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The effect of the steel-concrete interface on chloride-induced corrosion initiation in concrete

Authors

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Abstract

The steel-concrete interface (SCI) is known to influence corrosion of steel in concrete. However, due to the numerous factors affecting the SCI – including steel properties, concrete properties, execution, and exposure conditions – it remains unclear which factors have the most dominant impact on the susceptibility of reinforced concrete to corrosion. In this literature review, prepared by members of RILEM technical committee 262-SCI, an attempt is made to elucidate the effect of numerous SCI characteristics on chloride-induced corrosion initiation of steel in concrete. We use a method to quantify and normalize the effect of individual SCI characteristics based on different literature results, which allows comparing them in a comprehensive context.

It is found that the different SCI characteristics have received highly unbalanced research attention. Parameters such as w/b ratio and cement type have been studied most extensively. Interestingly, however, literature consistently indicates that those parameters have merely a moderate effect on the corrosion susceptibility of steel in concrete. Considerably more pronounced effects were identified for i) steel properties, including metallurgy, presence of mill scale or rust layers, and surface roughness, and ii) the moisture state. Unfortunately, however, these aspects have received comparatively little research attention. Due to their apparently strong influence, future corrosion studies as well as developments towards predicting corrosion initiation in concrete would benefit from considering those aspects. Particularly the working mechanisms related to the moisture conditions in microscopic and macroscopic voids at the SCI is complex and presents major opportunities for further research in corrosion of steel in concrete.

Keywords

Steel-concrete interface; Interfacial transition zone; Durability; Corrosion; Inhomogeneity; Variability

This article has been prepared within a framework of RILEM TC 262-SCI. The article has been reviewed and approved by all TC members.

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1 Introduction

The steel-concrete interface (SCI) has, in the literature, frequently been claimed to play a major role in initiation of chloride-induced reinforcing steel corrosion in concrete [1-11]. These studies generally focused on one particular characteristic of the SCI, such as the influence of a lime-rich layer at the steel surface or the presence of interfacial voids. However, given the complexity of the SCI and the variety of characteristics that may or may not occur locally [12], it remains unclear which are the most dominant influencing factors at the SCI that govern corrosion initiation. Here, we summarize available literature documenting the effect of numerous characteristics of the SCI on chloride-induced corrosion initiation in concrete, and an attempt is made to study these individual effects in a comprehensive context. The aim is to elucidate the dominating characteristics and to identify areas where further research is needed.

This paper was prepared by members of RILEM TC 262-SCI, and is closely linked to a recent publication [12] that presented a systematic approach to describe the SCI in terms of local characteristics and their physical and chemical properties. Here, we focus on the effect of SCI characteristics on chloride-induced corrosion of carbon steel reinforcement in concrete. Coated, alloyed or high-strength steels as well as influences of corrosion inhibitors and electrochemical techniques (e.g. cathodic protection) are not considered. Note that we focus on *corrosion initiation*, which we consider as the transition of the steel from passivity to stable localized active corrosion. Literature on corrosion propagation (factors influencing the corrosion rate, corrosion-induced concrete cracking, etc.) are beyond the scope of this paper. We are aware that the community has struggled to agree on a definition of “corrosion initiation”. Nevertheless, to avoid being forced to exclude a significant portion of the available literature, we have considered all relevant studies that, according to their authors, were designed to investigate chloride-induced corrosion initiation in concrete.

2 Methodology of reviewing the literature

2.1 Considered literature studies

To quantify the impact of SCI characteristics on the initiation of chloride-induced corrosion, reported experimental results were collected which allow the corrosion susceptibility of steel in an alkaline system to be assessed in the absence or presence of certain SCI characteristics. This objective imposes a number of requirements to the studies in order to be considered suitable for our review. Here, we considered studies in which the steel sample was exposed in an alkaline solution or in a cementitious system, and where the chloride concentration was systematically varied (added at different concentrations or increased over time). In order to quantify the susceptibility to corrosion initiation under these conditions, we looked for studies that either reported pitting potentials (E_{pit}), critical chloride contents (C_{crit}), or that measured times to corrosion initiation in solutions (t_{ini}). The vast majority of the reviewed literature quantified the susceptibility to corrosion in terms of C_{crit} . This parameter represents the chloride concentration in the concrete (or in a solution) at which the transition from passive steel to active corrosion occurs. In the literature, different methods are used to determine C_{crit} , and it is well-known that the chosen methodology affects the results [13]. Nevertheless, we consider C_{crit} a suitable parameter for our review, mostly because of the broad acceptance of this concept to characterize “corrosion initiation” of steel in concrete [13, 14]. In the literature, C_{crit} is reported in terms of the free (dissolved) chloride ion concentration in the electrolyte (liquid phase) or in terms of total (bound and free) chlorides in concrete, mortar, or cement paste.

2.2 Evaluation to quantify the effect of SCI characteristics

We used the following equation to quantify the effect of a particular characteristic, x , of the SCI:

$$\text{Effect of } x: \quad E_{x,j} = \frac{C_{crit}^{x,j} - C_{crit}^{Ref,j}}{\min\{C_{crit}^{x,j}, C_{crit}^{Ref,j}\}} \quad (1)$$

Here, $C_{crit}^{x,j}$ is the critical chloride content measured for the SCI characteristic x reported in study j , and $C_{crit}^{Ref,j}$ is the critical chloride content of the reference case in the same study j . As an example,

consider the case of the influence of mill-scale on the steel surface: Here, $C_{crit}^{x,j}$ would be C_{crit} in the presence of mill-scale in a certain study j , and $C_{crit}^{Ref,j}$ would be C_{crit} in the absence of mill-scale (reference case; e.g. removed by sandblasting) *in the same study* j .

Because C_{crit} values in the literature are known to be significantly affected by the test method used [13], we chose to normalize the effect of x by adopting the minimum of $C_{crit}^{x,j}$ and $C_{crit}^{Ref,j}$. With this normalization, the application of eq. 1 to literature data allows the effect of different SCI characteristics (x) determined in different studies (j) to be compared. Additionally, the sign of $E_{x,j}$ illustrates if the effect is positive or negative, and the minimum in the denominator of eq. 1 ensures that negative and positive effects are equally scaled.

Consider that n is the number of parallel samples (replicates) used in a study. For a study with $n=1$ (following the example of mill-scale: one reported result for C_{crit} with mill-scale, and one reported result for C_{crit} without mill-scale), eq. 1 would produce one single value for the particular study. In many studies, however, replicate samples were tested ($n>1$), and thus, various results for both $C_{crit}^{x,j}$ and $C_{crit}^{Ref,j}$ were reported. The average effect of x in a given study j becomes:

$$\bar{E}_{x,j} = \frac{\bar{C}_{crit}^{x,j} - \bar{C}_{crit}^{Ref,j}}{\min\{\bar{C}_{crit}^{x,j}, \bar{C}_{crit}^{Ref,j}\}} \quad (2)$$

Here, $\bar{C}_{crit}^{x,j}$ is the arithmetic mean of all $C_{crit}^{x,j}$ reported within one study j , and $\bar{C}_{crit}^{Ref,j}$ is the arithmetic mean of all $C_{crit}^{Ref,j}$ within study j . Eq. (3) was used to express the standard deviation of the effect of x in study j [15]:

$$SD_{E_{x,j}} \approx \sqrt{\left(\frac{\bar{C}_{crit}^{x,j}}{\bar{C}_{crit}^{Ref,j}}\right)^2 \cdot \left[\left(\frac{SD_{x,j}}{\bar{C}_{crit}^{x,j}}\right)^2 + \left(\frac{SD_{Ref,j}}{\bar{C}_{crit}^{Ref,j}}\right)^2\right]} \quad (3)$$

Here, $SD_{x,j}$ is the standard deviation of all (n) $C_{crit}^{x,j}$ reported within one study j , and $SD_{Ref,j}$ is the arithmetic mean of all $C_{crit}^{Ref,j}$ within study j .

Fig. 1 schematically shows how eqs. 1–3 were used to graphically represent the effect of a certain characteristic within different studies. This evaluation helped in assessing if there is agreement or disagreement concerning the effect of a certain characteristic x within the available literature, both in a qualitative (positive vs. negative effect) and a quantitative manner (numbers on the ordinate).

2.3 Remarks on the chosen methodology

Note that we assumed that all the examined characteristics of the SCI, x , are independent. We are aware of the fact that this may not entirely be the case. However, it should be mentioned that most experiments were designed to study a particular parameter (e.g. the cement type) keeping all other parameters constant. Since it is practically impossible to experimentally provide systematic data with sufficiently high number of combinations of the wide variety of characteristics (full factorial experimental design), we believe there are currently no better alternatives to the above-presented approach. Valuable conclusions can be drawn from this review, *but it is important to bear the limitations in mind and consider the outcome primarily as a guidance for further research rather than being directly applicable in engineering.*

We did not exclude any studies because of potential weaknesses in experimental methodology, because an assessment of the quality of the different studies would be far from straightforward, especially considering the controversial discussions in the literature regarding criteria to detect (stable) corrosion initiation in concrete, or how to measure critical chloride content experimentally [13, 16]. In other words, for the sake of a systematic and transparent methodology in evaluating the literature, we considered all available studies as equally relevant.

Note that the order of presenting information in the following sections is not according to their relevance, but follows the structure and conceptual treatment of the SCI presented in Ref. [12].

3 Documented influences

3.1 Reinforcing steel type

3.1.1 Metallurgy

In the 1970s, thermomechanical strengthening processes (quenching and self-tempering) were developed and this rapidly phased out cold-work hardened reinforcing steel bars from the construction markets in many countries. Nowadays, quenched and self-tempered steels with a tempered martensite (TM) periphery and a ferrite-pearlite (FP) core (contributing to the desired strength and ductility) are widely used in practice. A number of investigations focused on the mechanical behavior of this new type of reinforcing steel. In one of the early of these works Rehm and Russwurm [17] indicated that “*it is absolutely necessary to carry out corrosion tests*”. However, this issue has received very little research attention.

There are various studies in the literature focusing on the influence of steel microstructure on corrosion (generally on corrosion rate) in CO₂ or in pH-neutral chloride solutions, but with somewhat contradictory results [18-20]. The effect of inclusions on pitting corrosion initiation has also been studied, mostly in stainless steels [21]. However, limited literature data are available on the effect of metallurgy on chloride-induced corrosion initiation in alkaline environments. Trejo and Pillai [22, 23] compared C_{crit} of conventional (ferrite-pearlite), microcomposite (ferritic-martensitic), and stainless steels embedded in mortar. The ferritic-martensitic steels exhibited significantly higher C_{crit} (approx. by a factor of 10) than conventional steels and similar chloride threshold as the 304 grade stainless steels. Angst and Elsener [24] compared reinforcing steels (in as-received condition) from 9 different countries with respect to their corrosion behavior in chloride-containing saturated Ca(OH)₂ solution. The steel microstructure was identified as one of the major influencing parameters. C_{crit} was statistically significantly lower (by a factor > 2) for reinforcing steel that underwent thermomechanical strengthening (TM surface layer) compared to cold-work hardened steel (FP microstructure). Kumar [25] cut coupon specimens of TM and FP from the periphery and core, respectively, of a quenched and self-tempered steel rebar and immersed them in simulated pore solutions (pH ≈ 13) with various chloride concentrations. The average C_{crit} for TM coupons was slightly higher (by 11–16%) than for FP coupons. Note, however, that the tested samples were sections of steel bars, which may not be representative of the reinforcing bar surface. Further studies [26-28] investigated influences of steel metallurgy on corrosion in concrete, including dual-phase reinforcing steels, however, these studies did not report C_{crit} or other quantified measures of corrosion initiation.

In short, little information on the effect of metallurgy on corrosion initiation of steel in concrete is available. Although the few available studies are contradictory, evaluating them [22-25] according to section 2 indicates an overall effect of the order of 3–4. At this stage, it can be concluded that the microstructure of the rebar surface is likely to play an important role in the corrosion initiation. However, further studies are needed in this area.

3.1.2 Rebar geometry

The geometry of rebars may influence corrosion initiation by modifying the SCI locally, namely due to the presence of ribs on the rebar surface, different diameters, or bent parts of the rebars. We expect that such effects would ultimately be related to microscopic level phenomena, including metallurgical influences (section 3.1.1), reinforcing steel surface (section 3.2), or interfacial voids (section 3.4). However, literature is not consistent. While Alonso et al. [29] reported negligible differences in C_{crit} between ribbed and smooth rebars, Zafar and Sugiyama [30] found an increase in C_{crit} by factor 2 between deformed and plain rebars. Moreover, it was noted that for deformed bars, the corrosion started at ribs, whereas for plain bars the trigger for corrosion initiation was always an air void. Michel and Angst [31] found that, on largely rust-free surfaces, corrosion tends to initiate close to or on ribs, whereas on rusty rebars, corrosion does not preferably initiate on ribs (corrosion tests in alkaline solutions). In samples taken from engineering structures, Boschmann et al. [32] reported preferential corrosion initiation close to and on ribs for one concrete structure, while this was not observed in two other structures. Sandberg [4] found that corrosion initiated in approx. 70% of the cases in the deformed

parts of reinforcing steel bars, particularly on the inside of the U-bend rebars. This was attributed to mechanical damage to oxide scales on the rebar during bending. In summarizing the literature, it is clear that the current level of documentation does not permit us to quantify the effect of the rebar geometry on corrosion initiation.

3.2 Reinforcing steel surface

3.2.1 Mill scale and pre-existing rust layers

Different researchers reported [33-40] the influence of steel surface modifications (removing mill scale through sandblasting, polishing or pickling) on C_{crit} . Some studies were undertaken in mortar or concrete [36, 37, 40] and some in alkaline solutions [33-35, 38]. Fig. 2 shows the evaluation of the data according to section 2.2. In most studies, the presence of these oxide scales favored corrosion initiation; the effect (quantified according to eqs. 1 and 2) was generally in the range 0 to -2. In one study [35] the effect was much more pronounced and approached -6. In two studies [33, 40], however, it was found that the presence of the mill scale raised C_{crit} . The mill-scale was reported to be particularly dense and continuous, which appeared to have a corrosion-protective effect. However, mill scale on most commercially available steel reinforcement is non-uniform, discontinuous, and contains cracks and crevices [41, 42]. Therefore, the presence of mill scale appears generally to promote corrosion initiation.

A number of studies [24, 37, 38, 42-44] suggested that the effect of pre-existing rust layers on corrosion initiation may be different from that of mill scale. Mohammad and Hamada [44] as well as Chen et al. [38] reported that C_{crit} was lower in the presence of mill scale with additional rust than with mill scale alone. Angst and Elsener [24] compared rebars from different manufacturers and in different pre-rusted condition and reported that the degree of rust did not permit any reliable prediction of the susceptibility to chloride-induced corrosion. The influence of pre-formed rust scales was considered a weaker influencing factor than the manufacturing process or the metal microstructure. Al-Tayyib et al. [43] reported that pre-rusting did not favour corrosion initiation of rebars embedded in concrete.

3.2.2 Passive film

Apart from possible effects of steel microstructure (e.g. inclusions) the passive film is the main barrier against pitting corrosion. The passive state of steel in alkaline environments such as concrete remains stable until chloride ions reach the steel surface. During prolonged exposure (ageing) of black steel to alkaline solutions or in concrete, the open circuit potential (OCP) asymptotically increases to more positive values, see [12] and Refs. cited therein. Several authors reported that ageing of the passive film (pre-passivation) of black steel in alkaline media results in an increase in the pitting potential, E_{pit} [45-48]. The pitting potentials were measured in alkaline, chloride-containing solutions upon a given time of pre-passivation in chloride-free solutions of equal pH. For sandblasted steel in synthetic pore solution an increase of the pitting potential of about 150 mV was reported for a pre-passivation time of 30 days (Fig. 3) both for mildly corrosive conditions ($Cl^-/OH^- = 1$) and for severe conditions ($Cl^-/OH^- = 25$) [46, 49]. It is worth noting that the statistical distribution of the pitting potentials was very narrow for aggressive environments (high Cl^-/OH^- ratio) and relatively broad for mild conditions. In the case of ground or mechanically polished steel surfaces [45, 50] the effect of pre-passivation time on the pitting potential was found much more pronounced than for sandblasted steel [46, 49]. Studies on pure iron showed that pre-passivation in chloride-free solution for 2 h resulted in a breakdown of the passive film after 60-80 h exposure to chlorides, while longer pre-passivation of 36 hours resulted in breakdown after 100-120 h [47]. Similarly, potentiostatic ageing (pre-polarization) under conditions where pitting does not occur ($E_{cor} < E_{pit}$) tends to improve the resistance against later pitting corrosion [51, 52].

According to established corrosion theory [53], increasing E_{pit} is associated with an increase in C_{crit} . According to Fig. 3, E_{pit} increases by roughly 40 mV when steel is pre-passivated for one month instead of 1 week. From studies [33, 35, 54], it is known that E_{pit} decreases by about 400 mV for an increase in chloride concentration by a factor of 10. Thus, as a first-hand estimate, it can be concluded that pre-passivating for 1 month instead of 1 week (or 1 week instead of 1 d) increases C_{crit} by a factor of 1.25.

The association of E_{pit} with C_{crit} is also reflected in the observation that, all other conditions staying the same, C_{crit} tends to increase when the steel potential prior to corrosion initiation is made more

negative, either by macrocell coupling with other parts of the system or by applied polarization current [55, 56].

3.2.3 Steel surface preparation

It has been widely reported that higher C_{crit} were observed for rebars with modified surfaces, that is, through sandblasting, polishing or pickling than those in as-received conditions [33, 34, 40, 50, 57-60], as discussed in section 3.2.1. Moreover, literature indicates that the susceptibility to corrosion decreases with the degree of surface preparation. For instance, Mammoliti et al. [33] found that with an increasing degree of polishing (from grit 240 to grit 600, thereby reducing the surface roughness), pitting potentials of carbon steel in $\text{Ca}(\text{OH})_2$ solution were shifted to more anodic values by at least 300 mV, occasionally even up to 600 mV, at various chloride concentrations. In similar tests, Figueira et al. [60] found a shift of E_{pit} by approx. 300 mV in anodic direction for carbon steel in various alkaline simulated pore solutions when moving from 500 grit SiC paper to 1 μm diamond polishing suspension. Consistent with this, comparable relationships between E_{pit} and degree of polishing (surface roughness) were reported from studies on stainless steels, e.g. [61], although the impact was generally less pronounced than in [33, 60]. Following the reasoning presented in section 3.2.2, it can be inferred that such shifts in E_{pit} due to decreased surface roughness have a dramatic influence on C_{crit} . With the data reported in [33, 60], polishing to grit 600 instead of 240 or polishing with 1 μm diamond polishing suspension instead of 500 grit, would raise C_{crit} by roughly a factor of 10. While it may be concluded that the steel surface preparation that goes beyond removing oxide scales has a strong effect on corrosion initiation in concrete, it should be noted, however, that this is based on only two studies for carbon steel.

As a final comment, we would like to highlight that the surface of steel reinforcement is typically not modified in practice. However, corrosion sensors such as steel coupons used in structural monitoring often have sandblasted or polished surfaces. Additionally, such surface treatments may also be used in laboratory studies on corrosion of steel in concrete. Results from such studies as well as results from steel coupons exhibiting modified surfaces, e.g. in monitoring systems, should thus be interpreted with caution.

3.3 Concrete microstructure and chemistry at the SCI

3.3.1 w/b ratio

The water/binder ratio (w/b) is known to be an important parameter in concrete. Its effects on compressive strength and transport, e.g. of chloride ions, are well documented. The effect of w/b on corrosion initiation, on the contrary, is less well documented. C_{crit} reported by [62-69], generally based on Portland cements, are compiled here and evaluated according to section 2.2 with the reference of w/b = 0.50 (Fig. 4). Some publications did not include data for w/b = 0.50 [66, 68]; in those cases, results were interpolated. Hansson & Sørensen [62] and Breit [65] exposed mortar (w/b 0.40–0.60) with embedded steel in chloride solutions under anodic polarisation, and, upon corrosion initiation, material from adjacent to the rebar was sampled and analysed. Pettersson [63] and Li et al. [67] exposed mortar (w/b 0.40–0.60) with embedded steel to chloride solutions, either permanent or cyclic, and sampled material close to the steel upon electrochemical detection of corrosion onset. Sandberg et al. [64] derived what they call “engineering values” for submerged concrete from experiments (w/b 0.30–0.75). Oh et al. [66] admixed various chloride levels to concrete (w/b 0.35–0.55) and monitored corrosion potentials; after 30 days they sampled material around the steel and expressed pore solution and recorded the corroded area. Polder [68] exposed concrete (w/b 0.40–0.55) specimens to drying and wetting with chloride solutions and detected corrosion initiation with electrochemical measurements.

Summarizing these studies, both for concrete and mortar, either in natural exposure, under anodic polarization or with mixed-in chloride, the influence of w/b on the susceptibility of steel to corrosion is relatively small. While there is general agreement that lowering w/b below 0.5 slightly improves the resistance against corrosion initiation (increase of C_{crit} up to approx. 30%), the results for the effect of w/b in the range above 0.5 are contradictory, with an overall negligible influence. Based on the currently available literature, it is concluded that the effect of w/b on the corrosion susceptibility of embedded

steel is much less pronounced (and less clear) than the effect of w/b on mechanical and transport properties of concrete.

3.3.2 Cement type

Similar to w/b, cement type is also known to be an important parameter in concrete. For the purpose of this review, cement types have been broadly categorized into: (i) Portland clinker (PC) blended with supplementary cementitious materials (SCMs) such as blast furnace slags (BFS), fly ash (FA) and silica fume (SF), and (ii) alkali-activated cements (AAC), with BFS and FA as the main precursors.

PC blended with SCMs: A number of studies [57, 62, 63, 66, 70-74] reported C_{crit} when using different SCMs as summarized in Fig. 5. It is apparent that the literature is ambiguous and even contradictory since for all investigated SCMs (FA, BFS, SF) both positive and negative effects have been reported. Given the ambiguous nature of reported data on the influence of cement type on C_{crit} , it is difficult to conclusively state whether the blending of SCMs with PC has a positive or a negative effect on corrosion initiation. The C_3A content of PC has a significant impact on the ability of the binder to interact with chlorides and form Friedel's salt. As a result, increasing the C_3A content in PC increases C_{crit} when the latter is expressed as total chloride content [63, 75] (Fig. 5). Fig. 5 shows that most studies generally agree that the overall effect of the cement type, including different SCMs and differences in C_3A content, is moderate (in most cases, the absolute effect is <1.5).

Alkali-activated cements: Corrosion initiation in AACs is a much less studied topic in comparison to OPC and blended cements. Studies investigating C_{crit} in alkali-activated FA [76] and FA/BFS blended systems [77] typically found C_{crit} to depend primarily on the ratio (SiO_2/Na_2O) of the activator used. In the case of alkali-activated BFS, several studies have reported reducing conditions at the SCI due to the presence of sulfides in the pore solution [78, 79], which is claimed to influence, at least for some time, the corrosion performance [80, 81]. Unfortunately, no studies were identified that allow to quantify the effect of AAC with the methodology used in this review, generally because no reference data (samples with OPC) were studied in the reviewed publications.

3.4 Macroscopic interfacial concrete voids at the SCI

3.4.1 Interfacial air voids

Interfacial air voids occur when entrapped or entrained air bubbles in fresh concrete rise up during compaction and become trapped beneath reinforcing steel bars. Most (if not all) studies have focused on the effects of entrapped rather than entrained air voids on corrosion initiation.

Various studies have noted that corrosion of embedded steel reinforcement tends to initiate at locations where air voids are present at the steel-concrete interface [3, 82-88]. However, in many publications it remained unclear whether corrosion initiated at the steel surface exposed in the actual air void or at the steel surface exposed in the cementitious matrix adjacent to it.

Several authors [83, 85, 88] found that interfacial air voids significantly reduced C_{crit} (Fig. 6). In these studies, the authors varied the amount of compaction voids (entrapped air) in 150 mm concrete cubes with a single rebar at 15 mm cover. The samples were sealed-cured for 28 days, then saturated with water and submerged in 4 M NaCl solution at 30°C. Upon initiation of corrosion, as indicated by a galvanic current rise, the samples were split and the steel surface observed. Visual examination of the SCI indicated that corrosion tends to initiate at locations adjacent to entrapped air voids. Furthermore, for a range of binder types, an increase in air void content from 0.2 to 2% led to a decrease in C_{crit} from 1.2 to 0.2% by mass of binder. The most pronounced change occurred below 1% air content. However, concrete with air contents below 1% are rare in engineering practice. Between 1 and 2% entrapped air content, which is the range in which most concretes in practice are expected to be, the impact of entrapped air void content on C_{crit} was found to be approx. a factor of 2. For entrapped air contents above approx. 1.5%, the influence on corrosion initiation was negligible. The results of these studies indicate that the effect of interfacial air voids is greater than that of the cement type, at least at relatively low air void contents. Similar findings were also reported in [87, 89].

However, there are a number of studies that have reported findings contrary to the above [10, 32, 90-92]. In [10, 32, 90, 91], samples were exposed to wet/dry cycles in chloride solution and corrosion was detected by monitoring the corrosion potential of the steel. The samples included lab-made reinforced concrete samples (with ribbed steel) and cores drilled from various 20–50 year old engineering structures. Upon corrosion initiation, the samples were split and visually examined. Relatively large interfacial air voids of up to several mm in diameter were frequently found at the SCI, but the location of corrosion initiation, as evidenced by presence of deposited corrosion products, rarely coincided with the location of voids. Corrosion was found to initiate at macroscopic (>0.5 mm) air voids in only 12% of the studied cases (Fig. 7) [32]. However, in these cases, C_{crit} was generally higher than in cases where corrosion initiated elsewhere (Fig. 7). An important observation made in these studies is that in many cases corrosion was found to initiate at locations with no particular visual irregularities, despite the presence of air voids at the SCI elsewhere within the specimen.

In conclusion, the impact of interfacial air voids on corrosion initiation in concrete cannot be generalized. The frequently held view that air voids *a priori* favor corrosion initiation is not always true, based on the reviewed data. The role of interfacial air voids is probably influenced by other conditions, particularly the moisture conditions in the concrete (see section 3.6).

3.4.2 Artificially created interfacial voids

In various studies, artificial defects, supposedly representing interfacial voids, were introduced by means of filter paper at the steel surface, e.g. [4, 6]. Sandberg [4] reported average C_{crit} values of 0.84% and 1.05% chloride by cement mass in the presence and absence of filter paper defects, respectively, from field exposure (splash zone) (standard deviation in both cases approx. 0.23% chloride by cement mass; based on 8 replicate samples). In submerged conditions and from laboratory tests, the impact of the filter paper was lower and not statistically significant. Nygaard [6] examined reinforced mortar samples containing artificial defects created by covering part of the rebar with a filter paper prior to casting. These samples were continuously immersed in chloride solution and it was found that the chloride concentration needed to initiate corrosion decreased with increasing area fraction of the filter paper.

It must be noted, however, that filter paper is likely to behave differently than natural voids at the SCI. This is because filter paper typically has pores that are coarser than those of the cement paste, but much finer than those of macroscopic interfacial voids. Thus, introducing filter paper to simulate macroscopic interfacial voids may keep the latter in a higher moisture condition compared to the case of natural interfacial voids. Therefore, care needs to be taken when applying conclusions drawn from filter paper studies to interfacial air voids.

3.4.3 Settlement and bleeding zones

Fresh concrete reduces its volume in the formwork due to its own weight (compaction of matter, removal of entrapped air) and early chemical shrinkage. This volume reduction is called settlement, and may give rise to (empty) cracks/zones, among others, at the reinforcement. If the concrete is unstable or the setting time is long, settlement might be accompanied by segregation and bleeding. In case of bleeding, bleed water may accumulate under larger obstacles, e.g. under the reinforcement and – if present – in the initially empty settlement zones. During hydration the bleed water will be (partly) absorbed into the surrounding paste due to chemical shrinkage causing self-desiccation. Settlement and bleeding zones may thus be empty (air-filled) or partly water-filled. Unless exposed to highly moist environments, it is unlikely that water will be present in large quantities in settlement and bleeding zones in the hardened concrete due to the size of these voids. An exception to this is the case when cracks in the concrete cover provide a link from the settlement and bleeding zones to the exposure environment. The size of settlement and bleeding zones increase with the height of concrete below the reinforcing steel. As a result, top-cast horizontal bars exhibit more defects at the underside perimeter than bottom-cast horizontal bars. This effect is well-known by structural engineers as the “top-bar effect”, which affects the bond properties. Settlement and bleeding zones were documented to occur particularly at the underside of horizontal rebars. This is important because depending on the orientation of the exposed concrete surface, chlorides may arrive to the steel from the underside or upper side [93].

For a more detailed review of the properties of settlement and bleeding zones, the reader is referred to [12].

A number of researchers found that settlement or bleeding zones may have an effect on corrosion [5, 9, 11, 44, 90, 94, 95], i.e. that corrosion occurred preferably on the underside of reinforcing steel bars, where settlement or bleeding zones exist. In most of these studies, the reported observations were made after considerable time had passed after corrosion initiation. The reported observations include visual inspection upon splitting the specimen such as determining the location of corrosion (e.g. on undersides of rebars) or the extent of the corroded steel area. However, these works do not permit quantifying the influence of settlement and bleeding zones on corrosion initiation of reinforced concrete, as the results may be influenced by the impact on corrosion propagation. Data on initiation is available from two laboratory studies [90, 95] that reported C_{crit} , distinguishing between initiation on the underside of rebars that were horizontally oriented during casting (presumably showing settlement or bleeding zones [96]) and initiation at vertically oriented rebars (presumably free of settlement or bleeding zones [96]), as summarized in Fig. 8. Moreover, the figure shows results [91, 97] from testing samples taken from two different engineering structures. While the literature is somewhat contradictory on the qualitative effect, overall, the influence of settlement or bleeding zones on C_{crit} is modest, both increasing or depressing C_{crit} by 40% at most (under the conditions of testing).

Although the data are limited, Fig. 8 indicates that the adverse impact of settlement and bleeding zones is slightly more pronounced on smooth rather than ribbed steel bars and in moisture states close to saturation (immersed) compared to lower moisture contents (wetting/drying).

3.4.4 Cracks, slip, and separation (mechanical damage at the SCI)

Cracking of the concrete may lead to formation of voids at the steel surface [12]. This happens both when cracks appear in early stages (e.g. plastic shrinkage cracks, temperature induced cracks) and due to mechanical loading of mature concrete. The morphology of cracks varies depending on the cause. Bending cracks are usually accompanied by slip and separation at the rebar. A main difference between other interfacial voids e.g. settlement zones and crack-related interfacial voids, is that cracks may extend through the entire concrete cover concrete and, thus, exposing the voids to the environment, while settlement zones are encapsulated in the concrete and, thus, typically have limited direct access to the exposure environment. This has significant consequences for exchange of moisture, oxygen, and ions with the exposure environment; the concentrations of these can dramatically and relatively quickly change in crack-related voids, while such changes are comparatively slow in settlement and bleed water zones. In structures under load, however, bending cracks could intercept top-cast tensile bars affected by the top-bar effect. In this case, cracks allow quick access of settlement zones to the exposure environment and pitting nucleation was reported to occur along the affected rebars and not only in zones close to the cracks [98].

The premature initiation of reinforcement corrosion due to presence of cracks in the cover of concrete structures is well-known and has been investigated for more than 50 years, e.g. reviewed in [99-102]. Many investigations have generally concluded that cracking in the concrete cover facilitates rapid chloride ingress and subsequently reduces the time to corrosion initiation (e.g. [103]). The presence of cracks, interfacial slip and separation caused by mechanical loading has been reported to cause relatively excessive cross-sectional reductions after 1-2 months of ponding with 10% chloride solution [104], and both laboratory studies [105-109] and in-situ observations [110-112] have noted an expedited corrosion initiation in cracked concrete compared to pristine concrete. In contrast to this, Zhang et al. observed that corrosion did not occur at all cracks in loaded beams exposed to salt spray [98].

While most studies focused on the role of cracks in accelerating moisture and ion transport through the concrete cover, very little is known about its actual effect on corrosion initiation. Studies indicated that defects at the SCI induced by mechanical cracking may also reduce C_{crit} [94, 113]. Available observations on the preferred sites for pitting corrosion initiation are inconsistent. Pacheco [114] reported that in areas with separation between steel and concrete, the prominent sites for corrosion pitting nucleation are where the separation decreases. However, Michel et al. [104] and Pease [101] observed corrosion only in areas with larger separation.

3.5 Other characteristics at the SCI

3.5.1 Spacers, tie wires, rebar intersections, and welding spots

While the review in [12] showed that the presence of spacers, tie wires and the location where rebars intersect can present local inhomogeneities, particularly in terms of concrete porosity at the SCI, we could not identify any literature addressing the influence of this on steel corrosion initiation. The few studies reviewed in [12] generally looked into concrete transport properties, e.g. chloride ingress, but no tests regarding the corrosion behavior of the reinforcing steel were reported. In addition to concrete microstructural features, locations where rebars are welded present metallurgical inhomogeneities, due to the effect of the welding heat. Jang and Iwasaki [115] reported that welded zones on rebars were more active than the base metal and acted preferably as anodes. Stefanoni et al. [116] also showed experimentally that welding spots on welded meshes are more susceptible to chloride-induced corrosion in alkaline environments than the remaining areas of the rebars.

3.6 Moisture content and temperature

Only very few studies reported the influence of the concrete moisture state and the temperature on corrosion initiation. Both Pettersson [117] and Song et al. [118] reported C_{crit} and observed a pessimum in the RH range 80–95%. Pettersson [117] conditioned concrete samples at various levels of RH and in capillary saturated state and exposed them briefly to saturated sodium chloride solution at intervals of 1–2 months [119]. Corrosion was considered to have initiated when the corrosion current density increased sharply to more than $10 \mu\text{A}/\text{cm}^2$, which we consider a criterion that may have required a very high amount of chlorides to be achieved particularly in the lower RH range. Song et al. [118] exposed concrete samples at different RH and in immersed state during a year. The criterion for corrosion initiation was a corrosion current density higher than $0.5 \mu\text{A}/\text{cm}^2$. The method for introducing chlorides is unclear. Boschmann & Angst exposed concrete samples to chloride solutions, where one series was continuously immersed and another series was exposed to daily wetting/drying cycles. The RH in the concrete was measured at the depth of the rebar and found at $95 \pm 2.5\%$ RH during the wetting/drying cycles. Corrosion initiation was detected by means of a drop in steel potential.

Evaluating these data according to the methodology presented in section 2.2 yields Fig. 9. While RH < 65% may be of little practical relevance for chloride-induced corrosion, the figure indicates that the moisture condition has a pronounced effect in the range from RH 80% to capillary saturation (grey and white symbols in Fig. 9). This effect becomes smaller between RH 95% and capillary saturation (black symbols in Fig. 9).

Repeated wetting and drying appears to create more critical conditions with lower C_{crit} than at constant moisture state [120-122]. This was apparent from the unpublished data by Boschmann and Angst where the mean C_{crit} was approx. 50% higher in continuously immersed condition compared to wetting/drying. In agreement with this, using custom-made corrosion cells developed by Arup and Sørensen [123] and exposing them to chloride by submersion, wetting and drying or in the field, Sandberg and Sørensen [122] observed that submersion generally resulted in higher C_{crit} . They concluded that the results supported the earlier suggested [124, 125] hypothesis of moisture and temperature variations at the steel surface promoting corrosion initiation. Very high C_{crit} are expected in concrete submerged for long periods [124].

4 Discussion

4.1 Dominating influencing factors

Corrosion initiation in concrete is a stochastic process. Even when steel surface is in pristine condition and most confounding factors at the SCI that affect corrosion initiation are eliminated, corrosion may initiate at random sites on the steel surface when sufficient amount of chloride is present in the electrolyte. Where and how corrosion might initiate in these idealized conditions would depend

mainly on the local imperfections of the passive film. Beyond this well-accepted stochastic nature of localized corrosion, the most significant factors affecting corrosion initiation of steel in concrete are related to the characteristics of the SCI, which are covered in this paper.

Fig. 10 summarizes the effects of the different local characteristics of the SCI according to the literature evaluated in this work. Note that due to the uncertainties in the data, as discussed in section 3, the focus of Fig. 10 lies on assessing the relative influences rather than a quantitative assessment of these effects on corrosion initiation. On this basis, we make the following striking observations:

- The degree to which steel is polished, the steel metallurgy, and the moisture content at the SCI are by far the most dominant influencing characteristics for chloride-induced corrosion initiation in concrete (among the characteristics for which we have sufficient data for a quantitative assessment). The effects of these characteristics are almost one order of magnitude higher than the effect of almost all the other characteristics of the SCI that were quantified. Further indication for the dominating effect of moisture content is apparent from results reviewed in the various cases of macroscopic interfacial concrete voids (section 4.2.2).
- We consider it of particular interest that factors such as cement type and w/b ratio, which are well-known to play major roles in concrete technology and concrete properties (e.g. strength, bulk transport properties, etc.), have comparatively small effects (Fig. 10).
- The different possible influencing factors have received highly unbalanced research attention. Cement type, w/b ratio, and mill scale or rust layer influences have been studied most extensively. However, the effect on corrosion initiation of characteristics such as moisture, steel metallurgy, air voids, rebar geometry, spacers, etc. have been addressed in very few or no studies (note that here, we are not considering studies focusing on chloride transport through the cover).
- The best agreement on the effect of SCI characteristics described in the literature was found for mill scale and rust layers, passive film ageing, and moisture. Mixed conclusions were reported for the effects of w/b ratio, cement type as well as certain interfacial voids, but the studies generally agreed that the effect is relatively moderate for these characteristics.

4.2 Hypotheses of working mechanisms

4.2.1 Steel metallurgy and surface condition

Section 3.1 presented the effect of various characteristics related to the steel side of the SCI on corrosion initiation. A number of hypotheses have been proposed, that may be classified as follows:

- *Crevice corrosion mechanism*: Ghods et al. [41] suggested that cracks in the mill scale provide access for the concrete pore solution to reach the steel surface. Numerical simulations supported the hypothesis that the chemical composition of the pore solution within the cracks differs from that of the bulk solution through a process similar to the suggested mechanism of typical crevice corrosion [126]. This may lead to conditions favoring depassivation of the steel at the crack tip. It may be noted that crevices and cracks in the oxide scales may likely be of over-capillary dimensions and thus the hypotheses presented in section 4.2.2 may also apply to these. Similarly to the reasoning related to cracked oxides, it has been suggested that a rough surface, exhibiting depressions in the metal surface, provides a geometrical configuration in which diffusional transport limitations can more easily produce a sufficiently aggressive anolyte to sustain further local pit growth [130].
- *Cathode effect sustaining early pit growth*: Several authors suggested that the presence of oxides may provide an additional oxidation/reduction couple (Fe(II)/Fe(III)) that can – at least for a limited time – enhance the cathodic current capacity by a factor of 2–5 [127, 128]. The higher cathodic current was also explained by different effective steel surface areas between clean and mill-scale surfaces. More effective cathodes in the vicinity of pit nucleation sites are likely to sustain early stage pit stabilization and pit growth, hence initiation of stable corrosion.
- *Inhibited passivation*: Li and Sagüés [34] suggested that in the absence of pre-existing oxides, a fresh, dense passive film can grow better on the steel surface than in the presence of oxides. Similarly, from results using potentiodynamic polarization (PDP) and electrochemical impedance

spectroscopy (EIS) it was in [7,8] concluded that the mill scale can inhibit the formation of a passive film on steel exposed to alkaline solutions or mortar.

- *Surface heterogeneity and size effect:* A general explanation can be found in the inhomogeneity of the surface [36], e.g. in terms of chemical composition of the different surface components. Local electrochemical studies have revealed the marked diversity in local electrochemical behavior of as-received surfaces as well as their mutual polarization when immersed in an electrolyte [31, 129]. Surface inhomogeneities have also been related to a size effect, meaning that larger exposed electrodes exhibit a higher probability of corrosion through a size effect [10, 130-132]. Since surface roughness is related to the effective area, the size effect has been used to explain the influence of surface roughness on corrosion initiation [130].

While the data reviewed in this work appear to suggest a strong influence of steel metallurgy (Fig. 10), no conclusive hypotheses are currently available to explain this.

4.2.2 Macroscopic interfacial concrete voids (MICV) and the importance of moisture state

Section 3.4 presented the effect of various *macroscopic interfacial concrete voids (MICV)* on corrosion initiation. It became repeatedly apparent that their role in corrosion initiation is significantly affected by moisture state, which was furthermore supported by the marked overall effect of moisture (Fig. 10). Here, we attempt to provide hypotheses that are generally valid for the different macroscopic concrete voids. These include entrapped and entrained air voids, settlement and bleeding zones, and perhaps cracks, slip, and separation as well as other voids, e.g. at intersecting rebars or at spacer/rebar or stirrup/rebar interfaces, for which no literature documenting their influence on corrosion initiation is available. An important common feature of these MICVs is that their dimensions are above the capillary range. However, they differ in geometry and initial moisture condition as well as possible access to the exterior (Table 1), which will affect the availability of reactants and the efficiency of the electrical circuit in the corrosion cell [133].

Additionally, an important feature may be whether or not the MICV exhibit a thin layer of cement hydration products coating the steel surface adjacent to the void. Such a layer is expected in all cases where the steel is initially embedded in concrete during casting, that is, before the formation of the MICV such as in bleeding and settlement zones, concrete cracks, slip, separation, and entrapped/entrained air voids. However, there are situations where such a cementitious layer is absent, e.g. an air bubble that is attached to the reinforcement from the very beginning and remained stationary during concrete casting and hardening. This may include entrapped air voids as well as gaps and crevices between intersecting rebars or between rebars and spacers [12].

Our hypothesis on the combined impact of moisture state and macroscopic interfacial concrete voids on corrosion initiation is illustrated in Fig. 11. In completely saturated state, steel is directly exposed to liquid water (containing chlorides [134]) within the void (Fig. 11a). In this case, the steel will be more prone to corrosion initiation than steel covered with cement paste. This is because of the absence of calcium hydroxide at the steel surface that locally buffers the pH during stages of early pitting corrosion. During this stage the competitive migration of chloride and hydroxyl ions towards the anodic site is decisive for establishing stable pit growth [135, 136]. Thus, a locally lowered pH buffer capacity significantly impairs the corrosion inhibiting mechanism associated with the presence of a calcium hydroxide rich layer at the SCI, as postulated in [1, 2].

In partially saturated MICVs, i.e. voids containing both liquid electrolyte and air (Fig. 11b), there is, in addition to the susceptibility to corrosion in the wet areas as mentioned above, an increased availability of oxygen from the air filled part that potentially aggravates the corrosion initiation process. The mechanism for this may be related to, on the one hand, generally raising the steel potential prior to pitting nucleation [55, 56] and, on the other hand, enhancing the supply of oxygen to sustain the cathodic activity that is needed to overcome the stage of early pit growth and to achieve stable pitting corrosion [136-138]. Thus, steel in partially saturated MICVs may be considered the worst case.

Fig. 11c shows an air-filled (non-saturated, “dry”) MICV. Here, supply of oxygen through diffusional transport across the film of adsorbed water at the steel surface is high. On the contrary, ionic transport within the film of adsorbed water is expected to be severely limited [139]. Thus, it is unlikely to achieve pitting corrosion in the parts of the steel covered with adsorbed water, as this requires the

formation of significant galvanic elements, which is impossible due to the strong ohmic control within the adsorbed water film. Electrical resistance measurements reported in [133] indicated that interfacial air voids at non-saturated conditions are indeed electrically (and electrochemically) inactive. However, for the reasons given in the previous paragraph, air-filled MICVs may promote corrosion initiation in cement-paste covered steel areas close to the MICVs, as indicated by the red cross in Fig. 11c.

Fig. 11d shows the case where a cement paste layer is present at the steel surface adjacent to the MICV. Here, the susceptibility to corrosion is expected to be lower than in the case of Fig. 11b, mainly due to the pH buffering and barrier effect of the cement paste layer. Depending on the thickness of this cementitious layer as well as the reservoir of calcium hydroxide available, corrosion may still preferentially start here, particularly in unsaturated condition due to the combination of increased supply of oxygen and high ionic conductivity enabling the formation of galvanic elements to stabilize early pit growth.

Summarizing the different cases illustrated in Fig. 11, corrosion may be expected to initiate in the MICV or next to it, depending on the saturation state. The location of corrosion initiation with respect to the location of MICVs was not always evidenced with full clarity from studies reporting visual inspections of rust at the SCI. This may also have been masked to some extent by the fact that increased oxygen concentrations close to and in voids may locally favour precipitation of corrosion products that diffuse towards voids from other, more deoxygenated locations (Fig. 11c).

The strong dependency of corrosion initiation on the moisture state of interfacial air voids could explain the seemingly inconsistent findings from different studies (cf. section 3.4.1). Entrapped air voids initially contain no moisture and if they remain dry, corrosion would not occur at these locations due to the lack of electrolyte to support initiation. In the work of Angst et al. [10, 32, 90, 91], MICVs were clearly not saturated [12] owing to the cyclic wet/dry exposure conditions applied. Thus, it is not surprising that they were rarely found to be preferential sites for corrosion initiation. In other studies [83, 85, 87, 88], however, concrete samples were closer to saturation due to being continuously submerged, and potential partial saturation of the MICVs could explain the observation of corrosion initiation close to or at air voids.

Concerning the local role of oxygen as above hypothesized for Figs. 11c-d, it may be mentioned that often a locally triggered anode is galvanically coupled with the rest of the reinforcing steel bar. Thus, unless the concrete electrical resistivity is extremely high, the local presence of oxygen may lead to more global rather than pronounced local differences in steel potential on the rebar surface. Hence, the cathodic current can be supplied from the comparatively large rebar surface and not critically depend on a local supply of oxygen. This may thus be another explanation for the experimental observations where corrosion was not found to preferentially occur at MICVs.

In engineering practice, fully saturated MICVs are considered unlikely to be present in uncracked concrete, as even in submerged conditions it can take very long time to fill pores in the over-capillary range [12]. The proposed mechanism leading to saturation is that dissolution of the air (held in a void) into the surrounding pore water leads to a gradual filling of the void with liquid water [140]. The process is estimated to be very slow [140], and thus MICVs may not be completely filled during the lifetime of a structure. However, depending on the moisture exposure, MICVs may be partially water-filled. Finally, it should be mentioned that in cracked concrete where the cracks provide a direct path between MICV and the exposure environment, saturation of MICV may be reached relatively fast.

The microstructure and the chemical composition of the binder as well as the geometric configuration (e.g. cover, SCI characteristics) and temperature affect the actual moisture content and the pore liquid composition at the steel surface. However, numerical simulations performed by Ryu et al [141] indicated that short-term drying and wetting have little effect on the internal moisture content in uncracked concrete. Variations in the range of 15–35 °C and 30–90% RH were found to affect moisture content in concrete with $w/b=0.3$ and 0.6 to depths of 10 and 30 mm, respectively. For continuous rainfall during 24 h, the predicted moisture change in $w/b=0.6$ concrete was evident to a depth of 30 mm. The moisture content at depths ≥ 50 mm from the surface was scarcely affected [141]. Using recorded meteorological data for simulating internal temperature and moisture conditions in a concrete, Flint et al. [142] found, for concrete ($w/b=0.5$) in northern climate (Lofoten, Norway), that the temperature change was “instantaneous” throughout the investigated depth of 50 mm, whereas the degree of saturation was significantly affected only in the outermost 20–30 mm. Relling [143] measured convection zones (zones with varying moisture state) in Norwegian marine bridges of approximately

10 mm. Similar results were also reported by Moro, based on monitoring of exposed concrete with sensors [144].

In summary, it can be expected that engineering structures in chloride-bearing environments are typically exposed to moistures in the range from 80% RH to direct contact with liquid water (splash, tidal and submerged zones). Depending on exposure and cover depth, the moisture state in the here discussed macroscopic (over-capillary) interfacial concrete voids discussed here can vary considerably, which in turn plays a major role on the effect of these voids on corrosion initiation in concrete.

4.2.3 Cement type and w/b ratio

Perhaps one of the more striking findings of this review is that the cement type and the w/b ratio, both being important parameters in concrete technology and durability design, have relatively small effects on corrosion initiation. Without doubt, these parameters will continue to be important in ensuring the durability of concrete structures. However, their primary role appears to be in controlling mass transport through the concrete cover. Their role in affecting the susceptibility of steel embedded in cementitious matrix to corrosion seems secondary.

Nevertheless, there are a number of hypotheses about how cement type and w/b may influence initiation of chloride-induced corrosion. Due to the observed moderate effect (Fig. 10), we decided to present here only a summary of these hypotheses and refer to the literature for more details. The cement type affects the pore solution composition, particularly pH and its buffer capacity, and the microstructure [145-148]. Moreover, SCMs can affect chloride binding capacity [146, 149]. Finally, cement type and w/b ratio have an influence on the stability and packing of particles and thus on the micro- and macrostructure at the SCI such as the morphology and width of settlement and bleeding zones [96].

5 Conclusions

This review of the literature on chloride-induced corrosion of steel in concrete has revealed that parameters well-known to be important in concrete technology – such as w/b ratio and cement type (at least PC-SCM blends) – have relatively small influences on the actual corrosion susceptibility of steel embedded in a cementitious matrix. Future scientific and engineering developments towards predicting corrosion initiation in concrete would benefit from considering other parameters, namely *steel properties* – including metallurgy, presence of mill scale or rust layers, and surface roughness – as well as the content and distribution of *moisture* in microscopic and macroscopic voids at the SCI. Although limited data are available in the literature, our review suggests that these characteristics of the SCI have a strong influence on the susceptibility to corrosion. Thus, we expect that considering these factors will improve the reliability of future predictive models for corrosion initiation in concrete.

Our findings imply that endeavors to develop experimental methods aiming at determining C_{crit} as a function of concrete mix proportions – utilizing standardized steel surfaces and concrete consolidation methods – may have little relevance in holistically assessing the corrosion performance of reinforced concrete structures.

Although w/b ratio and cement type seem to have a surprisingly small effect on the corrosion susceptibility, these parameters will continue to play an important role in service life modeling, durability design, and condition assessment, since they have a strong impact on bulk concrete transport properties and on concrete cracking, and thus on the transport of chloride, moisture, oxygen, and carbon dioxide through the concrete cover, as well as on the propagation of corrosion.

We recommend that further studies focus on those characteristics that have so far received the least research attention or where the literature is inconclusive, but indicate potentially dominant effects. These include the moisture conditions at the SCI, interfacial air voids, reinforcing steel properties including metallurgy as well as macroscopic and microscopic geometry of the rebar and its surface, spacers, rebar intersection, welding points, and mechanical damage (cracks, slip, separation). On the contrary, parameters such as w/b ratio, cement type (with the exception of alkali-activated cements and other novel cements), and steel passivity may not need the highest priority in future studies related to

the corrosion susceptibility of steel embedded in a cementitious matrix because literature have consistently indicated merely moderate effects.

A final comment should be made on the quality of the available literature data. In the course of reviewing the extensive number of scientific publications, we found that important experimental details (such as the method of saturating concrete or introducing chlorides, or the number of replicate samples used) were often not reported, which significantly impeded the interpretation of the data. We consider it unsatisfactory that, after more than half a century of research on the issue of steel corrosion in concrete, many questions remain open, partly due to a simple lack of adequate reporting.

Acknowledgements

We acknowledge the input provided by all TC members attending the discussions of the TC meetings on 22 October 2017 (Philadelphia, U.S.), 7 April 2017 (Paris, France), 14 September 2017 (Zurich, Switzerland), 14 October 2017 (Anaheim, U.S.), 27 March 2018 (Sheffield, UK), 28 August 2018 (Delft, the Netherlands), 21 February 2019 (Zurich, Switzerland), and 19 March 2019 (Rovinj, Croatia). Additionally, we acknowledge the so far unpublished results that were provided by Raoul François from Université de Toulouse, France, (Fig. 5), and by Carolina Boschmann from ETH Zurich, Switzerland, (Figs. 8 and 9).

Compliance with ethical standards

Conflict of interest

The authors declare that they have no conflict of interest.

Figures and Tables

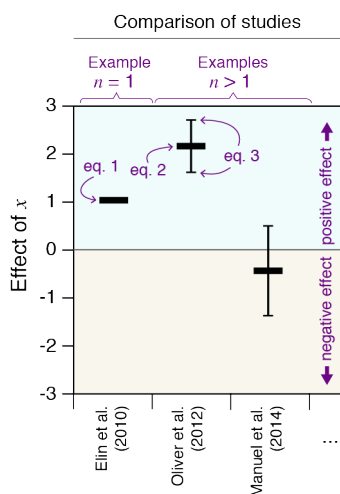


Fig. 1. Methodology used to quantify and compare the effect of an individual SCI characteristic x on corrosion initiation in concrete according to different literature studies; n = number of replicate samples in a study. See section 2.2 for more explanations.

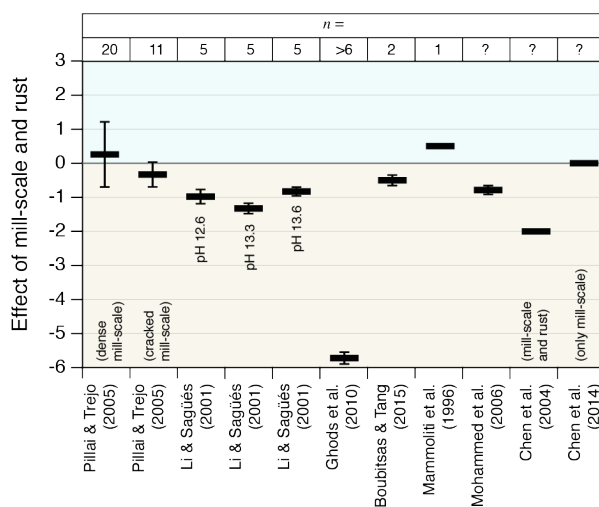


Fig. 2. Effect of mill scales and pre-existing rust layers on corrosion initiation [33-36, 38, 40, 44]. The presence of these scales generally favored corrosion initiation, as indicated by most data being located in the yellow area. The reference used to quantify the effect was the case without scales.

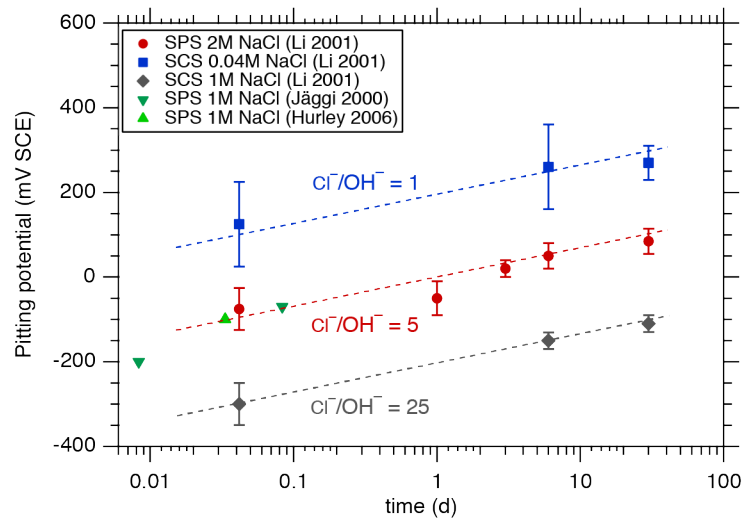


Fig. 3. Effect of the pre-passivation time on the pitting potential E_{pit} (steel pre-passivated at the open circuit potential in alkaline synthetic pore solutions (SPS) and in saturated calcium hydroxide solution (SCS)). Experiments of Li (2001) pre-passivation in chloride-free solution, then measurement of E_{pit} in corresponding chloride-containing solution [46]; experiments of Jäggi (2000) both pre-passivation and E_{pit} measurement in chloride-containing solution with Cl^-/OH^- ratio = 4 [45]; experiments of Hurley (2006) both pre-passivation and E_{pit} measurement in chloride containing solution for 1 h, $Cl^-/OH^- = 0.286$ [50].

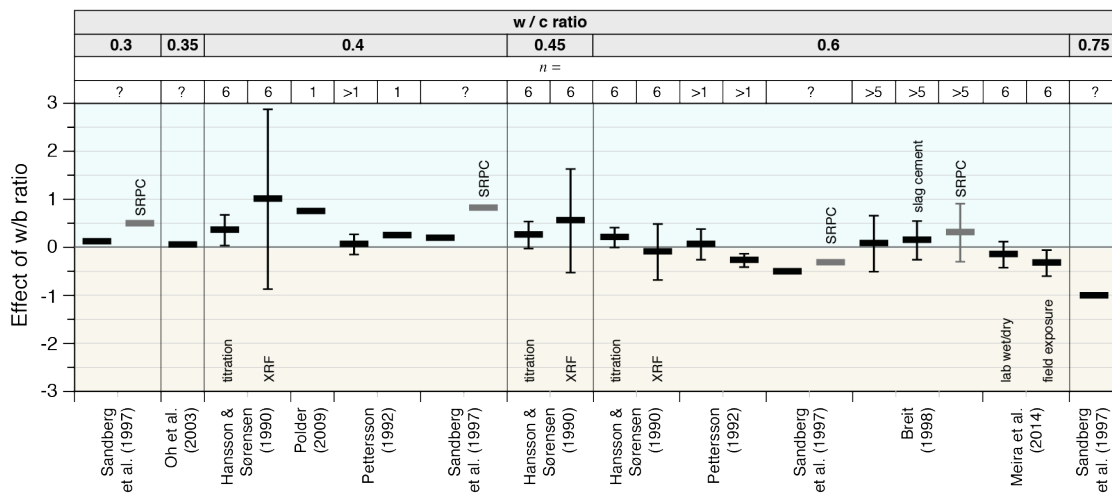


Fig. 4. Effect of w/b ratio on corrosion initiation based on Refs. [62-69]. The reference case for this evaluation of the literature data was $w/b = 0.5$. For $w/b < 0.5$ there is a slight tendency of improved resistance against corrosion initiation (blue area), while for $w/b > 0.5$ the results are mixed with no clear effect.

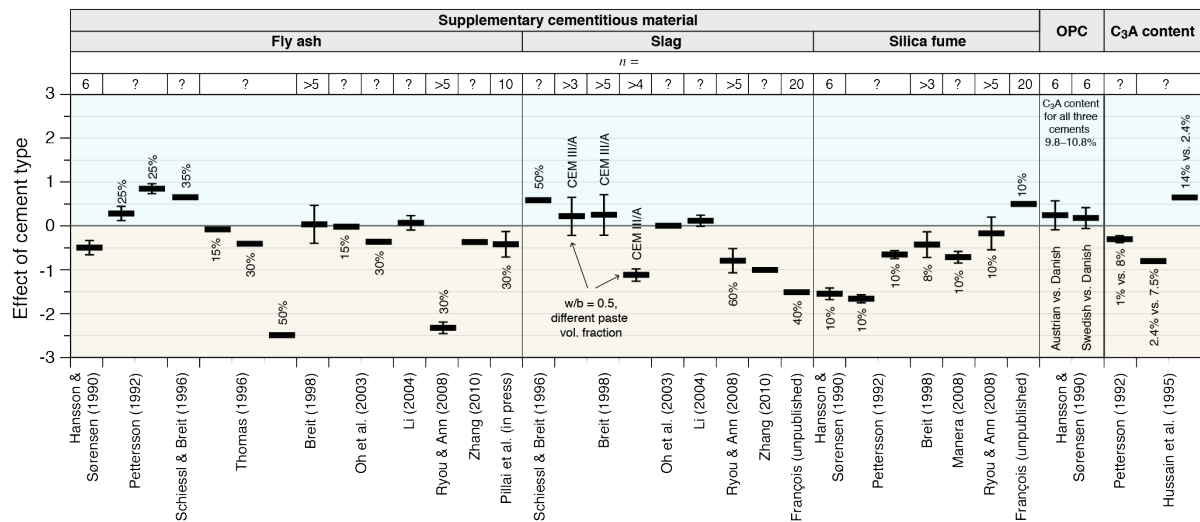


Fig. 5. Effect of cement type on corrosion initiation based on total C_{crit} reported in [57, 62, 63, 66, 70-75]. The reference case was always PC; for the effect of C_3A content, the reference C_3A content was ~8%. The percentages given indicate the amount of SCM or the amount of C_3A in the binder, respectively. The results are ambiguous, but in general the absolute effect of cement type appears to be <1.5.

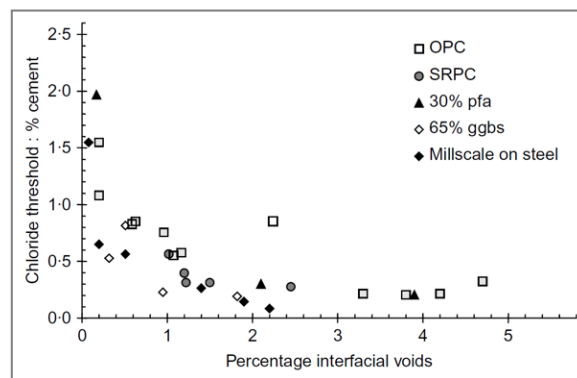


Fig. 6. Relationship between C_{crit} and entrapped air void content at the SCI for a range of binder types and steel surface conditions [83, 88]. Specimens were “saturated with water” and submerged in chloride solution during test.

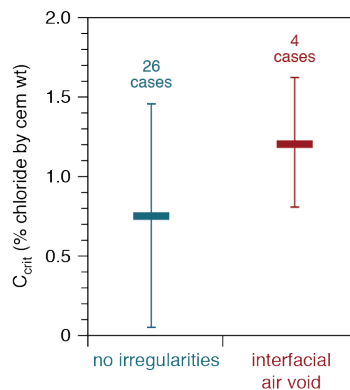


Fig. 7. C_{crit} observed in 30 samples taken from various engineering structures in Switzerland [32]. In the majority of samples (26 cases) visual inspection of the SCI upon splitting the samples after corrosion initiation did not reveal any visually apparent irregularities at the SCI. Only in 4 cases did corrosion coincide with interfacial air voids. Symbol = average, whisker = standard deviation.

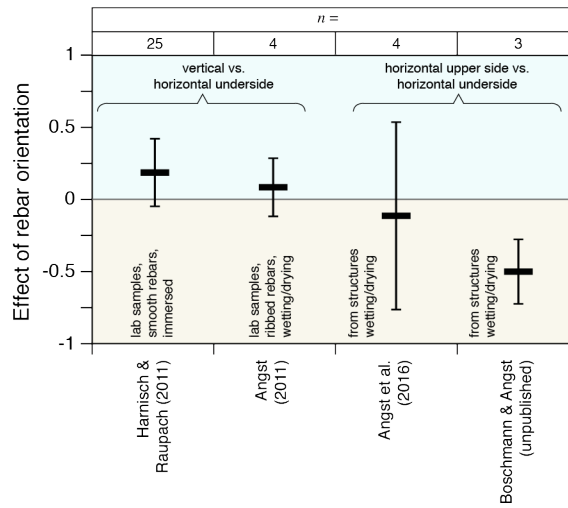


Fig. 8. Effect of rebar orientation on corrosion initiation (C_{crit}) [90, 91, 95, 97]. Some data indicate that vertical rebars were less susceptible to corrosion initiation than horizontal bars, and others suggest the upper side of horizontal rebars exhibit a better resistance against corrosion initiation than the underside of horizontal rebars (reference case). The literature results are somewhat contradictory on the impact of settlement and bleeding zones.

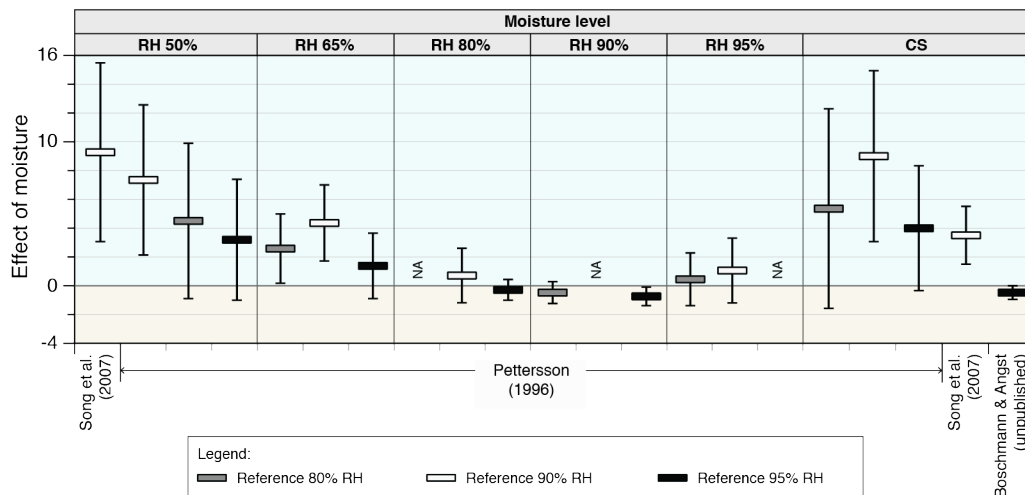


Fig. 9. Effect of concrete moisture state on corrosion initiation (C_{crit}), based on Pettersson [117], Song et al. [118], and unpublished data by Boschmann & Angst, all reporting C_{crit} as a function of the moisture state. CS = capillary saturated. In both the work of Pettersson [117] and Song et al. [118] the number of replicate samples (n) was not reported, Boschmann & Angst studied 3 replicate samples.

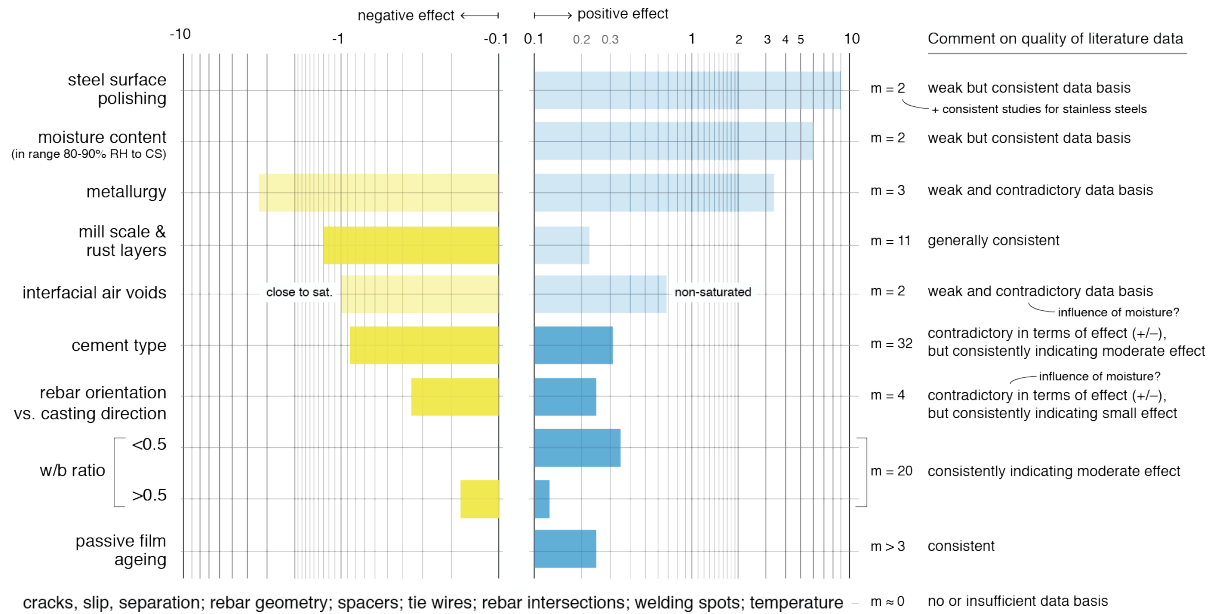


Fig. 10. Compilation of the influencing factors evaluated in this work and their effects quantified according to the methodology described in section 2.2. The blue and yellow bars show the average of positive effects and the average of negative effects, respectively. Light bar colors indicate a low number of reviewed studies (m). Note the logarithmic axis.

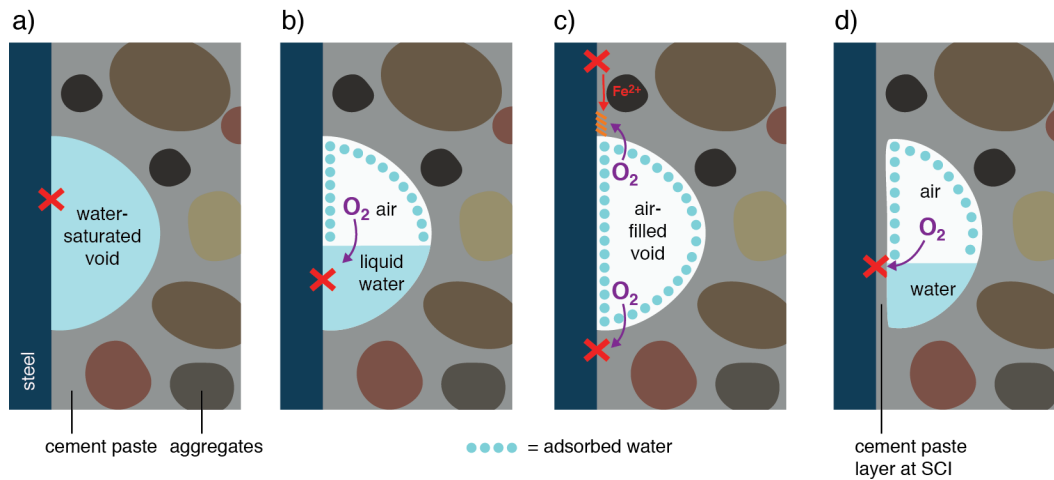


Fig. 11. Schematic sketches illustrating possible roles of macroscopic interfacial concrete voids (MICV) in corrosion initiation. Here, we use the example of an air void, but the same processes may occur in other MICVs. The red crosses indicate the location of corrosion initiation. a) water-saturated MICV with limited pH buffered liquid due to the relatively long diffusion path for hydroxyl ions released by CH; b) partially saturated MICV promoting corrosion in the liquid part, amongst other reasons due to the oxygen supply from the air-filled part; c) non-saturated (air-filled) void acting as oxygen reservoir for nearby zones, either promoting corrosion initiation (red cross shown at the bottom) or precipitation of corrosion products diffusing towards the void from an initiation site elsewhere (red cross shown at the top). Finally, d) illustrates a case where the steel adjacent to the void is coated with a thin cement paste layer, where moisture, oxygen, and a pH buffer ($(CaOH)_2$) may be abundant at the steel surface to influence corrosion initiation.

Table 1. Geometry, moisture state, and access from exposure environment in different macroscopic interfacial concrete voids (MICVs)

	Entrained air voids	Entrapped air voids	Settlement zones	Bleeding zones	Cracks, slip and separation
Length scale, perpendicular to steel	μm-mm	μm-cm	μm-mm	μm-mm	μm-mm
Length scale, along steel	μm-mm	μm-cm	cm-m	cm-m	μm-cm
Initially water saturated	No	No	No	Yes	Depends on exposure
Access to exterior (exposure environment)	No, unless linked to exterior through cracks in the cover	No, unless linked to exterior through cracks in the cover	No, unless linked to exterior through cracks in the cover	No, unless linked to exterior through cracks in the cover	Yes

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