Evaluation of the Diffusive Capacity of Hydrogen through 9Cr-1Mo Ferritic Steel in the Dual Environment of a Furnace: a Simulation Study in COMSOL Multiphysics®

David A. Méndez, Brayan E. Uribe, Juan Orozco-Agamez, Dario Peña-Ballesteros, Anibal Alviz-Meza

*Grupo de Investigaciones en Corrosión, Universidad Industrial de Santander, Parque Tecnológico Guatiguará, Piedecuesta 681011, Colombia

**Grupo de investigación en Deterioro de Materiales, Transición energética y Ciencia de Datos DANT3, Universidad de Sipán, Facultad de Ingeniería, Arquitectura y Urbanismo, Chiclayo, Pimentel 14001, Peru

alvizanibal@crece.uss.edu.pe

Hydrogen transport and storage represent a challenge due to its high diffusivity capacity, which causes the pitting of pipes. This study focused on addressing this problem by simulating the effect caused by the hydrogen generated during the heating pre-processing step of crude oil before addressing it to the distillation unit. For this purpose, the software COMSOL Multiphysics® was employed to determine how the hydrogen diffusion across ferritic steel containers increases the corrosion rate in a furnace. Then, a dual corrosion process was assessed at 650 °C, simulating the flow of water vapor at the outer part of the pipe whereas hydrogen sulfides -as contamination from crude oil- at the inner part. The ferritic steel selected for this research was the 9Cr-1Mo steel due to its usual implementation in processes developed at high temperatures in industry. Besides, we also studied the corrosion rate of both pipe’s faces independently, omitting the effect of the other fluid on the other side of the pipe -singular environments-. The influence of hydrogen sulfide on the corrosion rate on the water vapor side was found negligible. Meanwhile, water vapor boosted the corrosion rate on the hydrogen sulfide side, making it 1.2 times greater compared to its singular estimation. The corrosion rate induced by the water vapor is higher due to the higher number of corrosion products -current density- that it delivers when reacting with the steel elements at high temperatures. These approaches aimed to quantify the hydrogen diffusion contribution to the corrosion rate of the ferritic steels used in oil refining furnaces.

1. Introduction

The influence of two corrosive media on metallic structures is common in equipment such as furnaces, boilers, heaters, super-heaters, turbines, etc. For instance, dual corrosion occurs in pipes of refinery furnaces: fireside corrosion by flue gases on the external face and internal corrosion at the inner tube face. A few researchers have studied dual environments, demonstrating that ferritic steels undergo accelerated corrosion rates (Alnegren et al., 2016; Chandra et al., 2014; Coelho, 2014). Dual corrosion has attracted the attention of the scientific community beginning from the Lobnig & Grabke (1990) research work, which studied the simultaneous oxidation-sulfidation of Fe-20Cr-12Ni and Fe-20Cr steels in $H_2-H_2O-H_2S/H_2S-H_2O$ mixtures between 700 °C and 1150 °C, finding out that both phenomena were simulated simultaneously, increasing the alloys' corrosion rates. In general, in studies based on ferritic steels with different chromium percentages (2-14%), temperatures (550-750 °C), and corrosive environments, it has been found that one environment influences the corrosion rate of the other by up to a factor of 3, compared with their deterioration rate in isolated -singular- environments (Alnegren et al., 2016; Alviz et al., 2020; Chandra et al., 2014; Coelho, 2014; Huenert et al., 2010; Mosquera, 2019; Nakagawa et al., 2001). This augmentation in corrosion rate has been linked to the permeability of hydrogen through the wall thickness of alloys, which enhanced mass transfer processes (Chandra et al., 2014; Huenert et al., 2010; Reisert et al., 2021).
Ferritic steels such as Fe-9Cr-1Mo (P91) are commonly used in the metallic infrastructure of diverse equipment due to their good mechanical properties and acceptable corrosion resistance at high temperatures (Alviz et al., 2018). P91 steel corrosion rates have been deeply studied in water vapor and H₂S-containing environments (Ehlers et al., 2006; Rebak, 2011) whereas under the dual effect of the dual environment of furnaces from oil refineries, where crude oil flows in the inner pipes’ face and flue gases in the outer face, it hasn’t been assessed. To sum up, the gap that covers this work is the lack of experimental and theoretical studies to determine the role of H₂ in the corrosion rate of P91 alloy in the dual environment of a furnace from the oil refining sector. The influence of H₂ is evaluated through simulation by COMSOL Multiphysics®, limiting the dual environment of the furnace to H₂O and H₂S as representative compounds from flue gas and crude oil, respectively.

2. Materials and methods

The mass transfer conservation model was stated based on Fick's law, Faraday's law, and Navier Stokes equations, including the required boundary conditions. The equations (1) and (2) were used to determine the corrosion rate induced by each environment. The complete model can be extracted from the thesis work of Uribe & Méndez (2021) and Auinger et al., (2011), which due to its extension could not be included in this paper. The geometrical model used in COMSOL Multiphysics® is based on the image shown in Figure 1.

\[
V_{\text{corr}}(\text{mpy}) = FD_{\text{H}_2\text{O}\text{--N}_2}[\frac{\partial C_{\text{H}_2\text{O}}(g)}{\partial y} + \frac{\partial C_{\text{N}_2}(g)}{\partial x}] \left(\frac{\text{mA}}{\text{cm}^2}\right) \times 129 \frac{M}{a}
\]  
(1)

\[
V_{\text{corr}}(\text{mpy}) = FD_{\text{H}_2\text{S}--\text{N}_2}[\frac{\partial C_{\text{H}_2\text{S}}(g)}{\partial y} + \frac{\partial C_{\text{N}_2}(g)}{\partial x}] \left(\frac{\text{mA}}{\text{cm}^2}\right) \times 129 \frac{M}{a}
\]  
(2)

Figure 1: The geometry of the Comsol Multiphysics ® model. The figure shows that in section (1) flows a mixture of H₂S(g) + N₂(g) while in section (2) H₂O(g) + N₂(g). Point (3) represents the P91 steel sample.

The conservation of momentum and continuity equations were used in the chemical species transfer model and solved using Comsol Multiphysics® Version 5.6 (COMSOL Multiphysics®, 2019). The velocity profile was obtained from the above equations, which combined with the diffusion coefficient allowed us to determine either the concentration gradient. The current density and the corrosion rate were obtained once defined the concentration gradient. The equations set were resolved using the finite element method, which involves the division of the geometric model into different geometric elements of different sizes, by employing the physics of dilute species transport and laminar flow.

Table 1 shows the operating values of an oil refinery’s furnace while Table 2 shows the elemental weight fraction of the studied alloy. The steam concentration was taken as 6.96 mol/m³ and its volumetric flow as 250 mL/min according to the previous work reported by Alviz et al., (2021). The H₂S concentration was selected as 0.146 mol/m³ based on data from the oil refining industry whereas its flow remained the same as the water vapor.

In order to include the effect of one environment on the other in the simulation, we assume that this influence is due to the diffusion of hydrogen in the alloy’s bulk, generating discontinuities in the material and accelerating corrosion due to greater participation of the presented chemical species. Thus, we first solve the system for calculating the hydrogen diffusion effect on each side, further adding it to the current density generated in each isolated singular environment.
The proposed model includes the following limitations:

- It only considers H₂S and water vapor as the chemical species involved in the corrosion process of each pipe side.
- The simulation was carried out taking into account a 2D geometric model due to its axial symmetry and less troubling simulation process.
- It was considered a transient state to contemplate the fluctuating concentration.
- The model is diffusion-controlled: the kinetic of reactions are omitted considering an instantaneous process (Mosquera, 2019).

### 3. Results and discussion

#### 3.1 Simulation of singular environments.

This section presents the graphs obtained after the simulation, for which a variation of the flow rates was taken into account. The shape observed in the steam and H₂S curves is explained by the concentration profile developed in the scheme shown in Figure 1, which follows the gas flow gradient on the x-axis, as observed in the thesis of Uribe & Méndez (2021). The concentration is higher at the borders of the coupon on the steam side, which explicates the higher corrosion rates shown in Figure 2 at the extremes of the x-axis. In the case of the H₂S side, the highest concentration and therefore corrosion rate is located at the origin of the x-axis in Figure 3. This particular behavior is related to the low concentration of H₂S in the inner duct gas flow. The maximum corrosion rate on the steam side is 390 mpy, doubling the minimum value obtained at the center of the coupon after 30 seconds of reaction when the steady state is reached (see Figure 2). The rapid stability of the system is linked to the large flow rates introduced, compared to the dimensions of the geometry, as well as the consideration of instantaneous reaction kinetics in the system.

![Figure 2: Corrosion rate in the steam singular environment at 650°C](image)

Regarding sulfidation corrosion, Figure 3 shows that the corrosion rate reached a maximum value of 87 mpy and a minimum of 37 mpy for a stabilization time of close to 1 second. In this case, the stabilization time was shorter due to a higher gas flow velocity due to the 5.5 times smaller diameter of the inner duct in comparison with the outer one.
Figure 3: Corrosion rate due to H\textsubscript{2}S at 650°C

3.2 Results in the dual environment.

As demonstrated in studies done by Chandra & Kranzmann, (2018), Alnegren et al., (2016) y Nakagawa et al., (2001), in environments hydrogen-rich, H\textsubscript{2} diffuses through ferritic steels creating discontinuities, favoring the diffusion of chemical species from one side to the other, and augmenting their corrosion rate, as we obtained in Figure 4 and Figure 5. The influence of H\textsubscript{2}S on the water vapor side multiply 1.008 times its corrosion rate in the dual environment, compared to its singular baseline effect at 650 °C. Otherwise, water vapor multiplies 1.2 times the corrosion rate on the H\textsubscript{2}S side in the dual environment, regarding the H\textsubscript{2}S behavior under singular conditions at 650 °C. These results are explained by the hydrogen diffusion throughout the alloy, as suggested by other researchers (Mosquera, 2019; Nakagawa et al., 2001). Chandra & Kranzmann, (2018) showed similar trends in a dual environment of flue gas and water vapor: the corrosion rate of the ferritic alloy increase about 3 times the value obtained in the singular flue gas environment. Likewise, Alviz et al., (2021) found a corrosion rate for the P91 ferritic steel 1.6 times higher in the dual environment than in the singular one of an industrial simulated boiler.

Figure 4: Corrosion rate due to the influence of water vapor on H\textsubscript{2}S 650°C and 6x10\textsuperscript{4} s

Figures 6 and 7 show the corrosion rate obtained by different authors under similar singular conditions, compared with values gotten for singular cases in the present work. The conditions for steam of each author's studies are as follows: Laverde studied T91 steel at 550°C for 250 h in a mixture of Ar+H\textsubscript{2}O; Ehlers a P91 steel at 650°C for 60 h in an environment of N\textsubscript{2}-O\textsubscript{2}-H\textsubscript{2}O; Alviz et al., (2021) a P91 steel at 650°C for 200 h in flue gas/steam; Alviz et al., (2020) a P91 steel at 650°C for 200 h in N\textsubscript{2}-O\textsubscript{2}-H\textsubscript{2}O-H\textsubscript{2}S. On the other hand, the conditions used in the H\textsubscript{2}S studies are the following: Gao tested an API 5L steel at 120°C for 504 hours, in H\textsubscript{2}S-N\textsubscript{2}, and Dong a T91 steel at 700°C for 70 hours in N\textsubscript{2}-(0.5,2.5)%H\textsubscript{2}S molar.
A comparison of results for each singular environment is separately presented below to determine how well our model matches the reported values. In the water vapor case, our corrosion rate was found higher than in the selected research works; being the most distant 2.75 times lower (Alviz et al., 2021) whereas the closest 1.22 times lower (Alviz et al., 2020). This is associated with the difference in the corrosive agents presented in reactants. On the other hand, the H₂S corrosion rate was 3.5 times lower in our work than the one found by Dong (2016) and 1.2 times higher than that of Gao (2017). The large difference with Duong’s research is explained by the 3.5 times higher concentration of H₂S employed by him.

4. Conclusions

Based on a proposed model for the P91 steels’ corrosion rate estimation in a dual environment, a simulation was carried out in Comsol Multiphysics®, using the physics of dilute species’ transport and a laminar flow. The flow rate employed was 250 mL/min for both environments, with concentrations of 6.96 mol/m³ and 0.146 mol/m³ for H₂O and H₂S, respectively. It was determined that the corrosion rate of the system is higher in water vapor (between 275-400 mpy) than in H₂S (between 32-87 mpy) at 650 °C. In the dual environment, the corrosion rate
obtained in the H₂S environment was 1.2 times higher due to the H₂ effect. Meanwhile, for the H₂O side, it was 1.008 times higher due to the same effect from the hydrogen released by H₂S. These results were compared with scientific literature, obtaining acceptable corrosion rate matchings, and agreeing with the explanation given to the hydrogen contribution: it enhances the diffusive processes through the alloy’s bulk. In short, the contribution of hydrogen in corrosive environments should be considered a key factor to select the best alloy in equipment working at high temperatures in dual environments.

Acknowledgments
The authors express their gratitude to professor Jefferson Oswaldo Ruiz Lizarazo for his support in the development of the simulation system in COMSOL Multiphysics®.

References