A New Updated Model of CO₂/H₂S Corrosion in Multiphase Flow

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ABSTRACT
Following the introduction of an integrated mechanistic model of CO₂ corrosion in multiphase flow a few years ago, new advances have been made in understanding internal corrosion of mild steel pipelines. The original model was mechanistic and covered the key underlying processes such as: kinetics of electrochemical reactions at the steel surface, dynamics of coupled transient transport of multiple species between the bulk solution and the steel surface, through the turbulent boundary layer and through a porous surface film, kinetics of chemical reactions including precipitation of iron carbonate, as well as growth and protectiveness offered by iron carbonate scales. A new updated model based on the latest experimental results covers previously unknown territory: corrosion at very low temperatures (as low as 1°C) and corrosion in very high salinity brines (up to 25%wt. NaCl). However the biggest breakthrough was the development of a mechanistic H₂S corrosion model which includes the kinetics of iron sulfide growth by solid state reaction. The overall model has been extensively calibrated and verified with a reliable experimental database.

KEYWORDS: model, prediction, CO₂, carbon dioxide, organic acids, H₂S, hydrogen sulfide
INTRODUCTION

Many different models for internal corrosion of mild steel multiphase pipelines carrying a mixture of oil, gas and water have appeared since the first worthwhile attempts of deWaard and Milliams\(^1\) in 1973. Recently Nyborg\(^2\) has reviewed the performance of a group of models and concluded that most of them successfully predicted the “worst case” pure CO\(_2\) corrosion rate, however did not perform well when more complex effects were included (e.g. protective scales, H\(_2\)S, etc). The main reason for this problem lies in the fact that most of these models were empirical derivatives of the original mechanistic model of deWaard and Milliams\(^1\) or some other more empirical versions that appeared later. When dealing with effects that complicate the pure bare-steel CO\(_2\) corrosion such as precipitating of protective layers, effects of H\(_2\)S, organic acids, etc., these models typically resort to an arbitrary series of multiplicative empirical correction factors which have little or no theoretical background and as such are unable to perform well when extrapolating outside the dataset used to “calibrate” them. Nyborg’s\(^2\) review did not cover the integrated CO\(_2\) corrosion – multiphase flow modeling package MULTICORP V3.1 from Ohio University (referred to as “original model” in the text below). The outline of this package has been presented in 2004 and 2005 NACE conference papers by Nesic et al.\(^3,4\)

This original model was very firmly rooted in theory and took into account the effect of most important variables and processes occurring in CO\(_2\) corrosion such as:

- Kinetics of electrochemical reactions at the steel surface.
- Transient transport of species between the bulk solution and the steel surface, through the turbulent boundary layer and through porous surface scales.
- Kinetics of chemical reactions including precipitation.
- Kinetics of scales growth and the effect on corrosion.
- Dynamics of multiphase flow.

The equations behind the model were faithful descriptors of the abovementioned physico-chemical processes underlying corrosion. This is in contrast with most of the other models review by Nyborg\(^2\), which are fully empirical or semi-empirical. By using this model it was possible to reliably predict the effects of key variables that affect internal pipeline corrosion such as:

- Effect of temperature (20-100\(^\circ\)C);
- Effect of CO\(_2\) partial pressure (0 – 2 MPa);
- Effect of organic acids (0 – 1000 ppm);
- Effect of H\(_2\)S traces (0 – 1 Pa);
- Effect of pH and brine chemistry (pH 3 – pH 7);
- Effect of steel type;
- Effect of multiphase flow (single-phase water flow, oil-water, gas-water and oil-gas-water);
- Magnitude and morphology of localized attack;

Besides providing immediate answers - e.g. the corrosion rate, the model allowed the users to get a deeper insight into the root causes behind the problem, thereby raising the user's confidence in the provided answer. The other (semi)empirical models based on arbitrary mathematical equations lack this capability, completely or in part. Due to the strong theoretical background of the original model, the user could extrapolate the predictions outside the calibration domain with much more confidence than can ever be achieved with the (semi)empirical models whose extrapolation capabilities are questionable. The model was built so that further extensions of the model to include new phenomena could be done relatively easily, in a logical fashion, and without changing most of the existing coding. This is in contrast with the extensions of (semi)empirical models which are cumbersome and often prohibitively difficult.
A new version of the original model: V4.0 (referred to as “new updated model” in the text below) has recently been released including many new features such as:

- Corrected model of CO₂ corrosion at low temperatures;
- Corrected model of CO₂ corrosion in high salinity brines;
- A full blown model of sour corrosion (covering pH₂S=0.001 mbar - 10 bar).

The new updated model has been extensively calibrated and verified with a reliable experimental database (800+ laboratory conditions and 100+ field cases). Most of the laboratory data came from large-scale corrosion and multiphase flow experiments. A reliable field corrosion database, provided by the major oil and gas companies, has been used to verify the performance of the model. In the text below, the foundations of the original corrosion model will be presented first, followed by the discussion of some of the new physics built into the new version of the model.

**THE NEW UPDATED VERSION OF THE CORROSION MODEL**

**Outline of the Original Model of CO₂ Corrosion**

The various physical, mathematical and numerical aspects of the model were presented in past publications⁸-¹⁰, however a brief outline is given below to facilitate the understanding of the new developments.

The corrosion model is built around coupled electrochemical and mass transfer equations that describe the electrochemical reactions at the metal surface and the movement of the species involved in the corrosion process between the metal surface and the bulk fluid. A one-dimensional computational domain is used, stretching from the steel surface through the pores of a surface scale and the mass transfer boundary layer, ending in the turbulent bulk of the solution. The concentration of each species is governed by a species conservation (mass balance) equation. A universal form of the equation which describes transport for species \( j \) in the presence of chemical reactions, which is valid both for the liquid boundary layer and the porous scale, is:

\[
\frac{\partial (\varepsilon c_j)}{\partial t} = \varepsilon \frac{\partial}{\partial x} \left( \varepsilon^{1.5} D_j^{\text{eff}} \frac{\partial c_j}{\partial x} \right) + \varepsilon R_j \tag{1}
\]

where \( c_j \) is the concentration of species \( j \) in \( \text{kmol m}^{-3} \), \( \varepsilon \) is the porosity of the scale, \( D_j^{\text{eff}} \) is the effective diffusion coefficient of species \( j \) (which includes both the molecular and the turbulent component) in \( \text{m}^2 \text{s}^{-1} \), \( R_j \) is the source or sink of species \( j \) due to all the chemical reactions in which the particular species is involved in \( \text{kmol m}^{-3} \text{s}^{-1} \), \( t \) is time and \( x \) is the spatial coordinate in \( \text{m} \). It should be noted that in the transport equation above electromigration has been neglected as its contribution to the overall flux of species is small when it comes to spontaneous corrosion. Turbulent convection has been replaced by turbulent diffusion as the former is difficult to determine explicitly in turbulent flow. A well-established statistical technique is used here: instantaneous velocity is divided into the steady and the turbulent - fluctuating components. Close to a solid surface, the steady velocity component is parallel to the surface and does not contribute to the transport of species in the direction normal to the metal surface. The turbulent convection term can be approximated by a “turbulent diffusivity” term¹¹, \(-D_t \partial c_j / \partial x\) and lumped with the molecular viscosity term to give \( D_j^{\text{eff}} \). The turbulent diffusion coefficient \( D_t \) is a function of the distance from the metal or solid scale surface and is given by¹¹:
\[ D_i = \begin{cases} 
 0 & \text{for } x < \delta_f \\
 0.18 \left( \frac{x - \delta_f}{\delta - \delta_f} \right)^3 \frac{\mu}{\rho} & \text{for } x > \delta_f 
\end{cases} \]  
(2)

where \( \delta_f \) is the thickness of the porous scale in m. The liquid boundary layer thickness \( \delta \) in m is typically a function of the Reynolds number. For pipe flow it reads:\(^{11}\)

\[ \delta - \delta_f = 25 Re^{-7/8} d \]  
(3)

where \( d \) is the hydraulic diameter in m, \( Re = \rho U d / \mu \) is the Reynolds number, \( U \) is bulk velocity in m s\(^{-1}\), \( \rho \) is the density in kg m\(^{-3}\), and \( \mu \) is dynamic viscosity in kg m\(^{-2}\) s\(^{-1}\). It is assumed that there is no fluid flow within the porous scale (for \( x < \delta_f \)).

When species \( j \) is involved \( k \) chemical reactions simultaneously, one can write for the source/sink term in (1):

\[ R_j = a_{jk} r_k \]  
(4)

where tensor notation applies for the subscripts, \( a_{jk} \) is the stoichiometric matrix where row \( j \) represents the \( j \)-th species, column \( k \) represents the \( k \)-th chemical reaction, and \( r_k \) is the reaction rate vector.

Chemical reactions of special interest are precipitation of iron carbonate and iron sulfide, which have been implemented in the model, as they take place at the steel surface and in the porous corrosion scale. The precipitation reactions act as a sink for Fe\(^{2+}\), CO\(_3^{2-}\), S\(^2-\) ions, influencing the fluxes and concentration gradients for both the ions and all other carbonic and sulfide species. The kinetics of scale growth is described in more detail in the sections below.

One equation of the form (1) is written for each species. They all have to be solved simultaneously in space and time. The boundary conditions for this set of partial differential equations are: in the bulk - equilibrium concentrations of species (which is also used as the initial condition), and at the steel surface - a flux of species is determined from the rate of the electrochemical reactions (zero flux for non-electroactive species). This is where the link between corrosion and species transport is established. The corrosion process determines the fluxes of the species at the metal surface and thus affects their transport into the solution, while in turn the transport rate and mechanisms affect the concentration of the species at the metal surface which determine the rate of corrosion. Hence a two-way coupling is achieved, what is physically realistic.

As the corrosion process is electrochemical in nature, the corrosion rate can be explicitly determined by calculating the rate of the electrochemical reactions underlying it such as: iron oxidation as well as reductions of hydrogen ion, carbonic acid, acetic acid, etc. The electrochemical reaction rate can be expressed as a current density, \( i \) (expressed in A m\(^{-2}\)), which is a function of the electrochemical potential at the metal surface, \( E \) (expressed in V):

\[ i = z \ i_0 \cdot 10^{- \frac{E - E_{\text{aux}}}{b}} \cdot \prod_{s=1}^{n} (1 - \theta_s) \]  
(5)

This equation is unique for each of the electrochemical reactions involved in a corrosion process such as hydrogen reduction, iron oxidation, etc. The “+” sign applies for anodic reactions while the “−” sign applies for cathodic reactions. \( \theta_s \) is the fraction of the steel surface where a given electrochemical reaction does not occur because the surface is covered by a species \( s \) which could be an adsorbed
inhibitor or a protective scale (such as iron carbonate or iron sulfide). The product sign \( \prod \) accounts for a compounding effect by more than one surface species. For each electrochemical reaction, equation (5) is different because of the unique parameters defining it: \( i_o \) - the exchange current density in \( A m^{-2} \), \( E_{\text{rev}} \) - the reversible potential in \( V \), and \( b \) - the Tafel slope in \( V \). These parameters have to be determined experimentally and are functions of temperature and in some cases species concentrations. An overview covering how these parameters are calculated in the present model is given elsewhere (reference\(^{12}\) for the cathodic reactions and reference\(^{13}\) for the anodic reaction). The unknown electrochemical potential at the metal surface \( E \) in (5), is also called corrosion potential or open circuit potential, which can be found from the current (charge) balance equation at the metal surface:

\[
\sum_{a=1}^{n_a} i_a = \sum_{c=1}^{n_c} i_c
\]  

(6)

where \( n_a \) and \( n_c \) are the total number of anodic and cathodic reactions respectively.

### The effect of low temperatures

The original model described above has successfully been used to predict the uniform \( \text{CO}_2 \) corrosion rate at temperatures between 20ºC and 80ºC (see Figure 1).\(^3\) However, the corrosion rate is poorly predicted when this model is used at lower temperatures, often seen in the field (see Figure 2). Similar malfunction of a much simpler empirical NORSOK M-506 model was also seen at low temperatures as was demonstrated by Olsen et al. (see Figure 3).\(^{14}\) In both cases, the models greatly overpredicted the corrosion rate at very low temperatures. It is not easy to adjust an empirical model without doing a full recalibration or by introducing another questionable correction factor. However, a more straightforward remedy can be found for the original mechanistic model.

Fang\(^{15}\) has conducted numerous experiments at low temperatures and identified the source of the problem with the mechanistic model. By performing potentiodynamic sweeps at low temperatures and using them to analyze the corrosion mechanisms, Fang\(^{15}\) was able to pinpoint where the problem lies and what needs to be done about it. He concluded that the main issue was with the standard activation energies used to describe the behavior of various electrochemical reactions with temperature. The activation energy which applies between 20ºC and 80ºC is dramatically increased as the water freezing point is approached. In other words, the rate of the various electrochemical reactions slows down much more rapidly than anticipated as the temperature approaches 0ºC. Interestingly, the standard activation energies for the mass transfer and homogenous chemical reactions in the model worked well across the whole temperature range and did not display this anomaly at very low temperature. The only explanation for the observed behavior of the electrochemical reactions is a change of the reaction mechanism or, what is more likely, a change in the rate determining step (rds) of a given heterogeneous reaction sequence. Therefore, in order to obtain more accurate predictions at low temperatures, Fang\(^{15}\) suggested that, as a first remedy, the activation energies for the four key electrochemical reactions underlying the \( \text{CO}_2 \) corrosion need to be adjusted at low temperature. The values he suggested for 5ºC and 1ºC are shown in Table 1.

The predictions obtained with the new updated model, using adjusted activation energies for the four electrochemical reactions, is shown in Figure 4. Clearly a much better agreement is obtained and the new updated model can be trusted at low temperature. The question that remains unanswered is: what is the reason for this unusual change of reaction mechanisms? Even if one accepts Fang’s\(^{15}\) suggestion that it is only a change in the rds for a heterogeneous multistep electrochemical reaction sequence, the question is: which step becomes rate controlling at low temperature and why? Clearly this is a challenge that will require a new effort and a more detailed mechanistic electrochemical study.
Table 1. Activation energy for different reactions at different temperatures suggested by Fang\textsuperscript{15}

<table>
<thead>
<tr>
<th></th>
<th>80ºC&gt;T&gt;20ºC</th>
<th>5ºC</th>
<th>1ºC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proton reduction</td>
<td>30 kJ/mol</td>
<td>100 kJ/mol</td>
<td>140 kJ/mol</td>
</tr>
<tr>
<td>Carbonic acid reduction</td>
<td>50 kJ/mol</td>
<td>100 kJ/mol</td>
<td>140 kJ/mol</td>
</tr>
<tr>
<td>Iron dissolution</td>
<td>37.5 kJ/mol</td>
<td>185 kJ/mol</td>
<td>215 kJ/mol</td>
</tr>
<tr>
<td>Water reduction</td>
<td>30 kJ/mol</td>
<td>120 kJ/mol</td>
<td>140 kJ/mol</td>
</tr>
</tbody>
</table>

The effect of high salinity

Another situation often seen in the field and rarely studied in the lab is the CO\textsubscript{2} corrosion of mild steel in high salinity brines (>>1\% by weight). Contradicting anecdotal evidence exists about the effect of very high chloride content, some suggesting that it is detrimental to survival of mild steel in CO\textsubscript{2} saturated solutions, others suggesting that it is beneficial. This confusion carried over into the models. The original model therefore did not include any effect of high salinity.

In his thesis work, Fang\textsuperscript{15} has looked at this effect as well, and found that the CO\textsubscript{2} corrosion rates of mild steel are severely retarded with high salt concentrations across the whole temperature range. It is seen in Figure 5 that the original mechanistic model, which work reasonably well up to 3\% NaCl in solution, deviates progressively as the concentration of salt increases.

Higher salt concentration changes the brine physico-chemical properties comprehensively. The density and viscosity are increased; however this effect is well understood and readily modeled. As the salt concentration increases well beyond a few percent, the brine becomes a non-ideal solution. This is also well understood and can be accounted for by using a water chemistry model based on Pitzer’s theory to calculate the activity coefficients and adjust the concentrations of the various ionic species in solution. The result is shown in Figure 6, where it is seen that high salt concentration affects mainly the activity coefficient of H\textsuperscript{+} ions in saturated CO\textsubscript{2} solutions. Consequently, the pH of a brine changes significantly with increasing salt concentration. Furthermore, high salt concentration will decrease the solubility of CO\textsubscript{2} in the corrosion solution, another effect that is well known and readily modeled. However, even when all these effects are taken into account, the original model could not capture the effect of high salinity on diminishing uniform CO\textsubscript{2} corrosion rate, as seen in Figure 5.

By conducting a detailed electrochemical analysis, Fang\textsuperscript{15} has discovered that many other processes underlying CO\textsubscript{2} corrosion are affected by salt concentration. The homogenous chemical reactions were predominantly influenced by the abovementioned change in activity coefficients and the decrease in the solubility of CO\textsubscript{2} in water. The heterogeneous electrochemical reactions at the steel surface were affected in the first place by competitive adsorption and surface reaction retardation. Mass transfer coefficients were also changed for reasons beyond those related to the increased density and viscosity of the brine. When all these effects were accounted for by using the appropriate theories where available,\textsuperscript{15} and introduced into the new updated model, a much improved fit with the experimental results is obtained across the whole temperature and salinity range (Figure 7).
The effect of H₂S

Internal CO₂ corrosion of mild steel in the presence of hydrogen sulfide (H₂S) represents a significant problem for the oil and gas industry. In CO₂/H₂S corrosion of mild steel, both iron carbonate and iron sulfide layers can form on the steel surface. Studies have demonstrated that surface layer formation is one of the important factors governing the corrosion rate in H₂S corrosion. Until recently there were no predictive models of H₂S corrosion of mild steel. Only a very rudimentary model of CO₂ retardation by traces of H₂S was built into the original model. In a recent study Sun has proposed a rather simple mechanistic model which is described below.

The corrosion of mild steel in H₂S aqueous environments proceeds initially by a very fast direct heterogeneous chemical reaction at the steel surface to form a solid adherent mackinawite layer. The overall reaction scheme can be written as:

\[ \text{Fe}(s) + H₂S \Rightarrow \text{FeS}(s) + H₂ \]  (7)

As both the initial and final state of Fe is solid, this reaction is often referred to as the “solid state corrosion reaction”. This film is very thin (<1μm) but dense and acts as a solid state diffusion barrier for the sulfide species involved in the corrosion reaction. The thin mackinawite film continuously goes through a cyclic process of growth, cracking and delamination, what generates the outer sulfide layer which thickens over time (typically >1μm) and also presents a diffusion barrier; this outer sulfide layer is very porous and rather loosely attached, over time it may crack, peel and spall, a process aggravated by the flow. Iron sulfide precipitation and transformation may occur in long exposures and effect not accounted for in the present model.

Due to the presence of the thin inner mackinawite film and the outer sulfide layer, it is assumed that the corrosion rate of steel in H₂S solutions is always under mass transfer control. Based on the discussions above, a schematic of the H₂S corrosion process is shown in Figure 8. According to the depiction shown one can write the flux of hydrogen sulfide due to:

convective diffusion through the mass transfer boundary layer:

\[ \text{Flux}_{H₂S} = k_{m,H₂S} \left( c_{b,H₂S} - c_{o,H₂S} \right) \]  (8)

molecular diffusion through the liquid in the porous outer layer:

\[ \text{Flux}_{H₂S} = \frac{D_{H₂S}}{\delta_{os}} \varepsilon \Psi \left( c_{o,H₂S} - c_{i,H₂S} \right) \]  (9)

solid state diffusion through the inner mackinawite film:

\[ \text{Flux}_{H₂S} = A_{H₂S} e^{\frac{B_{H₂S}}{RT_s}} \ln \left( \frac{c_{i,H₂S}}{c_{s,H₂S}} \right) \]  (10)

where:

- \( \text{Flux}_{H₂S} \) flux of H₂S expressed in mol/(m²s),
- \( k_{m,H₂S} \) is the mass transfer coefficient for H₂S in the hydrodynamic boundary layer, \( k_{m,H₂S} = 1.00 \times 10^{-4} \) in nearly stagnant condition, in m/s,
\( c_{b,H_2S} \) is the bulk concentration of \( H_2S \) in the liquid phase in \( \text{mol/m}^3 \),
\( c_{o,H_2S} \) is the interfacial concentration of \( H_2S \) at the outer layer/solution interface in \( \text{mol/m}^3 \),
\( D_{H_2S} \) is the diffusion coefficient for dissolved \( H_2S \) in water, \( D_{H_2S} = 2.00 \times 10^{-9} \), in \( \text{m}^2/\text{s} \),
\( \varepsilon \) is the outer mackinawite layer porosity,
\( \Psi \) is the outer mackinawite layer tortuosity factor,
\( c_{i,H_2S} \) is the interfacial concentration of \( H_2S \) at the inner layer/film interface in \( \text{mol/m}^3 \).
\( \delta_{os} \) is the thickness of the mackinawite layer \( \delta_{os} = m_{os} / (\rho_{FeS} A) \) in m,
\( m_{os} \) is the mass of the mackinawite layer in kg,
\( A \) is the surface area of the steel in \( \text{m}^2 \),
\( A_{H_2S}, B_{H_2S} \) are the constants in the Arrhenius equation,
\( A_{H_2S} = 2.0 \times 10^{-5} \text{mol/(m}^2\text{s}) \) and \( B_{H_2S} = 15,500 \text{ J/mol} \),
\( T_k \) is the temperature in K,
\( c_{s,H_2S} \) is the “near-zero” concentration of \( H_2S \) on the steel surface and is set to \( 1.00 \times 10^{-7} \) in \( \text{mol/m}^3 \).

In the *new updated model* the fluxes through the liquid boundary layer (8) and the outer sulfide scale (9) are already accounted for by the general transport equation (1). The solid state diffusion flux through the very thin mackinawite film (10) was implemented as a boundary condition on the steel surface side of the computational domain and replaces the electrochemical boundary condition (5).

The prediction for \( \text{Flux}_{H_2S} \) depends on a number of constants used in the model which can be either found in handbooks, calculated from the established theory or are determined from the experiments. The unknown thickness of the outer sulfide layer changes with time and was calculated as described below.

It is assumed that the quantity of the sulfide layer retained on the metal surface at any point in time depends on the balance of:

- layer formation kinetics (as the layer is generated by spalling of the thin mackinawite film underneath it and by precipitation from the solution), and
- layer damage kinetics (as the layer is damaged by intrinsic or hydrodynamic stresses and/or by chemical dissolution)

\[
\text{sulfide layer retention rate} = \text{sulfide layer formation rate} - \text{sulfide layer damage rate}
\]

where all the terms are expressed in \( \text{mol/(m}^2\text{s}) \). It is here assumed that in the typical range of application \((4<\text{pH}<7)\), precipitation and dissolution of iron sulfide layer do not play a significant role, so it can be written:

\[
\text{sulfide layer retention rate} = \text{CR}_{H_2S} - \text{SDR}_{\text{mech}}
\]

Where \( \text{CR}_{H_2S} \) is the corrosion rate measured in \( \text{mol/(m}^2\text{s}) \). The implicit assumption is that all the corroded iron from the steel ends up as solid iron sulfide, which may adhere to or spall from the steel surface. Experiments have shown that even in stagnant conditions about half of the sulfide layer that forms is lost.
from the steel surface due to intrinsic growth stresses by internal cracking and spalling, i.e.
\[ SDR_n \approx 0.5 \text{ CR} \]. More experimentation is required to determine how the mechanical layer damage is affected by hydrodynamic forces. Once the layer retention rate \( SRR \) is known, the change in mass of the outer sulfide layer can be easily calculated as:

\[ \Delta m_{os} = SRR \ M_{FeS} \ A \ \Delta t \]  

(13)

where \( M_{FeS} \) is the molar mass of iron sulfide in kg/mol, \( \Delta t \) is the time interval in seconds. The porosity of the outer mackinawite layer was determined experimentally to be very high (\( \varepsilon \approx 0.9 \)), however due to its layered structure the tortuosity factor was found (by trial and error) to be very low (\( \Psi = 0.003 \)).

Similar equations were derived by Sun\(^{24}\) for diffusion of \( \text{CO}_2 \), \( \text{H}^+ \) and organic acids through the sulfide layers, thus completing the model of mixed \( \text{H}_2\text{S} /\text{CO}_2 /\text{organic acid} \) corrosion. This model was implemented into the \textit{new updated model} and extensively verified showing a remarkable agreement with the various experimental results in which the partial pressure of \( \text{H}_2\text{S} \) varied from a fraction of a mbar to 10 bar. Over this large range the predictions were never more than a factor 2 different from the measured values.

A sample of the comparison is shown in Figure 9 for a case of \( \text{CO}_2 \) corrosion with traces of \( \text{H}_2\text{S} \), i.e. at extremely low \( \text{H}_2\text{S} \) partial pressures ranging from 0.0013 – 0.32 mbar (experiments conducted by Lee\(^{23}\)). This corresponds to 1 – 250 ppm\( m \) (by mass) in the gas phase at 1 bar \( \text{CO}_2 \) and is a \( \text{CO}_2 \) dominated corrosion scenario (\( p\text{CO}_2/p\text{H}_2\text{S} \) ratio is in the range \( 10^3 – 10^6 \)). However the \( \text{H}_2\text{S} \) controls the corrosion rate even when present in such minute amounts. The pure \( \text{CO}_2 \) (\( \text{H}_2\text{S} \)-free) corrosion rate is reduced by 3 to 10 times, due to traces of \( \text{H}_2\text{S} \), because of the formation of a mackinawite film. The model successfully captures this effect as shown in Figure 9.

At the other extreme, corrosion experiments at very high temperature (120°C), high partial pressures of \( \text{CO}_2 \) (\( p\text{CO}_2=6.9 \) bar) and \( \text{H}_2\text{S} \) (\( p\text{H}_2\text{S}=1.38 – 4.14 \) bar) were recently reported by Kvarekval et al.\(^{25}\) In exposure lasting up to 16 days, a steadily decreasing corrosion rate was observed due to buildup of a protective iron sulfide layer. The effect of various \( \text{H}_2\text{S} \) concentrations on the corrosion rate was very small. This effect was readily captured by the model with very good accuracy as seen in Figure 10. This is a situation where the \( \text{H}_2\text{S} \) is the dominant corrosive species. At the highest \( p\text{CO}_2/p\text{H}_2\text{S} \) ratio of 5 (\( p\text{CO}_2=6.9 \) bar, \( p\text{H}_2\text{S}=1.38 \) bar) \( \text{H}_2\text{S} \) generated approximately 70% of the corrosion rate. At the lowest \( p\text{CO}_2/p\text{H}_2\text{S} \) ratio of 1.67 (\( p\text{CO}_2=6.9 \) bar, \( p\text{H}_2\text{S}=4.14 \) bar) \( \text{H}_2\text{S} \) generated 82% of the overall corrosion rate.

**CONCLUSIONS**

New advances have been made in modelling the internal corrosion of steel pipelines which are based on the latest experimental results. The biggest advances are:

- \( \text{CO}_2 \) corrosion at very low temperatures (as low as 1°C), where the corrosion rates are greatly reduced;
- \( \text{CO}_2 \) corrosion in very high salinity brines (up to 25%wt. NaCl), where strong retardation of corrosion rates is seen;
- \( \text{CO}_2 \) corrosion in the presence \( \text{H}_2\text{S} \) where the corrosion rate is governed by formation of a iron sulfide layer by solid state reaction.

The overall model has been extensively calibrated and verified with a reliable experimental database and very good agreement was found.
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Figure 1. Comparison between experimental results (points) and predictions (line) for the original model. Conditions: 1 bar CO$_2$, 1% NaCl solution, single-phase flow, $v=1$ m/s, in a 15 ID pipe. Data taken from the flow loop experiments of Nesic et al.$^3$

Figure 2. Comparison between experimental results (points) and predictions (line) for the original model at low temperature. Conditions: 1 bar CO$_2$, 3% NaCl solution, pH 4, single-phase rotating cylinder flow. Data taken from the glass cell experiments of Fang.$^{15}$
Figure 3. Comparison between experimental results and predictions for the NORSOK M-506 model at low temperature. Taken from Olsen et al.\textsuperscript{14}

Figure 4. Comparison between experimental results (points) and predictions (line) for the new updated model at low temperature. Conditions: 1 bar CO\textsubscript{2}, 3\% NaCl solution, pH 4, single-phase rotating cylinder flow. Data taken from the glass cell experiments of Fang.\textsuperscript{15}
Figure 5. Comparison between experimental results (points) and predictions (lines) for the original model at various NaCl concentrations. Conditions: 5°C and 20°C, 1 bar CO₂, pH 4, single-phase rotating cylinder flow, 1000 rpm. Data taken from the glass cell experiments of Fang.15

Figure 6. Calculated activity coefficient change with salt concentration at 20°C at 1 bar total pressure.
Figure 7. Comparison between experimental results (points) and predictions (lines) for the new updated model at various NaCl concentrations. Conditions: 5°C and 20°C, 1 bar CO₂, pH 4, single-phase rotating cylinder flow, 1000 rpm. Data taken from the glass cell experiments of Fang.¹⁵

Figure 8. A schematic of the iron sulfide controlled corrosion process.
Figure 9. Corrosion rate vs. partial pressure of H$_2$S; experimental (points), predictions using *new updated model* (lines); conditions: total pressure $p=1$ bar, CO$_2$ partial pressure 0.1 bar, H$_2$S gas partial pressure from 0.0013 – 0.32 mbar, $T=20^\circ$C, reaction time 24 hours, pH 5, 1000 rpm. Data taken from Lee.$^{23}$

Figure 10. Corrosion rate vs. pH$_2$S/pCO$_2$ ratio; experimental (points), predictions using the *new updated model* (lines); conditions: total pressure $p=7$ bar, CO$_2$ partial pressure 6.9 bar, H$_2$S gas partial pressure from 1.38 – 4.14 bar, $T=120^\circ$C, experiment duration up to 16 days, pH 3.95 – 4.96, liquid velocity 10 m/s. Experimental data taken from Kvarekval et al.$^{25}$