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Cathodic protection criteria for buried carbon steel structures

Joint EFC/CEOCOR Working group

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1 Scope

In recent years a number of key factors have significantly influenced cathodic protection and the assessment of effectiveness. State of the art three-layer polyethylene coatings and up to date construction techniques in combination with an appropriate quality control system result in extraordinary high coating resistances and only limited numbers of small coating defects. Unless specific measurement techniques are used (e.g. coupon measurement), on such structures the assessment of effectiveness of cathodic protection according to EN ISO 15589-1 is no longer possible since the measurement at the metal / electrolyte phase boundary according to EN 13509 is only applicable on large individual coating defects with relevant ohmic drops in the soil. Additionally, the increasing ac interference observed on many structures is significantly impacting the assessment of effectiveness with conventional methods, since the separation of the decoupling devices is no longer possible for safety reasons and conventional survey techniques may become inapplicable. Moreover, ac corrosion has become a key concern with respect to the integrity of the structures and the thresholds in EN ISO 18086 (limited maximum tolerated protection current density) are conflicting with the requirements in EN ISO 15589-1 and EN 50162 (minimum required IR-free potential). Currently, demonstrating compliance to all these standards based on measurements on the protected structures may be difficult to achieve. The complexity of the interactions and the importance of the integrity of the protected structures calls for the assessment of the current state of knowledge, an assessment of the involved mechanisms and the corresponding interactions as well as the elaboration of suitable procedures that allow for using possible new protection criteria for the assessment of the effectiveness of cathodic protection. The scope of the present document elaborated in a joint EFC/CEOCOR working group is the summary of the theoretical background, the assessment of the corresponding model concepts and an outline on important issues that remain open for future discussions. These aspects could be used for the upcoming revision of EN ISO 15589-1.

2 Introduction

Cathodic protection (CP) has been systematically applied on pipelines since 1928 [1]. Despite the long and successful application of this technology, there is still significant controversy about the applicable protection criteria, the corresponding threshold values and the underlying mechanisms. While there is general agreement that the protection is achieved through electrochemical polarization, there was only limited consideration to the type of polarization and the respective characteristics.

Within this document the argumentation is illustrated by means of experimental data plotted into Pourbaix diagrams, as shown in Figure 1. It is relevant to note that these are constructed based on thermodynamic data. The Pourbaix diagrams are valid only in the absence of substances, which can form soluble complexes of insoluble salts with Iron [2]. The predominant species relevant for buried structures responsible for deviation from the shown behaviour are carbonates (as illustrated in Figure 10), sulphides and organic acids. In the last years various investigations with respect to cathodic protection have confirmed the understanding, that the pH is an important parameter in providing the corrosion protection [3-6]. In many cases relatively small protection current densities result in a significant increase of the pH-value at the steel surface of coating defects of pipelines. This increase of the pH favours the formation of a passive film, which significantly limits any further corrosion [4-6]. By means of model calculations, that are based on thermodynamic and kinetic data, all currently used protection criteria in CP can readily be explained when taking into account the relevance of the pH and the formation of a passive film [7-10]. Based on these considerations the bedding conditions and the associated mass transport at the steel surface is a key determining factor in the effectiveness of CP.



Figure 1: Polarization of steel with cathodic current according to Leeds [11]. Green: Concentration polarization of a well bedded coating defect; Red: Activation polarization of a poorly bedded coating defect. Solid lines delineating the thermodynamic stability of the various areas assume the conventional concentration of 10⁻⁶ M Fe²⁺.

An increase of the pH-value at the steel surface, however, can only occur, when the pipeline is bedded in fine sand and soil, or if the precipitation of calcareous deposits from water with increased hardness is taking place. Under these circumstances the convection of water at the coating defect will be limited and the hydroxide ions that are formed at the steel surface by the cathodic current entering the steel surface can only be transported by means of diffusion and migration. Hence, they will accumulate and the pH-value at the steel surface will increase. Under these conditions relatively small protection current densities are required to cause a change in concentration of hydroxide ions at the steel surface. This type of polarization corresponds to a concentration polarization as shown by the green line in Figure 1. In contrast, the presence of streaming and especially soft water, as it can be expected in rough bedding material, will result in a rapid dilution of the generated hydroxide ions. Alternatively, the increase of the pH can be limited by the activity of sulphate reducing bacteria (SRB) [3]. Note that recent RNA studies have resulted in many sulphate reducing species being re-classified as Achaea. So, instead of sulphate reducing bacteria, the technically correct term is sulphate reducing prokaryotes. The increase of the pH-value and the formation of the passive film will not occur under these circumstances. As a consequence, corrosion protection can only be achieved by activation polarization into the immunity domain. Since the immunity domain is cathodic of the hydrogen equilibrium potential line, the polarization towards or into immunity is only possible with highly increased current densities. This case is illustrated by the red line in Figure 1.

The discussion of the two extreme cases of polarization ignores the fact that any activation polarization will give rise to some concentration polarization and all concentration polarization will have some activation polarization. In Figure 1 the extreme cases of these two types of polarization are shown. For further differentiation the key differences between them are discussed below:

Limited mass transport due to bedding in soil and sand, or formation of calcareous deposits on coating defects resulting in concentration polarization:

- The required current density for the increase of the pH-value and the formation of a passive film is in the range of 1 $m\text{A}/\text{m}^2$
- After passivation of the steel, often a much higher current density will establish that is determined by external parameters, such as the on-potential, the diffusion limited oxygen reduction current and the spread resistance of the coating defect
- In the presence and maintenance of passivity an IR-free potential more positive than any conventional protection criterion (e.g. more positive than -0.85 V_{CSE}) does not have to be interpreted as corrosion
- In the presence of passivity, limited anodic interference has only a minor impact on the corrosion behaviour

In case of fast mass transport due to contact with streaming and especially soft water or neutralization of the formed hydroxide ions due to microbial activity, the polarization is achieved through activation polarization:

- A reduction of corrosion rate is achieved by shifting the IR-free potential in the negative direction, i.e. in the immunity domain
- The required current densities for achieving the immunity domain are in the range of 1 A/m^2 or higher due to the kinetics of the hydrogen evolution reaction
- IR-free potentials more positive than the protection criterion of -0.95 $V_{\mbox{\scriptsize CSE}}$ indicate corrosion

- In the case of activation polarization, anodic interference has a dramatic impact on the corrosion behaviour due to the potential being consistently in the area where soluble Fe²⁺ is thermodynamically stable.
- Under typical operation conditions of CP, it can be difficult to achieve protection of larger coating defects in higher resistive soil

The above list is not exhaustive. A more detailed discussion is given in [12]. These parameters demonstrate the importance of mass transport and hence the bedding of the pipeline. Effective CP is comparably easy to achieve on a well bedded pipeline where limited mass transport favours concentration polarization. In contrast, in the case of poor bedding conditions and fast mass transport, it can be very difficult to achieve sufficient corrosion protection by means of activation polarization. For larger coating defects in high resistivity soil unrealistically negative on-potentials may be required to achieve sufficient corrosion protection. This makes it questionable whether or not achieving immunity by means of activation polarization is possible in all cases.

For the design, control and effectiveness of CP systems and the requirements for coatings and backfill of pipelines it is of the utmost importance to have an understanding of the predominant and relevant controlling parameters. This is even more important for interference conditions. In order to define a generally accepted understanding of the relevant processes for the next revisions of the relevant standards this document summarizes the current state of the art.

3 Definitions and abbreviations

Some of the following definitions deviate from those in the current standards and codes. They are required for the discussion in this document to clearly distinguish time and place dependent effects.

Coating defect: Defect in the coating of the structure consisting of at least one electrode.

Earth: Conductive mass of the earth, whose electric potential at any point is conventionally taken as equal to zero

Homogeneous Electrode: Metal surface with homogeneous potential distribution. In this document for simplification the term "electrode" was used instead of "homogeneous electrode".

Electrode potential (E): A potential resulting from two or more electrochemical reactions occurring simultaneously on one electrode. The assessment may require sampling rates of up to 1 kHz in ac interference conditions.

IR-free potential (E_{IR-free}): Electrode potential free from any voltage errors caused by the *IR*-drop due to protection current or any other dc current. This must be measured with a frequency of less than 10 Hz. The IR-free potential (E_{IR-free}) is linked to the on potential (E_{on}) through the spread resistance (R), the electrode surface (A) and the current density (J_{dc}) according to equation (1).

$$J_{dc} = \frac{E_{IR-free} - E_{on}}{R \cdot A} \tag{1}$$

Note that the measured IR-free potential value will vary with the method of measurement; in particular with the sampling rate, which can significantly affect the measured value depending on the environment and especially in the presence of ac interference. Hence, the measurement procedure must be clearly documented. See CEOCOR WG D "off potential" report.

On-potential (E_{on}): Structure potential where the sum of all dc-currents on all the electrodes is different from zero (under effective CP the sum of the current is cathodic). The measurement is taken with a rate of less than 10 Hz and requires the following information: reference electrode position (e.g. earth, above structure, at relevant position), connection point to the structure and time.

Instant off-potential (E_{off}): Structure potential measured within 0.1 to 1 seconds after synchronized interruption of all the cathodic protection current sources .The potential value requires the following information: Time delay for taking the reading, reference electrode position (e.g. earth, above structure, at relevant position), connection point to the structure and time. In this condition, stray currents and equalizing currents flow on and off the structure. The sum of all these dc currents flowing on and off the structure is zero. However, for a structure with several coating defects this measured value will not be IR-free.

Spread resistance (R): Ohmic resistance through a coating defect to earth.

Structure: Coated metal surface consisting of more than one electrode.

4 Existing standards and procedures

4.1 Introduction

The discussion of the existing standards and procedures in this section is based on [13]. Currently the effectiveness of cathodic protection is assessed based on the following internationally accepted standards:

EN 12954:2001, "Cathodic protection of buried or immersed metallic structures - General principles and application for pipelines"

EN 14505:2005, "Cathodic protection of complex structures"

EN ISO 15589-1:2015, "Petroleum and natural gas industries - Cathodic protection of pipeline transportation systems - Part 1: On-land pipelines"

EN 15112:2006, "External cathodic protection of well casing"

NACE SP0169:2013, "Control of External Corrosion on Underground or Submerged Metallic Piping Systems"

The corresponding criteria and the associated protection criteria are discussed in the following.

4.2 IR-free potential

The IR-free potential (or polarized potential) has been widely accepted as a protection criterion and was incorporated in EN 12954 [14]. IR free potential ($E_{IR-free}$) is the structure-toelectrolyte potential measured without the voltage error caused by the IR drop due to the protection current or any other current. Threshold values are shown in Table 1. Note that all potential values in the relevant standards and in this document are in reference to saturated Copper/Coppersulfate electrode (CSE).

	Conditions	E _{IR-free} [V _{CSE}]
Aerobic conditions (Oxygen pre- sent)	Normal conditions: $\rho < 100 \ \Omega m; T < 40^{\circ}C$	-0.85
	Normal conditions: $\rho < 100 \ \Omega m; T > 60^{\circ}C$	-0.95
	Aerated sandy soils $100 \ \Omega m < \rho < 1000 \ \Omega m;$ T < 40°C	-0.75
	Aerated sandy soils $\rho > 1000 \ \Omega m, T < 40^{\circ}C$	-0.65
Anaerobic conditions (no oxygen present)	Aggressive soils, SRB	-0.95

Table 1:	Threshold values applicable for the IR-free potential
	····· •••···· • • • • • • • • • • • • •

The resistivity of the soil is indicated by the symbol, ρ . It is important to note that there are various threshold values for various conditions. The -0.95 V_{CSE} [2] and -0.85 V_{CSE} [4] thresholds are based on thermodynamic considerations. In contrast, the threshold values of -0.75 V_{CSE} and -0.65 V_{CSE} for soils of increased soil resistivity were determined empirically [15].

These values are identical with those in EN ISO 15589-1. Also, it is important to note the fact that these values apply to the IR-free potential of the steel of individual electrodes or coating defects only. It may not be confused with the instant off-potential. The determination of the IR-free potential according to EN 13509 is only possible by means of the so called intensive measurement or with coupons. A detailed description of the associated problems is given in the section below titled, "Problems associated with heterogeneous polarization".

4.3 Instant Off-potential

The instant-off potential is the structure-to-electrolyte potential measured immediately after interruption of all sources of applied cathodic protection current. It is often used as an approximation of the IR-free potential. However, it must be clearly stated that this simplification according to EN 13509 may only be made when there are no equalizing currents (such as in the case of a coupon). The source of the equalizing currents involves multiple factors. For example, different geometry and different soil conditions at various coating defects will result in different current densities and, therefore, different IR-free potentials of the individual electrodes. When interrupting all sources of cathodic protection current, equalizing currents will establish due to the different IR-free potentials at the various coating defects. These currents will cause IR-drops that are often significant. A detailed description of the associated problems is given in the section below titled, "Problems associated with heterogeneous polarization".

The instant-off potential of a structure only corresponds to the IR-free potential if it can be assumed that all electrodes of that structure are identically polarized. Naturally, this assumption is generally not fulfilled. Although used in many countries, the applicability of the instant off-potential, as an approximation of the IR-free potential, is still subject to debate, because it represents a relevant deviation from the requirements of EN 12954, EN ISO 15589-1 and EN 13509.

4.4 On-potential

The on-potential has been widely used for assessing the effectiveness of cathodic protection. It is still the primary protection criterion in EN 14505. According to EN 14505, a threshold value of -1.2 V_{CSE} is required for a soil resistivity less than 100 Ω m. The on-potential is also included in NACE SP0169 with respect to a threshold value of -0.85 V_{CSE}; however, with the comment that the IR drop needs to be considered. The on-potential was originally introduced by Kuhn [1] in combination with a threshold value of -0.85 V_{CSE}. Kuhn's theoretical considerations were that the natural potential of steel electrodes in soil cannot be more negative than -0.85 V_{CSE}. Applying an on-potential more negative than -0.85 V_{CSE}, therefore, results in a current entering into all electrodes.

4.5 100 mV cathodic polarization

The shift of 100 mV cathodic polarization criterion is described in NACE SP0169 and EN ISO 15589-1. The criterion is based on the fact that a shift of the IR-free potential to more negative values decreases the corrosion rate. With a shift of 100 mV a decrease of the corrosion rate by a factor of 10 can be expected. This effect is empirically demonstrated by von Baeckmann [14]. This concept requires homogeneous electrodes, which is also the case for the instant off-potential criterion.

This effect is considered in EN ISO 15589-1 by limiting its application to homogeneously polarized cases. This will be discussed further in Section 4.8.

According to EN ISO 15589-1 the application of the 100 mV polarization criterion should be avoided at higher operating temperatures, in soils containingSRB, or with interference currents, equalizing currents and telluric currents. These conditions should be characterized prior to using this criterion. Furthermore, the criterion should not be used on structures connected to or consisting of mixed metal components. Many countries do not measure this criterion because the expectation and assumptions indicated above are not likely met and sometimes unfeasible. This is the reason for the absence of this criterion in the EN 12954.

4.6 Tafel criterion

The Tafel criterion or log "I vs. E" criterion is based on increasing the protection current and recording the corresponding structure potential. At low current densities the oxygen reduction will occur, while at increased current densities hydrogen evolution will take place as discussed in more detail in Section 5.2.6 and illustrated with the blue lines in Figure 15. With the onset of hydrogen evolution, a change in slope is observed in the log I vs. E plot. From the technical point of view this criterion is ensuring that the CP system polarizes many electrodes of the structure to the hydrogen evolution. In EN 15112 this method is described for assessing the effectiveness of external cathodic protection of well casings.

4.7 Other criteria

The 300 mV polarization criterion is described in EN 14505 and referenced in NACE SP0169. It is the negative shift of the on-potential caused by the application of the protection current. The application of this criterion does not necessarily result in an on-potential more negative than -0.85 V_{CSE} . Therefore, it cannot be assumed that current will be entering into all coating defects. The on-potential criterion is therefore a more suitable criterion.

4.8 **Problems associated with heterogeneous polarization**

A key problem associated with instant-off potential and 100 mV shift in cathodic polarization measurements is the heterogeneous polarization of electrodes and coating defects. This is caused by heterogeneous soil resistivity, heterogeneous aeration, and by the different sizes of the coating defects. These differences among the various coating defects on a structure result in different current densities and, therefore, different IR-free potentials of all individual coating defects. This situation is described in Figure 2. When the protection current sources are interrupted (figure right), the differences of IR-free potentials among the various coating defects cause the flow of equalizing currents. The coating defects with better polarization (i.e., more negative IR-free potentials) will act as anodes that provide protection current to the less polarized coating defects. This is illustrated in Figure 2. The resulting equalizing currents in the case of interrupted protection current cause a potential value that is composed of the IR-free potential of all the coating defects plus the associated IR-drops due to the equalizing currents. After interruption of the external cathodic protection current the individual electrodes of the structure still experience polarization. As a consequence, they do not fulfil the definition for the IR-free potential. The instant off potential is therefore only the special case of an on-potential measurement where the external current flow to the system is zero. This very effect is the basis of the intensive measurement as described in EN 13509.

It must be clearly stated that a structure never has an IR-free potential, unless all electrodes exhibit the same IR-free potential. In this case there is not equalizing current and the instant off potential may according to EN 13509 be used for assessing the IR-free potential based on an instant off potential measurement, since identical IR-free potentials will result in zero equalizing currents.

As a consequence, the E_{off} of a pipeline with two coating defects with the individual IR-free potentials ($E_{IR-free}1$ and $E_{IR-free}2$) and spread resistances (R1 and R2) can be calculated according to equation (2) [16-19]

$$E_{off} = \frac{E_{IR-free} \mathbf{1} \cdot R\mathbf{2} + E_{IR-free} \mathbf{2} \cdot R\mathbf{1}}{R\mathbf{1} + R\mathbf{2}}$$
(2)

The consequence of the equalizing currents is the overestimation of the polarization of the less polarized coating defects in the instant off-potential measurement. Moreover this effect will significantly affect the 100 mV shift in cathodic polarization.



Figure 2: Effect of coating defect size and heterogeneous polarization on the offpotential. The situations with (left) and without (right) protection current are shown.



Figure 3:Data collected on a PE-coated pipeline. Both the 1 cm² and the 7 cm² coupon
consist of a rod shaped coupon with a circular coating defect.

The equalizing currents will cause a current to enter or leave the coating defects at interrupted protection current. Since there is a current flowing, the measurement does not represent an

IR-free potential. Due to this effect there are various countries that are not using instant offpotential and 100 mV polarization measurements. This is illustrated by the fact that the EN 12954 specifically states IR-free potentials and does not even mention the 100 mV potential shift criterion. Moreover, related limitations are stated in EN ISO 15589-1. This clearly shows that the current standards are consistent within themselves and that their competent application provides correct results. The related effects are described in full detail in [14].

As a consequence of the equalizing currents the IR-free potential of the coating defects on a structure can only be determined with the intensive measurement technique. According to EN ISO 15589-1 this is the simultaneously measured pipe-to-electrolyte potentials and associated perpendicular potential gradients. Hence, the intensive measurement is a concurrent close interval potential survey (CIPS) and direct current voltage gradient (DCVG) measurement. The IR-free potential is determined from collected data based on an extrapolation to the situation with zero current and consequently zero IR-drop. Since this method requires a relevant ohmic drop (typically larger than 25 mV) caused by the current entering the coating defect, it is not applicable in the case of increased pipeline depths (e.g. larger than 5m), at moderate on-potentials (e.g. more positive than -1 V CSE) and on modern state of the art structures coated with 3 layer polyethylene.

The problems associated with equalizing currents influencing the measured instant off-potentials are illustrated (Figure 3) by a field survey on a pipeline equipped with numerous coupons. The coupons consisted of bare surfaces of 1 and 7 cm². The IR-free potentials detected on the coupons are in dramatic contrast to the instant off-potential measured at the very same test post with the reference electrode positioned directly above the coupons. Despite electrical connection to the pipeline the instant off-potential measurement did not reflect the actual IRfree potential of the coupons.

This confirms that the IR-drops caused by the equalizing currents can be significant and significantly influences the conclusion of the entire survey. While it is often observed that IRfree potentials measured on coupons exhibit significantly different values than the instant offpotential measured on the pipeline, the results in Figure 3 represent an extreme case. This is due to the fact that the instant off-potential is dominated by largest and least resistive coating defects [16-19] according to equation (2). The obtained value for the instant off-potential is, therefore, to a large degree arbitrary and may be very misleading. It especially leads in many cases to a severe overestimation of the polarization of the coating defects in more resistive environment that generally receive smaller current densities. For this reason, EN 13509 does not allow the use of the instant-off potential measurement for assessing the IR-free potential in presence of equalizing currents.

4.9 Summary

The discussion of the various criteria has shown the considerable problems associated with the IR-free potential measurement. The full compliance with the required measurement of the IR-free potential of EN ISO 15589-1 is only possible with the intensive measurement or with coupons. Since the intensive measurement is limited to structures with relevant coating defect sizes, compliance of modern pipeline systems with EN ISO 15589-1 can only be demonstrated with coupons. However, if the bedding conditions and the mass transport exhibit the relevance as discussed in section 1 the installation of the coupons predominantly influences the result. In this view also the use of coupons needs to be reconsidered and the applicability of the EN ISO 15589-1 in the current form has to be fundamentally questioned.

5 The mechanism of cathodic protection

5.1 Introduction

The fact that the requirements of the current standards for assessing the effectiveness of cathodic protection cannot be fulfilled on modern pipeline systems the relevant difference between concentration and activation polarization as discussed in Section 1 have severe consequences on the application of cathodic protection and the assessment of its effectiveness. If there is a lack of understanding of the involved processes, it will not be possible to develop appropriate protection criteria.

5.2 Literature review

5.2.1 Introduction

In the light of these concepts it is therefore relevant to revisit literature and compare those results with the proposed concepts. The following discussion will be based on the various aspects of the current understanding [20].

5.2.2 Increase of pH and passivation with concentration polarization

All the relevant processes described in Figure 1 for concentration polarization were essentially discussed by Evans in 1923 for the corrosion of steel in heterogeneous aeration [21]. In order to illustrate his concept, the relevant effects are discussed by means of the Evans droplet test as shown in Figure 4:

A piece of steel with a water droplet on top will develop within a few minutes a localized corrosion process based on the reduction of oxygen according to reaction (3) and the oxidation of iron according to reaction (4).

$$O_2 + 2H_2O + 4e^- \to 4OH^-$$
 (3)

$$2Fe \to 2Fe^{2+} + 4e^{-}$$
 (4)

The heterogeneous aeration due to the different diffusion distances for oxygen causes an increase of the pH at the edges of the droplet while oxygen depletion in the centre will result in increased corrosion. This mechanism originally reported by Evans was summarized by Pourbaix [2] in 1974 as follows:

"When a piece of iron is immersed in a practically neutral non-buffered solution, which is aerated in one region (edges of the droplet) and not aerated in another (centre of the droplet), it is noticed that this differential aeration produces an increase in the corrosion rate in the non-aerated regions, and a decrease in the corrosion rate in the aerated regions, with a flow of electric current between these regions. On account of the increase of the pH due to the reduction of oxygen, the aerated regions will be passivated, and the non-aerated regions will not be passivated."



Figure 4: Schematic illustration of the Evans droplet test causing local increase of pH due to heterogeneous aeration and subsequent passivation (yellow).

This description by Pourbaix is in good agreement with the effects illustrated in Figure 1 for concentration polarization. In other words, the Evans droplet test is a case of cathodic protection by means of a galvanic anode in the form of a poorly aerated steel surface. This very concept of increase of pH and formation of a passive film is confirmed by the Max Planck Institute für Eisenforschung in the case of cut edge tests of galvanized steel [22]. It was found that the good corrosion protection caused by the limited cathodic current densities delivered by zinc can only be explained based on a pH increase and subsequent passivation.

In 1928 Kuhn introduced cathodic protection as a means to control corrosion on pipelines. He introduced the value of -0.85 V_{CSE} for the on-potential and stated for the current entering on the cathodic areas [1]:

"It causes a film of hydroxide to form, which protects these areas from corrosion"

The process of accumulation of hydroxide ions and increase of pH is most effective in nonbuffered solutions with hindered mass transport, since otherwise the increase of the pH is limited and passivation will be delayed. Evans indeed observed that the current density required for achieving cathodic protection is dependent on the convection of the electrolyte. At increased flow rates higher current densities are required to compensate for the dilution of the hydroxide ions, as Evans stated in 1946 [23].

Based on these observations it can be concluded that some of the first discussions of the electrochemical corrosion processes of steel already took into account the effects of change of concentration at the steel surface and the resulting passivation.

5.2.3 The problems associated with activation polarization

The key problem associated with activation polarization is the fact that the immunity domain of iron is cathodic of the hydrogen evolution equilibrium line and hence outside the stability domain of water. Polarizing into the immunity, therefore, requires significantly increased protection current densities. Achieving cathodic protection by means of activation polarization is, therefore, difficult and requires increased current densities. This problem was recognized by von Baeckmann in the first edition of his Handbook in 1971 [24]. The discrepancy between the theoretically expected high current densities with the comparably small ones observed in the practical application led him to the conclusion that the hydrogen evolution is kinetically hindered by an over-potential:

"The existence of such an over-potential, i.e. a kinetic inhibition for the hydrogen evolution, is an important precondition for the applicability of CP. Otherwise the hydrogen evolution

would represent an important potential barrier. Otherwise the required negative potentials for immunity could not be reached."

However, investigations have clearly shown that there is no relevant over potential for hydrogen evolution on steel [25]. Consequently in the third edition of the Handbook 1988 [26] this explanation is no longer found. Instead the mechanism of cathodic protection is described as follows:

- Immunity cannot be reached
- Hence the protection criteria can only be determined empirically
- The strong decrease of the corrosion rate with decreasing potential in oxygen containing solutions is caused by the formation of oxide covering layers with the involvement of OH^- and O_2

This description is in good agreement with the discussed effects in Figure 1. The polarization close to or into immunity is practically not possible and the change in concentration of hydroxide ions is considered to be at least relevant in certain cases. However, the fact that no model is proposed for the mechanism of cathodic protection has severe implications. If the criteria are purely empirical, they are only applicable for the very conditions under which they were determined. It follows immediately that the thresholds for the IR-free potential of -0.75 and -0.65 V_{CSE} of EN 12954 and EN ISO 15589-1 may only be applied in the case of coating defects that are well bedded in fine sand, since those were the very conditions under which these thresholds were determined [15]. Interestingly, for well bedded conditions in combination with concentration polarization these values can readily be confirmed by recent model calculations [7-10]. Therefore, these criteria in EN 12954 implicitly require optimized mass transport conditions and concentration polarization. The same applies for the current densities stated in the EN ISO 15589-1.

5.2.4 The IR-free potential and the pH-value

One of the first investigations with respect to the relevance of the threshold of $-0.85 V_{CSE}$ was performed by Schwerdtfeger and McDorman [27] in 1951. They observed that the corrosion potential of steel in anaerobic electrolytes lays between the equilibrium line for reaction (2) at $-0.95 V_{CSE}$ and the hydrogen evolution line (Figure 5). The values gradually decrease with increasing pH, reaching $-0.85 V_{CSE}$ at pH-values of 9. These results are in agreement with the expected corrosion potential of iron under hydrogen evolution [7-10]. They are moreover consistent with the line shown in Figure 1 for concentration polarization, since their data demonstrate that an IR-free potential of steel more negative than $-0.85 V_{CSE}$ is linked to a pH larger than 9. Moreover, these data demonstrate that the steel does not exhibit a relevant over potential for hydrogen evolution, since a kinetic inhibition for hydrogen evolution would result in more negative potentials. This has relevant technical implications: The electrodeposition of iron by polarization close to or into immunity is limited to few specific applications due to the required current densities well above 10 A/m². Hence, the very conditions that prevent an extensive application of electrodeposition of iron make the cathodic protection of steel by activation polarization close to or into immunity impractical [26].



Figure 5: Corrosion potentials of steel in anaerobic electrolytes as a function of pH replotted from [27]



Figure 6: The pH values obtained after polarization in mass transport limited electrolytes replotted from [28]. All corrosion rates were below 0.01 mm/year. Red: anaerobic; blue: aerated; green: oxygenated.

Another investigation related to the IR-free potential and the pH at the steel surface was performed by Thompson and Barlo in 1983 [28]. Steel samples were polarized in anaerobic, aerated and oxygenated electrolyte to either -0.92 or -1.12 V_{CSE}. The mass transport was limited by a porous ceramic, allowing for accumulation of hydroxide ions at the steel surface. Based on these measurements, the applied IR-free potentials resulted in a relevant increase of the pH and a limitation of the corrosion rate below 0.01 mm/year in all cases (Figure 6). Based on these data the corrosion protection was achieved by either passivity or immunity at increased pH.

They evaluated their data with respect to protection current density and pH at the steel surface [28] as shown in Figure 22. Additionally, data obtained in artificial soil determined at higher current densities are shown [4]. Current densities as low as 1 mA/m^2 are sufficient to provide an increased surface pH and passivation of the steel surface under conditions of hindered mass transport. Moreover, it is found that protection current densities in the range of 0.1 A/m² result in pH values around 12 under these conditions and, therefore, allow for concentration polarization.

In 1988 Freiman et al. [29] recognized the discrepancy between concentration and activation polarization and the lack of differentiation between these two cases:

"Modern theories of cathodic protection for underground steel structures assume that the metal always corrodes in the active state, and the role of cathodic polarization is reduced to a reduction in potential to those values with which the anodic dissolution rate becomes below the technical permissible value or it almost equals zero. As a rule no consideration is given to an increase in pH of the layer near the electrode at the metal surface (pHs) due to a reduction in the rate of formation of acid corrosion products with retention or even some acceleration of the cathodic discharge of O₂ molecules. However, with pHs \geq 10.7 and quite low concentrations of active ions (sulphates and chlorides) in the soil the metal should be converted into the passive state not despite cathodic polarization, but due to it."



Figure 7: Correlation between the protection current density and the surface pH under conditions of restricted mass transport and hence concentration polarization. Green: Thompson and Barlo [28]; red: Büchler and Schöneich [4].



Figure 8: Representation of the corrosion rate from field tests [30] in the Pourbaix diagram. The corrosion rates for the green dots are smaller than 0.03 mm/year, while the red dots exhibit higher corrosion rates.

This conclusion is in line with the effects discussed in Figure 1. They are confirmed by the extensive field investigations of CP and the related protection criteria performed by Barlo in 1994. Coupons were cathodically protected under defined conditions during 5 years in Australia, Canada and the USA [30]. In Figure 8 the IR-free potentials and the pH values calculated from the relationship in Figure 7 are plotted. The pH of the corroding coupons was estimated to be in the neutral domain. The results for the coupons with small corrosion rates (green dots) show a sufficiently increased pH for passivation. This explains the small corrosion rates even at IR-free potentials more positive than -0.85 V_{CSE}. Based on these data it is clear that the corrosion protection can readily be determined based on the pH value rather than on the IR-free potential. Interestingly, applying the same approach to the data collected in Germany between 2011 and 2013 [31] resulted in an even better correlation between pH and corrosion protection. As long as the average current density was cathodic (larger than 1 mA/m²) and the average on-potential was more negative than $-0.85 V_{CSE}$, no corrosion was observed. This good agreement between the field test [31] with the model concepts in Figure 1 can readily be explained by the special care that was taken with respect to bedding of the coupons: they were oriented upward by an angle of 45° to prevent loss of contact with the soil in the case of soil settling. Hence, the bedding conditions were ideal, the mass transport was highly limited and the build-up of an increased pH with concentration polarization was possible even at small current densities. This confirms the relevance of the mass transport for the corrosion behaviour.

The problems associated with a limited pH increase are nicely illustrated by the investigation of Kajiyama and Okamura in 1999 [3] as shown in Figure 9 The acid produced by the SRB required activation polarization in order to achieve protection, while in absence of SRB an increase of pH and protection through passivation was readily possible. These data are again

in good agreement with the proposed model concept in Figure 1 and can readily be explained by means of concentration and activation polarization, respectively.



Figure 9: The pH and IR-free potential dependence determined for soils with (red) and without (green) SRB according to [3].

The results of Schwerdtfeger and McDorman (Figure 5), of Thompson and Barlo (Figure 6) and Kajiyama and Okamura (Figure 9) demonstrate a link between the pH and the IR-free potential as suggested by Leeds in 1992 (Figure 1). Kasahara, Sato and Adachi [32] also demonstrated a similar relationship between the IR-free potential, current density and pH. In the case of limited mass transport (sand bedding) and absence of SRB the protection current as small as 1 mA/m² causes an increase of surface pH and subsequent passivation. IR-free potentials more negative than -0.85 V_{CSE} indicate an increase of the pH above 9 and the possibility of passive conditions under these circumstances.

5.2.5 The IR-free potential in the case of activation polarization

Most literature data can readily be explained based on the concept for concentration polarization. However, there are relevant factors affecting the corrosion behaviour that are often overlooked. Pourbaix has pointed out in 1960 [33] that the cathodic polarization of steel in buffered solution can initiate or accelerate corrosion. The naturally formed protective passive film or rust layer can be cathodically dissolved and, therefore, cause an increase of corrosion rates under insufficient cathodic protection. The investigation of Schwenk [34] has indeed confirmed strongly increased corrosion rates in the range of 0.3 mm/year caused by polarization of steel to IR-free potentials of -0.85 V_{CSE} in carbonate buffers of pH 9.5. In his test IR-free potentials more positive than -0.65 V_{CSE} limited the corrosion rate to irrelevant levels. This effect is schematically illustrated for a solution of pH 7 in Figure 10. It is relevant to note that in the presence of carbonates the corrosion domain is further extended

to higher pH values well above 10 as indicated with the blue domain for 1 M CO_2 [35]. Schwenk used concentrations in this range and confirms that no active domain is observed at pH-values of 11 [34].



Figure 10: Possible acceleration of corrosion due to cathodic activation polarization (red arrow) from the passivity into corrosion domain in a buffered near neutral electrolyte according to [33, 34]. In yellow the pH dependent protection criteria proposed by Pourbaix are shown [2, 33]. In blue the expected additional corrosion domain in 1 M CO₂ is shown [35]. The green circles represent the expected pH values based on the IR-free potential values of - 0.85 and -0.95 V_{CSE} in anaerobic conditions in the case of concentration polarization.



Figure 11: Polarization scans (cathodic currents with positive sign) in electrolytes with various pH values redrawn according to [36]

It must be assumed that the possible risk of corrosion acceleration due to activation polarization from the passivity into the corrosion domain has motivated Pourbaix to propose pH dependent protection criteria as indicated with the yellow line in Figure 10 [2, 33]. Pourbaix's approach is in important contrast to the requirements of EN 12954 and EN ISO 15589-1. Even the most negative threshold of -0.95 V_{CSE} for the IR-free potential can only be accepted according to his concept, if the pH increase is limited. In case of a pH increase above values of 11, which has to be expected at typical current densities of 0.1 A/m² based on the data in Figure 7, a threshold of -1.05 V_{CSE} would be required (according to the yellow line in Figure 10). Based on Pourbaix's concept, the application of the current standards would result in significant corrosion risks.

The absence of an active domain anodic of the immunity in the pH range of 9 to 14 is, therefore, of utmost importance for the application of cathodic protection. Based on the results in Figure 7, it has to be expected that all well bedded coating defects exhibit pH values from 9 to 13. Moreover, the EN 12954 and EN ISO 15589-1 require the IR-free potential to be in the range of -0.85 to -1.2 V_{CSE}. This happens to be the transition domain between immunity and passivity with a possible presence of an active passive transition. There is no literature presenting any relevant active dissolution in the corresponding potential and pH-domain, with the exception of carbonate solutions. Typical results from Heusler et al from1958 [36] are presented in Figure 11. Clearly no relevant anodic dissolution is observed in the pH range from 9.5 to 13. At pH 15 however, significant anodic dissolution is observed. These results are in line with Schmuki et al. [37] for measurements at pH 13.

In contrast, active dissolution has to be expected in the case of increased carbonate concentrations as shown in Figure 10. This thermodynamically expected domain is experimentally confirmed by literature [34, 38]. A certain level of carbonates has to be expected in any soil and a consumption of the carbonates by precipitation of siderite (FeCO₃) is expected. This may contribute to the formation of covering layers that will enhance the increase of pH and contribute to a concentration polarization. Nevertheless, in the case of streaming soft and especially buffered water any threshold more positive than -0.95 V_{CSE} for the IR-free potential under activation polarization cannot be justified based on the available literature.

In contrast, under mass transport limited conditions on well bedded coating defects a limited concentration of carbonates may be expected. The precipitation of siderite will rapidly decrease their concentration and favor the formation of a passive film. This is confirmed by the data of Schwenk that show a rapid decrease of the corrosion rate in his test with a limited electrolyte volume [34]. In the case of concentration polarization, the application of a threshold of -0.95 V_{CSE} for the IR-free potential indicates a pH increase to 11. Under these conditions protection even in increased carbonate concentration according to the green circle in Figure 10 may be expected.

5.2.6 Electrochemical description of the processes

The effects taking place under cathodic protection are discussed in the following by means of a calculation model [39, 40] that includes the kinetic and thermodynamic data [2] as well as the mass transport processes [4, 41, 42] in order to illustrate the relevant electrochemical factors.

The calculations are performed for a well bedded electrode surface of 10 cm² allowing for concentration polarization. Furthermore, a diffusion limited oxygen reduction current density of 0.1 A/m² was assumed. In Figure 12 the calculated corrosion potential of -0.78 V_{CSE} for steel in a neutral soil with a resistivity of 60 Ω m is shown. Based on the polarization scans,

the potential, and pH value in the Pourbaix diagram [2] it is clear, that corrosion will occur and that the potential of steel is controlled by the hydrogen evolution.

When applying a cathodic protection with an on-potential of $-0.82 V_{CSE}$ the IR-free potential and the pH-value at the steel surface are changed according to Figure 13. The cathodic current of significantly less than 1 mA/m² is not sufficient to result in corrosion protection. The potential and the pH are within the corrosion domain of the Pourbaix diagram. Decreasing the on-potential to a value of $-0.84 V_{CSE}$ results in a significant modification of the situation (Figure 14). In this case the increase of the pH is sufficient for the formation of a protective passive film, resulting in corrosion protection. It is found that a slight change in the on-potential of 20 mV results in an increase of the cathodic current to $0.1 A/m^2$. The detailed analysis of this effect shows that this current is not the cause of the corrosion protection, but the consequence of the passivity. This means that a slight increase in the protection current as shown in Figure 13 causes a slight increase of the pH at the steel surface above 9, which is sufficient to allow for the formation of the passive film. This formation of the passive film in return causes the control of the potential by the diffusion of oxygen rather than by hydrogen evolution.

This observation is crucial for the discussion of the significance of the protection criteria. The cathodic current density that is established after reaching the corrosion protection by means of passivity is a consequence of the passivity, the spread resistance, the diffusion limited oxygen reduction current and the on-potential. This explains why until today no agreement for a critical current density can be found in literature. Based on the model calculation the required cathodic current density for passivation under the given mass transport conditions is in the range of 1 mA/m^2 . However, the subsequently establishing current density is arbitrary, since it depends on external parameters.



Figure 12: E_{IR-free} calculated for a soil resistivity of 60 Ω m. Left: Pourbaix diagram with the red dot showing the conditions at the steel surface. The dashed horizontal line represents the protection criterion of -0.85 V_{CSE} according to EN ISO 15589-1 and the solid horizontal line the corrosion potential of -0.78 V_{CSE} that is expected under the given circumstances. Right: Polarization curves for anodic (red) and cathodic reactions (blue) on steel at the corresponding pH value based on published literature data [39, 40].



Figure 13: E_{IR-free} calculated for a soil resistivity of 60 Ω m. Left: Pourbaix diagram with the calculated values shown as a red dot. The dashed horizontal line represents the protection criterion of -0.85 V_{CSE} according to EN ISO 15589-1 and the solid horizontal line the on-potential of -0.82 V_{CSE}. Right: Polarization curves for anodic (red) and cathodic reactions (blue) on steel at the corresponding pH value. The length of the green line is showing the magnitude of the current density.



Figure 14: E_{IR-free} calculated for a soil resistivity of 60 Ωm and an on-potential of -0.84 V_{CSE}. Details to the plot are described in Figure 13.



Figure 15: $E_{IR-free}$ calculated for a soil resistivity of 60 Ω m and an on-potential of -1.3 V_{CSE}. Details to the plot are described in Figure 13.

The further decrease of the on-potential to -1.3 V_{CSE} causes an additional increase of the current density and the pH-value as can be seen in Figure 15. In this case the current of 0.5 A/m² is controlled by the hydrogen evolution. The IR-free potential of the steel surface is still in the passivity domain.

The discussion shows that the protection criterion in EN ISO 15589-1 for normal conditions can be explained with the equilibrium line for hydrogen evolution that is reaching the passivity domain in the Pourbaix diagram at a potential value of -0.85 V_{CSE} according to [4]. Hence, potentials more negative than -0.85 V_{CSE} are bound to exhibit a sufficient increase in pH to provide passivity. At increased current densities it is possible that the immunity range is reached at elevated pH-values. The IR-free potential in the discussed cases was either controlled by the diffusion controlled oxygen reduction (Figure 14) or hydrogen evolution (Figure 15). Interestingly, in the calculated configuration an on-potential more negative than -0.85 V_{CSE} results in corrosion protection, which is in line with the observations of Robert Kuhn [1, 43].

5.2.7 Conclusion

Based on literature achieving corrosion protection by activation polarization close to or into immunity is difficult if not impossible. In contrast, there is wide agreement that cathodic protection is achieved by an increased pH and subsequent passivation. The second edition of the standard CP textbook by Peabody [44] from 2001 nicely summarizes this observation:

"The concepts presented for CP (activation polarization) are fundamentally correct at the instant that CP is applied but are too simplistic to consider the time-dependent behaviour of a cathodically protected underground structure..... This pH increase (concentration polarization) is beneficial because the corrosion rate of steel decreases with increasing pH, even under freely corroding conditions. The decrease in corrosion rate is the result of the formation of a protective oxide film on the metal surface in the elevated pH environment, a process referred to as passivation."

This view is confirmed by more recent work [45-47]. Based on the review of literature the requirements in EN 12954 and EN ISO 15589-1 can only be explained by means of concentration polarization. If the cathodic protection would be based on a shift of the IR-free potential into or towards immunity, no threshold for the IR-free potential more positive than -0.95

 V_{CSE} could be tolerated. More positive values would represent a significant corrosion risk. The fact that the more positive threshold values -0.85, -0.75 and -0.65 V_{CSE} have been successfully applied in the past years, confirms the fact that corrosion control is readily possible at these values and that concentration polarization is typically the predominant process. Denying the relevance of the pH-increase and concentration polarization implicitly puts the technical relevance of the current standards into question.

5.3 Consequences on criteria

5.3.1 Introduction

Based on the above discussion of the literature, the effect of concentration polarization, the increase of pH and the subsequent passivation for the corrosion protection of steel was demonstrated. The discussion of the threshold of $-0.85 V_{CSE}$ and the relevance of the onpotential versus an IR-free potential has continued since Kuhn's original publication in 1928. The corresponding literature is revisited with the proposed concept of activation and concentration polarization. This discussion is not only relevant for the future work on the corresponding standards, but significantly affects the protection criteria for stray current corrosion.

5.3.2 The significance of the on-potential

Kuhn has proposed in 1928 [1] an on-potential (E_{on}) criterion of -0.85 V_{CSE}. He identified the galvanic corrosion among differently aerated coating defects as the reason for the high corrosion rates. He found that applying a sufficiently negative on-potential ensures the compensation of these galvanic couples and a strong limitation of the corrosion process. Based on the above discussion his conclusion was technically sound and correct. The work of Schwerdtfeger and McDorman [27] has clearly shown that the IR-free potential ($E_{IR-free}$) of a steel surface corroding even under anaerobic conditions is always more positive than -0.85 V_{CSE} unless the pH is increased above 9. In most electrolytes such an increase of pH is sufficient for passivation. Based on equation (5) it must therefore be concluded that any on-potential more negative than -0.85 V_{CSE} is bound to result in a cathodic current I on an individual coating defect. This conclusion is independent on the spread resistance R and hence the soil resistivity.

$$I = \frac{E_{IR-free} - E_{on}}{R}$$
(5)

Taking into account the results in Figure 7 the current density for achieving pH 9 and hence passivation is as small as 1 mA/m^2 in the case of hindered mass transport and concentration polarization. At this current density negligible ohmic drop will be obtained even with large coating defects and increased soil resistivity. This explains the good success of Kuhn's criteria based on an on-potential and underlines its technical relevance.

It is argued that the very low soil resistivities in the Mississippi Delta, where Kuhn ran his experiment, have resulted in negligible ohmic drops [48]. This argument ignores the fact that Kuhn was working with coal tar dipped coated pipelines. The high current demand required by the comparably large metal surface area negates the effects upon the magnitude of the ohmic drop caused by the low soil resistivity in the range of 6 Ω m. However, when considering the small current densities required for protection, Kuhn's success with the on-potential criterion can readily be explained on the basis of concentration polarization. It might be relevant to consider that he performed his work on cast iron pipes. The limited mechanical properties

of this material required careful bedding of the pipelines and it may be assumed that this has favoured a limited mass transport and the build-up of an alkaline environment.

It follows that the on-potential is sufficient as a protection criterion in the case of a well bedded coating defect with limited mass transport allowing for concentration polarization in the case of homogeneous current distribution within the individual coating defects. This was indeed confirmed in a field test in Germany, where no relevant corrosion was found if the on-potential averaged over 24 hours was more negative than -0.85 V_{CSE} in the case of coating defects of 1cm^2 surface [31]. Consequently, the average current density was cathodic, and the pH could increase on the specifically well bedded probes with limited mass transport allowing for pH increase and passivation.

5.3.3 The significance of the IR-free potential

While the on-potential is a suitable method to ensure current entering into the coating defects, the IR-free potential in the case of concentration polarization corresponds to a pH measurement. This was already demonstrated by Schwerdtfeger and McDorman [27]. It could be argued based on Figure 1 that under concentration polarization (green line) an IR-free potential of -0.85 V_{CSE} corresponds to a pH 9 and IR-free potential of -0.95 V_{CSE} corresponds to a pH 9 and IR-free potential of -0.95 V_{CSE} corresponds to a pH 9 and IR-free potential of -0.95 V_{CSE} corresponds to a pH 9 and IR-free potential of -0.95 V_{CSE} corresponds to a pH 9 and IR-free potential of the green circles. Hence, these two empirically observed criteria for the IR-free potential that were already reported by 1957 [49] can readily be explained. Within this context also the maximum negative value of -1.2 V_{CSE} for the IR-free potential stated in EN ISO 15589-1 can be explained. This corresponds to a pH value above 13, where issues with cathodic disbondment have to be expected. It is relevant to note that this threshold has limited relevance with respect to activation polarization, since activation polarization is characterized by a lack of increase in pH. Limited issues with disbondment of the coating have to be expected in the case of activation polarization.

Within this discussion it is crucial to understand that the threshold values of -0.85, -0.95 and -1.2 V_{CSE} are only applicable for an estimation of the pH value at the steel surface, if the IRfree potential is controlled by hydrogen evolution as indicated by the green line in Figure 1. In contrast, if the IR-free potential is a result of the activation controlled oxygen reduction, as has been discussed in [7-10], significantly more positive values in the range of -0.65 or - $0.75 V_{CSE}$ may be expected as observed by Funk et al. in 1987 [15] in well aerated soils. This explains the corresponding values in EN 12954 and EN ISO 15589-1 for high resistive well aerated soils. However, under these conditions the IR-free potential may not be used to determine the pH with respect to protection levels or over-protection. In the case of concentration polarization, where CP is resulting in an increased pH and the formation of a passive film, it must be concluded that the IR-free potential is irrelevant for judging the corrosion protection of a passive surface [7-10]. This is best illustrated in Figure 8: The judgment of the corrosion situation is best done based on the pH value rather than the IR-free potential. In low resistive or poorly aerated soils the IR-free potential depends on the pH and may, therefore, be used to demonstrate the protection. Under well aerated conditions the IR-free potential does no longer correlate with pH and it is, therefore, irrelevant to draw any conclusion with respect to the corrosion situation.

Whenever this build-up of an increased pH is not possible due to a high removal rate of the formed hydroxide ions, such as in soft streaming water or in presence of SRB (see Figure 9), an activation polarization is required at significantly increased current densities. In this case only an IR-free potential of at least -0.95 V_{CSE} can assure sufficient corrosion protection by polarization close to or into immunity. Not meeting this threshold in the case of activation polarization is inherently linked to corrosion.

5.3.4 Consequences

The discussion of the IR-free potential (required in EN 12954 and EN ISO 15589-1) as well as the on-potential (required in EN 14505) demonstrates the relevance of these two criteria under the respective circumstances. Kuhn's approach with the on-potential is technically sound and valid until today in the case of concentration polarization for well bedded structures. However, it was bound to fail in the cases where the pH could not increase. Corrosion damages during the 1980's in Germany on a poorly bedded pipeline caused by SRB resulted in the introduction of the IR-free potential criterion in Europe in combination with a threshold value of -0.95 V_{CSE}. Based on the above discussion this is technically correct and justified.

While the approaches developed in the past are well justified, they lead to a key problem when it comes to determine the effectiveness of cathodic protection: As long as the bedding and the mass transport conditions of individual coating defects are not known it is very difficult to justify a protection criterion other than an IR-free potential in combination with a threshold value of $-0.95 V_{CSE}$. This assures corrosion protection in case of activation polarization to immunity or concentration polarization to a pH value of 11. However, not meeting this criterion in the case of concentration polarization does not indicate a corrosion risk, in the case of activation polarization it does.

5.4 Consequences on interference conditions

5.4.1 Introduction

In the above discussion the presence of ac and dc interference has been excluded. However, for the practical application their effect on the corrosion process is of utmost importance. It is beyond the scope of this document to treat these issues in detail. For simplification, the following assumptions were made:

- The reference electrode is placed above the structure
- The structure is coated with a state of the art three-layer PE coating with a coating resistance of at least 1 $M\Omega m^2$ exhibiting individual coating defects that are more distant than the structure depth
- The distance between the interfering source and the pipeline is significantly larger than the structure depth. As a consequence, the distance from the reference electrode placed above the coating defect is smaller than the distance between two neighbouring coating defects.

In the following only the most relevant effects of the individual parameters will be addressed. They are based on the available literature and on the proposed model concepts.

5.4.2 AC interference

Based on EN ISO 18086 there is general agreement that ac interference is caused by increased ac and dc current densities. This is especially critical in the case of concentration polarization, since the buildup of an increased hydroxide concentration at the steel surface and its distribution into the surrounding soil causes a relevant decrease of the spread resistance of the coating defects. Under given interference conditions the increase of pH and the drop of the spread resistance leads to increased current densities and, therefore, to an increased corrosion risk. The influencing processes according to the principles given in EN ISO 18086 are described in [50] in more detail. In the current understanding of the process it is the alternating formation and dissolution of the passive film within the transition between immunity and passivity at pH values around 13 that are responsible for the ac corrosion process [51]. So far there is no known case of ac corrosion under activation polarization conditions, which may be explained as follows: Based on theoretical considerations a short term decrease in cathodic protection current density and especially anodic currents caused by ac interference are expected to cause relevant corrosion damage as well. The key advantage in the case of activation polarization is the lack of pH increase and the absence of a decrease in spread resistance, which limit the risk of increased ac current densities and, hence, the risk for corrosion damages. This is indeed confirmed for the rails in the wet Simplon tunnel atmosphere. This was in absence of CP and therefore lack of pH increase where the ac traction return current was required to cause relevant corrosion of the rails [52].

Due to the relevant effect of pH and spread resistance, the corrosion rate is strongly controlled by the on-potential. More negative on-potentials result in higher pH at the steel surface, lower spread resistance and therefore higher ac corrosion rates. This makes the ac corrosion process for unknown mass transport conditions critical in the case of both, anodic and cathodic, dc interference.

5.4.3 Constant DC interference

In the case of constant dc interference, as it may be caused by a foreign CP system, the gradients in the soil will, depending on the polarity of the interference, either decrease (anodic interference) or increase (cathodic interference) the protection current density on a given coating defect. In the case of an anodic interference, the above discussion allows for some conclusions depending on the mass transport conditions and the predominant type of polarization:

In the case of concentration polarization, the on-potential measured above the struc-• ture must be at least more negative than -0.85 V_{CSE}. For practical reasons it was found that a safety margin is necessary and that a value of $-1 V_{CSE}$ should be used [53]. Under these conditions a cathodic current will be flowing towards the coating defect, resulting in an increased pH and passivity if concentration polarization is sufficient. If the ideal conditions of bedding, leading to adequate concentration polarization are not in place, corrosion remains possible. In the case of activation polarization any shift of the on-potential more positive than the minimum required value will cause an inherent risk of corrosion. With the Tafel slope of less than 100 mV/dec for the iron dissolution reaction (2) even a small shift of the on-potential to a less negative level can result in relevant corrosion [54]. Corrosion has to be expected already in the case of a decreased protection current density in the coating defect of less than 1 A/m^2 [30].As a consequence, anodic interference under activation control must be compensated by shifting the on-potential more negative, with all the negative consequences on ac corrosion risk on the corresponding well bedded coating defects exhibiting concentration polarization.

In the past not much attention was paid to cathodic interference. However, in combination of ac interference a cathodic interference (e.g. caused by an anode) can dramatically increase the corrosion risk of the structure as follows from the requirements in EN ISO 18086.

5.4.4 Time variant DC interference

In case of interference caused by dc traction systems, typically both, anodic and cathodic potential excursions are observed when the potential is monitored against a reference electrode placed above the structure. In a first approach the time variant stray current interfer-

ence could, therefore, be considered as low frequent ac interference. This has important consequences that are ignored when only focusing on anodic potential excursions as it is currently required according to EN 50162.

In the case of concentration polarization with an increased pH in the coating defect and a subsequently formed passive film, the shift of the IR-free potential in the anodic direction does not represent an imminent corrosion risk as follows immediately from Figure 8. In the first edition of Peabody's "Control of Pipeline Corrosion" from 1967 it is stated: "When steel is immersed in a sufficiently caustic solution with a pH around 11 and higher, it can be made to discharge current without appreciable metal loss" Similar effects are reported by [6, 28, 55-57]. Nagayama and Kawamura [58] have shown how current may be passed through a steel surface by means of the redox system Fe²⁺/Fe³⁺ in the presence of a passive film with a very high efficiency. Schmuki et al. [37] have also shown how the repetitive and reversible Fe²⁺/Fe³⁺/Fe²⁺ transition cycle can occur. The data demonstrates that in the presence of a passive film it is possible to observe anodic activity with no oxidation of Fe provided the potential does not become sufficiently cathodic to dissolve the passive film through electrochemical reduction.

Figure 16: Influence of the chloride ion concentration on the pitting potential. Chloride concentration from 0.001 M (green), 0.01 M (yellow), 0.1 M (orange) to 1 M (red) [59].

In chloride containing environments the initiation of pitting corrosion could in principle occur, but detailed investigations have shown that the redox system Fe²⁺/Fe³⁺ does not only consume large amounts of the passed anodic charge [4], but also limits the anodic excursion of the potential to values within the grey area in Figure 16. This makes the initiation of pitting corrosion less probable. Notably, this grey area is cathodic to the most negative possible pitting potential in solutions with pH-values above 11 even with very high chloride concentrations of 1 M [59] as shown in Figure 16. The chloride ion concentration will usually always be far less than 1 M under all practical

conditions (e.g. seawater has an approximate molarity of 0.48 M) and hence the pitting potential will in most cases be more positive than the gray area. Much lower chloride ion concentrations will usually occur in soils and the pitting potential becomes more positive in Figure 16 as the chloride ion concentration decreases from 10^{0} to 10^{-3} M. Hence the anodic excursion required for the initiation of pitting at low chloride concentrations or high pH-values is often insufficient.

In the case of concentration polarization, the only key requirement is a sufficiently increased pH value at the steel surface. This is achieved by a current passing to the steel surface that is on average cathodic (e.g. averaged over 24 hours). Important anodic potential excursions can be tolerated, if their duration does not substantially decrease the surface pH. In a first approach a net cathodic current can be assured with an average on-potential that is more negative than -1.2 V_{CSE}. Shifting this value more cathodic will increase alkalinity and increase the tolerance with respect to prolonged anodic potential excursions.

• In the case of activation polarization cathodic excursion cannot compensate for anodic current discharge. Furthermore, a rust layer containing the redox buffer Fe²⁺/Fe³⁺ cannot form due to the solubility of Fe²⁺. Hence, even decreasing the protection current density can cause an imminent corrosion risk and anodic current discharge is not acceptable, since this will result in iron oxidation to dissolved Fe²⁺. Even IR-free potentials more negative than -0.85 V_{CSE} and the absence of on-potentials more positive than this value can already cause relevant corrosion. Based on [54] even minor anodic shifts of the IR-free potential result in relevant acceleration of the corrosion process. In this respect anodic excursions must be strictly limited in the case of activation polarization.

In the case of time variant DC interference, the expected corrosion behaviour is significantly dependent on the type of polarization and the mass transport conditions at the steel surface.

5.4.5 Conclusions

The above discussion clearly demonstrates the importance of the bedding conditions and the resulting mass transport at the coating defect surface. In the case of concentration polarization on a well bedded coating defect severe ac and dc interference conditions are manageable and controllable. In contrast, in poor bedding conditions under activation polarization interference is critical. In the case of combined ac and dc interference, mitigation of the corrosion risk may become very challenging.

5.5 Consequences of the model concepts

The above discussion has highlighted some of the relevant processes taking place under cathodic protection. The discussion of the processes has a focus on individual coating defects. The presence of different coating defects with strongly different mass transport characteristics on the same structure has important consequences that need to be addressed. It may technically be achievable to ensure that all coating defects are well bedded at pipeline construction. Alternatively, some defects in poor bedding conditions will be in electrolytes that enable calcareous deposits to form and these will result in and enhance concentration polarization. However, it cannot be certain that all coating defects in poor bedding will develop calcareous deposits. Therefore, in typical cases for a pipeline it is possible that a large number of coating defects will exhibit concentration polarization, but a smaller number can only be protected by means of activation polarization. The relevant implications are discussed below: **Cathodic disbondment:** Achieving sufficiently high current densities for activation polarization on the poorly bedded coating defects is inevitably going to cause over-protection on at least some of the well bedded coating defects. It is technically not possible to ensure IR-free potentials more negative than -0.95 V_{CSE} on the poorly bedded ones and guarantee IR-free potentials more positive than -1.2 V_{CSE} on the well bedded ones. In this respect the requirements of EN ISO 15589-1 cannot be fulfilled in the case of concentration and activation polarization on the same structure.

The IR-free potential to measure the surface pH: The above discussion has shown that the IR-free potential measurement is a suitable method for assessing the surface pH of a coating defect exhibiting concentration polarization as long as the IR-free potential is controlled by hydrogen evolution. This is explaining the relevance of this criterion. Problems occur in the case of increased aeration, increased coating defect size and increased soil resistivity, when polarization to the hydrogen evolution is not possible under typical operation conditions of CP. Often this leads to a shift of the on-potential to more negative values by increasing the current output of CP-rectifiers. This will further shift the IR-free potential to more negative values, increase current density and surface pH resulting in a risk of over-protection. This is due to the fact that the threshold of -1.2 V_{CSE} represents a pH measurement as well, which is not applicable in the case of well aerated conditions. As a consequence, the IR-free potentials can be well more positive than -0.85 V_{CSE}, but CP has led to critical pH values at the steel surface resulting in cathodic disbondment.

The application of -0.85, -0.75 and -0.65 V_{CSE}**:** These values specified by EN 12954 and EN ISO 15589-1 are only applicable in the case of concentration polarization. They will not provide corrosion protection in the case of activation polarization. Under unknown mass transport conditions, it is difficult to justify their use.

The on-potential: The on-potential does not deliver any information with respect to an individual coating defect. However, average values more negative than -1.2 V_{CSE} , as required by EN 14505, ensures an average cathodic current to enter every individual coating defect. This can be considered a first minimum requirement for an effective CP. In the case of limited mass transport, build-up of an increased pH and passivation this will ensure corrosion protection in many cases. This explains the collectively good experience with CP, since most structures fulfil this on-potential requirement and have limited mass transport at most of their coating defects.

Interference conditions: Based on the above discussion, combined ac and dc interference can only be managed if all the coating defects can be protected by means of concentration polarization. Otherwise, severe anodic and cathodic interference in combination with high ac interference become practically impossible to mitigate.

6 New concepts for protection criteria

6.1 Introduction

The application of cathodic protection (CP) is generally statutory for high pressure gas- and oil-pipelines. Moreover, CP is frequently used for water pipelines and buried containers. The relevant underlying principles and protection criteria are stated in EN ISO 15589-1:2017. In the last years various investigations with respect to CP have raised the understanding, that the pH is an important parameter in providing the corrosion protection [3-6, 60, 61]. In many cases relatively small protection current densities in the range of a few tens of mA/m² result in an increase of the pH-value at the steel surface of coating defects of pipelines. This increase of the pH favours the formation of a passive film, which significantly limits any further corrosion [4-6, 55, 60, 61]. By means of model calculations, that are based on thermodynamic and kinetic data, all currently used protection criteria in CP can readily be explained when taking into account the relevance of the pH and the formation of a passive film [7-10]. Based on these considerations the accumulation of alkalinity and hence the mass transport at the steel surface are key determining factors in the effectiveness of CP. These aspects and the associated implications are discussed in detail by Angst et al [62]. Today's understanding of the mechanisms involved in CP provide a new view on the protection criteria and the processes taking place during interference conditions. This far reaching understanding has influenced the criteria in EN ISO 18086 for the assessment of AC corrosion and the work on ISO 21857 with respect to anodic stray current interference.

The relevant aspects involved in cathodic protection and the associated implications are presented. The physical and chemical significance of the protection criteria in EN ISO 15589-1 are explained and the implications with respect to the assessment of the effectiveness of CP are highlighted. Furthermore, the processes taking place under anodic stray current interference are discussed. These aspects are currently integrated in ISO 21857 and will provide a simple and straightforward approach to the assessment of the corrosion risk of pipelines under time variant stray current interference. Additionally, the implications of over polarization and the resulting AC corrosion risk will be discussed in the context of EN ISO 18086.

The far-reaching understanding of the mechanisms taking place during cathodic protection have led to conclusive models with respect to interference conditions. Based on these concepts it is today possible to determine the effectiveness of CP based on a new approach that is presented in the following. This approach is based on the documents DVGW GW 21, GW 28 and GW 28 B1. In Germany these documents have the status of a recognized state of the art. The main advantage of this documents is their applicability on all types of pipelines independent on their interference conditions. Based on them it is possible to assess the effectiveness of CP of AC and DC interfered pipelines based on the readily measurable on-potential and AC voltage, rather than the IR-free potential that can typically only be determined by means of coupon measurements on most modern pipelines.

Within this section a description of the underlying mechanism and a discussion of the contribution of the interference conditions is presented. Additionally, a description of the associated controlling parameters and the influencing factors is presented. Furthermore, a proposal for their quantification is given and examples for the field application is presented. These aspects form the basis for future discussion and elaboration.

6.2 The underlying mechanism

There is general agreement that the effectiveness of CP is based on polarization of the steel surface [1, 63, 64]. However, the actual chemical process that causes the polarization is often not further discussed despite of its important consequences on various aspects of the effectiveness. According to EN ISO 8044 the polarization and the associated the change of potential of a steel surface is caused by the cathodic current. This polarization can be achieved through activation polarization. The cathodic current flow inadvertently results in a change of the soil composition as a result of the electrochemical reactions taking place. This change of potential caused by the concentration change is called concentration polarization. According to von Baeckmann the activation polarization is achieved within milliseconds as a result of the current flow through the steel surface and the concentration polarization is built up within a fraction of a second, minutes or even hours [65]. Based on the present model concepts discussed in this document a more differentiated understanding is available today with respect to the involved electrochemical processes. The concentration polarization at the steel surface as a result of the CP current changes the composition of the soil not only in the immediate vicinity of the steel surface but also in the surrounding environment. This effect provokes a shift of both the pH and the potential along the hydrogen equilibrium line (blue arrows in Figure 17). The increase of the pH at the steel surface and in the surrounding soil as well as the passivation are illustrated in Figure 19a. Since the hydrogen equilibrium potential, which correlates with the cathodic limit of water stability, depends on the pH (hydrogen evolution line in Figure 17), the pH-value at the steel surface can directly be determined based on the IR-free potential. This has been discussed in detail by Angst et al [62] and is confirmed by the data of Yan et al [66] as reported in [67]. For this reason, the protection potentials according to EN ISO 15589-1 can readily be explained:

- The $E_{IR\text{-}free}$ of -0.85 V_{CSE} corresponds to a pH of 9, which is, according to Figure 17, sufficient for achieving passivity
- The $E_{IR-free}$ of -0.95 V_{CSE} corresponds to a pH of 10,5 (yellow circle in the middle of Figure 17), which allows for passivity even in aggressive soils
- On the other hand, the $E_{IR-free}$ of -1.2 V_{CSE} corresponds to a pH of 13 (yellow circle to the right in Figure 17), which, according to EN ISO 15589-1, may cause problems with the adhesion of coatings.

It should be noted in this context, that a pH value of 13 is typical for concrete. In concrete the formation of a protective passive film and the resulting corrosion protection for the steel reinforcement are well accepted.

The dependence of the pH-value at the steel surface on the CP current density has been investigated by several authors. The results are shown in Figure 18. It can be concluded that cathodic current densities in the range from 20 to 1000 mA/m², which are typical for the CP the exposed steel within the coating defects on buried pipelines, result in pH-levels from 10.5 to 12.5. According to Figure 17 these levels correlate with IR-free potentials in the range from -0.95 to -1.2V_{CSE}. Correspondingly, it is readily possible to plot the IR-free potential associated with the achieved surface pH value as shown on the right vertical axis in Figure 18. This consideration shows the relevance of the current density, the surface pH, the generation of an environment conducive to passivation as well as the protection criteria based on IR-free potentials for the practical implementation of CP on buried pipelines.

Figure 17: Pourbaix-diagram (Fe/H₂O) illustrating the cathodic polarization of steel [2]. The blue arrow shows the effect of concentration polarization and the resulting enhancement of the pH-level as well as the passivation. The yellow circles indicate the position of the protection potentials according to EN ISO 15589-1.

Based on this consideration the IR-free potential is associated with a surface pH measurement rather than the actual corrosion protection of the steel. Since an increase in pH is associated with passivation, corrosion protection is achieved. This view was emphasized by Freiman [29] and is today widely accepted. The standard CP textbook by Peabody [44] on CP describes the process as follows:

"The concepts presented for CP (activation polarization) are fundamentally correct at the instant that CP is applied but are too simplistic to consider the time-dependent behaviour of a cathodically protected underground structure..... This pH increase (concentration polarization) is beneficial because the corrosion rate of steel decreases with increasing pH, even under freely corroding conditions. The decrease in corrosion rate is the result of the formation of a protective oxide film on the metal surface in the elevated pH environment, a process referred to as passivation."

Figure 18: The pH-value at the steel surface as a function of the cathodic current density according to different literature sources summarized in [68]. On the right vertical axis, the expected IR-free potentials associated with the hydrogen equilibrium line are shown.

The relevance of passivation reveals another highly relevant aspect that immediately follows from Figure 17: The IR-free potential is not suitable for assessing the corrosion situation of a passive steel surface under cathodic protection. Passivity is present at increased pH over a wide potential range extending from the hydrogen evolution line all the way to the oxygen evolution line. In well aerated resistive soils, it may not be possible to achieve a sufficiently high current density to polarize the steel surface to the hydrogen evolution line. The resulting IR-free potential may well be in the range of -0.65 and -0.75 V_{CSE}. As long as the pH is increased, passivity and hence corrosion protection is achieved under such conditions [9]. This immediately explains the corresponding protection criteria determined by Funk et al [15] that are stated in EN ISO 15589-1.

It immediately follows that no correlation between the corrosion rate and the IR-free potential may be expected. Therefore, without consideration of soil resistivity and aeration single IR-free potential criteria can be associated with corrosion protection as emphasized by Table 1 in EN ISO 15589-1. This is because the IR-free potential may only be used as a pH measurement when the IR-free potential of the steel is controlled by the hydrogen electrode. At increased aeration of the soil this is not the case and the IR-free potential becomes an irrelevant number with respect to the assessment of the achieved pH and hence the level of corrosion protection.

This consideration raises the question with respect to an appropriate protection criterion. Based on Figure 18 a current density in the range of 1 mA/m^2 is sufficient to increase the pH to a level that generates conditions conducive to passivation. At such low current densities, the IR-error is negligible in many soils and even increased coating defect sizes. An example of a model calculation of such a configuration is shown in Figure 27. For sufficiently low soil resistivity and smaller coating defects corrosion protection is achieved as soon as the on-potential is more negative than -0.85 V_{CSE}. This is fully in line with the concepts and protection criterion proposed by Kuhn [1]. A more recent analysis of the two most extensive field investigations performed with respect to CP protection criteria performed so far has clearly shown

that the on-potential is at least as reliable as the IR-free potential with respect to the assessment of corrosion protection on coupons [69] with sizes below 20 cm², which is fully in line with the above conclusion. This raises the question with respect to the appropriate level of on-potential that provides enough pH increase for the generation of conditions conducive to passivation. While this current density of 1 mA/m2 may be sufficient for well bedded coating defects smaller than 20 cm², higher current density may be necessary in other conditions. The required level of current density required for achieving corrosion protection predominantly depends on the bedding conditions (see [62]) but also the coating defect size. The key parameter is the so-called reference current density J_{ref} that is required to increase the pH and establish effective CP. The reference on-potential that is required to ensure J_{ref} on all coating defects is called E_{ref} according to DVGW GW 21. For well bedded pipelines in sand and soil with small coating defects E_{ref} may be as positive as -0.85 V_{CSE}. Based on Angst et al [62] a value of -1.0 V_{CSE} is suggested for such conditions, since such an on-potential ensures current entering into all coating defects.

6.3 The effect of anodic stray current interference

It follows from the above consideration that the cathodic protection of steel is based on the increase of the pH-value and the resulting passivity. This discussion illustrates that the assessment of stray current interference of cathodically protected pipelines needs to be reassessed in the light of today's knowledge. The present mechanistic models, the available literature as well as available empirical data allow to develop new interference assessment concepts and corresponding thresholds. These will be presented and discussed in the following.

The cathodic protection based on passivity has far reaching consequences for the assessment of time variant stray current interference of a cathodically protected pipeline. The IR-free potential is a pH measurement rather than a controlling parameter for the assessment of the level of corrosion protection. This has immediate consequences on the assessment of the corrosion risk under an interfered situation: The temporary anodic shift of the IR-free potential of a passive steel surface in the positive direction as a result of anodic stray current interference, is not necessarily linked to a corrosion process. The IR-free potential more positive than the protection criterion of e.g. $-0.95 V_{CSE}$ is only the result of an influenced and hence irrelevant pH-measurement. In the case of well aerated soils IR-free potentials of -0.65 and -0.75 V_{CSE} are required for addressing this influencing effect of oxygen in well aerated soils in EN ISO 15589-1. Similar to the case of the oxidizing action of oxygen in the case of well aerated soils where IR-free potentials of -0.85 and -0.95 VCSE are no longer applicable significantly more positive IR-free potentials may well be associated with corrosion protection under anodic stray current interference. In both the well aerated soil and the anodic stray current interference condition the IR-free potential is no longer controlled by hydrogen evolution and therefore becomes a meaningless number.

Time variant interference is typically caused by DC-operated railways or trams. In these cases, the anodic interference has by experience in Holland typically a maximum duration of about 300 seconds. In contrast, time variant anodic stray current discharge from coating defects on pipelines under AC interference occurs at much higher frequency resulting anodic current discharge for only about 10 ms in the case of 50 Hz interference. Moreover, telluric interference is well known to cause time variant interference at very low frequency with anodic interference duration in the range of up to several hours. It is evident from the above discussion that a time limited anodic current discharge does not necessarily result in corrosion. In fact, the anodic polarization will only shift the potential in the positive direction within the passive domain, which usually does not result in corrosion. The anodic charge will rather

be consumed for the oxidation of ferrous ions (Fe(II)) within the oxide layers on the steel surface as described in [4]. As a consequence, the repeated cathodic and anodic interference caused by DC-operated railways or trams will cause a relevant charge transfer through the steel surface. This electric charge, however, will not be consumed by the corrosion reaction, but will be used for the redox system Fe(II)/Fe(III)/Fe(II). Bette observed this phenomenon already in 2005 [70] and described it with capacitive charging effects. In fact, the abovementioned redox system acts as a pseudo-capacity and may readily be compared to the charging/discharging of an accumulator. Consequently, the combination of passivity with this redox system explains the empirical observation that a time limited anodic current will neither result in major anodic potential shifts nor in corrosion. All more recent investigations of stray current induced corrosion on cathodically protected pipelines confirm that the electric charge will not be consumed for the corrosion reaction as long as the steel surface is passive [6, 28, 55, 56, 71, 72]. Hence, anodic current discharge is not inevitably related to a corrosive dissolution of the metal following Faradays Law, i.e. an anodic current does not always result in corrosion. This is due to the fact that the anodic charge is consumed for the reaction Fe(II)/Fe(III) on the passive surface, rather than the reaction Fe(0)/Fe(II) in the case of time variant stray current interference.

The criteria specified in EN 50162:2004 are, however, based on the assumption of a time constant anodic stray current load. Hence it is required to discuss the effects of a time constant anodic current on the corrosion processes. In case of a time constant anodic current it is not possible to achieve an increase of the pH at the steel surface. In fact, a time constant anodic current results in a decreasing pH-value and thus in an acidification (cause by concentration polarization) at the steel surface leading to corrosion according to Figure 19b. Moreover, the effect of a decreasing pH-value is illustrated with a red arrow in Figure 20. The formation of an acidic electrolyte provokes corrosive conditions and passivation will be impossible. Under such circumstances it is to be expected, that 100% of the anodic current is consumed by the redox system Fe(0)/Fe(II), i.e. for the corrosion reaction. This underlines, that EN 50162:2004, which had been elaborated for time variant interference from DCoperated railways or trams, is based on incorrect assumptions: The relevant increase of the pH-value as a result of effective CP, which is crucial for the effectiveness of cathodic protection, the resulting passivation, the relevance of the redox system Fe(II)/Fe(III) and also the consequences of a CP current that is cathodic when averaged over a representative period of time, are ignored.

Figure 19: Schematic illustration of cathodic concentration polarization (a) and anodic concentration polarization (b). The cathodic current entering the steel surface, results in a depletion of oxygen, an increase of the pH-value in the soil and passivation (a). The anodic current leaving the steel surface results in a lowering of the pH-value and provokes corrosion (b). The intensity of colouring indicates the extent of pH change.

Based on the above discussed aspects various influencing factors can be identified, which relevantly control the corrosion under time variant stray current interference:

Anodic current will decrease the pH, while cathodic current will result in an increase of pH at the steel surface. It immediately follows that the average current over a full interference cycle (e.g. 24 hours) represents a relevant parameter. This is due to the fact that the current efficiency for alkalinity formation is at least as good as the one for acidity formation in the case of steel. An average cathodic current is therefore bound to result in a pH increase over time irrespective of temporary anodic excursions. Hence the polarity of the average electrical charge controls the direction of pH change at the steel surface. If the excess of cathodic charge is associated with a permanent and sufficient increase of the steel surface pH, corrosion protection is achieved. This effect of concentration polarization can best be illustrated based on an analogy: In absence of anodic current discharge or in case of cathodic interference a reservoir of alkalinity is accumulated at the steel surface. This reservoir will be consumed during anodic current discharge. Saving alkalinity provides, therefore, corrosion protection during the hard times under anodic current discharge. As long as more alkalinity is saved than consumed, an increased surface pH and hence corrosion protection is ensured even under severe time variant anodic stray current interference.

This analogy reveals important implications: The duration of the anodic interference is bound to be relevant, since the decrease of pH is not only a result of anodic current discharge. Diffusion and migration into the surrounding soil, neutralisation with carbon dioxide as well as convection and streaming water will contribute to a loss of the accumulated alkalinity. These effects are negligible with respect to the pH at the steel surface in the case of short anodic interference during AC interference of 50 Hz or short telluric interference of up to 3 seconds. This is illustrated by means of the green arrows in Figure 20. Indeed, experience confirms that there is no detrimental effect with respect to corrosion caused by loss of surface pH under such conditions. However, based on the argumentation in EN 50162 they would have to result in relevant metal loss as a result of Faradays law and an assumed reaction of Fe(0)/Fe(II).

In contrast, a duration of the anodic interference of 30 seconds causes a relevant change of the surface pH during the anodic excursion, although the time averaged current is cathodic. (blue arrows in Figure 20). Such durations are typical for tram operation. A further increase of the duration of the anodic interference to 300 seconds, as they are characteristic for DC train operation, can cause a major loss of surface pH resulting in temporary initiation of corrosion (orange arrows in Figure 20). While the subsequent cathodic interference will re-establish passivating conditions according to Figure 20, a non-negligible corrosion rate will be observed in this interference situation. Based on the above discussion the efficiency of the anodic current with respect to the reaction Fe(0)/Fe(II) will be smaller than 100 % since only a fraction of the anodic charge is consumed for corrosion. This effect is indeed experimentally observed in the case of time variant anodic interference [70].

Figure 20: Polarisation of steel under stray current interference illustrated in the Pourbaix diagram [2]. The red arrow shows the concentration polarization under time constant anodic interference (in case of ineffective CP). A time variant interference in case of effective CP is illustrated with the green, blue and orange double arrows. The different duration of the anodic interference of 3 (green), 30 (blue) and 300 (orange) seconds results in different levels of concentration polarisation at a given cathodic charge.

Theoretically, the duration of the anodic interference is expected to be relevant with respect to the corrosion behaviour. However, there are additional aspects that require consideration in the practical application. The current will flow through individual coating defects as illustrated in Figure 19, which typically have various sizes and geometries and are in contact with various soils with different aeration and resistivities. It is therefore necessary to consider these aspects in the case of the practical application of the above concepts.

Based on the above discussion the current density is relevant with respect to the surface pH that is established (see Figure 18). With decreasing defect size and decreasing soil resistivity increased current densities are expected. As a consequence, in case of a time constant interference (red arrow inFigure 20) highest corrosion rates have to be expected on smallest coating defects in least resistive soil. However, in the case of time variant interference (e.g. caused by a DC traction system) it is observed that also the cathodic current densities are increased in this configuration. It follows that the effect of defect size and soil resistivity during time variant interference is essentially eliminated: High anodic current densities will be followed by high cathodic current densities. This will maintain an average cathodic current and hence corrosion protection irrespective of defect size and soil resistivity.

Based on this consideration only the following influencing parameters need to be considered:

• A reference current density (J_{ref}) for establishing effective CP at the steel surface in accordance with EN ISO 15589-1 is required. Typical values are between 20 to 200 mA/m².

- The cathodic charge must be larger than the anodic charge resulting in a time averaged pH increase at the steel surface.
- With increasing duration of the anodic excursion, the cathodic charge needs to be proportionally larger compared to the anodic charge. The excess in alkalinity is required to compensate for the contribution of migration and diffusion of alkalinity in the surrounding soil. This will ensure a durably increased pH at the steel surface and prevents the conditions shown with the orange arrows in Figure 20.

This consideration is illustrated in Figure 21 by means of schematic measuring data collected on a coupon on a cathodically protected pipeline under DC traction interference. The J_{ref} is required to establish effective CP. This J_{ref} is measured in the non-interfered situation, for example during the night in absence of train operation. During the day the train operation is causing anodic and cathodic current densities on the coupon, which are often a multiple of J_{ref} .

Figure 21: Schematic evolution of the current density in the coating defect (or a coupon) on a stray current interfered pipeline. The cathodic (blue) and anodic (red) charges are obtain based on the surface integral of the respective current densities. The duration t_a corresponds to the longest anodic excursion.

The ratio of the anodic and cathodic charge is relevant with respect to concentration polarization and hence the pH at the steel surface. The assessment of the corrosion risk can, therefore, be performed based on equation (1) with the anodic interference charge Q_a (total red surface in Figure 21) and cathodic interference charge Q_c (total blue surface in Figure 21) over the representative period.

$$Q \leq \left(\left| Q_{c} \right| - \left| Q_{a} \right| \right) / \left| Q_{a} \right|$$

$$\tag{1}$$

The value on the right-hand side of equation (1) describes the ratio of anodic and cathodic charge. For identical charges a value of zero is obtained. When the cathodic charge is double the anodic charge, it equals 1. Experimentally it was found that for artificial soil and artificial soil solutions with increased hardness the dependence of Q from t_a can be described by equation (2) [73]:

$$Q = t_{a}*0.004 \ s^{-1} \le (|Q_{c}| - |Q_{a}|)/|Q_{a}|)$$
(2)

Based on Ohm's law it is, however, possible to translate the current density consideration into a potential consideration under the following assumptions:

- The polarization resistances for anodic and cathodic reactions are small compared to the spread resistance. This is indeed the case for both the hydrogen evolution as well as the redox system Fe(II)/Fe(III).
- The spread resistance does not relevantly change during the cathodic and anodic interference. This might not be the case during extended interference durations, but the cathodic interference will always result in lower spread resistance (and hence smaller potential change) compared to the anodic interference. This assumption is, therefore, conservative.

Figure 22: Schematic potential evolution of a cathodically protected pipeline. The anodic interference $\Delta E_{a,avg}$ is the result of the average anodic shift of the potential relative to E_{ref} . The duration t_a corresponds to the largest anodic excursion.

Under these conditions for a typical DC traction interference DVGW GW 21 states with respect to the potential analysis equation (3) as the minimum requirement with respect to stray current protection:

$$E_{on,avg} \leq E_{ref} - \Delta E_{a,avg} \tag{3}$$

The on-potential averaged over 24 hours ($E_{on,avg}$) must accordingly be more negative than the reference potential (E_{ref}) minus the average anodic interference ($\Delta E_{a,avg}$). Similarly to the charge consideration in equation (1), a stronger anodic interference can hence be compensated by larger cathodic charge, which is obtained at a more negative average on-potential. In Figure 22 the analogy between the current density (see Figure 21) and the potential evolution is evident. E_{ref} corresponds to the minimal required on-potential in absence of DC interference for establishing effective CP. E_{ref} correspondingly ensures J_{ref} . $E_{on,avg}$ is the time average of all on-potential values recorded over a representative period of time. $\Delta E_{a,avg}$ corresponds to the average of all red potential values minus E_{ref} . Equation (3) was first published in 2009 and has today the status of a recognized state of the art in Germany. It is applicable for DC-traction interference where typical maximum values of t_a of about 250 seconds are observed, which results according to equation (2) in a value for Q of 1. Indeed, it can be demonstrated that equation (1) and equation (3) are mathematically identical for Q=1 [73]. It is hence readily possible to extend the application range of equation (3) based on the conditions in equation (2) and consider the effect of various values of t_a according to equation (4).

$$E_{on,avg} \le E_{ref} - Q^* \Delta E_{a,avg} = E_{ref} - 0.004 s^{-1} * t_a * \Delta E_{a,avg}$$
(4)

According to [73] and equation (4) the equivalence of the charge and potential consideration is evident as shown in equation (5):

$$Q \leq \left(\left| Q_{c} \right| - \left| Q_{a} \right| \right) / \left| Q_{a} \right| = (E_{ref} - E_{on,avg}) / \Delta E_{a,avg}$$
(5)

The relevance and the correctness of the requirements of DVGW GW 21 and the underlying concepts are confirmed by the independent validation in [73] based on an electrical charge consideration. The crucial advantage of equation (4) is the expansion of the application range of the DVGW GW 21 to the entire frequency domain of anodic stray current interference. This domain ranges from AC frequencies (t_a =0.01 s) over DC traction interference (t_a =300s) and tidal interference (t_a =22'000s) to time constant interference (t_a =infinite).

The validity of equation (4) and (5) is confirmed in [73] as well as the requirement of DVGW GW 21 for values of t_a in the range of a few minutes. For infinitely small values of t_a Q is approaching zero according to equation (2). Hence the anodic charge (or anodic potential excursion) may be approximately identical to the cathodic charge (or cathodic potential excursion). Based on equation (4) $E_{on,avg}$ must only be more negative than E_{ref} in this case. This is indeed confirmed by extensive field investigations in Germany [74]. Based on these data no corrosion is expected even in the case of AC current densities as high as several hundred A/m² at sufficiently negative average on-potentials (e.g. more negative than -1 V_{CSE}).

Equation (4) is also confirmed for very large values of t_a , as they are observed in case of time constant stray current interference. In this case Q becomes infinite and equation (5) can only be satisfied when the anodic charge (or $\Delta E_{a,avg}$) approaches zero. This is indeed only possible if equation (6) is fulfilled:

$$E_{on} \le E_{ref} - \Delta E_{a,max} \tag{6}$$

The on-potential must be shifted in the cathodic direction by the maximum anodic shift $(\Delta E_{a,max})$. This eliminates any anodic interference and Q_a becomes zero satisfying equation (1). Indeed, this is a common procedure for mitigation of time constant anodic stray current interference.

This consideration confirms the relevance of equation (4). It is well established by practical application in the last 10 years in Germany and can be expanded to cover the entire frequency domain of anodic stray current interference ranging from AC interference to time constant anodic DC interference. It allows for an assessment of the stray current corrosion risk as well as the identification and control of mitigation measures based on readily measurable potential data followed by an averaging process. The above procedure is introduced in ISO 21857.

6.4 The effect of cathodic stray current interference

The above discussion of anodic stray current interference has highlighted the relevant processes taking place under anodic stray current discharge. It has been shown that these concepts can be expanded to explain all the effects taking place over the entire frequency domain including the AC interference condition that is included in the scope of ISO 21857. This raises the question with respect to the relevance of EN ISO 18086 that also covers criteria with respect to AC corrosion. This apparent conflict can readily be resolved when taking into account the relevant mechanism of AC corrosion: The above discussion with respect to anodic interference has clearly shown that AC corrosion as a result of anodic interference cannot occur as long as effective CP is established by applying an average on potential more negative than E_{ref} (e.g. -1.0 V_{CSE}). Based on this consideration AC corrosion as a result of anodic current discharge is effectively excluded by establishing effective CP, which is based on an increase of the pH value at the steel surface and the establishing of a protective passive film. Anodic current discharge is therefore irrelevant as long as the passivating conditions are maintained which is indeed the case for the very short anodic excursions in the case of AC interference.

In contrast, EN ISO 18086 does not cover the corrosion aspects of anodic stray current interference, but those caused by excessive cathodic polarization. Indeed, the first leaks on cathodically protected pipelines occurred in over polarization conditions at on-potentials more negative than -2 V_{CSE} [75, 76]. Correspondingly, EN ISO 18086 limits the level of AC current density (J_{ac}) to less than 30 A/m² or the DC current density (J_{dc}) to less than 1 A/m². This implies that AC corrosion is an over polarization problem since limiting the level of cathodic current density is an effective measure that eliminates the AC corrosion risk. It has been demonstrated that AC corrosion can readily controlled by means of limiting J_{dc} to values below 1 A/m² [50].

The complexity of these effects requires a more detailed discussion, in order to highlight the relevant processes taking place under AC interference conditions. For illustrating the effect of AC interference on the actual IR-free potentials as a function of the current density (J) were recorded by Bette in artificial soils at a data acquisition rate of more than 1 kHz [77]. These results for two coupons are shown in Figure 23. Despite of significantly increased J_{DC} the E_{IR}free is temporarily anodic of the protection criterion of -0.85 V_{CSE} on both coupons. Based on the concurrently recorded corrosion rate data a metal loss is only observed when a temporary polarization cathodic of $-1.2 V_{CSE}$ occurs. As discussed above, the polarization cathodic of -1.2 V_{CSE} only occurs at J_{DC} larger than 1 A/m². Special attention must be paid to the time dependence of E_{IR-free}, which shows a variation in the range of 0.4 V as a function of the polarity of the current density (J). Based on the data in Figure 23 this time dependence can be assessed with a data acquisition rate of at least 1 kHz. Slower measuring rates of less than 10 Hz, as they are usually applied in CP for assessing the E_{IR-free}, do not show this time dependence due to the filter systems in the measuring instruments. Hence, the question regarding the physical significance of the various E_{IR-free} in Figure 23 rises. In this context only the most relevant influencing factors and the corresponding conclusions will be addressed. A more detailed discussion of the involved processes is given in [78].

Figure 23: IR-free potentials determined on an ER coupon as a function of the current density (J) under ac-interference of 16.7 Hz (cathodic currents with a positive sign). Green: No corrosion at J_{dc} 1 A/m², J_{ac} 128 A/m²; Red: Corrosion at J_{dc} 11 A/m², J_{ac} 309 A/m² according to [77]. The full circles correspond to $E_{IR-free}$ and the empty circles to the most negative cathodic excursion called E_{H} .

Based on Figure 23 it is not possible to determine a single value for the $E_{IR-free}$. It is, however, possible to determine an average of all the recorded values, which in a first approach corresponds to the classical slowly measured (about 100 ms after interrupting J_{dc}) $E_{IR-free}$. This value is illustrated with the full circles in Figure 23. Additionally, in Figure 23 also the most negative potential excursion limited by hydrogen evolution can be determined, which according to the model concept controls the ac-corrosion process. This parameter is in the following context described as E_H shown by means of the empty circles in Figure 23. These data confirm the relevance of the J_{dc} and J_{ac} given in EN ISO 18086. They furthermore demonstrate that it is the excessive cathodic polarization rather than the anodic polarization that initiates the AC corrosion process. Based on this discussion and in particular based on the potential variation in Figure 23 it is possible to further elaborate the relevant processes taking place under AC interference. It follows from the above discussion that in absence of AC-interference equation (7) applies:

$$E_{IR-free} = E_H \tag{7}$$

In presence of AC interference equation (8) applies with the contribution of the so called faradic rectification ΔE_F [79].

$$E_{IR-free} = E_H + \Delta E_F \tag{8}$$

This consideration represents a rough simplification of the processes taking place at the steel surface under exclusion of all time-dependent contributions. However, it provides a physical description for the empirically observed effects caused by the rectification of J_{ac} . The size and the polarity of ΔE_F are dependent on the ratio of the Tafel slopes of the anodic and cathodic

activation-controlled reactions. It is characteristic for passive systems that ΔE_F has a positive sign. This effect was used for so called "wet rectifiers" (or electrolytic rectifiers) in the past. Also, the DC decoupling devices based on Nickel plates in KOH solution (the so-called Kirk cells) are based on this effect. The key requirement for the electrodes used for wet rectifiers (and decoupling devices) was their passivity (see e.g. [80-82]). This results in a large non-symmetric current-potential behavior and a strong rectification.

The J_{dc}, which passes through a coating defect with a metallic surface of A, is the result of the difference between E_{on} and E_{IR-free} as well as the spread resistance R according to equation (9). For E_{on} more positive than -1.2 V_{CSE} it was demonstrated that J_{dc} can approach zero [50]. Based on this concept it is possible to limit ac-corrosion and satisfy the protection criterion of J_{dc} < 1 A/m² in EN ISO 18086 through the control of E_{on} even at very high levels of AC-interference.

$$J_{dc} = \frac{E_{IR-free} - E_{on}}{R \cdot A} \tag{9}$$

The J_{ac} passing through the metal surface causes a shift of the $E_{IR-free}$ in positive direction according to equation (8) and as illustrated with the full circles in Figure 23 with respect to the empty circles. For the determination of J_{dc} the $E_{IR-free}$ (average $E_{IR-free}$) is relevant, which is a result of E_H and ΔE_F . The evaluation of the literature data [4, 83] with respect to J_{ac} and the resulting anodic shift of $E_{IR-free}$ under assumption of a linear behaviour allows describing ΔE_F with a factor f according to equation (10) [40, 84, 85].

$$\Delta E_F = f \cdot \frac{U_{ac}}{R \cdot A} \tag{10}$$

$$U_{ac} = R \cdot A \cdot \left(J_{dc} \cdot R \cdot A - E_H + E_{on} \right) / f \tag{11}$$

The combination of equations (9) and (10) results in equation (11), which is a description of the E_{on} and the allowable U_{ac} as a function of the critical J_{dc} in the case of effective cathodic protection.

The further consideration of the thermodynamic [2] and kinetic [37, 41] parameters with a mathematical description of the decrease of the spread resistance caused by the increase of the pH-value at the steel surface [4] as a result of J_{dc} and the resulting spread of alkalinity as shown in Figure 19a allows for a more detailed description of the corrosion process under AC-interference on the basis of equation (11) [40, 84, 85]. This detailed model concept allows for explaining the relevant discrepancy between the actual damages on pipelines and the high corrosion rate on coupons as described in [85].

Based on equation (11) the acceptable U_{ac} reaches a value close to zero for decreasing defect sizes. In contrast, an increase of the metallic surface A in the coating defect will increase the acceptable U_{ac} . This has a relevant implication, since any corrosion process will cause an increase of the metallic surface as demonstrated in [40, 84, 85]. These considerations and the subsequent validation of the model have revealed that any AC corrosion process will reach negligible values once a certain corrosion depth has been reached. This is the result of the increase of the steel surface caused by the corrosion process and a direct consequence of the geometrical morphology of the corrosion site and equation (11). If the soil resistivity (ρ), the average E_{on} and U_{ac} , the original coating defect size and the allowable corrosion depth are known, an assessment of the acceptable interference conditions is possible. The key conclusions for ac-corrosion are therefore as follows:

- At small coating defects AC-corrosion cannot be prevented
- AC-corrosion will reach negligible rates at a certain depth
- This depth is higher on large coating defects than on small coating defects

These qualitative conclusions are in good agreement with the empirical observation of the last decades. Leaks due to AC interference were primarily observed on pipelines with small pipe wall thickness. Moreover, very high corrosion rates were found on thin coupons while only corrosion depths in the range of 1 to 2 mm were found on the associated pipelines.

Based on laboratory and field investigation a numerical description of all influencing parameters was developed. In particular the effect of pH increase at the steel surface and the migration of alkalinity into the surrounding soil (as shown in Figure 19a) was numerically described. The individual parameters were first calibrated in laboratory investigations and then validated in field tests [39, 86]. The applicability of the model was thus demonstrated. Moreover, the expected influence of the metallic surface as a function of corrosion depth on the corrosion rate was confirmed. This validated model confirms the empirical experience collected in the past 30 years. It allows, therefore, predicting the critical conditions and optimizing mitigation measures in the practical operation of pipelines. The resulting numerical model and the associated parameters were 2018 introduced into the document DVGW GW 28 B1 that represents the recognized state of the art in Germany.

This model for AC corrosion can explain the discrepancy between the high corrosion rates observed on coupons and the very limited number of damages on pipelines. The increase of the steel surface due to the corrosion process results with increasing corrosion depth in decreasing current densities. When they reach the threshold current densities stated in EN ISO 18086 the AC-corrosion process is expected to reach negligible rates. Based on the important relevance of the corroding steel surface and hence the corrosion depth, the AC-corrosion rate is strongly time dependent and of limited relevance. The very high corrosion rates in the early stages of AC-corrosion decrease rapidly with progressing depth. Hence, it is impossible to extrapolate a corrosion rate determined over a limited exposure time to the operation time of the pipeline.

All the available data demonstrate that the assessment of the acceptable AC interference level can only be based on an acceptable corrosion depth. The discussion of the critical coating defect surface demonstrates, that the assumption of a critical coating defect surface of 1 cm² already implies an acceptable corrosion depth in the range of one to two millimetres. The meeting of the requirements of EN ISO 18086 on coupons with 1 cm² defect surface cannot exclude higher current densities on smaller coating defects and hence corrosion. As predicted by the model calculation, these small coating defects never lead to perforation of pipelines. It is expected that they corrode rapidly in the early stages, but then the corrosion rate reaches negligible values within 1 to 2 mm depth. These considerations clearly show that the present standard has an implicitly accepted maximum corrosion depth, which confirms that the occurrence of AC-corrosion in the range of a few millimetres cannot be excluded on any pipeline even when all applicable standards are applied.

The validation of the model allows for the assessment of the AC-corrosion risk on pipelines caused by cathodic interference. The numerical description of the relevant influencing factors offers the possibility to correctly address them and optimize mitigation measures. Examples of this consideration for a maximum corrosion depth of 5 mm are shown in Figure 24. Clearly, an important dependence of the admissible U_{ac} on ρ and E_{on} is found, demonstrating that it is impossible to define generally valid interference levels for U_{ac} . The requirements in Figure 24 allow for determining critical sections of the pipeline system as well as the development of mitigation strategies. Experience shows that depending of the various factors the highest

corrosion risk is not necessarily associated with the highest U_{ac} induced on the pipeline, but rather low soil resistivity in combination with negative on-potentials.

Figure 24: Admissible average U_{ac} as a function of the average E_{on} for various soil resistivities ρ and an acceptable I_{max} of 5 mm calculated with the parameters in [39] for: a) PE and b) FBE coatings.

With this approach shown in Figure 24 it is possible to demonstrate relevant differences between fusion bonded epoxy coatings (FBE) with thickness in the range of 0.5 mm and threelayer polyethylene coatings (PE) with thickness in the range of 2 mm. In the case of FBE the corrosion products fracture the coating and the lateral extension of sthe corrosion process underneath the FBE coating is very limited. This characteristic behaviour of FBE is usually described as "fail safe" since the access of CP current is ensured by the fracturing of the coating. In the case of AC-corrosion, however, the defect diameter is increasing with the diameter of the corrosion site. This has relevant implications with respect to damage mechanism according to the described model. Small coating defects result in high current densities and high corrosion rates. Instead of a fast decrease of the corrosion process to negligible values (as expected for PE), the coating defect diameter grows with increasing corrosion depth and allows for further extension of the corrosion process and the reaching of larger depths. This is expected to result in faster perforation of the pipeline in the case of FBE compared to PE coating, since the corrosion process is always running at the most critical condition as described in [39].

In contrast, the PE coating is mechanically more robust. The experience shows that it will be lifted off the steel surface through the mechanical pressure of the corrosion products, while maintaining the original coating defect diameter. The shielding of the activation-controlled AC corrosion process is, therefore, an advantage of PE coatings. Based on these considerations different parameters are required for addressing the AC-corrosion risk on PE and FBE coated pipelines. This is readily possible with the two different parameter sets proposed in [39] and the resulting thresholds shown in Figure 24.

This concept has formed the basis for the DVGW GW 28-B1. No leaks have been reported in Central Europe for PE coated pipelines with wall thicknesses of more than 5 mm when the 24-hour average values of U_{ac} and E_{on} given in Figure 24a for the different values of the soil resistivity ρ are not exceeded.

This concept allows the assessment of the AC corrosion risk based on readily available potential values and significantly facilitates the decision with respect to mitigation measures. Based on this approach, the critical sections on the pipeline are readily identified. The additional installation of coupons in these specific areas allows for demonstrating the correctness of the numerical analysis and the determination of the current densities required by EN ISO 18086. In the evaluation of these data the relevance of processes described in equation (11) and in [39] still need to be taken into account.

6.5 Establishing E_{ref} and J_{ref}

6.5.1 Introduction

Based on the present discussion it is evident that the assessment of J_{ref} and E_{ref} are of key relevance with respect to protection criteria. A possible procedure for identifying these values is presented in the following. The assessment of J_{ref} and E_{ref} requires the identification of a number of relevant influencing parameters. They are discussed in the following.

6.5.2 Concentration polarization and the resulting J_{ref}

An increase of the pH value on the steel surface and hence sufficient concentration polarization is always possible, if sufficiently high current densities are applied. The following relevant influencing factors affect the required level of J_{ref} :

- **Bedding:** A sufficient quantity of fine bedding material leads to a more homogeneous current distribution due to a smaller shielding of large stones. In addition, there is a greater retention capacity for water. Fine bedding of the pipeline contributes to smaller levels of J_{ref} and less negative E_{ref}.
- **Water hardness:** An increased water hardness favours the formation of calcareous deposits on coating defects and can thus compensate a coarse bedding. Soft water cannot form a chalk-bed and also prevents the precipitation of salts of organic acids and carbonates. Soft water hence increases J_{ref} and shifts E_{ref} to more negative values.
- Flow: Flowing water can dilute the hydroxide ions and leads to higher required protective current densities. Extreme cases are changing groundwater levels or rainfall flowing through the pipe trench, which are then empty again. In combination with coarse bedding materials and good aeration, heterogeneous conditions and increased corrosion rates are to be expected. Flow hence increases J_{ref} and shifts E_{ref} to more negative values.
- Aeration: Increased aeration makes it difficult to achieve a homogeneous current density distribution due to the possibility of heterogeneous aeration and favours partial passivation and the formation of galvanic elements within individual coating defects. On larger coating defects an increased aeration may increase J_{ref} and shift E_{ref} to more negative values.
- **Coating defect size:** Smaller coating defects favour homogenous conditions and therefore require smaller current densities. Small coating defects requires smaller levels of J_{ref} and in particular less negative E_{ref}.

As a result of these boundary conditions, requirements for J_{ref} can be derived. Depending on these conditions, J_{ref} can vary between 1 mA/m² and 3 A/m² within individual coating defects on buried structures. While it is possible to identify influencing parameters, there is presently no concept available with respect to quantifying J_{ref} . In the following a conservative approach with respect to the quantification of J_{ref} as a function of the influencing parameters is given in equation (a).

$$J_{ref} = a + b + c \cdot (d + e) \tag{a}$$

The equation (a) as well as the values for the parameters "a" through "e" shown in Table 1 are based on semi empirical estimations so far. Their application must be calibrated with practical experience. In case of doubt it may be useful to apply a value of the 0.1 A/m^2 in absence of any further information as a first conservative approximation.

Defect surface	[cm ²]	<1	1-10	10-100
а	[A/m²]	0.01	0.03	0.1
Aeration (J ₀₂)	[A/m²]	<0.01	0.01 - 0.1	0.1 - 1
b	[A/m²]	0.01	0.1	1
Hardness	[°fH]	0-15	15-25	>25
с	[-]	1	0	0
Flow	[m/day]	<0.1	0.1 - 1	>1
d	[A/m²]	0	0.1	1
Bedding size	[mm]	< 1	1-10	> 10
e	[A/m²]	0	0.1	1

 Table 1:
 Relevant factors influencing J_{ref}

The quantifying of these influencing parameters allows for determining a value for J_{ref} that is based on the presently known influencing parameters. Furthermore, this procedure assists in an improved risk assessment and the identification of the potentially critical pipeline sections. Hence, the present model concepts allow for the first time the identification and consideration of the relevant influencing factors in the assessment of the corrosion risk based on a physical chemical concept that is passed on well recognized concepts. Therefore, this evaluation is expected to provide an improved risk assessment of pipelines.

6.5.3 Determining E_{ref}

With J_{ref} it is possible to determine E_{ref} based on the spread resistance of individual coating defects and Ohms law (equation (9)). A generally accepted approach for determining the spread resistance R for a disc shaped coating defect it given in equation (b) for the defect diameter d and the soil resistivity ρ :

 $R = \frac{\rho}{2 \cdot d} \tag{b}$

It is evident that the spread resistance is controlled by the soil resistivity and the defect size. Practical experience shows that it is normally not possible to determine the spread resistance of a coating defect based on the soil resistivity and the coating defect size. This is a consequence of the reduction of the local soil resistivity at the coating defect due to the increased OH⁻ ion concentration as well as their distribution into the adjacent soil. Consequently, effective CP and the associated concentration polarization results in a reduction in the spread resistance, a further increase of the current density and an increase of the potential gradient measured at the soil surface. This interaction between concentration polarization, spread resistance and DCVG indications usually makes it impossible to determine the actual coating defect size based on voltage gradients. The associated effects and the validated corresponding mathematical model are described in detail in [39, 87]. While it is not possible to determine the actual spread resistance and assess the defect size based on DCVG data and the

pipeline depth, it is possible to estimate the largest possible spread resistance and the associated largest possible coating defect size for the worst-case assumption of absence of any far reaching concentration polarization into the adjacent soil. This concept is discussed in the following.

Figure 25: Dependence of E_{ref} from defect diameter d (and defect surface A) and soil resistivity for a J_{ref} of 0.02 A/m².

Based on the assumed limitation of the pH increase to the very steel surface it is possible to calculate E_{ref} required to achieve J_{ref} for various coating defect sizes and soil resistivities. Combining equation (b) with equation (9) immediately allows to calculate the required E_{ref} to achieve the J_{ref} for any given coating defect diameter d as a function of the soil resistivity as shown in equations (f) and (g).

$$J_{ref} = \frac{8 \cdot (E_{IR-free} - E_{ref})}{\rho \cdot \pi \cdot d} \tag{f}$$

$$E_{ref} = E_{IR-free} - J_{ref} \cdot \frac{\pi \cdot d \cdot \rho}{8}$$
(g)

Assuming a J_{ref} of 20 mA/m² and making a conservative assumption for $E_{IR-free}$ of -1 V_{CSE}, it is immediately possible to calculate the required E_{ref} for various coating defect sizes. Again, these are worst-case assumptions, since any effect of concentration polarization on the spread resistance is ignored. The results are shown in Figure 25. It is evident that higher soil resistivities and larger coating defects require more negative on potentials to achieve the current density of 20 mA/m². This current density is expected to result in a surface pH of 10.5 and an IR-free potential of -0.95 V_{CSE} in the case of good bedding conditions according to Figure 18.

In the practical application the defect size on the pipeline is usually not known and it is generally accepted that it is not possible to determine it based on DCVG surveys due to unknown level of spread of the concentration polarization into the soil and the resulting decrease of the soil resistivity in the direct vicinity of the coating defect. Despite of unknown defect size distribution on the pipeline it is common practice to install coupons for demonstrating the effectiveness of CP. Based on the data in Figure 25 it is evident that demonstration of CP effectiveness is only possible, if the coupon is installed in the highest possible soil resistivity and if the coupon represents the largest possible coating defect. If it is accepted that the coupons are suitable for demonstrating the effectiveness of CP, then the choice of coupons size can be used to determine the required level of E_{ref} based on Figure 25. For a 100 cm² coupon in soil resistivity of up to 300 Ω m an on potential of -1.28 V_{CSE} would be required to achieve 20 mA/m², a pH of 10.5 and an IR-free potential of -0.95 V_{CSE} at the steel surface.

Figure 26: Worst-case coating defect diameter in absence of far reaching concentration polarization calculated for various DCVG indications as well as pipeline depths x.

This argumentation can be further extended. The use of this coupon is only justified if it is confirmed that there are no bigger coating defects on the pipeline. It is common practice to perform DCVG on pipelines and it is justified and appropriate to profit from these data for the assessment of the effectiveness of CP. According to [26] the diameter d of a coating defect in absence of any concentration polarization can readily be determined based on the %IR (or U/U_0) obtained in DCVG surveys for a given distance x of the reference electrode from the disc shaped defect according to equation (c):

$$d = 2x \cdot \tan\left(\frac{U \cdot \pi}{\cdot 2}\right) = 2x \cdot \tan\left(\frac{\% IR \cdot \pi}{200}\right)$$
(C)

The calculation of the worst-case size coating defects associated with the obtained DCVG indications according to equation (d) is shown in Figure 26. The following conclusions are possible based on the worst-case consideration assuming absence of any spread of alkalinity from the steel surface into the adjacent soil:

- a) The largest possible coating defect can be determined based on the %IR indications. This allows to verify the appropriateness of the chosen coupons size for the assessment of the effectiveness of CP. For the example of the 100 cm² coupon it follows from Figure 26 that all coating defects with more than 3 %IR at a pipeline depth of 1.5 m or more need to be repaired. Otherwise the choice of coupons size cannot be justified.
- b) Alternatively, it is possible to recalculate the required E_{ref} to ensure effective CP based on the determined worst-case defect size according to equation (c) based on equation (g) or Figure 25.
- c) If neither item a) or b) are possible the actual current density for a worst-case consideration can be calculated and compared to the parameters in Table 1 and equation (a). This procedure allows for identifying areas with increased risk that will justify further consideration.

This simple argumentation highlights the possibilities associated with the use of generally accepted concepts in CP. Their combination with well-established chemical and physical concepts allows for determining effectiveness of CP, optimize the protection or perform a risk assessment. Since this approach is based on readily accessible CP parameters the problems associated with the determination of an IR-free potential are overcome and an assessment of actual cause and effect is performed.

The acceptable threshold with respect to %IR is often discussed. Some companies use %IR indication thresholds for the integrity management of pipelines. Based on the present discussion it immediately follows that it is readily possible to perform an assessment of effectiveness of CP based on DCVG for the special case of a worst-case consideration. Combining equations (b), (c) and (9) immediately provides equation (d) that describes the current density achieved for a given DCVG indication at a given on-potential. This argumentation can be further expanded and the required E_{ref} can be calculated to ensure a required J_{ref} based on equation (e).

$$J_{ref} = \frac{4 \cdot (E_{IR-free} - E_{ref})}{\rho \cdot \pi \cdot x \cdot \tan\left(\frac{\% IR \cdot \pi}{200}\right)} \tag{d}$$

$$E_{ref} = E_{IR-free} - J_{ref} \cdot \tan\left(\frac{\% IR \cdot \pi}{200}\right) \frac{\pi}{4} \cdot \rho \cdot x$$
 (e)

An application example of equation (e) is shown in Figure 27 for a J_{ref} of 2 mA/m². According to Figure 18 this will result in a surface pH higher than 9 and an IR-free potential more negative than -0.85 V_{CSE}. According to Figure 27 this current density of 2 mA/m² is exceeded for soil resistivities smaller than 300 Ω m and for on-potentials more negative than -1.2 V_{CSE} even at 30% IR indications. At 1000 Ω m this current density and hence this level of corrosion protection could be ensured up to 10% IR indications. This is well in line with the field experience on numerous FBE coated pipelines that are not allowed to be operated at on-potentials significantly more negative than -1.2 V_{CSE} in order to avoid overprotection is achieved even at increased %IR-indications, provided the soil conditions according to EN ISO 15589-1 allow for the use of an E_p of -0.95 V_{CSE}.

In Figure 28 the same consideration is shown für a J_{ref} of 20 mA/m². According to Figure 18 this will result in a surface pH of 10.5 and an IR-free potential more negative than -0.95 V_{CSE}. This E_p and hence this current density are required to ensure corrosion protection in anaerobic aggressive soils according to EN ISO 15589-1. Based on Figure 28 this current density of 20

mA/m² is exceeded for soil resistivities smaller than 30 Ω m and for on-potentials more negative than -1.3 V_{CSE} even at 30% IR indications. At 300 Ω m this current density can only be ensured at the on-potential of -1.3 V_{CSE} for %IR indications of up to 3%.

Figure 27: Dependence of E_{ref} from the %IR indication (U/U₀) and soil resistivity for the following conditions: J_{ref} of 0.002 A/m²; x of 1.5 m and $E_{IR-free}$ of -0.85 V_{CSE}.

Figure 28: Dependence of E_{ref} from the %IR indication (U/U₀) and soil resistivity for the following conditions: J_{ref} of 0.02 A/m²; x of 1.5 m and $E_{IR-free}$ of -0.95 V_{CSE}.

Figure 29: Dependence of E_{ref} from the %IR indication (U/U₀) and soil resistivity soil resistivity for the following conditions: J_{ref} of 0.1 A/m²; x of 1.5 m and $E_{IR-free}$ of -1.0 V_{CSE}.

In Figure 29 the same consideration is shown für a J_{ref} of 100 mA/m². According to Figure 18 this will result in a surface pH above 11 and an IR-free potential more negative than -1.0 V_{CSE}. According to Figure 29 this current density of 100 mA/m² is exceeded for soil resistivities smaller than 30 Ω m and for on-potentials more negative than -1.4 V_{CSE} only at 7% IR indications. At 300 Ω m this current density can only be ensured at an on-potential of -1.4 V_{CSE} for %IR indications of up to 0.7%. According to Figure 26 this corresponds to a coating or coupon defect size of 10 cm². The application of an Eon of -1.4 VCSE in soils of 300 Ω m will ensure current densities of 100 mA/m² on all coupons and coating defects smaller than 10 cm².

Note that this consideration applies to the worst-case consideration where no spread of alkalinity into the surrounding soil has occurred. With a spread of alkalinity, a significantly increased current density of 350 mA/m² would be expected when calculated on the validated numerical model presented in [39]. In contrast, the formation of a calcareous deposit layer that covers 90% of the coupon would result in a current density of 26 mA/m² calculated with an assumed defect surface of 10 cm². However, this is only a result of an erroneous defect surface. In fact, the formation of the calcareous deposit would have caused an effective current density on the remaining steel surface of 1 cm² of 264 mA/m² when the real defect surface would have been considered. This 1 cm² is the remaining surface of a 10cm² defect with 90% coverage of calcareous deposits.

This discussion highlights the relevance of the influencing parameters, the possibility of assessing the worst-case coating defect size and calculating the required E_{ref} for ensuring a minimum current density with respect to providing effective CP.

6.5.4 Soil resistivity

The above discussion has highlighted the relevance of the soil resistivity in assessing effectiveness of CP. While the procedures for measuring the resistivity are well established the number of data points and their treatment needs further consideration. In the last six years, the soil resistivity in Switzerland has been systematically recorded at all test points.

Possible reference soil resistivity val-	Measured example values over 3 measuring points $[\Omega m]$ and the correspond- ing minimum and maximum reference values			
ues [Ωm]	Measured value		Resulting reference value	
3	max.	min.	Max.	Min.
10	75	35	100	30
30	210	35	300	30
100	400	35	1000	30
300	400	25	1000	10
1000	400	6	1000	3

Table 2:Estimation of the soil resistivities

Thus, at least some general soil resistivity values are available with respect to local conditions. Of course, a variation of the soil resistivity along the pipeline has to be expected. In addition, the values depend on temperature, weather and seasonal influences. Regarding these considerable influencing factors, a procedure was established that takes into account the resistivity values of the previous as well as the following test post. Based on these three data sets an evaluation for the lowest (with respect to the AC corrosion risk) and the highest (with respect to effective CP) soil resistivity value is performed. The highest value of the three is rounded up and the lowest rounded down to the next reference resistance value according to the first column in Table 2. Table 2 also shows some examples. The resistance distribution over several test posts is represented by the reference values. If knowledge of locally further reduced soil resistivity is available (for example, a horizontal drilling in bentonite), these values must also be considered in addition to the actually measured values. In critical cases, a further limitation of the soil resistivity values can be achieved by geoelectrical probing on the affected section.

6.6 Implementation of the protection criteria

Based on the presented model concepts on the mechanism of cathodic protection in combination with the effects taking place during anodic as well as cathodic stray current interference a comprehensive description of all aspects of the associated relevant corrosion processes has been developed. For the first time a unique procedure for addressing all influencing parameters in a single approach is has been developed. Independent on the level of interference, the type of CP system, the type of coating, the presence of bonding of the pipeline and the possibility to determine instant off-potentials, a straightforward assessment of the effectiveness of CP is available. This approach is not limited to addressing the effectiveness of CP but takes into account also the conflicting requirements for mitigating DC interference and AC interference: DC interference with the risk of corrosion due to anodic polarization has to be addressed by shifting the on-potential more negative and AC interference with the risk of excessive cathodic polarization has to be mitigated by shifting the on-potential more positive. Based on this new approach a straightforward analysis of all corrosion risks and an optimized setting of the CP operation conditions are readily possible. Additionally, the identification of possible mitigation measures even in combined and complex interference situations is possible. This new approach that is presently implemented in Switzerland will be shortly discussed in the following based on a hypothetical example. A PE coated pipeline with a wall thickness of 5 mm in soil with a resistivity between 30 to 100 Ω m is assumed. This process requires three steps:

- 1. First E_{ref} needs to be determined. This value corresponds to the on-potential that is required to ensure J_{ref} and hence effective CP on all coating defects. Typical values for E_{ref} are between -1.0 to -1.6 V_{CSE}. The assessment of E_{ref} can for example be based on $E_{IR-free}$ values potentials in absence of DC interference on the pipeline, $E_{IR-free}$ values determined on coupons or current density considerations taking into account Figure 18. In the given example an E_{ref} of -1.05 V_{CSE} is determined as shown in Figure 30a. In absence of stray current interference effective CP is ensured when the E_{on} is more negative than the red line shown in Figure 30a.
- 2. Secondly the average anodic interference relative to E_{ref} is recorded over a representative period of time. For DC traction interference this is typically 24-hours. In the present example $\Delta E_{a,avg}$ was found to be about 0.4 V. The largest anodic interference period t_a is determined to be 250 seconds. According to equation (4) a required $E_{on,avg}$ of about -1.45 V_{CSE} is calculated. This corresponds to the blue line in Figure 30b. Effective CP under anodic stray current interference is ensured when the E_{on} averaged over 24 hours is more negative than the blue line shown in Figure 30b.
- 3. Thirdly the applicable interference line of Figure 24a for a PE coated pipeline for the smallest soil resistivity is selected. In the present example the soil resistivity ranges from 30 to 100 Ω m. Correspondingly, the orange line for 30 Ω m is selected. These threshold values for the acceptable average U_{ac} are plotted in Figure 30c in orange. Effective corrosion protection is ensured when the 24 h average U_{ac} is below the orange line and the 24 h average E_{on} is more negative than the blue line.

Based on this approach a straightforward assessment of effective corrosion protection is possible based on the present model concept of the mechanisms involved in cathodic protection. The example shown in Figure 30c emphasizes the conflicting requirements with respect to mitigating anodic stray current interference and AC interference. The mitigation of DC interference requires the shifting of the $E_{on,avg}$ to -1.45 V_{CSE}. It follows from Figure 30c that effective corrosion protection in this combined AC and DC interference situation can only be managed by lowering the average U_{ac} to values in the range of 5 V. In contrast, in absence of DC interference the risk of AC corrosion could effectively have been eliminated by applying a 24-hour average on-potential in the range of -1.2 V_{CSE}. In this case even significantly increased levels of U_{ac} can be tolerated and mitigation of AC corrosion is possible based on a simple adjustment of the rectifiers rather than the installation of expensive earthing installations.

Figure 30: Admissible average U_{ac} and average E_{on} for ensuring effective CP: a) E_{ref} required to ensure effective CP in absence of any interference; b) E_{on,avg} for effective corrosion protection in case of anodic DC interference; c) Requirements for effective CP in case of combined DC and AC stray current interference.

This example illustrates that the new understanding of the mechanisms involved in cathodic protection provides a comprehensive model that describes all influencing parameters. This model has some key consequences on a number of aspects that have not been appropriately addressed in the past:

- Cathodic protection is achieved by applying a cathodic current to a steel structure. While there is some contribution of an activation polarization at the very moment the current is applied, the relevant polarization that provides corrosion protection is a result of the concentration polarization. This change in concentration with respect to oxygen and hydroxide concentration at the steel surface requires time to build up.
- Due to the time constants involved in concentration polarization, the corrosion protection is not lost when CP is interrupted or when anodic current discharge occurs. The slow diffusion processes associated with the loss of concentration polarization explain the relevance of 24-hour average values and the irrelevance of anodic interference during time variant stray current interference.
- The IR-free potential is a measure of concentration polarization. It detects the depletion of oxygen and the level of pH increase at the steel surface as a result of the cathodic current. It corresponds to a pH measurement based on the hydrogen electrode on the steel surface. IR-free potentials (measured 0.1 to 1 second after interrupting the CP current) more negative than $-0.85 V_{CSE}$ are only possible at pH values higher than 9. At such high pH values conditions for the formation of a protective passive film are given in many soils.
- In aggressive soil conditions in presence of carbonates, sulphides and humic acids, as they are typically found under anaerobic conditions, a pH of 10.5 corresponding to an IR-free potential of -0.95 V_{CSE} is required for passivation and hence corrosion protection.
- The IR-free potential may not be used to assess corrosion protection in case of anodic interference or in aerated soils. The pH electrode only provides reliable data with respect to the pH-value when the steel potential is controlled by the hydrogen reduction. Oxygen and stray current turn these IR-free potentials and the associated pH-measurements into meaningless numbers.
- Anodic stray current interference is irrelevant with respect to the integrity of pipelines as long as the concentration polarization (i.e. the increased pH at the steel surface) is maintained. Longer anodic interference can hence be compensated by means of an increased level of CP.
- At very low levels of CP (e.g. at on-potentials of -0.9 V_{CSE}) no anodic current discharge can be tolerated based on the presented concept due to the absence of a enough concentration polarization.
- The new approach with respect to the assessment of effective CP under anodic stray current interference covers the entire frequency range of stray currents including AC interference, DC traction interference, tidal interference, telluric interference and even time constant anodic interference.
- While anodic AC interference has no relevance with respect to the integrity of pipelines in case of effective CP, high corrosion rates occur due to AC voltages in the case of over-polarization. Hence AC corrosion is a problem associated with over polarization resulting in cathodic interference.

- The numerical modelling of all relevant processes in CP provides a simple approach with respect to the assessment of the AC corrosion risk. The consideration of all influencing parameters allows for translating the requirements in EN ISO 18086 into potential based threshold values.
- The assessment of the stray current corrosion risk is readily possible based on DC and AC potential measurements on pipelines in normal operation conditions. This methodology is applicable to all pipelines independent on their interference condition and the possibility of synchronous interruption of all DC current sources, such as rectifiers, DC decoupling devices, galvanic anodes, drainages and bonds. For the first time the corrosion protection of all pipelines can be assessed based on the same concept.
- The effectiveness of CP can be assessed even in case of combined AC and DC interference. Based on the presented concepts the planning of mitigation measures and the assessment of their effectiveness are readily possible. Areas with high corrosion risk and possibly inacceptable conditions are identified. Based on this risk analysis the locations for the installation of coupons or probes can be identified. Having coupons installed in the highest risk areas significantly increases the value of their readings.
- The presented concepts have the status of a recognized state of the art in Germany based on the documents DVGW GW 21 and DVGW GW 28 B1.

The various aspects presented in this paper provide a complete description of cathodic protection and the relevant influencing parameters. Based on these concepts a new approach with respect to the assessment of the effectiveness of CP is possible. The effectiveness is determined based on the readily accessible average values of E_{on} and U_{ac}. This concept is applicable to all pipelines independent on the coating system, the interference conditions, the possibility to interrupt all CP current sources, the presence of galvanic anodes and drainage bonds. Since all relevant influencing parameters are considered, an objective assessment of corrosion protection possible. This approach allows for identifying areas with increased risk and the evaluation of mitigation measures. In areas with conflicting requirements it may not be possible to find ideal solutions with respect to corrosion protection. Based on these concepts it will, however, be possible to operate the CP under optimized conditions while minimizing the corrosion risks caused by conflicting requirements. Considering the relevant associated costs of mitigation, the use of objective, technically correct and justifiable assessment criteria is highest importance.

7 Summary

This document summarizes the outcome of the discussion performed within the joint EFC and CEOCOR working group "protection criteria" with respect to the processes taking place under cathodic protection, the relevance of the present protection criteria in EN ISO 15589-1 and the possible solutions with respect to future criteria. The document was elaborated from 2016 to 2021 in a total of 11 working group meetings at Ceocor and Eurocorr congresses. The document contains a description of relevant influencing parameters and proposes a possible new procedure of assessing effectiveness of cathodic protection that is based on the DVGW documents GW10, GW 21, GW27, GW 28 and GW 28 B1. These documents have the status of a recognised state of the art within Germany. The benefit of this proposed procedure is that it is based on on-potentials and AC voltages. As a consequence, it has the potential to be applied to all cathodically protected buried steel structures irrespective of their size, their interference condition, the possibility to determine instant off potentials and their size. The goal of this document is to illustrate a possible procedure for assessing the optimized operation conditions of cathodic protection within the conflicting requirements given in EN ISO 15589-1, EN ISO 18086, EN 50162 and ISO 21857. This procedure not only allows for demonstrating effective corrosion protection but also identifying areas of lowest level of corrosion protection and optimal locations for the installation of coupons. These identified locations are expected to represent the most severe conditions on the pipeline and the installed coupons are, therefore, representative of these worst-case conditions. It is suggested that the installation of coupons in these locations can justify the use of a limited number of coupons for demonstrating effectiveness of cathodic protection.

Within the working group agreement has been achieved on the following aspects of this document:

- Cathodic protection is achieved by applying a cathodic current to the buried steel structure.
- The cathodic current results in an increase of the surface pH and this process is of high importance for the corrosion protection to be achieved. Beneficial aspects of the pH increase are the formation of calcareous deposits and the generation of an environment that is conducive to the formation of protective oxide films.
- The protection criteria given in EN ISO 15589-1 with respect to the IR-free potential are relevant for assessing the effectiveness of cathodic protection.
- The instant-off potential may only be used as an approximation to the IR-free potential and will not provide relevant information in the case of DC and AC interference conditions or in case of exchange currents. As a consequence, on modern well-coated structures assessing the effectiveness of CP is only possible with coupon measurements.
- The on-potential cannot be used as an approximation to the IR-free potential. However, the on-potential is a parameter that can be assessed on all pipeline systems independent on the level of AC and DC interference, the ability to synchronously switch rectifiers and decoupling devices as well as the size of the pipeline network.
- The on-potential affects the current density on individual coating defects. At a given on-potential decreasing current densities are obtained with increasing coating defect size and increasing soil resistivity. In contrast shifting the on-potential to more negative values will result in higher current densities on a given coating defect in a given soil resistivity.

 The present standards have conflicting requirements: EN ISO 15589-1 and EN 50162 may require the shifting of the on-potential in the negative direction to meet the protection criteria associated with the IR-free potential. In contrast the meeting of the limiting DC current density of 1 A/m² in EN ISO 18086 as well as limiting over-polarisation according to EN ISO 15589-1 may require shifting the on-potential in the positive direction.

While there was agreement on these important aspects, a number of important issues remain open for future discussion:

- No agreement could be achieved whether protective oxide films will spontaneously form in the alkaline environment formed at the steel surface. While one view was that it was sufficient for the protective oxide film formation to keep the IR-free potential in the passivity domain of the Pourbaix diagram, another view was that an oxidizing action caused by an anodic current discharge during AC or DC interference was required for its formation.
- The properties of the protective surface oxide films, their composition as well as their stability as a function of soil composition and formation conditions were not further detailed within the working group.
- While there was agreement that the soil parameters listed in this document are relevant with respect to the cathodic current density to achieve corrosion protection, no agreement was achieved whether this list is final or whether the proposed quantification of the influence of these parameters is correct.
- While soil resistivity and soil aeration are important parameters for assessing the protection criteria with respect to the IR-free potential according to EN ISO 15589-1, no final proposal with respect to the assessment method and the lateral measurement resolution for these parameters was agreed on.
- The assessment of the effectiveness of CP based on 24-hour average values of onpotential and AC voltage was not considered relevant by all participants of the working group.
- The applicability of electrochemical equilibrium thermodynamics presented in this document are put into question as detailed in [88]. Instead, the relevance of electrode kinetic consideration is considered more relevant as discussed in [89].

Despite of the open aspects it was agreed to finalize this document. It will be relevant to apply these concepts and test the obtained results in the field application. The assessment of most of the parameters proposed in this document are anyways required for a competent application of the present EN ISO 15589-1, since the protection criteria based on IR-free potentials depend on soil resistivity. Furthermore, the performing of DCVG investigations, the determination of the on-potential and the AC voltage are common practice for most of the operators. Since the most relevant data are anyways collected by most pipeline operators on the basis of their function and effectiveness control of cathodic protection, the proposed procedures can readily be applied and tested. First promising results are meanwhile available [90]. The collected experience in the next years will allow for a revisiting and further elaboration of this document.

8 Referenced standards and codes

- DVGW GW 21:2014 Stray current interference of buried metallic structures caused by direct current systems (Beeinflussung von unterirdischen metallischen Anlagen durch Streuströme von Gleichstromanlagen)
- DVGW GW 28:2014 Assessment of the corrosion risk of cathodically protected pipelines caused by alternating current (Beurteilung der Korrosionsgefährdung durch Wechselstrom bei kathodisch geschützten Stahlrohrleitungen und Schutzmassnahmen)
- DVGW GW 28 B1:2018 Assessment of the corrosion risk of cathodically protected pipelines caused by alternating current: Annex 1: Modified criteria for AC voltage and on-potential (Beurteilung der Korrosionsgefährdung durch Wechselstrom bei kathodisch geschützten Stahlrohrleitungen und Schutzmassnahmen: Beiblatt 1: Modifizierte Kriterien für Wechselspannung und Einschaltpotential)
- EN 12954:2001 Cathodic protection of buried or immersed metallic structures. General principles and application for pipelines
- EN 13509:2003 Cathodic protection measurement techniques
- EN 14505:2005 Cathodic protection of complex structures
- EN 50162:2004 Protection against corrosion by stray current from direct current systems
- EN ISO 15589-1:2015 Petroleum, petrochemical and natural gas industries Cathodic protection of pipeline systems - Part 1: On-land pipelines
- EN ISO 18086:2019 Corrosion of metals and alloys Determination of AC corrosion Protection criteria
- ISO 21857:2021 Prevention of corrosion on pipeline systems influenced by stray currents
- NACE SP0169:2013 Control of External Corrosion on Underground or Submerged Metallic Piping Systems

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