Impact of saline aquifer water on surface and shallow pit corrosion of martensitic stainless steels during exposure to CO$_2$ environment (CCS)

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Abstract. Pipe steels suitable for carbon capture and storage technology (CCS) require resistance against the corrosive environment of a potential CCS-site, e.g. heat, pressure, salinity of the aquifer, CO$_2$-partial pressure. Samples of different mild and high alloyed stainless injection-pipe steels partially heat treated: 42CrMo4, X20Cr13, X46Cr13, X35CrMo4 as well as X5CrNiCuNb16-4 were kept at T=60 °C and ambient pressure as well as p=100 bar for 700 h - 8000 h in a CO$_2$-saturated synthetic aquifer environment similar to possible geological on-shore CCS-sites in the northern German Basin. Main corrosion products are FeCO$_3$ and FeOOH. Corrosion rates obtained at 100 bar are generally much lower than those measured at ambient pressure. Highest surface corrosion rates are 0.8 mm/year for 42CrMo4 and lowest 0.01 mm/year for X5CrNiCuNb16-4 in the vapour phase at ambient pressure. At 100 bar the highest corrosion rates are 0.01 mm/year for 42CrMo4, X20Cr13 (liquid phase), X46Cr13 and less than 0.01 mm/year for X35CrMo4 and X5CrNiCuNb16-4 after 8000 h of exposure with no regard to atmosphere. Martensitic microstructure offers good corrosion resistance.

1. Introduction
During the compression of emission gasses e.g. from combustion processes of power plants into deep geological layers of a geological on-shore saline aquifer CCS-site (CCS Carbon Capture and Storage [1]), the corrosion of injection pipe steels may become an issue when CO$_2$ is compressed [2], [3]. CO$_2$-corrosion is sensitively dependent on alloy composition, contamination of alloy and media, temperature, CO$_2$ partial pressure, flow conditions and protective corrosion scales [4]-[10]. Suitable heat treatment [8], [11] and martensitic micro structure [12] decrease the pitting potential. Corrosion phenomena are reason for early failure of materials under cyclic load [13], [14]. Surface corrosion layers grow slowly and locally corroded samples usually show the same corrosion products as surface layers [7], acting as corrosion catalyster. Generally siderite FeCO$_3$ [2] is formed on steels exposed to CO$_2$-environment due to the low solubility of FeCO$_3$ in water (pK$_{sp}$ = 10.54 at 25 °C [14]) leading to anodic iron dissolution. The initial formation of transient Fe(OH)$_2$ [6], [11] possibly leads to an increase of the local pH near the hydroxide film. As a consequence an internal and external ferrous carbonate film precipitates [14]. In this paper we are presenting extended and compiled data of former work [3], [7], [13], [15]-[18].
2. Experimental Procedure

2.1. Steels
Static corrosion tests at ambient pressure as well as at high pressure (100 bar) were carried out using samples of mild steels, martensitic and duplex stainless steels (Table 1-5):
1. AISI 420 (X20Cr13, 1.4021) (Table 1)
2. AISI 420C (X46Cr13, 1.4043) (Table 2)
3. No AISI (X35CrMo17, 1.4122) (Table 3)
4. AISI 630 (X5CrNiCuNb 16-4, 1.4542) (Table 4)
5. AISI A182 F51 (329LN) SAF 2205 (X2 CrNiMoN 22-5-3 (UNS S31803) 1.4462) (Table 5)

In order to confirm the material’s chemical composition, samples were analyzed via spark emission spectrometry SPEKTROLAB M and by Electron Probe Microanalyzer JXA8900-RLn (Table 1-5).

| Table 1. Chemical composition of 1.4021 (X20Cr13, AISI 420, (in mass-%)). |
|-----------------|----------|----------|--------|------|----------|--------|------|-------|
| Elements        | C        | Si       | Mn     | P     | S       | Cr     | Ni   | Fe    |
|                 | 0.22     | 0.39     | 0.32   | 0.007 | 0.006   | 13.3   | 0.123| rest  |

| Table 2. Chemical composition of 1.4043 (X46Cr13, AISI 420C), (in mass percent). |
|-----------------|----------|----------|--------|------|----------|--------|------|-------|
| Elements        | C        | Si       | Mn     | P     | S       | Cr     | Mo   | Ni    |
|                 | 0.46     | 0.25     | 0.45   | 0.018 | 0.003   | 13.39  | 0.03 | 0.13  |

| Table 3. Chemical composition of 1.4122 (X35CrMo17), (in mass percent). |
|-----------------|----------|----------|--------|------|----------|--------|------|-------|
| Elements        | C        | Si       | Mn     | P     | S       | Cr     | Ni   | Co    |
| acc standard a  | 0.33-0.45|<1.00     |£1.00  |£0.045|£0.03    |rest   |      |       |
| Elements        | Cr       | Mo       | Ni     | Co   | Fe      |
| acc standard a  | 15.5-17.5|0.8-1.3   |£1.00  |0.20-0.45|        |
| a) elements as specified according to DIN EN 10088-3 in % |

| Table 4. Chemical composition of 1.4542 (X5CrNiCuNb16-4, AISI 630), (in mass percent). |
|-----------------|----------|----------|--------|------|----------|--------|------|-------|
| Elements        | C        | Si       | Mn     | P     | S       | Cr     | Mo   | Ni    |
|                 | 0.03     | 0.42     | 0.68   | 0.018 | 0.002   | 15.75  | 0.11 | 4.54  |
|                 |          |          |        |       |         |        |      | 3.00  |
|                 |          |          |        |       |         |        |      | 0.242 |

Duplex steel 1.4462 (X2 CrNiMoN 22 5 3, (table 5) manufactured by means of continuous casting followed by an appropriate tempering and water quenching allowing for an equilibrium of ferrite and austenite phase. Due to its percentage of austenite 1.4462 reveals a much better corrosion resistance resulting and a PREN number (35.1) twice as high as 1.4542 [16].

| Table 5. Chemical composition of 1.4462 X2 CrNiMoN 22 5 3 (UNS S31803) (in mass percent). |
|-----------------|----------|----------|--------|------|----------|--------|------|-------|
| phases          | C        | Si       | Mn     | Cr   | Mo       | Ni     | N    |
| α & γ **        | 0.023    | 0.48     | 1.83   | 22.53| 2.92     | 5.64   | 0.15 |

*PREN α =37.4, γ ≈32.4 ** P=0.024, S=0.008

2.2. Aquifer Water
To simulate in-situ geothermal condition the geothermal aquifer water (as known to be similar to the Stuttgart Aquifer [19], [20]: Ca²⁺: 1760 mg/L, K⁺: 430 mg/L, Mg²⁺: 1270 mg/L, Na⁺: 90100 mg/L, Cl⁻: 143,300 mg/L, SO₄²⁻: 3600 mg/L, HCO₃⁻: 40 mg/L) was synthesized in a strictly orderly way to avoid precipitation of salts and carbonates.
2.3. Corrosion Experiments

Laboratory scale exposure tests in CO₂-saturated aquifer brine and water saturated CO₂ were carried out using coupons of the steel qualities that may be used as injection pipe. Therefore steel samples made of as received and thermally treated specimens with 8 mm thickness, 20 mm width, 50 mm length were used. To gain martensitic microstructure with sufficient hardness and toughness prior to exposure heat treatment was done following commonly used protocols [7], [15], [17].

A hole of 3.9 mm diameter was used for sample positioning. Samples of each base metal were positioned within the vapour phase and within the liquid phase. Flow control (3 NL/h) of the technical CO₂ (purity 99.995 vol.-%)) into the brine at ambient pressure was done by a capillary meter GDX600_man by QCAL Messtechnik GmbH, Munich. The exposure of the samples between 700 h to 8000 h was disposed in reaction vessels according to the conditions at the geological site at 60 °C at ambient pressure – each material in a separated reaction vessel [3] and at 100 bar [7], [13], [15] (Figure 1).

![Figure 1. Reaction vessels and experimental set up [7], [17].](image)

Before corrosion tests the surfaces of the steels were activated by grinding with SiC-Paper down to 120 μm under water. After the corrosion tests, the samples were cut partly for scale analysis with the corrosion layer and prepared partly for kinetic analysis after the scale was etched. Descaling of the samples was performed by exposure to 37% HCl. Then parts of the samples were embedded in a cold resin (Epoxicure, Buehler), cut and polished first with SiC-Paper from 180 μm to 1200 μm under water and then finished with diamond paste 6 μm and 1 μm.

2.4. Analysis

Different light optical and electron microscopy techniques were performed on specimens to investigate the layer structures and morphology of the samples. X-ray diffraction was carried out in a URD-6 (Seifert-FPM) with CoKα-radiation with an automatic slit adjustment, step 0.03° and count 5 sec. Phase analysis was performed by matching peak positions automatically with PDF-2 (2005) powder patterns. Mainly structures that were likely to precipitate from the steels were chosen from the ICSD and refined to fit the raw-data-files using POWDERCELL 2.4 [21] and AUTOQUAN ® by Seifert FPM. Kinetics of the corrosion were determined by the corrosion rates which were calculated via mass change of the samples before and after corrosion testing according to DIN 50 905 part 1-4.

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\text{corrosion rate} \left[ \frac{\text{mm}}{\text{year}} \right] = \frac{8760 \left[ \frac{\text{hours}}{\text{year}} \right] \cdot 10 \left[ \frac{\text{mm}}{\text{cm}} \right] \cdot \text{weight loss} \left[ \frac{g}{\text{cm}^2} \right]}{\text{area} \left[ \frac{\text{cm}^2}{\text{cm}} \right] \cdot \text{density} \left[ \frac{g}{\text{cm}^3} \right] \cdot \text{time} \left[ \text{hour} \right]} \tag{1} \]

3. Results and Discussion

As expected due to the high PREN (53.1 (Pitting Resistance Equivalent (PRE = %Cr + 3.3% Mo + 16% N)) for the duplex stainless steel X2 CrNiMoN22-5-3 is reason for no corrosion phenomena at all, even after long exposure times in Stuttgart Aquifer water [19] and Northern German Basin [20] with
or without flowing technical CO\textsubscript{2}. Neither ferrite nor austenite phase showed corrosive deterioration (Figure 2). Molybdenum specially increases the resistance to pitting and crevice corrosion.

![Figure 2. Microstructure of experimental alloy prior to testing; (a) transversal and (b) longitudinal sections respecting the casting direction. Etching: Beraha II [19].](image)

In general corrosion rates are lower at 100 bar compared to ambient pressure. This applies for mild steel 42CrMo4 (Figure 3) as well as for martensitic stainless steels X46Cr13 (Figure 4), X35CrMo17 (Figure 5) and X5CrNiCuNb 16-4 (Figure 6). Corrosion rates are higher at ambient pressure than at 100 bar, which may directly be related to an open capillary system within the corrosion layer. Under pressure these are not present suppressing fast mutual diffusion of ionic species that is necessary for scale growth [18]. The corrosion rates of 42CrMo4, X20Cr13 and X46Cr13 increase with time at ambient pressure and decrease with time at 100 bar. X5CrNiCuNb16-4 and X35CrMo17 show increasing corrosion rates at ambient pressure as well as at 100 bar [18].

One reason for the decreasing corrosion rates lower gas tightness of the low pressure system with excess oxygen in the test vessels at ambient pressure leading to higher corrosion rates as a function of exposure time. Decreasing growth rates at 100 bar may be related to the fast formation of initial hydroxides [2], [6], [14], [18] then precipitating to a passivating siderite-layer with increasing thickness as a function of exposure time. This corrosion scale acts as a diffusion barrier towards mutual diffusion of ionic species to the metal surface. The corrosion rate may also decrease in the presence of iron carbonate precipitates and explains lower corrosion rates at longer exposure times. In general corrosion rates are severely dependent on the exposure time for specimens kept in the vapour phase. For example the corrosion rate is twice as high for 42CrMo4 after 8000 h (0.8 mm/year) in comparison to 4000 h (0.45 mm/year) when exposed at ambient pressure. This effect has less impact on specimens kept in the liquid phase or at 100 bar with no regard to atmosphere [18]. However, the atmosphere (liquid, vapour or supercritical atmosphere) does not significantly influence the general corrosive behavior.

The pressure has little influence on the corrosion rates. For 42CrMo4, X20Cr13 and X35CrMo4 the corrosion rates at ambient pressure are much higher than at 100 bar. For X20Cr13 and X5CrNiCuNb16-4 the difference is not as significant. In general the higher corrosion rates at ambient pressure may be explained by higher diffusion kinetics. The corrosion rates in the vapour phase (water-saturated CO\textsubscript{2}) exceed those measured in the liquid phase (CO\textsubscript{2}-saturated water) by a factor of 3–8. But, at high pressures (100 bar) corrosion rates in the liquid phases can be higher or lower than in the vapour suppercritical phase.
Figure 3. Comparison of corrosion rates of 42CrMo4 after 8000 h of exposure to aquifer brine water at 60 °C and ambient pressure [3] as well as at 100 bar [18].

Figure 4. Comparison of corrosion rates of X20Cr13 in the liquid and vapour/supercritical phase after 8000 h of exposure to aquifer brine water at 60 °C and ambient pressure as well as at 100 bar.

Figure 5. Comparison of corrosion rates of X46Cr13 in the liquid and vapour/supercritical phase after 8000 h of exposure to aquifer brine water at 60 °C and ambient pressure [3] as well as at 100 bar [18].

Figure 6. Comparison of corrosion rates of X35CrMo17 in the liquid and vapour/supercritical phase after 8000 h of exposure to aquifer brine water at 60 °C and ambient pressure as well as at 100 bar.

However, the corrosion rates for 42CrMo4 are higher in the liquid/vapour phase and also higher at ambient pressure compared to high pressure. This corresponds with the thickness of the precipitation layer that is greater in vapour/supercritical than in the liquid phase (Figure 3 and Figure 7).
The multi-layered corrosion scale precipitated on the steels at ambient pressure is complicated and has been described in detail [7], [17], [18]. This characteristic carbonate/oxide structure consists of siderite FeCO$_3$, goethite α-FeOOH, Mackinawite FeS and akaganeite Fe$_8$O$_8$(OH)$_8$Cl$_{1.34}$ and in addition spinel-phases of various compositions. Carbides, such as Fe$_3$C or chromium-rich iron carbides were visible within the corrosion layer.

4. Conclusion
Immersion tests at 60°C at ambient pressure and at 100 bar were performed in saline aquifer water (Stuttgart Aquifer and Northern German Basin) at 60°C using technical CO$_2$ to model the CCS-environment. According to DIN 6601 42CrMo4, X20Cr13 and X46Cr13 would be unsuitable for pressure vessel application, when being surrounded by the CO$_2$-saturated brine at ambient pressure. The following conclusions for the corrosion for corrosion phenomena of different steels may be stated:

i. Main corrosion products of a continuous corrosion scale as well as pits after static corrosion tests at ambient pressure as well as 100 bar were FeCO$_3$ and FeOOH for all steel qualities. Corrosion reactions are diffusion controlled with slower diffusion in the liquid phase leading to increased siderite crystal sizes than in the vapour/supercritical phase (smaller crystals due to higher diffusion rates).

ii. Corrosion rates obtained at 100 bar are generally much lower after 8000 h of exposure than those measured at ambient pressure (depending on the atmosphere (vapour or liquid) by a factor of ca. 10–80 for 42CrMo4, a factor of ca. 1.5–30 for X46Cr13, a factor of ca. 10–100 for X20Cr13 and a factor of ca. 10 for X35CrMo17-1, exception for X5CrNiCuNb16-4 with similar surface corrosion rates in the liquid phase).

iii. Highest surface corrosion rates are 0.8 mm/year for 42CrMo4 (1% Cr), 0.3 mm/year for X46Cr13 (13% Cr), 0.3 mm/year for X20Cr13 (13% Cr), 0.1 mm/year for X35CrMo17-1 (17% Cr) and 0.01 mm/year for X5CrNiCuNb16-4 (16% Cr) in the vapour phase at ambient pressure after 8000 h of exposure. At 100 bar the highest corrosion rates are 0.01 mm/year for 42CrMo4, X20Cr13 (liquid phase) X46Cr13 and less than 0.01 mm/year for X35CrMo4 and X5CrNiCuNb16-4 after 8000 h of exposure with no regard to atmosphere.

iv. The carbon content does not show significant influence on corrosion rate. But higher chromium content results in lower surface corrosion rates and therefore better corrosion resistance. Due to high Cr, Ni and Mo content X2CrNiMoN 22-5-3 shows now surface corrosion at all under the conditions given.
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