ± 0.05 in a test of up to 30 m of sliding distance, which did not occur for the plasma nitrided samples (with the exception from plasma nitrided samples at 550 °C for 6 and 9 h). The results show that the nitriding treatments were responsible for the reduction of the COF, Figure 6a-d, this can be attributed to the increase in hardness (Table 2) and the contact area between ball and surface, which reflects in smaller deformations in contact surfaces and depends on elastic deformations. The lower COF at first occurs due to the ceramic characteristics of the compound layer. The increase in COF, over time of wear is due to the presence of the third body of the ball and of the compound layer that was wearing out. As shown in Figure 7a-c, the sudden growth of COF is related to the wear of the surface layer that exposes the diffusion zone. When the diffusion zone is reached, the friction rises and may even reach or exceed the friction of the non-nitried samples. Some local damage, results in the propagation of microcracks or delamination of the surface layer..

Figure 7. Friction coefficient obtained at the time of microcracks formation or delamination of the surface layer in reciprocating wear tests on non-nitried samples and plasma nitrided samples for 3, 6 and 9 h at (a) 400 °C, (b) 450 °C, (c) 500 °C and (d) 550 °C.

Table 3. P-value obtained through a two-way analysis of variance.

| Input Factor   | Case depth | Surface hardness | Sliding distance** |
|----------------|------------|------------------|--------------------|
| Temperature    | 0.00*      | 0.00*            | 0.00*              |
| Time           | 0.00*      | 0.03             | 0.01               |
| Temperature*Time| 0.68       | 0.01             | 0.16               |

*The Minitab software package displays the P-value = 0 when the results tend to zero. **Sliding distance at the time of microcracks formation or delamination of the surface layer.
As inferred from the Table 3 and case Figure 12. The interaction plot demonstrates how the $-\text{Value}$ samples at 550 °C for 6 and 9 h. In general, all plasma nitrided from the zirconia ball on the worn tracks of plasma nitrided and cross-section profiles was the deposition of particles of the plasma nitrided samples at 550 °C, see the softer hardness gradient between the surface and core. Another factor that must be acting strongly in this case is the compound layer, preventing the beginning formation of microcracks or delamination, as seen in Figure 11a, b. However, the test was interrupted before the complete spalling of the plasma nitrided surfaces (Table 2). In general, the higher nitriding temperatures promoted an increase in the compound layer thickness. This increase associated with the $\varepsilon$-Fe$_2$N(C)N phase, which according to Doan et al. has a lower friction coefficient, Figure 2a-c, provided the bearing capacity of the compound layer, preventing the beginning formation of microcracks or delamination, as seen in Figure 11a, b. Another factor that must be acting strongly in this case is the softer hardness gradient between the surface and core of the plasma nitrided samples at 550 °C, see Figure 3a-c.

Another fact observed, in EDS analyses by Figure 10c, d and cross-section profiles was the deposition of particles from the zirconia ball on the worn tracks of plasma nitrided samples at 550 °C for 6 and 9 h. In general, all plasma nitrided samples showed zirconia deposition on the worn tracks. As only slight wear occurred in the plasma nitrided samples at 550 °C for 6 and 9 h, the SEM/EDS analysis, Figure 10a-d detected only particles of third body generated by the wear of the zirconia ball.

Figure 9c, d, Figure 10c, d and Figure 11c, d also show the EDS analyses carried out on the worn tracks after reciprocal sliding wear to identify the chemical elements present in the surface of the plasma nitrided samples, in addition to the zirconia transferred from balls to the tracks. The peaks associated with the core steel lose intensity in samples with higher compound layer thickness, that is, in plasma nitrided samples at higher temperatures (500 and 550 °C).

Studies developed by Dutrey et al. show that the plasma nitriding with a nitrogen rich gas composition on a low-alloy steel (AISI 4140) formed intergranular fracture mode in the diffusion zone, related to nitride precipitation at grain boundaries. Despite the gas mixture of 76 vol.% N$_2$ can be used to improve the surface properties of the DIN 18MnCrSiMo6-4 steel, the nitrogen rich gas composition showed precipitation of nitrides, carbo-nitrides or carbides at grain boundaries for plasma nitriding of M2 high-speed steel. It is known that grain boundary precipitation increases with time and temperature. The presence of precipitation at grain boundaries results in brittleness of the diffusion zone, and this can lead to the delamination of the surface layer, Figure 9a, b and Figure 11a, b, or even spalling.

The pile-up of tracks were analyzed from the cross-sections, according to Figure 12. As inferred from the cross-section profiles illustration (Figure 12) and case depth (Table 2), tracks did not reach the substrate in the plasma nitrided surfaces. The friction and wear results can be associated to the operating wear mechanism found in the plasma nitrided samples. Observing a cross-section profile view of tracks revealed in the optical interferometry, one can note the absence of pile-up for the group of plasma nitrided samples at 550 and 500 °C for 6 and 9 h and 450 °C for 9 h. For the group of plasma nitrided samples at 550 and 500 °C for 3 h, 450 °C for 3 and 6 h and 400 °C for 3, 6 and 9 h, it was verified of pile-up, resulting in the predominance of plastic deformation. As mentioned before, samples presenting higher case depths, exhibited lower plastic deformation, since the diffusion zone provided the bearing capacity of the compound layer.

In order to evaluate the statistical significance of treatment temperature and time on the case depth, surface hardness and sliding distance on the time for microcracks formation or delamination of the surface layer, the collected data results were analyzed using the Minitab 16 software. The results of the interaction plot by analysis of variance are shown in Figure 13a-c. The interaction plot demonstrates how the relationship between a categorical factor and a continuous response depends on the value of the second categorical factor. The interaction strength is greater when the lines in the graphs are non-parallel.

From a statistical point of view (Table 3), it is noticed that the case depth was influenced by the temperature ($P_{\text{Value}} = 0.00 < \alpha = 0.05$) and time ($P_{\text{Value}} = 0.00 < \alpha = 0.05$) parameters, but there was no synergistic effect between temperature and treatment time ($P_{\text{Value}} = 0.68 > \alpha = 0.05$).

![Figure 8](image_url)
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Figure 13a, because there is a competition between sputter yield and nitrogen uptake. Tukey’s-b post hoc test shows that the case depth has a significant difference between the times of 3 and 6 h, with a technical tie between 6 and 9 h. As for the case depth, the temperature that showed the best response was the nitriding performed at 550 °C, confirmed by the Tukey’s-b post hoc test.

As can be seen in Figure 13b, the surface hardness behavior of the samples remained relatively close. Both factors studied temperature and time showing statistical relevance ($P_{\text{Value}} = 0.01 < \alpha = 0.05$) and were presented in Table 3. Tukey’s-b post hoc test showed that although the temperature is significant in increasing the surface hardness, the difference between the results obtained in the time variation was not greater than three times the standard deviation. Tukey’s-b post hoc test between the different temperatures showed that there was a significant increase between 400 and 500 °C. Regarding the hardness profile, the plasma nitrided samples at 550 °C did not show any significant difference with nitriding performed at 500 °C, but in terms of surface hardness, the best temperature was 500 °C. The time should be determined according to the need for the case depth. However, as previously discussed, Figure 13a, low case depths (compound layer + diffusion layer) provided the pile-up formation. Therefore, as there was a statistical tie on the wear of the plasma nitried samples at 6 and 9 h at 550 °C, Figure 13c, the use of 6 h is recommended, due to the process cost.

Figure 13c shows that the sliding distance at the time of microcracks or delamination of the surface layer had no synergistic effect ($P_{\text{Value}} = 0.16 > \alpha = 0.05$), and that the main effects were more important (Table 3). However, the input parameters: temperature ($P_{\text{Value}} = 0.00 < \alpha = 0.05$) and time ($P_{\text{Value}} = 0.01 < \alpha = 0.05$) had significant effects on the sliding distance until the abrupt increase in COF, Figure 7a-d. By Tukey’s-b post hoc test analysis the nitriding time did not show differences greater than three times the standard deviation in the sliding distance. However, low nitriding times generated smaller case depth and favored the pile-up formation which, in some cases, contributed to delamination of the surface layer. The temperature variation analysis showed that the ideal temperature for increasing the sliding distance before the abrupt increase in COF and the consequent increase in wear resistance occur at 550 °C. The best performance for reciprocating dry sliding wear tests was found in plasma nitried samples at 550 °C with nitriding time of 6 h.

3.4. Final discussion

The compilation of case depth, surface hardness and sliding distance at the time of microcracks formation or delamination of the surface layer of plasma nitried samples are given in Table 2. Time and temperature can influence not only the compound layer thickness, but also its composition. The X-ray analysis, Figure 2a-c, showed the formation of biphasic compound layer, $\varepsilon$-Fe$_3$,(C,N) and $\gamma'$-Fe,N, for in all treatment conditions. It might normally be expected that
Figure 10. (a, b) SEM images and (c, d) EDS analysis of the wear tracks after reciprocal sliding wear of plasma nitrided samples at 550 °C for 9 h (as a representative result of plasma nitrided samples at 550 °C for 6 and 9 h).

Figure 11. (a, b) SEM images and (c, d) EDS analysis of the wear tracks after reciprocal sliding wear of plasma nitrided samples at 450 °C for 9 h (as a representative result of plasma nitrided samples at 400, 450 and 500 °C for 3, 6 and 9 h).
the compound layer containing predominant ε-Fe₂₋₃(C)N has higher hardness than the compound layer monophasic containing γ'-Fe₄N. For nitriding temperatures up to 500 °C, hardness increases with treatment temperature and time. The observed behaviour seems to be governed, by the increase of the compound layer thickness. For 550 °C on the other hand, the surface hardness is lower than for 500 °C, which happened due to the overheating of the substrate, as core hardness decreased for this temperature. Concerning the sliding distance at the time of microcracks formation or delamination of the surface layer, as nitriding temperature and time increased to 500 °C, the compound layer also increased. The same happened to the sliding distance and the worn volume, as expected. For the group of plasma nitrided samples at 550 °C for 6 and 9 h did, no delamination of the surface layer (even at a sliding distance of 58 m), was observed. Actually, a marked wear of the zirconia balls took place for these testing conditions. The plasma nitrided surfaces presented a lower COF again zirconia ball than non-nitrided steel. When the surface layer starts to delaminate, the friction coefficient quickly exceeds the substrate values.

The statistical analysis showed, with 95% of statistical confidence, that the best setup to resist dry sliding reciprocal wear on the tested configuration was plasma nitriding at 550 °C for 6 h. However, Dalcin et al. showed that the core hardness of samples nitrided at 550 °C was impaired. Therefore, for practical applications in components such as gears, the use at 500 °C is recommended, due to temperatures at 400 and 450 °C would require too long treatment times to produce the appropriate case depth.

In this work, it was found that the gas mixture of 76 vol.% N₂ can be used to improve the surface properties of the DIN 18MnCrSiMo6-4 steel, but this nitrogen rich gas composition can cause embrittlement of the diffusion zone, mainly in high carbon steels. The embrittlement is associated with the formation of precipitates in the grain boundary, but for steels with a low carbon, the embrittlement does not seem to be as intense. In order to avoid the embrittlement of the diffusion zone, a new study using a gas mixture with a lower nitrogen composition would be indicated. This issue will be addressed in future work.

4. Conclusions

Figure 12. Cross-section profiles illustration of tracks produced in reciprocating wear tests on the plasma nitrided surfaces.

Figure 13. Analysis of variance. Interaction plot of temperature and nitriding time with: (a) Case depth; (b) Surface hardness; (c) Sliding distance at the time of microcracks formation or delamination of the surface layer.
Plasma nitriding of DIN 18MnCrSiMo6-4 continuous cooling bainitic steel is viable, since it demonstrated good response to the process with excellent results regarding layer depth and surface hardness. All plasma nitriding treatments were able to form a compound layer and generate a significant increase in surface hardness, raising it from 329.8 HV0.1 in the material core to values above 1000 HV0.1.

Plasma nitriding treatments were responsible for the reduction of the friction coefficient. The lower friction at first occurs due to the ceramic characteristics of the compound layer. The increase in COF, over time of wear is due to the presence of the third body of the ball and of the compound layer that was wearing out. The sudden growth of COF is related to the wear of the surface layer that exposes the diffusion zone. When the diffusion zone is reached, the friction rises and may even reach or exceed the friction of the non-nitrided samples.

If assessing the wear mechanisms, only the samples that have greater case depth (compound layer + diffusion zone), the diffusion zone supports the compound layer. This causes the wear to be slower, precisely because the beginning wear is in the hardest region of the compound layer until it reaches less hard regions. For this reason, this transition is smoother and takes longer before microcracks or delamination appears. In the case of plasma nitrided samples at 550 °C for 6 and 9 h, the wear test would have to be extended for a longer time until the damage mentioned above appears;

Since that the plasma nitrided samples at 550 °C have a thicker compound layer and diffusion zone higher than the other conditions investigated, these samples showed better wear performances in the ball-on-flat reciprocal dry sliding than the other groups investigated. The nitrided layer formed in the treatment at 550 °C for 6 h present the best performance for the tested conditions, with 95% of statistical confidence.

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