Corrosion behavior of low carbon steel processed by ECAP in neutral and alkaline environment

Jorgimara Oliveira Braga 1
Jefferson Fabricio Lins 2
Gabriel Pessoa de Castro 2
Helder Alves de Almeida Junior 1
Igor Cuzzuol dos Santos 1 *
Gustavo Henrique Sousa 1
Izabel Lima Criscuolo 1
Cláudio Marcio Santana 1
Ursula Cid Pereira 1

Abstract

The purpose of this study was to evaluate the corrosion behavior of 1010 steel processed by Equal Channel Angular Pressing (ECAP), a technique that induces severe plastic deformation (SPD) in materials. The samples were processed by one and two passes of ECAP through routes A and C. The corrosion behavior was evaluated using the potentiodynamic polarization and electrochemical impedance spectroscopy, using NaCl (pH = 6.5) and sodium hydroxide/sodium phosphate (pH = 10.5) solutions. The polarization tests showed better performance for the samples processed by ECAP than in the as cast samples both in alkaline and neutral solutions. The results showed that alkaline solution presented higher corrosion rate for the as cast and one ECAP pass materials. It is seen that samples processed through routes A and C develop a film stabilization on the surface of these samples. In addition, the values of polarization resistance obtained from the electrochemical impedance spectroscopy in NaCl environment were higher for sample processed in route C.

Keywords: Low carbon steels; SPD; ECAP; Corrosion.

Comportamento de corrosão de aço com baixo carbono processado por ECAP em ambiente neutro e alcalino

Resumo

O objetivo deste estudo foi avaliar o comportamento à corrosão do aço 1010 processado por por Preensagem em Canais Equiangulares (ECAP), uma técnica que induz deformação plástica severa (DPS) em materiais. As amostras foram processadas em um e dois passes nas rotas A e C. O comportamento em relação à corrosão foi estudado por meio da polarização potenciodinâmica e espectroscopia de impedância eletroquímica em soluções de NaCl (pH = 6.5) e hidróxido de sódio/fosfato de sódio (pH = 10.5). Os testes de polarização apresentaram melhores desempenhos para as amostras processadas por ECAP do que nas amostras sem processamento, e isso ocorreu tanto na solução alcalina quanto na neutra. Os resultados mostram que os materiais no estado bruto de fusão e em apenas um passe de ECAP apresentaram maior taxa de corrosão em solução alcalina. Foi observado que as amostras processadas nas rotas A e C promoveram uma estabilização do filme em suas respectivas superfícies. Ademais, os valores de resistência de polarização obtidos por espectroscopia de impedância eletroquímica em ambiente de NaCl foram maiores para a amostra processada na rota C.

Palavras-chave: Aços baixo carbono; DPS; ECAP; Corrosão.
1 Introduction

Low carbon steels account for about 70% of the world’s ferrous alloy production. These materials have been used in a range of industrial applications, from immense structural components to small parts of mechanical devices. In order to meet the market’s demand, it is important to develop materials with special mechanical characteristics, such as high mechanical strength allied with high toughness. For example, some requirement mechanical properties of low carbon steels used for structural purposes are fatigue strength, wear resistance, mechanical strength associated with good toughness, weldability and corrosion resistance [1].

One of the methods applied to improve the mechanical properties of low carbon steels is the reduction of the material grain size using severe plastic deformation techniques. Severe plastic deformation (SPD) can be define as a group of techniques in which a high plastic deformation is apply in the material in order to produce ultrafine microstructures, also known as ultrafine grains (UFG) [2]. Materials with UFG are of great interest, since the grain refinement promotes an increase in the material mechanical properties, enabling the production of components and structures that demand high mechanical resistance and high corrosion resistance [3].

ECAP was the SPD technique chosen in the present study. In this technique, the sample is place and press through an angular channel using a plunger. The nature of the imposed deformation is simple shear, which occurs as the sample passes through the die. The ECAP process is know as one of the most promising SPD techniques [4]. One characteristic of ECAP is the multiple-step, which means that various strain paths can be followed according to the processing routes defined by the billet rotation between successive passes. In route A, the billet is not rotated between consecutive passes, whereas in route B the billet is rotated 180° between passes. In one pass, the billet extend over only once through the channels [5].

The electrochemical behavior of materials with deformed grains is an important factor to be evaluate mainly in steels, because it has great applicability in industries, buildings and marine environments. However, it is observed that this kind of study has received limited attention and some results are incoherent [6]. Some studies [7,8] have showed that the basic electrochemical characteristics of ultra-fine grains do not vary significantly compared to their coarse grains. On the other hand, refined grain materials generally promote uniform corrosion rather than microscopic corrosion, which minimizes catastrophic results by localized corrosion attack on coarse grain materials.

Then, the purpose of this study is to obtain more information about corrosion behavior of low carbon steel processed by ECAP in different solutions, since there are many controverses about the type of corrosion and electrochemical studies in steel that undergo severe plastic deformation (SPD).

2 Methodology

A commercial low carbon steel with nominal composition in Table 1 was used in the current study. The material was provided by Companhia Siderurgica Nacional (CSN-Volta Redonda/Brasil) in the form of as cast ingots. Billets having 10 × 10 × 50 mm were machined out from the as cast ingots.

It is widely known that ECAP induces severe plastic deformation in metallic materials, which the processing consists of pressing the billet through two equal cross-section channels, intercepting at an angle Φ. Since the billet does not change the cross section, the process can be repeated several times. The ECAP system of Federal Fluminense University (UFF) consists of an universal test machine (EMIC DL-60) with a maximum load capacity of 600 KN and a die with two-part tool steel H13, dimensions of 10x10 mm² and channel angle of 90° (Φ). The die and samples were lubricated with a calcium sulfonate grease prior to each assay. The samples were processed by one pass (no defined route) and two passes using routes A and C at room temperature.

The ECAP-processed samples were mechanically ground using abrasive papers to #4000 SiC and polished using 1.0 µm and 0.06 µm silica colloidal suspension solution. The microstructural surface was examined by optical microscope (Leica DM ILM) coupled with Nikon Digital Sight camera and NIS element D.30 software. In addition, the microstructure was analyzed by the scanning electron microscope (SEM), model EVO MA10 from ZEISS with LaB6 filament. The SEM was operated by backscattered electron mode, acceleration voltage of 25 kV and working distance (WD) of 8 mm. After acquiring the image by optical microscope and SEM, the grain sizes were measured using the linear intercept method according to the standard ASTM E1112 [9].

The electrochemical tests were carried out using potentiostat/galvanostat (SOLARTRON ANALYTICAL XM ECS/FRA) connected by three electrodes: 1010 steel as working electrode, Ag/AgCl as reference electrode and platinum as counter electrode. The electrochemical tests consisted of immersing the electrodes in 100 ml of the electrolyte in a glass cell. The solutions were prepared from analytical purity reagents (Sigma-Aldrich, 99.9%) and distilled water. The-solutions used in the tests were NaCl (0.25 mol. L⁻¹ - pH 6.5) and buffer solution of Na₂HPO₄ (0.1 mol.L⁻¹ - pH 10.5).

| Element | C   | Mn | Ti  | Al  | P* | Cu* | Mo* | Fe  |
|---------|-----|----|-----|-----|----|-----|-----|-----|
| Composition (wt%) | 0.104 | 0.164 | 0.069 | 0.061 | 0.012 | 0.018 | 0.006 | remain |

*Impurites elements.
The polarization curves were performed in order to evaluate the corrosion resistance of the deformed material compared to the as cast. The tests were conducted after potential stabilization and were performed in triplicates to achieve reproducibility in the results and in different solutions. The electrochemical impedance spectroscopy was realized to evaluate the corrosion behavior in 0.25 NaCl mol.L\(^{-1}\) at immersion time of 48 hours using a frequency range of 10 kHz to 10 mHz, signal amplitude of 10 mV in relation to the open circuit potential (OCP) and scan rate of 5 mV/s. The results obtained through the Electrochemical Impedance Spectroscopy (EIS) test were plotted on the Nyquist diagram, the representation is made by \(Z'\) and \(Z''\), which means real and imaginary part of the impedance, respectively. In other words, the abscissa is the real part (resistive terms, represented by R) and the ordinate is the imaginary part (capacitive or inductive terms). In EIS tests, it is important to correlate the physical and chemical processes that occur in the system with electrical circuit elements to better understand the phenomena and interpretation of the results. The Zview \(^{®}\) software was applied to data extrapolation and to obtain the equivalent circuit of the system.

In order to facilitate the understanding of the experimental procedure in the text, the as-cast sample was designated as A. The processed materials were labeled as B for one ECAP pass and C and D for two passes, using routes C and A, respectively (shown in the Table 2).

### 3 Results

#### 3.1 Microstructural analysis

After processing by ECAP, the material presented very elongated grains, as shown in Figure 1b. The grain size (Figure 1a) evaluated was about 48 ± 2.7 μm for the sample A and 23.9 ± 3.1 μm for sample B (Figure 1b). This means a significant reduction of approximately 77% of the grain size after ECAP. The evaluated grain size for the samples C and D were 11.2 ± 5.5 μm and 8.6 ± 7.8 μm, respectively. The high error values can be attributed to the heterogeneity induced by variation of grain size in the material, which makes the linear intercept method more difficult to be evaluate.

| Sample | Test Condition                  |
|--------|--------------------------------|
| A      | As cast                        |
| B      | One pass                       |
| C      | Two passes route C             |
| D      | Two passes route A             |

**Table 2. Processed samples used in this study**

![Figure 1. Optical Microscopy of 1010 steel in the cross section: (a) sample A; (b) sample B; (c) sample C and (d) sample D.](image-url)
3.2 Electrochemical behavior in neutral and alkaline environment

Figure 2a shows the electrochemical behavior of 1010 steel in NaCl solution (all results were obtained in triplicate). From these results, it is possible to observe an increase in the current density with a rise in the potential values. In Figure 2a, the increase in the current density begins at the pitting potential (Epitting) and the pitting corrosion occurs due to the rupture of the passive film. The potential values (Epitting) for the sample A was evaluated as -0.43 V and the value for the processed materials was -0.41 V; -0.34 V; and -0.36 V for samples B, C and D, respectively. These data confirm a greater stability of the passivation film in the samples C and D.

The tests conducted in the alkaline solution with a sodium hydroxide 0.1 mol.L⁻¹/sodium phosphate 0.1 mol.L⁻¹ (pH 10.5) are presented in Figure 2b. It is observed that the passive current density (i_pass) for the samples A and B are higher when compared to samples C and D. In other words, the material processed by two ECAP passes presented a passive film more stable when compared to the as cast and the sample processed by only one pass.

Analysis from Figure 3 confirms a greater number of pits that occurred in the samples A and B (Figure 3a and 3b) in comparison with C and D:

To confirm the behavior of the samples in NaCl, measurements of electrochemical impedance were carried out after 48 hours immersion. Figure 4 and Table 3 present the equivalent circuit and fitting values, in which the results were extrapolated by the Zview program. R1 is equivalent to the electrolyte resistance (NaCl), CPE1 is a constant phase element and considered a non-ideal capacitor due to the heterogeneity of electrode surface and double layer characteristics. Polarization resistance was represented in the electrical circuit by R2.

Figure 5 shows the EIS results by Nyquist Plot representation, where the tests were conducted after 48 hours immersion in NaCl solution. The values of polarization resistance (R2) means that the larger the diameter of the semicircle the higher the resistance, and hence, the lower the corrosion rate. Therefore, the values obtained for polarization resistance were R2 = 413.5 Ω.cm² for the sample A and 964.6 Ω.cm², 1770 Ω.cm², 1322 Ω.cm² for samples B, C and D, respectively.

4 Discussion

After one ECAP pass (Figure 1b), it can be observed that the grains in the material became very elongated and presented a fragmented structure. In sample A, the perlite

![Figure 2](image_url)

**Figure 2.** (a) Anodic polarization curve in NaCl 0.25 mol.L⁻¹; (b) anodic and cathodic polarization curve in Na₂HPO₄ 0.1 mol.L⁻¹.

**Table 3.** Simulation of the impedance data by the Zview program for immersion time 48 hours

| Samples | Immersion time (h) | R₁ (Ω.cm²) | CPE₁-T (µF.cm⁻²) | CPE₂-P (µF.cm⁻²) | R₂ (Ω.cm²) | χ² (10⁻³) |
|---------|--------------------|------------|------------------|------------------|------------|-----------|
| A       | 48                | 14.6       | 1154             | 0.75             | 413        | 0.94      |
| B       | 48                | 17.6       | 2185             | 0.78             | 964        | 1.76      |
| C       | 48                | 25.1       | 2284             | 0.84             | 1770       | 1.15      |
| D       | 48                | 19.4       | 2541             | 0.85             | 1322       | 6.54      |
has precipitated in the ferrite grain boundaries. After the first ECAP pass (sample B), it was noticed dark contours inside the ferrite grains. Probably, this is an indicative of the perlite phase fragmentation caused by intense deformation.

For the sample C (Figure 1c), the morphology became equiaxed, which is a typical characteristic of this route. This phenomenon is possible due to a restoration of the cubic element after rotation at 180° [7]. As said earlier, the perlite contours were visible after the first ECAP pass. This observation agrees with Zrník et al. [10] and can be proved by the micrographs of low carbon steel in Figure 1c.

Figure 3. Optical microscopy after polarization test in NaCl 0.25 mol.L⁻¹: (a) as cast; (b) one pass; (c) route C and (d) route A.

For the specimen D (Figure 1d), the grains have became extensively elongated with a very elongated ferritic and perlite structure. This is explained by a continuous and intense processing, which is favorable to a strong rolling texture formation [10]. In addition, it is seen a greater change

Figure 4. Equivalent circuit used to simulate the impedance.

Figure 5. Nyquist diagram for 1010 steel in immersion for 48 hours in NaCl solution.
Table 4. Summary of the main results obtained in this study

| Sample | Test condition | Grain size (µm) | Epitting (V) | ipass (mA.cm⁻²) | R_p (Ω.cm²) |
|--------|----------------|----------------|--------------|----------------|-------------|
| A      | As cast        | 48 ± 2.7       | -0.43        | 0.08           | 413.5       |
| B      | One pass       | 23 ± 3.1       | -0.41        | 0.07           | 964.6       |
| C      | Two passes route C | 11.2 ± 5.5   | -0.34        | 0.03           | 1770        |
| D      | Two passes route A | 8.5 ± 7.8    | -0.36        | 0.04           | 1322        |

in the microstructure of the material processed by route A rather than C. This is explained by the fact that route A developed grain substructures. Elongated and small grains were also detected in route A while a more homogeneous microstructure with equiaxed grains are noticed in route C.

According to the diagram of Fe-H₂O (diagram of Pourbaix-E-pH), it is concluded that the iron is passivated by Fe(OH)₃, for a pH of 6.5 [11]. As shown before (Figure 2a), the increase in the current density begins at the pitting potential (Epitting) and pitting corrosion occurs due to the rupture of the passive film. The formation of pitting in a carbon steel have been studied by Eletre et al. [11] in NaCl solution and the authors have proved that chloride ion leads to a sudden and noticeable increase in the current density, which promotes a destruction of the passive film and, consequently, a beginning of the corrosion by pitting. The same analysis was reported by Bi et al. [12]. In this work, the passive film was destroyed, and the 1010 steel surface was exposed to an active dissolution point. From Figure 2a, it is observed that the samples A and B have larger currents with a increasing in the potential, especially above -0.35 V. This means that the specimens C and D presents greater resistance to pitting comparing to the samples A and B.

Previous study by Hadzima et al. [8] evaluated the corrosion resistance of an IF steel processed by two passes of ECAP via route C in NaCl solution. This work confirmed that the Epitting increased from + 70 mV for the as cast up to + 250 mV for the deformed material. The authors confirmed an increase in the Epitting values between the imperfect passivity grain boundaries and the pitting region. This supports the observation that passive film is more stable for the materials with elongated grains than for those with coarse grains. As seen in Figure 2b, samples C and D have formed a passive film more stable when compared to A and B. Once again, this can be explained by the stability of the film, which is more stable for the deformed samples, due to the high surface energy created after ECAP processing.

It has been proved that in the alkaline environment the film basically corresponds to a double layer model, consisting of magnetite in the inner region and a ferric oxide in an external region, according to the Fe₃O₄/Fe₃O₃ structure [13]. Using the polarization curve showed in Figure 2b it was possible to obtain i_pass values. The values for the specimens A and B were measured at 0.08 mA cm⁻² and 0.07 mA cm⁻². For C and D the values were 0.03 mA cm⁻² and 0.04 mA cm⁻², respectively. The passivation of steel in the alkaline solution is due to the formation of a very thin, but highly protective oxide/hydroxide layer [14].

Some authors found EIS values different from this study [15,16]. This might occur due the fact that the initial conditions were different, despite the same equivalent circuit for data simulation were used. In summary, samples with higher deformation (C and D) presented better performance than the less deformed ones (A and B), as proved in the Nyquist plots (Figure 5). These are also in agreement with the results obtained by the polarization technique, as shown in the Figure 2. The summary of the main results, such as grain size and electrochemicals values are shown in Table 4.

5 Conclusion

From the results obtained in this work, it was possible to evaluate the corrosion behavior of the samples before and after the deformation in different environments. It can therefore be concluded that:

- Sample A presented very coarse grains compared to the samples processed by ECAP. Large elongation grains and fragmented structures was observed in the deformed samples;
- Higher values of Epitting were found for the deformed samples in C and D using NaCl solution (pH 6.5). Furthermore, higher current densities with the increase potential was found for the specimens A and B;
- It was observed a greater value of polarization resistance (immersion time of 48 hours), for the material processed by routes A and C using ECAP. This means a presence of a better resistance to the localized corrosion;
- In the alkaline sodium hydroxide/sodium phosphate pH 10.5 environment, the formed passive film is more stable for higher deformed samples - C and D (lower passivation current - i_pass).

Acknowledgements

The authors would like to thank CAPES (Coordenação de Aperfeiçoamento de Pessoal Ensino Superior) for their financial support.
References

1. Abdalla AJ, Hashimoto TM, Pereira MS, Anazawa RM. Formação da fase bainítica em aços de baixo carbono. Revista Brasileira de Aplicações de Vácuo. 2008;25(3):175-181.

2. Kawasaki M, Langdon TG. Producing ultrafine-grained materials through severe plastic deformation. Emerging Materials Research. 2014;3(6):252-260.

3. Azushima A, Kopp R, Korhonen A, Yang DY, Micari F, Lahoti GD, et al. Severe plastic deformation (SPD) processes for metals. CIRP Annals. 2008;57(2):716-735.

4. Valiev RZ, Langdon TG. Principles of equal-channel angular pressing as a processing tool for grain refinement. Progress in Materials Science. 2006;51(7):881-981.

5. Figueiredo RB, Pinheiro IP, Aguilar MTP, Modenesi PJ, Cetlin PR. The finite element analysis of equal channel angular pressing (ECAP) considering the strain path dependence of the work hardening of metals. Journal of Materials Processing Technology. 2006;180(3):30-36.

6. Dobatkin S, Zrnik J, Mamuzic I. Ultrafine-grained low carbon steels by severe plastic deformation. Metalurgija. 2008;47(3):181-186.

7. Shin DH, Kim BC, Kim YS, Park KT. Microstructural evolution in a commercial low carbon steel by equal channel angular pressing. Acta Materialia. 2000;48(9):2247-2255.

8. Hadzima B, Janeček M, Estrin Y, Kim HS. Microstructure and corrosion properties of ultrafine-grained interstitial free steel. Materials Science and Engineering A. 2007;462(1-2):243-247.

9. American Society for Testing and Materials. ASTM E112-13: standard test methods for determining average grain size. West Conshohocken: ASTM International; 2015.

10. Zrnik J, Mamuzić I, Dobatkin SV, Stejskal Z, Kraus L. Low carbon steel processed by equal channel angular warm pressing. Metalurgija. 2007;46(1):21-27.

11. Eletre AE, Abdallah M, Soliman MG, Mabrouk EM. Inhibition of pitting corrosion C-steel in NaCl solution by some inorganic compounds. Commun. Fac. Sci. Univ. Ank. Ser. B. 2000;46:25-31.

12. Bi H, Li Z, Liu J, Cheng Y, Toku-Gyamerah I. Study on pitting corrosion of storage tank bottom steel in acidic condition using acoustic emission. International Journal of Electrochemical Science. 2015;10:4416-4427.

13. Nagajama MI, Cohen M. The anodic oxidation of iron in a neutral solution. Journal of the Electrochemical Society. 1962;109(9):781-795.

14. Yang Y, Xiao L, Zhao Y, Wang F. Hydrothermal synthesis and electrochemical characterization of a-MnO2 nanorods as cathode material for lithium batteries. International Journal of Electrochemical Science. 2008;3(1):67-74.

15. Hu J, Cao SA, Xie J. EIS study on the corrosion behavior of rusted carbon steel in 3% NaCl solution. Anti-Corrosion Methods and Materials. 2013;60(2):100-105.

16. Nady H, El-Rabiei MM, Samy M. Corrosion behavior and electrochemical properties of carbon steel, commercial pure titanium, copper and copper-aluminium-nickel alloy in 3.5% sodium chloride containing sulfide ions. Egyptian Journal of Petroleum. 2017;26(1):79-94.

Received: 11 Apr. 2020
Accepted: 19 Oct. 2020