Indirect Galvanostatic Pulse in Wenner Configuration: Numerical Insights into Its Physical Aspect and Its Ability to Locate Highly Corroding Areas in Macrocell Corrosion of Steel in Concrete

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Abstract: The use of indirect electrical techniques is gaining interest for monitoring the corrosion of steel in concrete as they do not require any connection to the rebar. In this paper, we provide insights into the physical aspects of the indirect galvanostatic pulse (GP) method in the Wenner configuration. Considering uniform corrosion, the instantaneous ohmic drop is decreased due to the presence of the rebar, which acts as a short-circuit. However, we observed that this phenomenon is independent of the electrochemical parameters of the Butler–Volmer equation. They are, however, responsible for the nonlinear decrease of the current that polarizes the rebar over time, especially for a passive rebar due to its high polarization resistance. This evolution of the resulting potential difference with time is explained by the increase of the potential difference related to concrete resistance and the global decrease of the potential difference related to the polarization resistance of the rebar. The indirect GP technique is then fundamentally different than the conventional one in three-electrode configuration, as here the steady-state potential is not only representative of polarization resistance but also of concrete resistance. Considering non-uniform corrosion, the presence of a small anodic area disturbs the current distribution in the material. This is essentially due to the different capability of anodic and cathodic areas to consume the impressed current, resulting in slowing down the evolution of the transient potential as compared to uniform corrosion. Hence, highly corroding areas have a greater effect on the transient potential than on the steady-state one. The use of this temporal evolution is thus recommended to qualitatively detect anodic areas. For the estimation of their length and position, which is one of the main current problematic issue when performing any measurement on reinforced concrete (RC) structures with conventional techniques, we suggest adjusting the probe spacing to modulate the sensitivity of the technique.

Keywords: reinforced concrete; indirect galvanostatic pulse; corrosion monitoring; finite-element modeling; uniform corrosion; non-uniform corrosion

1. Introduction

Reinforced concrete (RC) is a durable material that allows the construction of safe and strong structures. Concrete has a high compressive strength and acts as a protective barrier to the steel reinforcement bar (rebar), which provides tensile strength and ductility to the structures. However, RC degrades over time, the main cause being corrosion of the steel rebars [1,2]. This is an electrochemical process that involves the anodic oxidation of iron and, generally, the cathodic reduction of
oxygen, whereby the concrete pore solution acts as electrolyte to close the electrical circuit. Corrosion is
generally triggered by the ingress of aggressive agents—mainly CO$_2$ and Cl$^-$—up to the rebar,
which degrade the protective layer formed on steel surface in sound and alkaline concrete [2,3].
Due to the inherent heterogeneity of concrete, all transport phenomena within the material are
globally non-uniform. Hence, irrespective of the depassivation mechanism, non-uniform corrosion
is expected to be the main process due to local variations in environmental exposure, or to the
presence of interconnected rebars with different properties [4–6]. The volume expansion of the
corrosion products can result in concrete cracking, spalling, and delamination, which progressively
decreases the performance of the structures and can lead to their collapse. To prevent such disasters,
we need monitoring devices that can detect an early stage of corrosion without altering the integrity of
the structures.

Due to the electrochemical nature of corrosion, several electrical methods have been developed for
monitoring the corrosion rate [2]. Conventional techniques are based on a three-electrode configuration
that require a connection to the rebar. They apply an external perturbation to the system for polarizing
the rebar, generally close to its natural steady state, to determine a polarization resistance $R_p$ that
can be converted into corrosion current density $i_{corr}$ and then into corrosion rate in terms of loss of
steel cross-section (generally expressed in $\mu$m year$^{-1}$). The main techniques are the linear polarization
resistance (LPR), the Tafel extrapolation, the galvanostatic pulse (GP) method, and the electrochemical
impedance spectroscopy (EIS). All these techniques provide an accurate estimate of the corrosion rate
of actively corroding rebar under laboratory conditions [7–9].

However, some drawbacks affect their application to RC structures. First, determining the
 corrosion current density from polarization resistance is still mainly based on using the Stern–Geary equation (Equation (1)):

$$i_{corr} = \frac{B}{A \times R_p} = \frac{\beta_a \beta_c}{\ln(10) \times (\beta_a + \beta_c)} \times \frac{1}{A \times R_p},$$

where $i_{corr}$ is the corrosion current density ($A \cdot m^{-2}$), $B$ is the Stern–Geary constant (V), $A$ is the polarized
area of steel ($m^2$), $R_p$ is the polarization resistance ($\Omega$), and $\beta_a$ and $\beta_c$ are the experimentally obtained
anodic and cathodic Tafel constants (V), respectively. However, this equation was defined for uniform
corrosion [10] and is fundamentally non-applicable to non-uniform corrosion, which is expected in RC
structures [11]. Recently, a new perspective has been proposed for calculating the rate of localized
corrosion from GP measurements. This considers the temporal evolution of the current in anodic
areas, which reduces the error related to the use of the Stern–Geary equation [12]. Thus, this technique
is robust to determine the corrosion current. However, a limitation still exists and concerns the
determination of the polarized area $A$ during the measurement, as it is required for calculating the
corrosion current density. Though the use of a guard ring with modulation has been proposed for
confining the polarization to a known length of rebar, many studies have shown that it often fails to
achieve this goal and even contributes to additional polarization of the rebar, thus compromising the
measurement [13–19]. Finally, the main drawback of the different techniques mentioned above is the
need of an electrical connection to the rebar, often requiring breakout of local parts of the concrete to
make the measurement.

In the late 1990s, Monteiro et al. proposed the use of a non-destructive method involving indirect
polarization of the rebar, without any connection to it, for monitoring the corrosion of steel in RC
structures. This was achieved by making indirect electrochemical impedance spectroscopy (EIS)
measurements in a four-electrode configuration directly on the concrete surface [20–25]. This technique
applies an alternating current between two electrodes and measures the resulting potential difference
between the other two electrodes, all electrodes being generally placed in Wenner configuration,
i.e., with the same probe spacing between each four electrodes. It was shown that the results obtained
with this configuration were qualitatively similar to those obtained using the conventional EIS technique,
with a similar capacitive loop at low frequencies [26–32]. In addition to this indirect EIS technique,
it was also proposed that an indirect GP technique can also be used to determine the corrosion rate of steel embedded in concrete using a similar four-electrode configuration [33,34]. The main difference with the EIS technique is that a direct current is applied in GP rather than an alternating current. As for the conventional GP technique in three-electrode configuration, it was proposed to use an electrical equivalent circuit (EEC) to fit the results in order to extract the values of the EEC component and then calculate the corrosion rate [20,28,30,31,34,35]. By using an appropriate EEC, Fahim et al. have obtained similar corrosion rate values when using conventional electrochemical techniques and their patented connectionless electrical pulse (CEPRA) technology, which typically apply a current in the range of 0.5 to 2 mA on concrete surface [9,35].

Despite their great practical interest, these indirect techniques have been largely ignored in recent reviews devoted to corrosion monitoring. This lack of consideration could be related to their complexity, which can hardly be elucidated only experimentally. However, in recent years, numerical simulations have gained much interest as they can provide further insight into the mechanisms of macrocell corrosion [6,36–41] and the corrosion monitoring in three-electrode [18,42] and four-electrode configurations [26,27,31,32,35,43]. For the latter case, Keddam et al. showed that the current distribution in the material is dependent upon the measurement frequency; the steel/concrete interface behaves essentially as a capacitance at high frequencies, whereas it acts mainly as an insulator at low (zero)-frequency [26,27]. Fahim et al. investigated numerically the effect of concrete resistivity, cover depth, and rebar diameter in uniform corrosion and showed that an increase in concrete resistivity, a decrease in cover depth and a decrease in rebar diameter result in an increase in the total current reaching the rebar [35].

As shown by Fahim et al., one advantage of the indirect polarization in Wenner configuration is its ability to naturally confine the polarization current to a well-defined and similar length of the rebar, irrespective of concrete resistivity, cover thickness, and rebar diameter [35,43]. Thus, it appears possible to more precisely estimate the polarized area than with the three-electrode configuration because the guard ring is not adapted, which is required for converting the corrosion current in effective corrosion rate. However, the authors did not systematically analyze the evolution of the potential response with time and did not address the sensitivity of the potential probes, which can be different than the polarized area. Alexander and Orazem have shown the ability of the indirect EIS technique to locate highly corroding areas of external post-tensioned tendons in a macrocell system using Wenner and Wenner-Schlumberger configurations [31,32]. They showed that the presence of an anodic area disturbs the signal, depending on its position relative to the monitoring device, as two distinct time constants can be observed for non-uniform corrosion, whereas only one time constant is observed in the case of uniform corrosion. This effect essentially occurs when the anode is located below one of the current-injecting electrodes. The sensitivity of the technique depends upon the length of the actively corroding area, the magnitude of corrosion rate, and the resistivity of the cement-based material [32]. However, they believe that methods based on the DC-limit response may not be adapted for detecting localized corrosion, as the value obtained at low-frequencies in non-uniform corrosion is very similar to the one observed in uniform corrosion.

The objective of this preliminary study was to provide insights into the ability of indirect GP for locating highly corroding areas in non-uniform rebar corrosion, as well as to determine the sensitivity of this technique. In this paper, we first provide experimental evidences on the ability of the technique for indirect polarization of the rebar and corrosion monitoring. We then provide numerical insights on the physical aspects of this technique, considering both uniform and non-uniform corrosion.

2. Materials and Methods

2.1. Experiments

Experiments were done on a CEM I 52.5R (Lafarge, France) mortar sample (≃55 × 34 × 13 cm³, w/c = 0.4) with one ribbed carbon steel horizontal rebar (L = 60 cm, Φ = 12 mm) embedded at a cover depth of 33 mm. The sample was air-dried cured to be more representative of conditions encountered
on field. Measurements were first done on the passive rebar and after introducing Cl\(^-\) (C = 0.5 M) by the upper part of the sample through wetting (2 days)/drying (5 days) cycles for 3 months. The sample was then allowed to equilibrate back to the laboratory environment for 1 month, so all measurements were done in comparable conditions, at constant relative humidity (\(\approx 30\)–\(40\)\% RH) and room temperature (\(\approx 22 \, ^\circ\text{C}\)).

A potentiostat-galvanostat (Princeton Applied Research PMC-2000, HTDS, Massy, France) was used to perform electrical measurements with VersaStudio software (version 2.59). For the indirect GP technique, a device with 10 electrodes made in stainless-steel or brass electrodes (\(\Phi_t = 3 \, \text{mm}, \Phi_e = 5 \, \text{mm}\)), with a probe spacing of \(a = 5 \, \text{cm}\), was placed on concrete surface and parallel to the rebar (Figure 1a). A spring present in each electrode allows a good physical contact with concrete even if irregularities are present on its surface. The electrolytic contact was made with sponges wet beforehand with a highly conductive solution (0.5 M NaCl) and placed inside the probes.

Measurements were done in Wenner configuration, in which the two external probes are the current probes and the two internal probes are the potential probes. All the results represent the average value obtained by 10 consecutive measurements when the self-potential variation was small (Figure 1c). The impressed current between \(C_1\) and \(C_2\) varied between 50 and 1000 µA, while the probe spacing varies between 5 and 15 cm. It was not possible to use properly a higher sampling rate due to the presence of some filters in the potentiostat-galvanostat. Hence, the instantaneous ohmic drop was here evaluated at 0.1 s (10 Hz), so errors are expected in the calculation of the apparent concrete resistivity.

Supplementary experiments were also done and are presented in Figure S1. Measurements were done in the conventional three-electrode configuration using the GP technique to compare qualitatively the results obtained with the two techniques. A mercury/mercurous sulphate located at the center of the sample was used as reference electrode, and a stainless-steel counter electrode (55 \times 10 \, \text{cm}^2) was...
placed on moist sponges on the sample surface directly above the rebar. The impressed current was 10 \( \mu \)A between the rebar and the counter electrode. Experiments were also done with the indirect EIS technique, from 10 to \( \approx 10^{-2} \text{--} 10^{-3} \) Hz with 10 points per decade and an AC perturbation of 200 \( \mu \)A, using a similar device.

2.2. Simulations

All finite-element simulations used COMSOL Multiphysics software (version 5.3). The 3D dimension of the concrete domain was \( 200 \times 60 \times 13 \) cm\(^3\). This length was selected in order to observe the supposed self-confinement of the current \([35,43]\) when varying the probe spacing, which was not possible using the same size as the experimental sample. The first numerical simulations used the AC/DC module on an unreinforced specimen in order to determine the geometric factor for considering the finite size of the element. The secondary-current distribution module was then selected for modeling the distribution of the current in the entire domain (concrete and steel/concrete interface). The rebar was modeled as a 2-m carbon steel cylinder (diameter \( \Phi = 6 \text{--} 25 \) mm, electrical conductivity \( \sigma_{\text{steel}} = 10^7 \text{ S m}^{-1} \)) centered in the xy-plane (Figure 2).

![Geometry and mesh of the finite-element model used in this study (200 × 60 × 13 cm\(^3\)).](image)

The two outer electrodes \( C_1 \) and \( C_2 \) act as current-injection electrodes, while the two inner electrodes \( P_1 \) and \( P_2 \) act as potential electrodes. Probe spacing \( \delta \) varies between 2.5 and 15 cm. The steel-concrete interface was modeled with a parallel contribution of the exchange current density and the double-layer capacitance using the Butler–Volmer equation.

The concrete cover depth varied between 20 and 60 mm in uniform corrosion, whereas it was fixed at 40 mm in non-uniform corrosion. The concrete domain acts as the electrolyte with uniform resistivity \( \rho \) and electrical isolation at the external boundaries of the domain. Concrete resistivity varies between 100 and 1000 \( \Omega \) m to consider different binder types, \( w/c \) ratios (concrete quality), and degree of water saturation \([44]\). If the complexity of concrete, such as its porosity or the distribution of coarse aggregates, is not considered, the concrete is commonly simplified to a resistance with the use of the Randles circuit in three-electrode configuration. Here, it is simplified as one global resistance that considers the resistance of the continuously and discontinuously connected porosity but neglects all capacitive elements of the concrete matrix \([45]\). This simplification is possible as the capacitive loops associated to concrete are mainly observed at high frequencies (>10 kHz), while we only focused on the low-frequency response of the steel/concrete interface when the reactance and phase shift associated to concrete are generally small (<50 mrad) \([46,47]\).

In the concrete volume, Ohm’s law (Equation (2)) and charge conservation (Equation (3)) govern the electrical phenomena:

\[
i = \frac{1}{\rho} \nabla E, \tag{2}
\]
where \( i \) is the local current density (A m\(^{-2}\)), \( \rho \) is the concrete resistivity (\( \Omega \) m, reciprocal of electrical conductivity \( \sigma \)), and \( E \) is the electrical potential (V). The corrosion reaction was modeled as a boundary condition at the steel/concrete interface using the general Butler–Volmer equation in COMSOL (Equation (4)):

\[
\dot{i}_{\text{tot}} = i_0 \left\{ \exp \left[ \frac{\alpha_a F}{RT} (E - E_{\text{eq}}) \right] - \exp \left[ - \frac{\alpha_c F}{RT} (E - E_{\text{eq}}) \right] \right\},
\]

where \( i \) is the total current density (A m\(^{-2}\)), \( i_0 \) is the exchange current density or corrosion rate (A m\(^{-2}\)), \( \alpha_a \) and \( \alpha_c \) are the anodic and cathodic charge transfer coefficients, respectively, \( F \) is the Faraday constant (96,485 C mol\(^{-1}\)), \( R \) is the universal gas constant (8.314 J mol\(^{-1}\) K\(^{-1}\)), \( T \) is the absolute temperature (K), \( E \) is the rebar potential (V), and \( E_{\text{eq}} \) is the equilibrium potential (V). Three main parameters are thus required for modeling the corrosion of steel in concrete: the exchange current density \( i_0 \), the Tafel slopes (i.e., the charge transfer coefficients \( \alpha_a \) and \( \alpha_c \)), and the rebar potential \( E \) [48,49].

The oxygen concentration was not implemented in the Butler–Volmer equation as its reduction is not the only possible cathodic reaction as water (or protons, depending on the pH) can also act as depolarizing agents [52,53].

Simulations were first run considering uniform corrosion under active or passive conditions. A same electrochemical state was thus imposed on the entire rebar. Generally, the rebar is considered to be in passive state when \( i_0 \) < 0.1 \( \mu \)A cm\(^{-2}\) (10\(^{-3}\) A m\(^{-2}\)) and in active state when \( i_0 \) > 1 \( \mu \)A cm\(^{-2}\) (10\(^{-2}\) A m\(^{-2}\)) [54]. Here, \( i_0 \) was fixed at 10\(^{-5}\) A m\(^{-2}\) for the passive rebar and at 0.1 A m\(^{-2}\) for the active rebar. We must remember that, experimentally, the corrosion rate is highly dependent upon the water content at the steel/concrete interface [51]. Maximum values are generally observed at high relative humidity when the material is wet. For carbonation-induced corrosion, values up to 20 \( \mu \)A cm\(^{-2}\) are expected in outdoor exposure, even if values up to 20 \( \mu \)A cm\(^{-2}\) have been reported under laboratory conditions [55]. For chloride-induced corrosion, values can be up to 1 mm year\(^{-1}\) (i.e., 86 \( \mu \)A cm\(^{-2}\)) [56]. In this study, the maximum value for the active state was set at 10 \( \mu \)A cm\(^{-2}\). Hence, higher values than those simulated in this study are expected for highly corroding areas in wet conditions in chloride-induced corrosion.

To simulate non-uniform corrosion, two different conditions were imposed on the rebar: a small part along the entire circumference of the rebar acts as the corroding area, with the remainder acting as the passive area. Table 1 lists typical values of electrochemical parameters of the Butler–Volmer equation found in the literature for modeling non-uniform corrosion of steel in concrete. The reported parameters are quite different, especially concerning the difference in the exchange current density in corrosion potential between anodic and cathodic sites. Here, the exchange current density of the passive area was fixed at 10\(^{-5}\) A m\(^{-2}\), while that of the active area was 10\(^{-4}\) A m\(^{-2}\) (same order of magnitude, as in References [43,44,57]), or 0.1 A m\(^{-2}\) (same order of magnitude as in References [18,58,59]). If these values affect any quantitative determination, they do not affect the qualitative insight obtained with numerical simulations.

**Table 1.** Typical values of electrochemical parameters found in the literature for modeling non-uniform corrosion of steel in concrete.

| Active Area (Anode) | Passive Area (Cathode) | References |
|---------------------|------------------------|------------|
| \( i_{eq,a} \) (A m\(^{-2}\)) | \( E_{eq,a} \) (V) | \( \beta_{eq,a} \) (V) | \( \beta_{eq,c} \) (V) | \( i_{eq,c} \) (A m\(^{-2}\)) | \( E_{eq,c} \) (V) | \( \beta_{eq,c} \) (V) | \( \beta_{eq,c} \) (V) |
| 1.875 \times 10\(^{-4}\) | -0.78 | 0.06 | - | 6.25 \times 10\(^{-6}\) | 0.16 | - | 0.16 | [44,57] |
| 3 \times 10\(^{-4}\) | -0.78 | 0.09 | 0.18 | 10\(^{-5}\) | 0.16 | 5 | 0.18 | [43] |
| 5 \times 10\(^{-5}\) | -0.65 | 0.09 | 0.15 | 2.5 \times 10\(^{-4}\) | -0.15 | 0.4 | 0.15 | [5] |
| 0.1 | -0.7 | 0.06 | 0.16 | 10\(^{-4}\) | -0.1 | 0.4 | 0.16 | [18,58] |
| 0.3 | -0.576 | 0.046 | 0.3 | 6 \times 10\(^{-5}\) | -0.11 | 0.24 | 0.3 | [59] |
When modeling non-uniform corrosion, an important parameter is the free macrocell current that flows spontaneously between anodic and cathodic sites without external polarization [18]. It depends upon the parameter of the Butler–Volmer equation used in this study, mainly the exchange current density and the corrosion potential difference between anode and cathode (Figure 3). The objective of our study was to assess the influence of the monitoring technique on an established corrosion cell. However, in the modeling, the distribution of the macrocell current lines in the domain is not initially established. Hence, the simulation results considered both the effect of the impressed current for corrosion monitoring and the establishment of a free macrocell current. Irrespective of the exchange current density of the anode, the free macrocell current lines are removed when \( E_{eq,a} = E_{eq,c} = 0 \) V. For the other conditions, the current varies with time but gradually tends towards a constant value, indicating that a steady state is reached (surface potential). Hence, to eliminate the influence of these spontaneous current lines on corrosion monitoring, both anodic and cathodic potentials were fixed at 0 V in this study. Even though this value is not representative of a real macrocell process and can affect any quantitative determination, it does not affect the qualitative insights obtained by our study. To use representative values for anodic and cathodic potentials, it would have been possible to use the steady-state obtained without external polarization as initial conditions to the simulations with external polarization to remove the effect of the free macrocell current of the potential difference.

**Figure 3.** Illustration of the free macrocell current (evaluated at 50 s) that flows spontaneously between anodic and cathodic areas without applying any external polarization \( (I_{cl} = 0 \mu A) \) for \( \rho = 100 \Omega \) m. The cover depth \( c \) was set at 40 mm and the anode length \( L_a \) at 1 cm. The inserts show the temporal evolution of the potential difference between \( P_1 \) and \( P_2 \).

We also used a corrosion potential value of 0 V for modeling uniform corrosion as the results are used as reference states for determining the difference between microcell and macrocell corrosion. The input parameters of the Butler–Volmer equation used in this study are summarized in Table 2. The \( \alpha \) values were selected in order to obtain \( B \) values (Stern–Geary constant, see Equation (1)) close to 26 mV for active corrosion and close to 52 mV for passive corrosion, which are the common recommended values [60]. In each section, we input one Tafel slope for the anodic reaction and one Tafel slope for the cathodic one, as both microcell and macrocell components are non-negligible values in the total current density [61–63].

Another input parameter for considering the time-dependency of corrosion is the double-layer capacitance \( C_{dl} \) [31,35], which notably influences the time required to reach the steady-state potential. This parameter was implemented as a boundary condition at the steel/concrete interface for both anodic and cathodic areas. Typical values of \( C_{dl} \) for steel in concrete range from 1 to 150 \( \mu F \) cm\(^{-2} \) [64], even if greater ranges have also been reported from laboratory work [65]. Some differences between \( C_{dl} \) values for active and passive states are expected because of variation in electrolyte concentration or changes in exposed metal surface. Here, to separate the contribution of each parameter, we first...
considered a same value for both active and passive areas, investigating first the effect of the exchange current density alone. Then, we investigated the effect of $C_{dl}$ by increasing its value in the anodic area.

| Active Corrosion | Passive Corrosion |
|------------------|-------------------|
| $i_{0,a}$ (A m$^{-2}$) | $E_{eq,a}$ (V) | $\alpha_a, a$ (V) | $\alpha_c, a$ (V) | $i_{0,c}$ (A m$^{-2}$) | $E_{eq,c}$ (V) | $\alpha_a, c$ (V) | $\alpha_c, c$ (V) |
| 0.1 | 0 | 0.5 | 0.5 | - | - | - | - |
| 0.1 or $10^{-4}$ | 0 | 0.5 | 0.5 | $10^{-5}$ | 0 | 0.012 | 0.5 |

Table 2. List of the Butler–Volmer parameters used in this study for uniform (lines 1 and 2) and non-uniform corrosion (line 3).

To model the indirect GP measurements, the four electrodes were modeled as perfect point objects on the concrete surface in Wenner configuration, i.e., with the same probe spacing between each electrode and aligned right above and parallel to the rebar. The assumption of point-shaped electrodes is suitable as the geometry of a four-point probe has only a limited effect on calculating the concrete resistivity [66]. The two outer electrodes C$_1$ and C$_2$ act as current electrodes to inject 100 $\mu$A and $-100$ $\mu$A, respectively, in order to virtually create their connection to the current generator, while the two inner electrodes P$_1$ and P$_2$ act as potential electrodes; probe spacing $a$ varies between 2.5 and 15 cm. To determine the sensitivity of the technique, the electrodes remain fixed at the center of the element, while the anodic site is moved along the rebar. Thus, the distribution of the current in the material is not affected by changes in border effects, which allows the comparison of all simulations. An extremely fine mesh, with 2.99 million tetrahedral elements ranging from 0.0004 to 0.04 m, was used for evaluating the temporal evolution (without further indication from 0 to 30 s, with time steps of 0.2 s) with MUMPS (multifrontal massively parallel sparse direct solver) and a relative tolerance of 0.001 in order to obtain an accurate solution.

From the simulations, we derived the potential difference between P$_1$ and P$_2$ ($V_{P1-P2}$), and the current that polarizes the rebar $I_{rebar}$ and the anode $I_{anode}$, by integrating for each time step the absolute normal current density over the total rebar surface or over the anode surface, respectively. The integrated current was divided by two to account for the two different currents impressed on the concrete surface. The results obtained for the non-uniform corrosion were then compared to the relevant passive state to investigate the sensitivity of the indirect GP technique for detecting localized corrosion.

3. Experimental Results

The experimental results are presented in Figure 4. Figure 4a and Figure S1C represent the influence of the probes position with respect to the rebar (0–12 cm; 0 cm indicates that measurements are done above and parallel to the rebar). Irrespective of the position of the device, a clear polarization is observed. This polarization is related to the rebar and not to the concrete, as only an ohmic drop is observed on unreinforced specimens (Figure S1A). Results indicate that the further the monitoring device is from the rebar, the higher the apparent concrete resistance (i.e., $V_{P1-P2}$ at 0.1 s) and the lower the apparent polarization resistance (i.e., $V_{P1-P2}$ at steady-state). Indeed, more current flows in the rebar at 0 cm, whereas more current flows in the concrete at 6 and 12 cm. This result can in part be comparable to what is expected when increasing the cover depth; increasing the cover depth increases the distance between the probes and the rebar, decreasing the part of the current that polarizes the rebar.

The influence of the impressed current ($I = 50–1000 \mu A$) is presented in Figure 4b. Results indicate that an increase in the impressed current increases $V_{P1-P2}$ for each time. Visually, it appears that a longer time is required to reach the steady state when increasing the impressed current. However, a linear evolution of $V_{P1-P2}$ with the impressed current is obtained at $t = 0.1$ s or $t = 30$ s (Figure 4c). Hence, comparable results are obtained irrespective of the impressed current and increasing the current can help to obtain a more significant variation on $V_{P1-P2}$. Even if the injected current values appear to
be high as compared to the values traditionally used in three-electrode configuration (generally a few µA), we must recall that only a fraction of the current applied on concrete surface will polarize the rebar in this configuration [35]. As discussed later in Section 4.2, this fraction decreases with time in the case of a passive rebar, so the polarization of the rebar is not as strong as expected from the values of $V_{P1-P2}$.

![Graphs showing experimental results](image)

**Figure 4.** Experimental results obtained on the mortar sample using the indirect galvanostatic pulse (GP) technique. (a) Influence of the distance of the probe from the rebar on the evolution of $V_{P1-P2}$ with $I = 100$ µA and $a = 15$ cm, all electrodes being parallel to the rebar, (b) influence of the injected current (50–1000 µA) on the evolution of $V_{P1-P2}$ with $a = 15$ cm, (c) evolution of $V_{P1-P2}$ as a function of the impressed current for $a = 15$ cm at $t = 0.1$ s and 30 s, (d) influence of the probe spacing on $V_{P1-P2}$ with $I = 200$ µA for the passive rebar (reproduced from Reference [2]), (e) influence of the probe spacing on $V_{P1-P2}$ with $I = 200$ µA after the introduction of chlorides from the concrete surface through wetting/drying cycles, and (f) evolution of $\Delta V$ with time for each probe spacing, showing the difference between the passive rebar and the rebar after Cl$^-$ introduction.
Figure 4d and Figure S1D show the influence of the probe spacing \((a = 5–15 \text{ cm})\) for the passive rebar. The initial value obtained at 0.1 s decreases when increasing the probe spacing. However, as the current distribution is impacted by the finite size of the sample, only a small difference is observed between 10 cm and 15 cm at 0.1 s. The results also indicate that increasing the probe spacing increases the value of \(V_{P_1-P_2}\) on the steady-state potential. Irrespective of the probe spacing, the time required to reach the quasi-steady state is largely reduced as compared to the results obtained in the three-electrode configuration (Figure S1B), which is in agreement with References [22,28,35].

The results obtained after the local introduction of Cl\(^{-}\) through wetting and drying cycles (Figure 4e) indicate that a longer time is required to reach the steady state as compared to the passive rebar, for each probe spacing. We then calculated the difference in \(V_{P_1-P_2}\) between the passive rebar and the rebar after Cl\(^{-}\) introduction for each time (Equation (5)):

\[
\Delta V(t) = V_{P_1-P_2}(t) \text{ for passive rebar} - V_{P_1-P_2}(t) \text{ for rebar after Cl}^{-} \text{ introduction.}
\]  

(5)

The results are presented in Figure 4f. An increase in \(\Delta V\) is observed before it reaches a maximum value. Increasing the probe spacing increases the time required to reach this maximum, but it is observed in less than 30 s in all cases. The magnitude is also higher when \(a = 15 \text{ cm (} \Delta V_{\text{max}} = 20.2 \text{ mV) as compared to } a = 5 \text{ cm (} \Delta V_{\text{max}} = 10.2 \text{ mV). After this maximum value, } \Delta V \text{ decreases progressively, indicating that the difference between the two electrochemical state is less important at the steady state. Hence, as stated by Reference [32], the DC-limit seems not adapted for detecting localized corrosion. However, it appears that comparing the evolution of the transient potential can provide insights on the different corrosion states of the rebar.}

To sum up, the experimental results demonstrate that (i) the rebar can effectively be polarized without any connection by applying a direct current between two probes located on concrete surface, (ii) the distance of the probes from the rebar affects \(V_{P_1-P_2}\), (iii) the size of the specimen affects the current distribution in the material, and (iv) the time required to reach the steady state appears related to the corrosion state of the rebar.

4. Numerical Results

4.1. Unreinforced Specimen: Geometric Factor

In Wenner configuration, the geometric factor is calculated according to Equation (7):

\[
k_{\text{Wenner}} = 2\pi a,
\]  

where \(a \text{ (m)}\) is the probe spacing. However, this equation was defined for homogeneous and isotropic material with a semi-infinite geometry, which in practice is not true for RC structures [66]. Hence, the applicability of Equation (7) was verified by calculating the geometric factor obtained with the input resistivity and the results of simulations (Equation (8)):

\[
k_{\text{COMSOL}} = \frac{\rho_{\text{input}}}{R_{\text{without rebar}}} = \frac{\rho_{\text{input}}}{\left(\frac{V_{P_1-P_2}}{I_{C1}}\right)_{\text{without rebar}}}.
\]  

(8)

The results obtained for five probe spacings are presented in Table 3. The conventional geometric factor \(k_{\text{Wenner}}\) is quite similar to the calculated \(k_{\text{COMSOL}}\) one when \(a = 2.5\) or 5 cm. However, increasing \(a\) up to 15 cm gives quite different values, with lower values obtained by the numerical simulations. This difference may be related to edge effects due to the finite size of the element. Hence, as already shown in other studies [67–70], the sample geometry is an important parameter when running experiments and numerical simulations, and an accurate geometric factor must be determined case by case, depending on specimen size and probe spacing.
Table 3. Comparison of the conventional geometric factor in Wenner configuration $k_{\text{Wenner}}$ (Equation (7)) and the geometric factor calculated in COMSOL $k_{\text{COMSOL}}$, which considers the sample geometry (Equation (8)).

| $a$ (cm) | $k_{\text{Wenner}} = 2\pi a$ (m) | $k_{\text{COMSOL}}$ (m) |
|----------|---------------------------------|------------------------|
| 2.5      | 0.157                           | 0.149                  |
| 5        | 0.314                           | 0.304                  |
| 7.5      | 0.471                           | 0.387                  |
| 10       | 0.628                           | 0.424                  |
| 15       | 0.942                           | 0.405                  |

4.2. Uniform Corrosion: Parametric Study

The input parameters investigated for uniform corrosion are listed in Table 4.

Table 4. List of input parameters used for the case of uniform corrosion.

| Input Parameters                        | Values                      |
|-----------------------------------------|-----------------------------|
| Current impressed on $C_1$ $I_{C1}$ ($\mu A$) | 100, 300 or 500             |
| Concrete resistivity $\rho$ ($\Omega$ m)     | 100, 200, 500 or 1000        |
| Probe spacing $a$ (cm)                   | 2.5, 5, 7.5, 10 or 15       |
| Cover depth $c$ (mm)                     | 20, 40 or 60                |
| Rebar diameter $\Phi$ (mm)              | 6, 12 or 25                 |

4.2.1. Instantaneous Ohmic Drop

We first focused on the instantaneous potential difference obtained at $t_0$. All the values reported in Table S1 were obtained considering both active and passive corrosion. Hence, the instantaneous ohmic drop appears independent of the electrochemical state of the steel/concrete interface (i.e., Butler–Volmer input parameters). For each resistivity, $V_{P1-P2}$ decreases nonlinearly when increasing the probe spacing $a$ (Figure 5a), decreasing the cover depth $e$ (Figure 5b), or increasing the rebar diameter $\Phi$ (Figure 5c).

![Figure 5](image_url)

*Figure 5.* Evolution of the instantaneous ohmic drop observed at $t_0$ for both active and passive conditions, for $I_{C1} = 100 \mu A$ as a function of: (a) probe spacing $a$, for each concrete resistivity, $\epsilon = 40$ mm and $\Phi = 12$ mm, (b) cover depth $e$, for each concrete resistivity, $a = 5$ cm and $\Phi = 12$ mm; (c) rebar diameter $\Phi$ for each concrete resistivity, $a = 5$ cm and $e = 40$ mm.

Only a fraction of the current impressed on the concrete surface polarizes the rebar. As shown in Table S1, this value is almost independent of concrete resistivity. However, higher values are observed when the probe spacing is increased, as more current penetrates deeper into the concrete volume. Higher values are also observed when the cover depth is decreased and when increasing the rebar diameter, in agreement with Reference [35]. Though the value of the impressed current $I_{C1}$ also influences the instantaneous ohmic drop, it has no effect on the calculated resistance, in agreement with Ohm’s law and the experimental results. As the total injected current at the concrete
surface remains constant, the other fraction of the current flows in the concrete without polarizing
the rebar. For each input resistivity, the highest $V_{P1-P2}$ values are observed when this fraction is
maximum. The instantaneous potential difference was used for calculating concrete resistivity using
the geometric factor $k_{\text{COMSOL}}$ determined numerically that accounts for potential border effects (see Table 3).
This value was then compared to the different input resistivities (100, 200, 500, or 1000 $\Omega$ m). In all
cases, the resistivity calculated using the total impressed current was lower than the input resistivity
(Figure 6).

![Figure 6](image_url)

**Figure 6.** Comparing input concrete resistivity with the concrete resistivity calculated with Equation (6),
considering the accurate geometric factor $k_{\text{COMSOL}}$ for each simulation. The calculated resistivity
decreases when increasing $\sigma$, decreasing $\varepsilon$ or increasing $\Phi$ as more current reaches the rebar.
A percentage of underestimation was calculated as (Equation (9)):

$$\% \text{ of underestimation} = \frac{\rho_{\text{input}} - \rho_{\text{calc}}}{\rho_{\text{input}}} \times 100.$$  (9)

As shown in Table S1, this percentage increases when increasing $\sigma$, decreasing $\varepsilon$ and increasing
$\Phi$, though it is independent from the input resistivity. The underestimation is related to the presence
of rebar that acts as a short-circuit due to its very low electrical resistivity at high frequencies.
Hence, the instantaneous ohmic drop results from the contribution of concrete resistivity but also of
the rebar. This is known as the rebar effect and is discussed in Section 5.1.

4.2.2. Transient Potential

We then focused on the transient potential. Some examples of the results are presented in Figure 7,
and all the results are presented on Figure S2. The ohmic drop is not shown on these figures as the
first point of the potential difference $V_{P1-P2}$ at $t_0$ was normalized to 0 mV for easier comparison of the
different simulations.

For active corrosion, the current polarizing the rebar slightly decreases before reaching a constant
value, which is highest for the highest resistivity (Figure 7, left). As there is more resistance to current
flow in the concrete in this condition, it becomes easier for the current to polarize the rebar. Compared
to the conventional GP technique in the three-electrode configuration, only a fraction of the injected
current polarizes the rebar in the indirect GP technique.

In parallel, $V_{P1-P2}$ rapidly increases before reaching a constant value (Figure 7, middle). The time
required to reach this steady-state potential depends upon the double-layer capacitance $C_{dl}$: for
$C_{dl} = 0.2 \text{ F m}^{-2}$, the steady-state potential is reached in less than 1 s, whereas more than 4 s are
required for $C_{dl} = 2 \text{ F m}^{-2}$. Though this steady-state potential value is independent of $C_{dl}$, it depends
upon the exchange current density $i_0$, and on concrete resistivity $\rho$, probe spacing $a$, cover depth $e$, and rebar diameter $\Phi$ (Figure 7 and Figure S2).

For passive corrosion, the current polarizing the rebar decreases nonlinearly over time before reaching very low values at the end of the simulations (Figure 7 left and Figure S2). After 30 s, the total current on the rebar is less than $1 \mu A$ for $\rho = 100 \, \Omega \, m$, irrespective of probe spacing, but it remains higher than $5 \mu A$ for $\rho = 1000 \, \Omega \, m$. As concrete resistivity decreases, the current decreases more...
rapidly. This is related to the high polarization resistance of the rebar and the different behavior of a capacitor at high and low frequencies [26]. Theoretically, a capacitor acts as a short-circuit at high frequencies (i.e., over very short times), whereas it acts as an open circuit at low frequencies (i.e., over a long period of time) [35]. Between these two opposite behaviors, the current polarizing the capacitor decreases at a velocity that is dependent on the time constant of the system (τ = RC). This means that at the start of injection, the current can easily flow through the rebar capacitance. In the conventional three-electrode GP technique, as the rebar is the only current sink in the system, the entire impressed current polarizes it each time, either in its capacitive or its resistance component, and the current flows progressively through the polarization resistance when increasing the duration of the injection. However, in the indirect GP technique, the current can simply flow between C1 and C2 through the concrete without polarizing the rebar. In passive condition, due to the high polarization resistance, the concrete progressively becomes the less resistive path in the system as the duration of the injection increases, especially for low resistive concrete. This behavior explains the very low current in the rebar after 30 s when ρ = 100 Ω m. As the total injected current remains constant in the system, the fraction of the current that flows in the concrete without polarizing the rebar increases over time.

In parallel to the decrease in the current on the rebar, \( V_{P1-P2} \) increases nonlinearly over time (Figure 7 right). A decrease in the probe spacing \( a \) decreases \( V_{P1-P2} \), while a decrease in the cover depth \( e \) increases \( V_{P1-P2} \). In fact, the higher the initial current on the rebar, the higher will be the potential difference \( V_{P1-P2} \), as it was observed experimentally. Increasing concrete resistivity also increases \( V_{P1-P2} \). If the transient potential was only related to the polarization resistance, a decrease in \( V_{P1-P2} \) would have been expected as the current that polarizes the rebar decreases over time. In fact, for each time, there is an increase in \( V_{P1-P2} \) related to the concrete resistivity—as more current flows in the concrete—and a decrease in \( V_{P1-P2} \) related to the polarization resistance of the rebar, as less current flows in the rebar. Overall, this results in an increase in \( V_{P1-P2} \) (Figure 8). This means that the indirect GP technique is fundamentally different from the conventional GP technique in three-electrode configuration, for which the transient potential is not dependent on concrete resistance.

![Figure 8](https://example.com/figure8.png)

**Figure 8.** Schematic representation of the physical aspect of the indirect GP technique in Wenner configuration. (a) Evolution of the distribution of the current in the concrete and in the rebar; (b) evolution of the potential difference attributed to concrete resistance, polarization resistance, and the global response.

A “quasi-steady-state” is reached after only 30 s for \( \rho = 100 \, \Omega \, m \). Indeed, as the polarizing current is already very low at this time, variations in \( V_{P1-P2} \) are almost insignificant. This is considerably shorter than the time required to reach the steady state in the conventional three-electrode configuration for passive rebar, as already noted in References [22,28,35]. However, the higher the concrete resistivity, the slower the current that polarizes the rebar decreases and the longer it takes to reach the quasi-steady-state potential. Hence, for highly resistive concrete, a longer polarization time is required to reach the steady state, though a quasi-steady-state is reached in about 300 s (Figure S3).

By comparing the values of \( V_{P1-P2} \) at steady state or “quasi-steady-state”, some practical recommendations can be formulated. For a highly corroding rebar, as \( V_{P1-P2} \) is low irrespective of concrete resistivity, the impressed current must be higher than 100 µA to improve the signal-to-noise
ratio. For a passive rebar in highly resistive concrete, $V_{P1-P2}$ values seem to indicate a strong polarization of the rebar as $V_{P1-P2} > 20$ mV. However, as already stated, this is not the case as most of $V_{P1-P2}$ is mainly related to concrete resistivity. As the current polarizing the rebar decreases over time, the strongest rebar polarization is obtained shortly after injection of the current. Hence, when the rebar condition is unknown, one can impress a high current (e.g., 300 or 500 $\mu$A) for improving the signal-to-noise ratio without causing irreversible changes to the steel rebar. If the resulting $V_{P1-P2}$ value is high enough, the value of the impressed current can be decreased to reduce potential damage to the concrete microstructure if the measurements will be repeated several times.

4.3. Non-Uniform Corrosion: Detection of the Actively Corroding Area

The input parameters for non-uniform corrosion are summarized in Table 5. In this section, we only focus on the transient potential and not on the instantaneous ohmic drop obtained at $t_0$.

**Table 5.** List of input parameters used for the non-uniform corrosion of steel in concrete.

| Input Parameters                      | Values                |
|---------------------------------------|-----------------------|
| Current impressed on C1 $I_{C1}$ ($\mu$A) | 100                   |
| Concrete resistivity $\rho$ ($\Omega$ m) | 100, 200, 500 or 1000 |
| Anode length $L_a$ (cm)               | 1 or 3 (5, 7.5, 10, or 15 in SI) |
| Probe spacing $a$ (cm)                | 2.5, 5 or 15          |
| Cover depth $e$ (mm)                  | 40                    |
| Rebar diameter $\Phi$ (mm)            | 12                    |
| Double-layer capacitance $C_{dl}$ (F m$^{-2}$) | 0.2 or 2              |

4.3.1. Influence of an Anode with a Low Anodic Exchange Current Density

The effect of the anodic exchange current density was first studied with $i_{0,a} = 10^{-4}$ A m$^{-2}$. To compare the results obtained for non-uniform corrosion with those for uniform corrosion in passive state, we calculated the potential difference $\Delta V$ obtained at each time with Equation (5). The greater this difference, the more important is the effect of the anode on potential and current distribution. This equation can be used if the impressed current is similar for all measurements and if the concrete resistivity profile is uniform in the investigated area.

Probe spacing was set at 5 cm, which is an industry standard for concrete resistivity meters. The anode length $L_a$ was first set at 1 cm ($S_a = 3.77$ cm$^2$), and the results are shown on Figure 9. Almost no difference with the reference state is observed when the anode is located at the center of P$_1$ and P$_2$ or at 7.5 cm and more from C$_1$ or C$_2$, as $\Delta V$ remains close to 0, especially when $\rho = 100$ or 200 $\Omega$ m. Only small differences are observed when the anode is below the current electrodes when $\rho = 500$ or 1000 $\Omega$ m, but $\Delta V_{\text{max}}$ remains lower than 5 mV in these cases.

In concrete and other porous media, the growth of anodic sites is more rapid close to anode/cathode boundaries than deeper down in the pit due to the non-uniform distribution of current densities [71]. Hence, the size of a pit is greater on surface than at depth. Thus, it is expected that anodes over 1-cm long can develop at the steel/concrete interface. However, even when the anode length is increased to 3 cm ($S_a = 11.3$ cm$^2$, Figure S4), the difference with a fully passive rebar remains very small and only significant for the highest resistivity ($\rho = 1000$ $\Omega$ m), for which $\Delta V$ can be up to 9 mV.

We must note that the exchange current density used in this first case for the anodic area is very low and is sometimes used for cathodic areas (as indicated in Table 1). A rebar is generally considered in the passive state if the corrosion current density is less than 0.1 $\mu$A cm$^{-2}$ [54], i.e., $10^{-3}$ A m$^{-2}$. Hence, the results of this simulation may rather represent a passive rebar with different electrochemical properties due to the heterogeneity of the passive film at the steel/concrete interface [72]. Considering this case, the low sensitivity of the indirect GP technique cannot be seen as a drawback as it seems non-essential to differentiate these two states as corrosion remains in passive
conditions. Further studies are required to determine if this small difference can still have a practical interest, for example, for locating areas with higher susceptibility to corrosion initiation [73].

![Diagram showing the evolution of V_{P1-P2} and ΔV over time in the presence of a slightly corroding area at different positions relative to the monitoring device.](image)

**Figure 9.** Evolution of V_{P1-P2} and ΔV over time in the presence of a slightly corroding area at different positions relative to the monitoring device. Input parameters: \( i_{0,a} = 10^{-4} \text{ A m}^{-2} \), \( i_{0,c} = 10^{-5} \text{ A m}^{-2} \), \( L_a = 1 \text{ cm} \), \( a = 5 \text{ cm} \), \( e = 40 \text{ mm} \), \( C_{dl,a} = C_{dl,c} = 0.2 \text{ F m}^{-2} \), \( I_{C1} = 100 \text{ µA} \).

4.3.2. Influence of an Anode with a High Anodic Exchange Current Density

To represent a highly corroding area, the anodic-exchange current density was increased to 0.1 A m\(^{-2}\) [18,32,58,59]. Even if this value is mainly expected in a low-resistive concrete with a high water-saturation level, we investigated the effects for all resistivity values as they depend on concrete composition. It has been shown that a steel rebar can corrode at relative similar rates when embedded in a low-resistive mortar prepared only with Portland cement, or in a high-resistive mortar prepared with a mix of Portland cement and fly ash [74].

The anode length was first set at 1 cm (Figure 10). When the anode is perfectly centered between \( P_1 \) and \( P_2 \), the difference with a fully passive rebar is small (\( \Delta V_{\text{max}} < 1.1 \text{ mV} \)). This is similar to the observation made with a low anodic-exchange current density (Figure 9), but, when the anode is below \( C_1 \) or \( C_2 \), the anode has a significant effect on the measured potential. In this position, the anode receives a higher part of the injected current and, thus, is more polarized, and the anodic area will have a significant effect on the evolution of \( V_{P1-P2} \) over time. The strongest effect on \( V_{P1-P2} \) is observed for the highest resistivity, as \( \Delta V_{\text{max}} \approx 14 \text{ mV} \).

It is important to note that the time required to reach \( \Delta V_{\text{max}} \) increases with increased concrete resistivity. After this maximum, \( \Delta V \) slowly decreases before reaching a constant value when the steady-state potential is reached. Hence, the maximum is not observed at the steady state but at the transient potential. The only exception is when the anode is located below \( C_2 \) for \( \rho = 1000 \Omega \text{ m} \), as \( \Delta V_{\text{max}} \) is observed at the steady-state potential.

The difference with passive rebar becomes small again when gradually moving the anode away from the current electrodes as shown by a decrease in \( \Delta V \) values (Figure S5A): \( \Delta V_{\text{max}} < 2.6 \text{ mV} \) at 7.5 cm from \( C_1 \) and \( \Delta V_{\text{max}} < 1.3 \text{ mV} \) at 15 cm from \( C_1 \) when \( \rho = 1000 \Omega \text{ m} \), which is close to the value observed when the anode is centered between \( P_1 \) and \( P_2 \). The sensitivity of the monitoring device is thus spatially limited as the anode is mainly observed when it is located below or close to one of the
current electrodes. These results show that it is possible to estimate the position of such a small but highly corroding anode if the measurement is repeated at several positions along the rebar.

![Graphs showing the evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0a} = 0.1$ A m$^{-2}$, $i_{0c} = 10^{-5}$ A m$^{-2}$, $L_a = 1$ cm, $a = 5$ cm, $e = 40$ mm, $C_{dl,a} = C_{dl,c} = 0.2$ F m$^{-2}$, $I_{C1} = 100$ µA.](image)

For practical considerations, it is important to note that, when the total current injected is 100 µA, $\Delta V_{\text{max}}$ is only 1.2 mV for $\rho = 100$ Ω m in the best case, i.e., when the anode is below one of the current electrodes. It is questionable whether this difference is experimentally significant because of the noise or of the variation in self-potential, which are expected during any measurement. To guarantee the ability of the technique to locate highly corroding areas, it is necessary to impress a higher current (300 or 500 µA) on the concrete surface to increase the signal for a low-resistive concrete (Figure S6).

The anode length was then increased to 3 cm (Figure 11). The stronger effect of the anode on $V_{P1-P2}$ is still observed for the highest resistivity, for which $V_{P1-P2}$ is clearly lower than the reference state. This is the case when the anode is below $C_1$ ($\Delta V_{\text{max}} \approx 23$ mV) or below $C_2$ ($\Delta V_{\text{max}} \approx 31$ mV at 30 s). The main difference with the small anode is observed when the anode is centered between $P_1$ and $P_2$, as significant value of $\Delta V$ are observed. The effect of the anode is, however, still lower than that observed when it is located below one of the current electrodes, as $\Delta V_{\text{max}} < 6$ mV when $\rho = 1000$ Ω m.

The middle of the current probes (and of the potential probes in Wenner configuration) corresponds to a point where the total current on the rebar is null (Figure S7 and Reference [35]). As a result, if a small anode is in this area, only a small part of the current will flow in it, as it was already observed for $L_a = 1$ cm in Figure 10, so the potential difference $V_{P1-P2}$ is almost as if there was no anode. With increasing anode length, more current will flow through it, so the presence of the anode has a significant effect on $V_{P1-P2}$.

It must be noted that $\Delta V$ remains constant close to its maximum values at the steady-state potential when the anode is centered between $P_1$ and $P_2$, which is different than what it is observed when the anode is below one of the current electrodes ($C_1$). It is thus possible to estimate the position of the highly corroding area by comparing both $\Delta V_{\text{max}}$ values and $\Delta V$ at the steady-state potential for each measurement position.

To obtain further insight into the physical aspects of monitoring non-uniform corrosion, the simulations were continued up to 800 s. We focused on the results obtained when the anode is below $C_1$ (Figure 12). The results show that the time to reach the steady state when the anode is below
one of the current electrodes increases with increasing concrete resistivity. Especially when the anode is below C1, it increases from about 50 s for $\rho = 100 \, \Omega \cdot m$ to over 600 s for $\rho = 1000 \, \Omega \cdot m$. In this last case, this is longer than the time needed for reaching the steady state for a passive rebar (Figure S3). Hence, the experimental results presented in Figure 4e indicate the non-uniform corrosion of the rebar, which is expected after locally introducing chlorides through wetting/drying cycles.

![Figure 11](image.png)

**Figure 11.** Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0,a} = 0.1 \, A \cdot m^{-2}$, $i_{0,c} = 10^{-5} \, A \cdot m^{-2}$, $L_a = 3 \, cm$, $a = 5 \, cm$, $\varepsilon = 40 \, mm$, $C_{dl,a} = C_{dl,c} = 0.2 \, F \cdot m^{-2}$, $I_{C1} = 100 \, \mu A$.

The presence of the highly corroding area disturbs the current distribution in the entire material. Right after the instantaneous ohmic drop, the total current in the rebar rapidly decreases, whereas the current at the anode slightly increases. The higher the concrete resistivity, the longer and the more the current in the anode increases. After reaching a maximum value, the current in the anode progressively decreases before reaching a constant value. With increasing injection time, it becomes easier for the current to flow within the concrete than in the rebar, especially when concrete resistivity is low, which explains the decrease of the current in the anodic area.

Hence, it seems that a fraction of the current polarizing the cathodic areas is first redirected to the anodic area, as the latter has the lowest resistance in the system, before being partly redirected to concrete domain, with a magnitude that depends on concrete resistivity. This behavior slows down the current decrease in the anodic area and in the entire rebar, which explains why the time required for reaching the steady-state potential is higher for non-uniform corrosion. Even when the current in the anode is stable, the current that polarized the rebar is still slowly decreasing. As indicated by the $I_{anode}/I_{rebar}$ ratio, over 90% of the current polarizes the anode in the steady state, with the highest values being obtained for $\rho = 100 \, \Omega \cdot m$.

To support this hypothesis, the anodic area was divided into six identical areas of 5-mm length each, to analyze the current distribution in each section. The results obtained for $\rho = 100 \, \Omega \cdot m$ and 1000 $\Omega \cdot m$ are shown on Figure 13. At $t_0$, the current is slightly higher for the central anodes (anodes 3 and 4) as they lie below C1, but the current increase in the first seconds is more important for the two external anodes (anodes 1 and 6), in agreement with a redirection of the current from adjacent cathodes. In addition, the decrease in current is slower for the most eccentric anode relative to the device (anode 1).
Another interesting result observed in Figures 10 and 11 is the difference in $V_{P1-P2}$ evolution obtained when the anode is below $C_1$ or $C_2$, notably when $\rho = 1000 \, \Omega \, m$. This signal dissymmetry can be related to the difference in current distribution. It is shown that a higher current, in absolute values, flows in the anode when it is located below $C_2$. This effect is attributed to the difference in the anodic charge-transfer coefficient $\alpha_a$ between the anode and the cathode, as the system is symmetric when the same $\alpha_a$ value is input in both anodic and cathodic areas (Figure S8). The current distribution at the steel/concrete interface is affected by the current direction, as both anode and cathode have a different capability for consuming anodic or cathodic current, depending on the value of this charge-transfer coefficient. Notably, as indicated by the $I_{\text{anode}}/I_{\text{rebar}}$ ratio, the percentage of the
current on the anode is clearly lower when the anode is below $C_2$. Reversing the polarity during the measurement may thus help in differentiating the electrochemical parameters of the rebar: the greater the dissymmetry, the more the charge-transfer coefficients are different between anode and cathode. However, this behavior is mainly observed for $\rho = 500$ and 1000 $\Omega$ m, which limits its interest for highly resistive concrete. For the lowest resistivities, as there is less resistance to current flow in the concrete, the total current polarizing the rebar decreases rapidly to values below 6 $\mu$A. Hence, irrespective of the polarity direction, there is almost no difference between the two sides as the potential difference induced by anode polarization rapidly becomes insignificant.

4.3.3. Influence of the Probe Spacing

If the sensitivity of the technique to localized corrosion was assessed by modifying the anode length, this parameter is in fact unknown in RC structures. In practice, only the probe spacing can be adjusted as the other parameters, such as cover depth, rebar diameter, or electrochemical condition of the rebar, are the main characteristics of the investigated structures. Thus, we had to determine if varying the probe spacing will modify the sensitivity of the technique.

The probe spacing was first reduced to 2.5 cm (Figure 14). The values of $\Delta V_{\text{max}}$ are less important when the anode is below $C_1$ or $C_2$ than those obtained for $a = 5$ cm, as $\Delta V_{\text{max}} \approx 14.7$ mV here. This can be explained by a decrease of the current that polarizes the rebar at $t_0$ when decreasing the probe spacing. The evolution of $\Delta V$ over time when the anode below $C_1$ or $C_2$ is also more similar, irrespective of concrete resistivity, indicating that the system is more symmetric.

As observed for $a = 5$ cm, the difference with the passive rebar becomes small again when gradually moving the anode away from the current probes (Figure S5C). But, here, $\Delta V$ is lower at 7.5 cm from $C_1$ as the anode is less polarized. Hence, reducing the probe spacing increases the “internal” sensitivity but also decreases the “external” sensitivity of the indirect GP technique.

The probe spacing was then increased to 15 cm (Figure 15). The results obtained with this probe spacing show some similarities with those obtained for $e = 5$ cm and $L_a = 1$ cm (Figure 10), despite that a longer time is required to reach the quasi-steady-state here. There is almost no difference from the reference passive state when the anode is centered between $P_1$ and $P_2$, notably for the lowest resistivities. In this case, the technique is not sensitive to the presence of a highly corroding area of
3 cm. A small difference is still observed for $p = 500$ or $1000 \, \Omega \, m$, but it remains lower than 1.2 mV. As shown on Figure S9, the anode length must be higher than 5 cm to observe a significant effect when the anode is centered between $C_1$ and $C_2$.

![Graphs showing $V_{P1-P2}$ and $\Delta V$ over time](image)

**Figure 15.** Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0,a} = 0.1 \, A \, m^{-2}$, $i_{0,c} = 10^{-5} \, A \, m^{-2}$, $L_a = 3 \, cm$, $a = 15 \, cm$, $c = 40 \, mm$, $C_{dL,a} = C_{dL,c} = 0.2 \, F \, m^{-2}$, $I_{C1} = 100 \, \mu A$.

The distribution of the current density along the rebar can help defining the sensitivity of the device according to probe spacing. As shown on Figure S7, there is a point of zero-current density on the rebar at the center of $C_1$ and $C_2$, which corresponds to the point of sign inversion of the current density, and to two points of maximum current density just below the current electrodes. In between, the current density increases from zero to its maximum value at a rate that increases with decreasing probe spacing. As a result, the current fraction that polarizes the central area under the device decreases when increasing the probe spacing. Conversely, polarization is important away from the current electrodes as a longer part of the rebar is polarized; even when the anode is 7.5, 15, or even 30 cm distant from $C_1$, it still has a significant effect on the measured $V_{P1-P2}$ (Figure S5D).

To sum up, an increase in the probe spacing decreases the “internal” sensitivity and increases the “external sensitivity” of the technique, as well as increases the time required to reach the steady-state potential. Nevertheless, irrespective of probe spacing, the most important difference with the reference passive state is observed when the anode lies below one of the current probes as it is the main polarized area.

### 4.3.4. Influence of the Double-Layer Capacitance of the Anode

The effect of the double-layer capacitance was investigated by considering a hypothetical case in which the exchange current density of the anodic and cathodic areas is equal ($i_0 = 10^{-5} \, A \, m^{-2}$). The objective was to separate the contribution of $C_{dL}$ from the other input parameters. We considered only one anodic length of 3 cm and one probe spacing of 5 cm. $C_{dL}$ was fixed at 0.2 F m$^{-2}$ for the cathode and at 2 F m$^{-2}$ for the anode, and the results were compared to the reference state at 0.2 F m$^{-2}$. As $C_{dL}$ is higher, a longer time to reach the steady-state potential is expected when the anode is polarized in the sensitivity area of the device.

The results for $a = 5 \, cm$ are presented on Figure 16. The most pronounced effect is observed when the anode is below one of the current electrodes, confirming once again that this corresponds to the
area of maximum sensitivity. Irrespective of concrete resistivity, the maximum difference \( \Delta V_{\text{max}} \) is observed before 10 s, when \( C_{dl,a} = 2 \, \text{F m}^{-2} \). The difference is lower for \( \rho = 100 \, \Omega \, \text{m} \) (\( \Delta V_{\text{max}} < 2 \, \text{mV} \)) than for \( \rho = 1000 \, \Omega \, \text{m} \) (\( \Delta V_{\text{max}} > 15 \, \text{mV} \)). After this maximum value, the difference progressively tends to 0, in agreement with the similar exchange current densities between anodic and cathodic areas.

In practice, both the exchange current density and the double-layer capacitance of the anodic area can be different than those of the passive areas. Thus, we verified the combined effect of \( i_0 \) and \( C_{dl} \) on \( V_{P1-P2} \) (Figure 17). These results are compared to those obtained in Figures 11 and 16. There is only a small increase in \( \Delta V_{\text{max}} \) (<1 mV) as compared to the results obtained with the same value of \( C_{dl} \) in the entire rebar (Figure 11). Thus, increasing \( C_{dl} \) by one order of magnitude has a limited effect on \( V_{P1-P2} \) for a small anode in highly corroding state.

**Figure 16.** Evolution of \( V_{P1-P2} \) and \( \Delta V \) over time in the presence of an anode with a high double-layer capacitance at different positions relative to the monitoring device. Input parameters:

- \( i_{0,a} = i_{0,c} = 10^{-5} \, \text{A m}^{-2} \), \( l_a = 3 \, \text{cm} \), \( a = 5 \, \text{cm} \), \( e = 40 \, \text{mm} \), \( C_{dl,a} = 2 \, \text{F m}^{-2} \), \( C_{dl,c} = 0.2 \, \text{F m}^{-2} \), \( I_{C1} = 100 \, \mu\text{A} \).

**Figure 17.** Evolution of \( V_{P1-P2} \) and \( \Delta V \) over time in the presence of a highly corroding area with a high double-layer capacitance at different positions relative to the monitoring device. Input parameters:

- \( i_{0,a} = 0.1 \, \text{A m}^{-2} \), \( i_{0,c} = 10^{-5} \, \text{A m}^{-2} \), \( l_a = 3 \, \text{cm} \), \( a = 5 \, \text{cm} \), \( e = 40 \, \text{mm} \), \( C_{dl,a} = 2 \, \text{F m}^{-2} \), \( C_{dl,c} = 0.2 \, \text{F m}^{-2} \), \( I_{C1} = 100 \, \mu\text{A} \).
Hence, a similar value of $C_{dl}$ can be applied in the entire rebar without causing a significant error in the determination of $\Delta V_{\text{max}}$. We do not exclude that another conclusion can be drawn in other conditions, for example, for a more uniform corrosion morphology in which the anode can be several centimeters long, with a lower exchange current density, as it can be expected for carbonation-induced corrosion.

5. Discussion

5.1. Rebar Effect on the Instantaneous Ohmic Drop

As shown in Section 4.2.1, the instantaneous ohmic drop obtained on reinforced concrete is not only related to concrete resistivity but also results from a rebar effect [75,76]. The latter originates from the fact that the measured resistivity in a four-probe configuration corresponds to an apparent resistivity, which is an integrated value of local resistivities from the concrete surface to a set depth. In the presence of rebar, the apparent resistivity represents a weighted contribution of concrete and steel rebar, the latter being much more conductive than concrete at high frequencies. Expressed as percentages, we showed that the rebar effect is not affected by changes in the injected current or in concrete resistivity. However, it is stronger when there is an increase in probe spacing $a$ or in rebar diameter $\Phi$ or a decrease in cover depth $e$. This agrees with other studies that showed that measurements of concrete resistivity in Wenner configuration are affected by cover depth, probe direction, probe spacing, and rebar spacing [77–83]. Concrete thickness is also a major factor affecting measurement in the case of thin slabs [80]. However, rebar diameter has only a limited impact.

Each measurement made in front of a rebar in four-electrode configuration must then be corrected to determine the true concrete resistivity $\rho_{\text{corr}}$ from the apparent resistivity $\rho_{\text{calc}}$. Garzon et al. [70] proposed to correct the conventional geometric factor used in Wenner configuration (Equation (7)) by adding two correction factors: a shape factor $f_s$ that accounts for the finite size of the element and a rebar factor $f_b$ that accounts for the presence of rebar (Equation (10)):

$$k = f_s f_b 2\pi a.$$  

They defined the values of $f_s$ and $f_b$ for some geometries, i.e., prismatic, cylindrical, and slab specimens with specific dimensions, in the absence and presence of one rebar. For other configurations, their parameters cannot be used, and the general procedure presented in our study must be reproduced. First, one must accurately evaluate the specimen size by determining the geometric factor. This is done by a numerical simulation of unreinforced concrete with the accurate dimension and probe spacing. Then, the percentage of underestimation in the presence of rebar must be determined by a numerical simulation of reinforced concrete with the accurate dimension, cover depth, rebar diameter, and probe spacing. The input values for concrete resistivity and electrochemical parameters do not matter as they do not influence this percentage. Finally, Equation (11) determines the corrected apparent resistivity:

$$\rho_{\text{corr}} = \frac{\rho_{\text{calc}}}{1 - \frac{\% \text{ of underestimation}}{100}}.$$  

We also observed in uniform corrosion that the instantaneous ohmic drop is independent of the electrochemical state of the steel/concrete interface, i.e., of $i_0$, $E_{\text{eq}}$, $a$, and $C_{dl}$ values. This aspect is important as it allows calculating the concrete resistivity without knowing the electrochemical condition of the rebar, which is the case when making measurements on RC structures. This finding is quite opposite from the results in Reference [84], where Nguyen et al. concluded, from both numerical simulations and experiments, that the electrochemical state of rebar affects the resistivity drop. Concerning their simulations, the time at which they evaluated the ohmic drop is not specified. As they observed almost no resistivity drop for a passive rebar in low-resistive concrete (100 $\Omega$ m), we believe that their study was done in stationary mode, i.e., at steady state. Consequently, they considered
not only the ohmic drop but also the temporal evolution of the current distribution related to the polarization of the rebar. Hence, the absence of resistivity drop can be related to the fact that almost all the current polarizes the concrete at steady state, as shown in Figure 7, as a passive rebar acts mainly as a current insulator at low frequencies. Still, it can be possible to differentiate active and passive corrosion by using an intermediate frequency (5 Hz in their study) as the current distribution is already different between the two cases, but the measured potential does not correspond only to the instantaneous ohmic drop.

Experimentally, the measurement frequency is a major parameter that must be defined for a correct determination of concrete resistance. Impedance spectroscopy is necessary for determining the frequency at which the imaginary part of the impedance (reactance) is least for correctly assessing the concrete resistance. When measurements are done with a direct current, further investigations must determine if this minimum reactance can be defined by analyzing the temporal evolution of $V_{P1-P2}$ acquired at high sampling rate.

5.2. Ability of Indirect GP to Detect Localized Corrosion in Wenner Configuration

The numerical simulations done in this study have demonstrated the ability of the indirect GP method to detect localized corrosion of steel in concrete with a uniform resistivity, by considering the difference in variation of the transient potential over time. Based on all results, we can estimate the length of a localized corroding area by considering the difference in “internal” sensitivity of this probe configuration depending on probe spacing. As for the half-cell potential technique [85], it seems more appropriate to consider the potential gradients and not the absolute values. Here, we propose using the maximum difference value observed over the injection time, $\Delta V_{\text{max}}$, for each position of the monitoring device.

For this, we must first determine the evolution of $\Delta V$ with time for each measurement position using Equation (5), which can be used if the impressed current is similar for all measurements and if the concrete resistivity profile is (almost) uniform in the investigated area. Otherwise, the data must be normalized by concrete resistivity to make them comparable. In practice, the reference state used in this study for calculating $\Delta V$, i.e., a fully passive rebar, will be unknown in most cases. Nonetheless, it is possible to use the signal with the highest values of $V_{P1-P2}$ for each measurement time as the reference value for calculating $\Delta V_{\text{max}}$. Afterwards, $\Delta V_{\text{max}}$ can be reported as a function of measurement position. Different scenarios are then expected from the different patterns obtained for $\Delta V_{\text{max}}$ along the rebar (Figure 18):

- **Case A:** If $\Delta V_{\text{max}}$ is close to 0 in the middle of two well-defined— but not necessarily symmetric—peaks, it is the result of a much smaller anode than the probe spacing ($L_a << a$);
- **Case B:** As $L_a$ increases, the value of $\Delta V_{\text{max}}$ in the middle of the two peaks also increases. As indicated by further simulations (Figure S10), it becomes similar to the value of the peaks when $L_a \approx 1.5-2a$ for $e = 40$ mm;
- **Case C:** If $\Delta V_{\text{max}}$ is high over a large section with a maximum value close to its center, it is the result of an anode larger than the probe spacing ($L_a > 2a$). This indicates a large actively corroding section of the rebar; and
- **Case D:** If $\Delta V_{\text{max}}$ remains around 0, it is the result of uniform corrosion along the investigated area. It can either represent a fully passive rebar, a passive rebar with an anode with a low exchange current density, or an actively corroding rebar if the considered reference state also corresponds to active corrosion. If $\Delta V_{\text{max}}$ is similar along the rebar and different than 0, this indicates an actively corroding rebar with a passive reference state.

These scenarios only consider the presence of one anode in the investigated area, and different results are expected in the case of multiple anodes, depending on their relative positions. If they are far away from the current electrodes, they will not have any influence on $V_{P1-P2}$. If they are located below or near the current probes, all of them will influence $V_{P1-P2}$. However, it will not be possible to
differentiate each anode as the results will only reflect an average value of the contributions of each anode. It will not be possible to estimate the length of each anode, but it will always be possible to locate highly corroding areas and to classify them according to the values of $\Delta V_{\text{max}}$ and, thus, to their relative corrosion state. It is expected that comparing the results obtained with several probe spacings will yield further insight into the distribution of the anodic areas along the rebar.

![Figure 18. Schematic representation of the main scenarios expected when monitoring a rebar with one highly corroding area in concrete with uniform resistivity for $e = 40 \text{ mm}$. The figure shows the evolution of $\Delta V_{\text{max}}$ values as a function of the measurement position. Depending on the charge transfer coefficients, the two peaks observed in cases A, B, and C can either be symmetric or not. Case A can be expected even if concrete resistivity is not uniform, as the anode effect is almost insignificant irrespective of concrete resistivity when $L_e << a$.]

5.3. Comparison of Indirect GP with Indirect EIS

Alexander and Orazem showed the ability of indirect EIS to detect localized corrosion in post-tensioned tendons with uniform resistivity [31,32]. They concluded that highly corroding areas can be located if they are below the current-injecting electrodes or directly adjacent to an electrode array. In this case, two distinct time constants can be observed for non-uniform corrosion instead of only one time constant for uniform corrosion [32], attributing the high-frequency response to the corroding section and the low-frequency one to passive regions. Otherwise, almost no sign of active corrosion was detected, especially when the corroding area lay below the potential probes in Wenner configuration. With the indirect GP technique, we cannot draw such a conclusion from only one measurement as there is no reactance for this DC method. However, even with indirect EIS, several measurements are needed to determine the corrosion rate along rebar as the system is heterogeneous. Hence, this aspect cannot be considered as a drawback of the pulse technique for the monitoring of RC structures.

As already stated, Alexander and Orazem also concluded that the sensitivity of the indirect EIS method for localizing non-uniform corrosion depends upon the length of the highly corroding area, the magnitude of the exchange current density, and the resistivity of the cement-based material [32]. These conclusions also are valid for the indirect GP technique, showing the similarity of the results obtained with these two techniques (also shown by comparing Figure 4 and Figure S1C,D). The indirect GP method has the advantage of being less time consuming than the indirect EIS one, which seems more practical for the monitoring of RC structures. However, in the conventional three-electrode configuration, EIS is a powerful technique for mechanistic investigations of the corrosion process [86,87]. It is then expected that more information could be obtained from the indirect EIS method that might improve our understanding of the macrocell corrosion mechanism. The development of both techniques is then important for improving the monitoring of corrosion of steel in concrete.

5.4. Towards a Quantification of the Corrosion Rate in Non-Uniform Corrosion?

Though our study aimed at determining the ability of indirect GP to qualitatively detect localized corrosion, this technique may also be useful for a quantitative determination of the corrosion rate from the steady-state potential value. Most of the previous studies have proposed the use of an equivalent electrical circuit (EEC) for modeling the system in the four-electrode configuration [20,28,30,31,34,35]. For example, one EEC considers several resistances in series and parallel for representing the contact resistances, the resistance to current flow from the concrete alone (parallel to the rebar,
i.e., the bulk resistance), and the resistance in the path through the concrete perpendicular to the rebar (i.e., the surface resistance) [30,35]. The steel/concrete interface is represented by one resistance and one capacitance in parallel, as is traditionally done for modeling $R_p$ and $C_{dl}$, respectively, using the traditional Randles circuit. In this case, only an average value of the polarization resistances of the rebar can be obtained. This approach is then suitable for uniform corrosion as only one time constant is observed, as shown in References [31,32] and in Figure S1C,D on a passive rebar.

However, in the case of non-uniform corrosion, it should be more appropriate to consider a Randles circuit with at least two time constants for differentiating the two different areas, in agreement with the results from the indirect EIS technique [31,32]. This approach can be helpful for the qualitative location of highly corroding areas, depending on which EEC better fits the results. If it is the Randles circuit with only one time constant, then the anode has no influence on the potential response and cannot be detected. Conversely, if a Randles circuit with more than one time constant better fits the results, this indicates that the anode has an influence on the potential response and could be detected.

One of the main difficulties in assessing the corrosion rate is related to the importance of the charge-transfer coefficient $\alpha$ on the current distribution at the steel/concrete interface over time and on the conversion of the corrosion current density. This aspect is not specific to the four-electrode configuration as it already exists in the three-electrode one. Here, we considered only one value of $\alpha$ for the anode and one for the cathode, based on the recommended $B$ values (26 mV for active corrosion and 52 mV for passive corrosion) [60]. However, the $\beta_a$ and $\beta_c$ values reported in the literature can be inconsistent with this recommendation [88]. Thus, we considered hypothetical cases of uniform corrosion with different $i_0$, ranging from $10^{-5}$ to 0.1 A m$^{-2}$, either with $\alpha_a = 0.5$ ($B = 26$ mV) or $\alpha_a = 0.012$ ($B = 50$ mV). As shown on Figure S11, the anodic-charge coefficient also influences the steady-state potential in uniform corrosion. This is especially the case when $i_0$ and $\rho$ are increased, as a greater part of the current polarizes the rebar. Without knowing the charge-transfer coefficients, an error will be introduced during calculation of the corrosion rate. However, according to experimental values of $\beta_a$ and $\beta_c$ in the literature [5,48,49,88,89], this error should be a factor of 10 at most when the charge-transfer coefficients are unknown.

5.5. Implication for Practical Applications on RC Structures

This study provides theoretical insights on the indirect GP technique based on numerical simulations, but some perspectives for its application on existing structures can be defined. The indirect GP technique could be included in a general procedure aiming at evaluating corrosion [2]. First, a visual inspection is required for defining the locations for testing. Before making any measurements, the position and diameter of the rebar must be determined for determining the position of the electrical measuring device. This is generally done with ground-penetrating radar (GPR) or other electromagnetic techniques [90,91]. In addition to the rebar position, it is possible to estimate simultaneously the rebar diameter and cover thickness with high accuracy [92].

Currently, the most common electrical methods used for the assessment of corrosion are based on measuring the half-cell potential and concrete resistivity. It has been shown the good ability of using both techniques in combination for assessing the probability of corrosion [93,94]. However, these techniques do not provide a direct quantification of the effective corrosion rate. If commercial devices and recommendation exist for measuring the polarization resistance in real structures [54], its determination is still very rare in practice, mainly because it requires sophisticated equipment [95]. The indirect GP technique could thus be integrated in the overall monitoring strategy as (i) it allows the detection of anodic areas without any connection to the rebar, (ii) it provides surface potential and concrete resistivity, and (iii) it can provide quantitative information on the corrosion rate, as already demonstrated for uniform corrosion [9,35].

However, some parameters must be accurately defined as they must affect the measurement. First, two types of fluctuation can be expected when using electrical methods. The first are related to self-potential, i.e., electrochemical potential and streaming potential generated by the movement of
ions in concrete. Ideally, the potential electrodes should be non-polarizable (solid-state Ag/AgCl or Cu/CuSO₄ electrodes) with a small diameter to limit such disturbances. The self-potential variation must be considered accurately as, otherwise, it could be difficult to differentiate a steady state from a ‘quasi-steady-state’. The second type of fluctuation is related to electrochemical noise, causing stochastic pulses associated with the corrosion process, but its effect is more limited as the associated signal amplitude generally is low. To evaluate experimental noise and increase data quality, measurements must be repeated several times in the same place to obtain an average value.

Contact resistances between the electrodes and the concrete surface may also influence the measurement [66]. This parameter must be evaluated before any measurement by determining the imposed voltage between C₁ and C₂. To reduce these resistances, a good electrolytic contact is required, but special attention must be paid to ensure that all measurements are performed under comparable conditions. Otherwise, additional errors may be introduced in the measured potential difference because of variation in water and ionic content.

Finally, we only considered the presence of one rebar, and we defined the “longitudinal” sensitivity of the potential probes according to probe spacing. However, RC structures are composed of a rebar mesh. As several rebars can be polarized simultaneously, the “lateral” sensitivity of the potential probes must be assessed, by investigating the effect of the rebar spacing, to define the distance at which a highly corroding area on an adjacent rebar can influence the measurement over another rebar. As proposed above, it will be appropriate to analyze the results by comparing the relative values of ΔVmax, in three dimensions, to gain more insight into the position and distribution of the anodic areas along the different rebars.

5.6. Necessity of Electrical/Electrochemical Tomography

As shown in the different simulations, concrete resistivity has a major influence on the current distribution within the concrete volume and at the steel/concrete interface. An accurate determination of concrete resistivity is thus needed for correctly interpreting the results for assessing steel corrosion. Though, in our study, concrete resistivity is assumed homogeneously distributed in the concrete domain, we must recall that concrete is inherently heterogeneous by nature. To consider this point, an electrical resistivity tomography (ERT) is necessary [96,97]. Such measurements are generally done with a multi-electrode device for determining first the apparent resistivities at specific depths in the concrete volume. Pseudo-sections of the true resistivities are obtained with an inversion algorithm, which is not presented here. Several commercially available geophysical software systems exist for determining true resistivity values from apparent ones. However, as many reconstruction algorithms exist in electrical tomography, further investigations of RC structures are required so that inversion models will converge toward realistic pseudo-sections of true resistivities [98].

Still, many advances have been done for characterizing transport properties in cement-based materials, as highlighted in details in Reference [98]. For example, ERT can be used for determining the ingress of water and aggressive ions into concrete [99–103], which can be helpful for assessing both initiation and propagation stages of corrosion. However, as many phenomena are susceptible to occur simultaneously on RC structures, further studies are required to determine first the influence of each phenomenon separately in order to separate the contribution of each parameter on the resistivity profile. Notably, the influence of the variability in the degree of water saturation needs to be elucidated as it would theoretically lead to an infinite number of solutions obtained from Wenner data. One advantage that has been highlighted recently is the fact that ERT directly accounts for the rebar effect as this is no longer observed on true resistivity values [68]. Hence, the correcting factors presented in Section 5.1 are no longer required with this approach, and concrete resistivity can be determined irrespective of the sample geometry.

A similar inverse method must be developed to quantify the corrosion rate from the surface potential \( V_{P1-P2} \). This would allow to spatially distribute the variation of the corrosion rate to consider accurately the localized aspect of corrosion of steel in concrete. As concrete resistivity can be determined
with the instantaneous ohmic drop using the indirect GP technique, a single multi-electrode device can be used for determining the distribution of both resistivity in concrete and of polarization resistance along the rebar. Considering the inherent heterogeneities of both concrete and steel will improve the diagnosis of the service life of RC structures. The use of a multi-electrode device has other advantages for monitoring the corrosion rate. For example, when large RC structures must be investigated, measurements can be very time-consuming, and a multi-electrode device can simultaneously measure the potential difference between pairs of electrodes to reduce the total duration of measurement and to limit the variation of environmental factors. In addition, as shown previously, varying the probe spacing and position may help in locating small anodes and their position and estimating their length.

In this study, we only considered the Wenner configuration for locating highly corroding areas as this is the conventional configuration for measuring concrete resistivity. Though this configuration has a good sensitivity to vertical changes, it has a low sensitivity to horizontal changes [104]. In that case, a dipole-dipole array can be a useful alternative due to its good lateral resolution; this configuration is especially interesting for decreasing the rebar effect observed on the apparent resistivity when the measurement is done parallel to the rebar, as the potential probes are not directly located in the effectively polarized area. The obtained potential difference will then be attenuated compared to the one obtained in Wenner configuration, which will provide further insight into the size and position of highly corroding areas. In addition to these traditional electrode arrays, recent studies (e.g., References [105,106]) have shown the interest of optimizing electrode positions to control the image resolution, in order to improve the reliability of the measurements and the quality of the reconstruction process. Hence, by using several probe configurations and spacings, complementary data will be obtained for increasing the data quality and spatial resolution of the results.

6. Conclusions

The main conclusions are summarized as follows:

(1) Sample geometry is an important parameter when performing any experiments or numerical simulations. An accurate geometric factor must be determined case-by-case, depending on specimen size and probe spacing, to convert concrete resistance into concrete resistivity.

(2) The instantaneous ohmic drop is largely affected by the presence of rebar that acts initially as a short-circuit. This rebar effect decreases the value of concrete resistivity, especially when the measurement is done right above the rebar. It mainly depends on the probe spacing, the cover depth, and the geometry of the slab. It was also shown that the electrochemical state of the rebar has no influence on the rebar effect; thus, it is possible to calculate a corrective factor for an accurate determination of concrete resistivity irrespective of the corrosion state, which is the case when making measurements on RC structures.

(3) Contrary to the conventional GP technique in three-electrode configuration, the steady-state potential obtained with the indirect GP technique is not only representative of the polarization resistance but also of concrete resistivity.

(4) In non-uniform corrosion, $V_{P1-P2}$ increases slower as compared to passive corrosion. This is essentially due to the different capability of anodic and cathodic areas of consuming the impressed current. Hence, the anode has a greater effect on the transient potential than on the steady-state potential. Thus, it is preferable to examine the temporal evolution of $V_{P1-P2}$ to qualitatively detect the presence of a highly corroding area.

(5) The effect of the anode differs depending on its position relative to the monitoring device. It was shown that (i) corroding areas can mainly be located when they are below or in the vicinity of the current electrodes, and (ii) the area below the center of the device is almost not polarized irrespective of concrete resistivity. Hence, by adjusting the probe spacing, highly corroding areas will be either detected or not, depending on their position. This specificity should be helpful for
estimating the position and length of highly corroding areas, which is one of the main problems when making any measurements on RC structures.

Further experiments are required to confirm the different scenarios observed numerically in this study and to determine the applicability and efficiency of the technique for everyday practice in real structures. Future studies should also be focused on the development of electrical tomography with inverse methods to quantify the corrosion rates from the surface potential $V_{P1-P2}$ in non-uniform corrosion, in order to spatially distribute both concrete resistivities and corrosion rates in the material to improve the diagnosis of RC structures.

Supplementary Materials: Supplementary data are available online at http://www.mdpi.com/2624-5558/1/3/18s1. Table S1: Initial values of the current that polarizes the rebar and the potential difference $V_{P1-P2}$ corresponding to the ohmic drop, for each input parameters, and the resistivity calculated from $V_{P1-P2}$ and the total impressed current ($I_{C1} = 100 \mu A$) using Equation (6), considering the accurate geometric factor. Similar results were obtained in both active corrosion ($i_0 = 0.1 \ A \ m^{-2}$) and passive corrosion ($i_0 = 10^{-5} \ A \ m^{-2}$), which indicates that the instantaneous ohmic drop is independent on the electrochemical state of the rebar. However, a small difference between the two electrochemical states (3.3%) was observed for $a = 2.5 \ cm$, Table S1bis: Effect of the impressed current on the ohmic drop and the calculated concrete resistivity. Initial values of the current that polarizes the rebar and the potential difference $V_{P1-P2}$ corresponding to the ohmic drop, for each input parameters, and the resistivity calculated from $V_{P1-P2}$ and the total impressed current using Equation (6), considering the accurate geometric factor. In agreement with Ohm’s law, there is no effect of the value of the impressed current on concrete resistance, and thus on concrete resistivity, Figure S1: (A) Evolution of $V_{P1-P2}$ with time in the case of a small unreinforced specimen showing only an ohmic drop, indicating that the polarization observed on reinforced specimen is related to the rebar. (B) Evolution of $V$ with time obtained in the three-electrode configuration using the GP technique for the passive rebar with $l = 5 \ \mu A$. The results indicate that the quasi-steady-state is far from being reached after 100 s in this configuration. (C) Influence of the distance of the monitoring device from the rebar with $a = 15 \ \text{cm}$ using indirect EIS technique for the passive rebar. 0 cm = measurements above and parallel to the rebar. The further the monitoring device is from the rebar, the higher the measured apparent concrete resistivity and the lower the measured apparent polarization resistance. (D) Influence of the probe spacing on the potential response using the indirect EIS technique for the passive rebar. The increase in the probe spacing decreases the measured apparent resistivity (at high frequency) and increases the measured polarization resistance (at low frequency). Only a small difference is observed between 10 cm and 15 cm for the apparent resistivity due to the small size of the element, which affects the distribution of the current in the material (border effects), Figure S2: Evolution of $I_{rebar}$ and $V_{P1-P2}$ when considering uniform corrosion, in active ($i_{0,a} = 0.1 \ A \ m^{-2}$) or passive states ($i_{0,c} = 10^{-5} \ A \ m^{-2}$) with $C_{dl} = 0.2 \ F \ m^{-2}$, Figure S3: Evolution of emph$V_{P1-P2}$ and $I_{rebar}$ during 800 s for a passive rebar. Input parameters: $i_0 = 10^{-5} \ A \ m^{-2}$, $a = 5 \ cm$, $e = 40 \ mm$, $\Phi = 12 \ mm$, $C_{dl} = 0.2 \ F \ m^{-2}$, $I_{C1} = 100 \ \mu A$. The time required to reach the steady state increases when increasing concrete resistivity, Figure S4: Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a slightly corroding area at different positions relative to the monitoring device. Input parameters: $i_0,a = 10^{-4} \ A \ m^{-2}$, $i_0,c = 10^{-5} \ A \ m^{-2}$, $L_0 = 3 \ cm$, $a = 5 \ cm$, $e = 40 \ mm$, $C_{dl} = 0.2 \ F \ m^{-2}$, $I_{C1} = 100 \ \mu A$, Figure S5: (A) Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_0,a = 0.1 \ A \ m^{-2}$, $i_0,c = 10^{-5} \ A \ m^{-2}$, $L_0 = 1 \ cm$, $a = 5 \ cm$, $e = 40 \ mm$, $C_{dl,a} = C_{dl,c} = 0.2 \ F \ m^{-2}$, $I_{C1} = 100 \ \mu A$, (B) Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: emph$i_0,a = 0.1 \ A \ m^{-2}$, $i_0,c = 10^{-5} \ A \ m^{-2}$, $L_0 = 3 \ cm$, $a = 5 \ cm$, $e = 40 \ mm$, $C_{dl,a} = C_{dl,c} = 0.2 \ F \ m^{-2}$, $I_{C1} = 100 \ \mu A$, (C) Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0,a} = 0.1 A m^{-2}$, $i_{0,c} = 10^{-5} A m^{-2}$, $L_0 = 3 cm$, $a = 5 cm$, $e = 40 mm$, $C_{dl,a} = C_{dl,c} = 0.2 F m^{-2}$, $I_{C1} = 100 \ \mu A$, (D) Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0,a} = 0.1 A m^{-2}$, $i_{0,c} = 10^{-5} A m^{-2}$, $L_0 = 3 cm$, $a = 15 cm$, $e = 40 mm$, $C_{dl,a} = C_{dl,c} = 0.2 F m^{-2}$, $I_{C1} = 100 \ \mu A$, Figure S6: Influence of the impressed current $I_{C1}$ on $\Delta V$ when the anode is below $C_1$ for $\rho = 100$ or $200 \ \Omega \ m$. $\Delta V_{max}$ increases when increasing $I_{C1}$, which allows a better detection of the highly corroding area, Figure S7: Effect of the probe spacing on the distribution of the normal current density, evaluated here at 0.2 s, along the upper ridge of the rebar (the most polarized ridge). Input parameters: $i_{0,a} = 0.1 A m^{-2}$, $i_{0,c} = 10^{-5} A m^{-2}$, $\rho = 200 \ \Omega \ m$, $L_0 = 3 cm$, $a = 40 mm$, $C_{dl,a} = C_{dl,c} = 0.2 F m^{-2}$, $I_{C1} = 100 \ \mu A$, the figure (b) is a zoom on the central part of the rebar, where the length $L_a$ and position of the anode are indicated, Figure S8: (A) Evolution of $V_{P1-P2}$ $I_{rebar}$, $I_{anode}$ and $I_{anode/rebar}$ over time when the anode is below $C_2$. Input parameters: $i_{0,a} = 0.1 A m^{-2}$, $i_{0,c} = 10^{-5} A m^{-2}$, $L_0 = 3 cm$, $a = 5 cm$, $e = 40 mm$, $C_{dl,a} = C_{dl,c} = 0.2 F m^{-2}$, $I_{C1} = 100 \ \mu A$, (B) Evolution of $V_{P1-P2}$, $I_{anode}$ and $\Delta V$ over time when the anode is below $C_1$ or $C_2$ considering the same anodic charge transfer coefficient for anodic and cathodic areas ($\alpha_a = 0.012$). Input parameters: $i_{0,a} = 0.1 A m^{-2}$, $L_0 = 3 cm$, $a = 5 cm$, $e = 40 mm$, $C_{dl,a} = 0.2 F m^{-2}$, $I_{C1} = 100 \ \mu A$; (C) Evolution of $V_{P1-P2}$ $I_{anode}$ and $\Delta V$ over time when the anode is below $C_1$ or $C_2$, considering the same anodic charge transfer coefficient for anodic and cathodic areas ($\alpha_a = 0.5$). Input parameters: $i_{0,a} = 0.1 A m^{-2}$, $L_0 = 3 cm$, $a = 5 cm$, $e = 40 mm$, $C_{dl,a} = 0.2 F m^{-2}$, $I_{C1} = 100 \ \mu A$, Figure S9: Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence
of a highly corroding area centred between $P_1$ and $P_2$, for several length $L_a$. Input parameters: $i_{0,a} = 0.1 \text{ A m}^{-2}$, $i_{0,c} = 10^{-5} \text{ A m}^{-2}$, $a = 15 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl,a} = C_{dl,c} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$, Figure S10: (A) Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0,a} = 0.1 \text{ A m}^{-2}$, $i_{0,c} = 10^{-5} \text{ A m}^{-2}$, $L_a = 5 \text{ cm}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$; (B) Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0,a} = 0.1 \text{ A m}^{-2}$, $i_{0,c} = 10^{-5} \text{ A m}^{-2}$, $L_a = 10 \text{ cm}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$; (C) Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0,a} = 0.1 \text{ A m}^{-2}$, $i_{0,c} = 10^{-5} \text{ A m}^{-2}$, $L_a = 15 \text{ cm}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$; (D) Evolution of $V_{P1-P2}$ and $\Delta V$ over time in the presence of a highly corroding area at different positions relative to the monitoring device. Input parameters: $i_{0,a} = 0.1 \text{ A m}^{-2}$, $i_{0,c} = 10^{-5} \text{ A m}^{-2}$, $L_a = 40 \text{ mm}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$; (E) Influence of the anodic charge transfer coefficient of the cathode $\alpha_{a,c}$ on the evolution of $V_{P1-P2}$ over time in uniform corrosion. Left: $\alpha_{a,c} = 0.012$; right: $\alpha_{a,c} = 0.5$. Input parameters: $i_0 = 10^{-5} \text{ A m}^{-2}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$; (F) Influence of the anodic charge transfer coefficient of the cathode $\alpha_{a,c}$ on the evolution of $V_{P1-P2}$ over time in uniform corrosion. Left: $\alpha_{a,c} = 0.012$; right: $\alpha_{a,c} = 0.5$. Input parameters: $i_0 = 10^{-4} \text{ A m}^{-2}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$; (G) Influence of the anodic charge transfer coefficient of the cathode $\alpha_{a,c}$ on the evolution of $V_{P1-P2}$ over time in uniform corrosion. Left: $\alpha_{a,c} = 0.012$; right: $\alpha_{a,c} = 0.5$. Input parameters: $i_0 = 10^{-3} \text{ A m}^{-2}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$; (H) Influence of the anodic charge transfer coefficient of the cathode $\alpha_{a,c}$ on the evolution of $V_{P1-P2}$ over time in uniform corrosion. Left: $\alpha_{a,c} = 0.012$; right: $\alpha_{a,c} = 0.5$. Input parameters: $i_0 = 10^{-2} \text{ A m}^{-2}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$; (I) Influence of the anodic charge transfer coefficient of the cathode $\alpha_{a,c}$ on the evolution of $V_{P1-P2}$ over time in uniform corrosion. Left: $\alpha_{a,c} = 0.012$; right: $\alpha_{a,c} = 0.5$. Input parameters: $i_0 = 0.1 \text{ A m}^{-2}$, $a = 5 \text{ cm}$, $e = 40 \text{ mm}$, $C_{dl} = 0.2 \text{ F m}^{-2}$, $I_{C1} = 100 \mu \text{A}$.

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