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CO2 corrosion resistance of carbon steel in relation with microstructure changes

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HIGHLIGHTS

- The effect of change in microstructure on CO2 corrosion resistance was evaluated.
- An API 5L X42 carbon steel was immersed in a 0.5 M NaCl solution saturated with CO2.
- Banded, normalized, quenched-tempered and annealed microstructures were considered.
- Electrochemical measurements were coupled with surface analysis.
- Morphology and distribution of undissolved Fe3C control corrosion kinetics.

ABSTRACT

The microstructural effects on the corrosion resistance of an API 5L X42 carbon steel in 0.5 M NaCl solution saturated with CO2 was investigated. Four microstructures were considered: banded (B), normalized (N), quenched and tempered (Q&T), and annealed (A). Electrochemical measurements (polarization curves and electrochemical impedance spectroscopy) were coupled with surface analyses (scanning electron microscope (SEM) and X-ray photoelectron spectroscopy (XPS)) to characterize the formation of the corrosion product layers. Electrochemical results revealed that corrosion resistance increased in the following order: B < N < Q&T < A. From the polarization curves it was shown that specifically, cathodic current densities were affected by microstructural changes. SEM images indicated that ferrite dissolved earlier than cementite and a thin layer of corrosion products was deposited on the steel surface. XPS analyses revealed that this layer was composed of a mixture of iron carbonate and non-dissolved cementite. It was also found that the quantity of FeCO3 content on the steel surface was greater for Q&T and A microstructures. These results, in agreement with the electrochemical data, indicate that the deposition mechanism of iron carbonate is closely related to the morphology of the non-dissolved cementite, determining the protective properties of the corrosion product layers.

1. Introduction

Carbon dioxide corrosion is an important degradation problem with major implications for oil and gas industries. Steel pipes in contact with production fluids are particularly susceptible to this kind of damage. Corrosion results from the dissolution of CO2 in the source water forming carbonic acid. Its aggressiveness is affected by interrelated parameters dependent on operational and metallurgical variables. The former are related to geological characteristics and operational conditions and the latter to the metal's characteristics and the processes used in its manufacture. Even though CO2 corrosion phenomena have been widely studied since the 70’s, only few works are devoted to investigating the influence of the steel’s microstructure on its corrosion resistance [1–10].

In early 2000, Lopez et al. [1] published a state-of-the-art appraisal concerning the influence of microstructure and the chemical composition of carbon and low alloy steels on CO2 corrosion. They found that metallurgical variables have a significant influence on the CO2 corrosion phenomena but concluded that it is
not possible to obtain general behavior patterns due to the complexity of the problem and the interrelations occurring among the various parameters involved in the occurrence of this phenomenon.

The chemical composition of carbon steel and/or the thermomechanical treatment applied during the manufacturing process determine its microstructure. These variables are not independent and the same microstructure can be obtained with different chemical compositions and/or heat treatments. Most previous works have been devoted to evaluating the effect of one parameter without taking into account that the others can also be modified. Thus, some authors have found that the corrosion resistance of carbon steels with a ferritic/pearlitic microstructure was higher than that reached with martensitic steels, whereas other studies have reported the opposite [1–4]. However, as pointed out by Lopez et al. [1], results from these studies are not comparable because the experimental conditions were different.

Although there is no general agreement as to how the microstructure affects the corrosion resistance of carbon steels, many authors consider that this parameter not only modifies the characteristics and the kinetics of corrosion products formation but also has a significant influence on the action of corrosion inhibitors. Some authors [2,3,5,11–16] have found that the steel microstructure plays an important role on the adherence of the corrosion products which mainly consist of iron carbonate (FeCO₃). Several authors have associated the protective properties of the carbonate films to the anchoring effect provided by iron carbide (Fe₃C). Indeed, during the exposure time, the ferritic phase dissolves whereas iron carbide (contained in the pearlite) are highlighted. This effect is a consequence of the galvanic coupling that occurs between these micro-constituents: ferrite acts as anodic site whereas cementite acts as a cathode [17,18]. However, the role of these two components on the stability and on the protective properties of the corrosion product layers is not well understood.

The aim of the present study is to reach a better understanding of the influence of the carbon steel microstructure on its corrosion resistance focusing on the formation of the corrosion product layers. For this purpose, heat treatments were performed on carbon steel API 5L X42 samples in order to modify the morphology and the distribution of the steel constituents while keeping the chemical decomposition constant. Afterward, corrosion tests were carried out in a CO₂ saturated brine using electrochemical measurements coupled with surface analyses in order to evaluate the effect of the steel microstructure.

### 2. Experimental

#### 2.1. Materials

API 5L X42 carbon steel samples were taken directly from a pipeline. Their chemical composition, determined by inductively coupled plasma – optical spectrometry (ICP-OES), is given in Table 1.

As received, the steel presents a banded microstructure that occurs between these micro-constituents: ferrite acts as anodic site whereas cementite acts as a cathode [17,18]. However, the role of these two components on the stability and on the protective properties of the corrosion product layers is not well understood.

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#### 2.2. Metallurgical characterization

The microstructures of the samples were characterized by optical microscopy (OM) following conventional metallographic procedures specified in ASTM E 3-01 and E 407-99 standards [19,20]. The samples were etched with 5% Nitral solution during 10 s. Phase fractions were determined using Image®J analysis software. The mechanical properties of the heat-treated steels were tested by Rockwell B measurements correlated to the ultimate tensile stress (UTS) using conversion tables presented in the ASTM E140-97 standard [21].

#### 2.3. Electrochemical measurements

A standard Pyrex three-electrode cell was used with a platinum grid as counter electrode, a saturated calomel electrode (SCE) as reference and rods of the steel as rotating disk electrodes. Working electrode rods consisted of cylinders 7 mm of diameter which were covered with a heat shrinkable sheath in order to leave only the tip of the steel electrodes in contact with the corrosive medium.

Prior to electrochemical measurements, the electrodes were ground with SiC papers down to grade 1200. Afterwards, they were cleaned with ethanol, rinsed with distilled water and dried in warm air. The electrode rotation rate was fixed at 200 rpm. The corrosive medium was a 0.5 M solution of NaCl (reagent grade), initially purged with nitrogen for 30 min and then CO₂ was bubbled through for 1 h before introduction of the sample into the cell. CO₂ bubbling was maintained during the experiment to limit the ingress of air. Experiments were carried out at atmospheric pressure and room temperature. The pH of the corrosive solution was around 3.5 and did not change during the experiments.

Polarization curves and electrochemical impedance diagrams were obtained after 6 h of immersion at the corrosion potential. Current-voltage curves were obtained under potentiodynamic regulation using a Solartron 1286 electrochemical interface. They were plotted from the cathodic to the anodic range, from −1 V/SCE to −0.4 V/SCE. The potential sweep rate was fixed at 10 mV min⁻¹.

Electrochemical impedance measurements were carried out using a Solartron 1286 electrochemical interface connected to a Solartron 1250 frequency response analyzer. Impedance diagrams were obtained under potentiostatic regulation, at the corrosion potential, over a frequency range of 65 kHz to 7 mHz with 8 points per decade, using a 8 mV peak-to-peak sinusoidal voltage. Three experiments were carried out to ensure reproducibility.

#### 2.4. Surface analyses

The morphology of the corroded steel electrodes was observed using a LEO 435 VP scanning electron microscope operating at

| Table 1: Chemical composition of API 5LX 42 carbon steel. |
|-----------------|---------|----------------|---------|---------|---------|---------|---------|---------|---------|-----------|
| C    | Si   | Mn  | P     | Cu    | Nb    | V      | Ti     | Fe     | Balance |
| 0.211 | 0.263 | 0.768 | 0.011 | 0.086  | 0.002 | 0.004  | 0.003  |        |           |

| Table 2: Characteristics of the heat treatments applied to the carbon steel samples. |
|-----------------|-----------------|
| Heat treatment | Characteristics |
| Normalized (N)  | Austenitizing at 1350 °C for 10 min followed by air cooling |
| Quenched and tempered (Q&T) | Austenitizing at 900 °C for 20 min followed by quenching in water. The steel was tempered at 650 °C for 60 min |
| Annealed (A)    | Heating at 710 °C for 24 h followed by furnace cooling |

lamellar aggregate of ferrite (z) and of cementite (Fe₃C) which results from the eutectoid decomposition of high temperature austenite. Four microstructures were thus considered: banded (B), normalized (N), quenched and tempered (Q&T), and annealed (A).
15 kV. The chemical composition of the corroded electrodes was determined using x-ray photoelectron spectroscopy (XPS) carried out on a VG Escalab 220i-XL spectrometer. For the surface analyses, steel samples were prepared by the same procedure as for the electrochemical experiments. After 6 h of immersion, the electrodes were removed from the solution after anodic polarization. They were rinsed with ethanol and dried in warm air before being placed in a fast-entry pre-chamber and then in the analysis chamber. The specimens were irradiated using a Mg Kα X-ray source with a power of 300 W. Measurements were made at a take-off angle \( \theta = 90^\circ \). The experimental resolution of the binding energy was 1 eV. After background subtraction, peak deconvolution was performed using XPSGRAF102 software. Binding energies were corrected for possible charging effects by referencing to the C1s (284.5 eV) peak. The chemical composition of the steel surfaces was analyzed monitoring the C1s, O1s and Fe 2p\(_{3/2}\) signals.

3. Results

3.1. Metallurgical characterization

Representative micrographs of the different steel microstructures are presented in Fig. 1. The microstructures present are seen to depend on the heat treatment applied with light and dark zones which have different distributions and morphologies. It can be noted that the banded microstructure of the reference sample disappeared after the heat treatments. The banded (B) microstructure is composed of alternating light and dark bands corresponding to equiaxed proeutectoid ferrite grains and coarse pearlite, respectively [22]. This microstructure results from a slow cooling rate from the austenitizing temperature and is the manifestation of the segregation of alloying elements during the steel solidification process [23,24]. Light and dark areas observed on the normalized (N) microstructure correspond to ferrite and fine pearlite, respectively [22]. In this case, proeutectoid ferrite located at the grain boundaries of a coarse former austenite grain has a polygonal morphology; ferrite situated within the grains grows as Widmanstätten ferrite plates [22]. This microstructure is frequently obtained when a short, high temperature heat treatment is performed on a low carbon steel [25,26]. The quenched and tempered (Q&T) microstructure consists of dark precipitates randomly dispersed in a light matrix which corresponds to isolated cementite globules and a ferritic matrix, respectively [22]. Globular cementite results from the transformation of martensite to ferrite and carbides [25]. The annealed (A) microstructure consists of light and dark grains whose sizes are relatively uniform and correspond to proeutectoid ferrite and colonies of pearlite, respectively [22]. In addition, coarse pearlite is also observed at ferrite grain boundaries. The long sub-critical annealing applied in this study leads to a redistribution of the cementite (Fe₃C) in pseudo-spherical pearlite agglomerates [22,26]. Volume fractions determined for the different steel microstructures revealed that all samples contain ferrite as major component while the pearlite was slightly different depending on the steel microstructures: B (25.9%), N (27.6%), Q&T (27.5%), A (26.3%). In the pearlitic fraction, the amount of ferrite was much larger than the amount of cementite. If the pearlitic fraction is considered constant and equal to 27% for all microstructures, the total Fe₃C quantity can be estimated from the Fe–Fe₃C equilibrium phase diagram using the lever rule at the eutectoid temperature. This implies a volumetric pearlitic fraction near to 3% but the exact percentage is slightly modified if we consider the effect of other alloying elements.

Correlations of the Rockwell B hardness measurements made on different samples to the ultimate tensile stress (UTS) indicated that the UTS values are between 579 and 482 MPa after the different heat treatments [21]. It can be mentioned that 414 MPa corresponds to the minimum UTS value specified in the API 5L Standard for X42 grade carbon steel pipes [27]. From an industrial point of view, the steel samples evaluated in this study fulfill the mechanical requirements specified in the API 5L standard.

Fig. 1. Optical micrographs of the different API 5LX 42 steel microstructures: (B) Banded, (N) Normalized, (Q&T) Quenched and Tempered and (A) Annealed.
an active electrodes was plotted as a function of immersion time (Fig. 2). It can be observed that $E_{\text{corr}}$ shifted slightly in the positive direction and then it slowly decreased to remain relatively constant at around $-0.670$ V/SCE, indicating that the electrochemical system reached a steady state after 6 h of immersion.

Electrochemical impedance diagrams obtained at the corrosion potential after 6 h of immersion in a 0.5 M NaCl solution saturated with CO$_2$ for the different steel microstructures.

3.2. Electrochemical characterization

The variation of the corrosion potential, $E_{\text{corr}}$, of the carbon steel electrodes was plotted as a function of immersion time (Fig. 2). It can be observed that $E_{\text{corr}}$ is independent of the steel microstructure: after 1 h of immersion, $E_{\text{corr}}$ shifted slightly in the positive direction and then it slowly decreased to remain relatively constant at around $-0.670$ V/SCE, indicating that the electrochemical system reached a steady state after 6 h of immersion.

Electrochemical impedance measurements were carried out at $E_{\text{corr}}$ for the different samples (Fig. 3). The spectra are characterized by two time constants: a capacitive loop in the middle frequency range and an inductive loop in the low frequency domain. The shape of the diagram in the low frequency domain is not exactly the same for all the spectra. For the Q&T and for the A microstructures, the inductive loop tends to close on itself giving rise to another capacitive loop. In previous works [17,18,30], the medium frequency loops are attributed to the charge transfer process on the ferritic surface that freely dissolves into a faulty layer of siderite (FeCO$_3$) as corrosion product. In this case, the charge transfer resistance ($R_T$) would also reflect differences in the uncorroded cementite (Fe$_3$C) area, revealed after 6 h of immersion for the different microstructures. Inductive behavior is generally associated with the existence of relaxation processes of adsorbed species at the metal/solution interface [27–31] and closure of the inductive loop is attributed to an active–passive transition [31,32].

Equivalent electrical circuits are frequently used to extract the parameters associated with the impedance diagrams where a constant phase element (CPE) is used instead of a capacitance to take the non–ideal behavior of the interface into account. The CPE is given by:

$$Z_{\text{CPE}} = \frac{1}{Q(j\omega)^\alpha}$$

where $\alpha$ is related to the angle of rotation of a purely capacitive line on the complex plane plots and $Q$ is a constant expressed in $\Omega^{-1}$ cm$^{-2}$ s$^\alpha$. In the present study, parameters were graphically obtained: the electrolyte resistance ($R_e$) and the resistance associated with the capacitive loop ($R_T$), physically related to the corrosion resistance, were directly measured on the impedance spectra; $\alpha$ and $Q$ were determined using the graphical method proposed by Orazem et al. [18]. The effective capacitance $C_{\text{eff}}$ (expressed in Farads), was calculated from the CPE parameters considering a distribution of the charge transfer resistances on the electrode surface [29]:

$$C_{\text{eff}} = \frac{Q^1/\alpha(R_eR_T/(R_e + R_T))^{(1-\alpha)/\alpha}}$$

The impedance parameters are reported in Table 3. It can be seen that the $R_e$ value depends on the heat treatment and increases in the following order: $B < N < $ Q&T < A. On the other hand, the capacitance values were higher than 50 $\mu$F cm$^{-2}$ and the product $R_T C_{\text{eff}}$ was not constant; therefore, the capacitive loops are thought to result from both a charge transfer process occurring on the ferritic phase, that freely dissolves and a film effect due to the corrosion products that precipitate on the steel surface. Thus, the impedance results indicated that the CO$_2$ corrosion resistance of the API 5L X42 steel was sensitive to the heat treatments performed. Furthermore, the shape of the impedance spectra in the low frequency range indicates different corrosion mechanisms when B/N and Q&T/A microstructures are compared.

To evaluate the influence of the microstructure on the electrode kinetics, polarization curves were obtained for the different heat

![Fig. 2.](image1.png)  
**Fig. 2.** Corrosion potential versus immersion time in a 0.5 M NaCl solution saturated with CO$_2$ for the different steel microstructures.

![Fig. 3.](image2.png)  
**Fig. 3.** Electrochemical impedance diagrams obtained at the corrosion potential after 6 h of immersion in a 0.5 M NaCl solution saturated with CO$_2$ for the different steel microstructures.

![Fig. 4.](image3.png)  
**Fig. 4.** Current-voltage curves obtained after 6 h of immersion in a 0.5 M NaCl solution saturated with CO$_2$ for the different steel microstructures.
treated steels after 6 h of immersion (Fig. 4). It can be observed that the shape of the current–voltage curves is similar. In the anodic domain, a Tafelian part for the low anodic overvoltage is observed. The anodic Tafel slopes are independent of the steel microstructure with values varying between 90 and 94 mV dec⁻¹. Besides, around −0.54 V/SCE, the anodic current densities systematically decreased recalling a pseudo-passivation plateau; nevertheless, current densities increase again for higher overvoltages. For the cathodic domain, a well-defined current plateau is observed which is ascribed to the mass transport of the hydrogen reduction reaction [33]. The values of the cathodic current densities on the plateau (i_{lim}) are reported in Table 4. The values decrease in the following order i_{lim~B} > i_{lim~N} > i_{lim~Q&T} > i_{lim~A}, in agreement with the increase of the R_T values obtained from the impedance spectra. A decrease of i_{lim} of about 30% is observed between the banded and the annealed samples. This result was reproducible and indicates that the heat treatment performed on the steel mainly affects the cathodic

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<td>153</td>
<td>122</td>
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<td>1.9</td>
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Table 4: Electrochemical parameters involved in the calculation of the corrosion rate for the API SLX 42 steel with different microstructures.

Fig. 5. SEM observations of the corroded steel surfaces after a preliminary hold time of 6 h in a 0.5 M NaCl solution saturated with CO₂; left and right images were obtained for two magnifications.

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reaction. Modification of the diffusion current densities could be linked to the precipitation of non-soluble corrosion products such as siderite that limits the cathodic reaction decreasing the surface area available for the reaction. This result is in agreement with the impedance data.

The corrosion rates (CR) of the steel with the different microstructures were calculated from the electrochemical parameters (Table 4) according to the ASTM G 102-04 Standard [34] with the following equation: \( CR = (K \times \text{corr} \times EW)/\rho \), where \( K \) is a constant \( 3.27 \times 10^{-3} \text{ mm g } \mu\text{A}^{-1} \text{ cm year} \), \( \rho \) is the density of iron \( (7.86 \text{ g cm}^{-3}) \) and \( EW \) is the equivalent weight of the carbon steel estimated at 28. Corrosion current densities \( (\text{corr}) \) were determined using the simplified Stern and Geary equation \( (\text{corr} = \beta_i/2.3 R_I) \) since it was shown that the cathodic reaction is under diffusion control [35]. The \( R_I \) values were obtained from the impedance spectra (Table 3) and anodic Tafel slopes \( (\beta_a) \) were measured from the polarization curves. The anodic Tafel slope value used for the calculation was 90 mV dec\(^{-1} \) for all microstructures. As expected, the corrosion rate decreased in the following order: B > N > Q&T > A. The corrosion rate of the steel with an annealed microstructure is about 35% lower than that of the banded steel but it is predictable that this difference will be greater for longer exposure times. This result is in agreement with those obtained by Al-Hassan et al. [5] who used weight loss measurements to study the effect of the microstructure on the corrosion of steels in aqueous solutions containing carbon dioxide. In their study, longer immersion periods (>400 h) were necessary to show differences in the corrosion resistance of the carbon steels with different microstructures; in our study, the same result was obtained after only 6 h.

3.3. Surface analyses

SEM images of corroded carbon steel surfaces after 6 h of immersion in the 0.5 M NaCl solution saturated with CO\(_2\) can be seen in Fig. 5. The features of light and dark zones imitate the steel microstructures presented in Fig. 1. Higher magnification images (right panel) reveal that the dark areas correspond to the selective dissolution of the ferritic matrix. The lamellar or globular morphologies of the light zones correspond to the non-dissolved cementite contained in the pearlitic structure. Selective dissolution of the ferritic phase with respect to cementite indicates that microgalvanic cells are formed between these constituents influencing the kinetics of the corrosion process. SEM images revealed that ferrite is the anode while iron carbide, contained in the eutectoid mixture, is the cathode, in agreement with its lower electrical conductivity [36]. Recently, Farelas et al. [17] confirmed that cementite forms preferential cathodic sites with lower overpotential favoring hydrogen evolution. These results are also in agreement with previous works reported in the literature [11,37]. In addition, it can be seen that the cementite surface area increases when selective dissolution of ferrite takes place and thus the galvanic coupling between the iron carbides and the ferritic matrix is enhanced. A very thin whitish layer can be observed on cementite, which is more visible on the lamellar structure of banded (B) and normalized (N) samples. For these two microstructures, it seems that a very thin layer of corrosion products forms on the outer part of the remaining cementite structure far from the metal substrate; conversely, in the case of quenched and tempered (Q&T) and annealed (A) microstructures, this layer could precipitate near the metal surface due to the different morphology of the iron carbide. This feature can be observed on the cross-sectional views of the N and Q&T samples (Fig. 6).

The chemical composition of the corroded steel surfaces was determined by XPS analysis. Fig. 7 shows a comparison of the high resolution spectra corresponding to C 1s, O 1s and Fe 2p regions obtained from the corroded steel surfaces with different microstructures. Deconvolution of C 1s region reveals peaks at 283.6, 284.6, 286.3, and 288.7 eV which correspond to binding energies of carbon in cementite (Fe\(_3\)C), adventitious species like graphitic carbon and ethanol (\( \text{CH}_2\text{CH}_2\text{OH} \)), and iron carbonate (Fe\(_3\)CO\(_3\)), respectively [38–40]. Peaks obtained at 528.2, 530.2, and 531.7 eV from deconvolution of the O 1s spectra correspond to the binding energies of \( \text{O}^2^- \), \( \text{OH}^- \), and \( \text{CO}_2^- \) ions in hematite (Fe\(_2\)O\(_3\)), iron oxy-hydroxide (FeOOH) and iron carbonate, respectively [3,37,38]. These signals are in agreement with those obtained from the Fe 2p\(_{3/2}\) spectra. Although the Fe signals are inherently less intense and more surface sensitive, deconvolution of iron spectra reveals different signals according to the steel microstructure. In the case of the banded (B) and normalized (N) samples, only a signal at ~711 eV was observed due to the binding energy of iron in Fe\(_2\)O\(_3\). Additionally, quenched and tempered (Q&T) and annealed (A) samples exhibited a signal at ~712.2 eV corresponding to iron carbonate. Iron oxy-hydroxide signals could be related to the partial decomposition of iron carbonate when CO\(_2\) corroded samples were exposed to air before being placed in the analysis chamber. According to several authors [39,40], iron carbonate decomposed into \( \alpha\text{-Fe}_2\text{O}_3 + \text{CO}_2 \) in air at temperatures below 100 °C. This could explain why the metallic iron signal at about 706.6 eV was not detected. However, the amount of iron carbonate at the metal surface detected by XPS analyses was higher for the less corrosion-prone samples (Q&T and A) and diminished with decreased corrosion resistance (N and B).

4. Discussion

The electrochemical results presented here show that the CO\(_2\) corrosion resistance of the API 5L X42 steel depends on its
microstructure. From the electrochemical measurements, it was found that the Q&T and A samples show the lowest corrosion rates compared with the B and N microstructures. Polarization curves revealed that this result was mainly linked to the differences in the cathodic reduction reaction. SEM images showed that when the corrosion process took place, the morphology of the undissolved cementite was revealed by the selective dissolution of the ferritic phase. Additionally, a very thin whitish layer deposits on the metal surface which is more or less continuous depending on the steel microstructure. From XPS analyses, the presence of FeCO3 on the steel surfaces was more abundant on Q&T and A microstructures. Joint analysis of electrochemical and surface characterizations indicated that the corrosion resistance of the API 5L X42 steel, is closely related to the morphology of non-corroded iron carbide which controls the iron carbonate deposition. It is determined by the heat treatment performed on the steel samples which also establishes the characteristics of the microgalvanic coupling between ferrite and cementite. When corrosion develops, the galvanic coupling could be accentuated by the precipitation of siderite on the steel surface.

Our results suggest that when the ferrite and pearlite areas were better differentiated, the strength of the galvanic coupling between ferrite and cementite increased. Thus, as the corrosion process evolves, a critical concentration of ferrous cations at the metal/solution interphase was reached; as a consequence, iron carbonate precipitated partially covering the steel surface. Precipitation of siderite is also favored by a local increase of the pH when the hydrogen evolution occurs on the cathodic sites. For steels with banded (B) and normalized (N) microstructures, iron carbonate forms on the outer part of the remaining cementite structure far from the metal substrate. This feature leads to the internal acidification between the non-corroded lamellae of Fe3C that further prevents precipitation of FeCO3 directly on the metal surface. In these cases, the resulting layers are non-protective and the CO2 corrosion resistance of the steel is low. Conversely, for samples with Q&T and A microstructures, the morphology and distribution of the undissolved cementite favor the formation of FeCO3 directly in contact with the metal substrate. In these cases, corrosion product layers are more continuous and protective; this explains the highest RT values, the lower effective capacitance, and the lowest cathodic current densities obtained for these samples.

5. Conclusions

Electrochemical measurements carried out after 6 h of immersion in a 0.5 M NaCl solution saturated with CO2 revealed that the corrosion resistance of API 5L X42 steel was dependent on the heat treatments performed. It was found that the corrosion rate decreased as a function of the microstructure in the following order: banded (B) > normalized (N) > quenched and tempered (Q&T) > annealed (A). This behavior was explained by the differences between the distribution and morphology of the non-corroded cementite which controls the intensity of the galvanic coupling between ferritic and pearlitic steel constituents and the formation of iron carbonate layers at long term.

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