

The accuracy of calcium-carbonatebased saturation indices in predicting the corrosivity of hot brackish water towards mild steel

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Synopsis

Industry has always relied on water's inherent ability to inhibit mild steel corrosion by virtue of its levels of calcium hardness and total alkalinity. This research seeks to verify the application of this principle to brackish water used in industrial systems at moderately elevated temperatures. A brief review is first given of the conventional calcium-carbonate-based scale or corrosion predictive indices. Laboratory corrosion tests were performed at various levels of calcium hardness and total alkalinity, resulting in the generation of an empirically derived nonlinear regression model. The newly developed model and the existing indices were then compared statistically in predicting the corrosivity of brackish water in contact with mild steel at 45°C. The accuracy, broader application, and relevance of the indices are also discussed.

Keywords

saline water, brackish water, cooling, corrosion prediction, computer modelling, Langelier Saturation Index, Ryznar Stability Index, mild steel, calcium carbonate saturation.

Introduction

Between 1960 and the mid-1990s the Langelier Saturation Index (LSI) (Equation [1]) (Langelier, 1936) attracted adverse commentary from the municipal potable water community. It was stated that the LSI had no correlation with corrosion rate (Stumm, 1960; Larson and Sollo, 1967; Singley, 1981; Schock, 1984; Piron et al., 1986; Pisigan and Singley, 1987), and based on the empirical evidence it was suggested that the use of the LSI for corrosion prediction should be abandoned (AWWARF and DVGW, 1996). The same sentiment applied to the Ryznar Stability Index (RSI) (Ryznar, 1944) (Equation [2]) and the various other 'corrosion prediction indices' that emerged during this period and that were also grounded on the same principle:

- ► Momentary Excess (ME) (Dye, 1958)
- Calcium Carbonate Precipitation Potential (CCPP) (Merrill and Sanks, 1978)
- Aggressiveness Index (AI) (Millette *et al.*, 1980)
- ► Driving Force Index (DFI) (Rossum and Merrill, 1983).

LSI (Langelier Saturation Index) = $pH - pH_s$ [1] where the variables and their range of applicability are as follows:

pH: measured pH (7.0–9.5), temperature: 25–80°C, total dissolved solids <800 mg/l. pH_s: the pH at which the cooling water will be saturated. This value is calculated based on the solubility product for calcite, the second dissociation constant for carbonic acid and calcium concentrations, and total alkalinity of the cooling water Positive LSI values indicate 'oversaturation' and a tendency for a protective CaCO₃ coating to form, whereas negative values indicate the tendency to dissolve an existing CaCO₃ coating.

- RSI (Ryznar Stability Index) = $2(pH_s) pH$ [2]
- PSI (Puckorius Scaling Index) = $2(pH_{eq}) pH$ [3] Puckorius uses an equilibrium pH for this index rather than the actual cooling water pH. The index applies for temperatures up to 93°C.

pHeq = $1.465 \times \log 10$ (total alkalinity) + 4.54 [4]

Total alkalinity: mg/l CaCO₃. RSI or PSI values 6 and higher indicate increasingly severe corrosive tendencies, whereas values 6 and lower indicate more CaCO₃ scaling tendencies.

The PSI (Puckorius and Brooke, 1990), calculated using Equations [3] and [4], is a refinement of the RSI, in which an empirical alkalinity function is derived to modify the calculated pH of saturation for calcium carbonate (pH_s). In large industrial and power station cooling systems, either the LSI of PSI is still used to control acid feed or the cycles of concentration for calcium carbonate scale

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control. This supports their continued role in predicting the impact of water-soluble species on industrial water systems.

Early research (Larson and Skold, 1957) confirmed that the precipitation/dissolution of calcium carbonate is not the only water quality parameter relevant to the corrosion of distribution systems. Other factors such as the ratios of anions, flow velocity, pH, and calcium concentration also contribute to corrosion rates, but perhaps the best researched is the Larson and Skold Index, also known as the Larson Ratio (LR) (Larson and Skold, 1957, 1958) (Equation [5]). This index includes the corrosive effects attributable to the chloride and sulphate concentrations.

Larson-Skold Index, Larson ratio (LR) or Corrosivity Index (CI) = ([Cl-] + $[SO_4^{2^-}]) / ([HCO_3^-])$ [5]

where

 $[Cl^-]$, $[SO_4^{2^-}]$ and $[HCO_3^-]$: meq/litre of chloride, sulphate, and total alkalinity respectively.

The water studied approximated the quality of the Great Lakes waters of North America. The higher the index the more corrosive the water.

Feigenbaum *et al.* (1978) demonstrated poor correlation between the already-mentioned calcium-carbonate-based indices and the saline waters of the Negev Desert, and therefore developed an empirical index that included the effect of calcium carbonate solubility and the ions of the LR.

The reported lack of a definite correlation between the LSI and corrosion rates evident in both the drinking water industry and laboratory-scale closed-loop experiments prompted Pisigan and Singley (1984) to embark on a series of jar tests. The results of the laboratory tests permitted them to empirically derive a four-variable model (Equation [6]) and an eight-variable model (Equation [7]) that could account for 98% of the variations in corrosion rate under the experimental conditions explored. The equations hypothesized indicated that the corrosion rate of mild steel was in fact influenced by factors beyond just the precipitation or dissolution of calcium carbonate.

Pisigan and Singley 4-variable equation:

 $CR4 = ((TDS)^{0.253} (DO)^{0.820})/((10^{S1})^{0.0876} (Day)^{0.373}) [6]$ Pisigan and Singley 8-variable equation: $CR8 = (Cl)^{0.509} (SO_4)^{0.0249} (Alk)^{0.423} (DO)^{0.780} / ((Ca)^{0.676} (\beta)^{0.0304} (Day)^{0.381} (10^{S1})^{0.107}) [7]$

where

TDS (total dissolved solids): mg/l, Ca (calcium): mg/l as Ca, Mg (magnesium): mg/l as Mg, Na(sodium): mg/l as Na, Cl (chloride): mg/l as Cl, SO₄ (sulphate): mg/l as SO₄, Alk (alkalinity): mg/l as CaCO₃, β (buffer capacity): mg/l as CaCO₃/pH, DO (dissolved oxygen): mg/l as O₂. Picigan and Singley's eight variable model (1984)

Pisigan and Singley's eight-variable model (1984) suggests that increasing chloride, sulphate, alkalinity, and dissolved oxygen levels would accelerate corrosion, whereas increases in calcium concentration, buffer capacity, saturation index, and exposure time would lead to decreasing corrosion rates. In this hypothetical equation, the alkalinity is declared to accelerate rather than reduce corrosion, as is commonly known. The authors attributed this contradiction to the overwhelming influence of the increased ionic strength over the effect of alkalinity with increasing dosages of sodium bicarbonate while attempting to raise the alkalinity during the laboratory experiments. The first use of the buffer capacity (β) appeared in the literature pertaining to the subject of corrosion prediction in the work by Stumm (1960). Laboratory tests with synthetic solutions helped explain the mutual interaction of corrosion-stimulating and -inhibiting factors of natural waters, namely: pH, buffer capacity, CaCO₃ deposition, and alkalinity. This study was thought to at least partially explain the increase in corrosion rates with increasing pH between the values of 7.0 and 8.5.

Several studies found that as the pH approached 8.4, either from a higher or lower pH value, the corrosion rate of cast iron increased with decreasing buffer intensity. It is presumed that this effect occurs as a result of there being fewer but larger cathodic and anodic areas, thereby encouraging the electrochemical cell (Stumm, 1960).

Based on the independent studies reported by Pisigan and Singley (1987) and Imran *et al.* (2005a), it was possible to propose a modified Larson ratio (LRM) that would compensate for the increase in total dissolved solids with the increasing alkalinity by including the sodium ion concentration.

Imran continued his work on potable water distribution systems and published an article (Imran *et al.*, 2005a) that included a wider range of parameters in an empirically derived nonlinear model. The model is based on the change in apparent colour (Δ C in cpu) as a measure of corrosion in distribution lines, as it was found to be a reliable surrogate measurement of total iron. Calcium and pH were not deemed significant during the statistical modelling, because all tests were performed in waters stabilized for CaCO₃ solubility. Alkalinity was the only variable that could be effectively controlled by chemical addition.

In the arena of oilfield brines, where the high salinity affects the ionic strength and influences the calcium carbonate solubility, the Stiff-Davis Index (SDI) (Equation [8]) has been used (Stiff and Davis, 1952) in place of the Langelier Index. Waters with total dissolved solids levels higher than 4000 mg/l require that the SDI is used.

SDI = pH - pCa - pAlk - K [8] where

pH: pH measured, pCa = -log (Ca in mg/l as Ca^{2+}), and pAlk = -log (M alkalinity in mg/l as $CaCO_3$), K = constant based on the total ionic strength and temperature.

 $\begin{array}{l} I_{s} \left(0 \ \& T \ scaling \ Index \right) = log \ (T_{Ca} \ A_{Ik}) + pH - 2.78 + \\ 1.143 \times 10^{-2} \ T - 4.72 \times 10^{-6} \ T^{2} - 4.37 \times 10^{-5} \ P - 2.05 \times I^{1/2} + \\ 0.727 \times I \end{array}$

where

I (ionic strength): moles/l, T_{Ca} (calcium): moles/l as Ca, A_{lk} (alkalinity): moles/l as HCO_3^- , T (temperature): ${}^{0}F$, P (pressure): psi.

As with the Langelier Saturation Index, positive SDI or I_s values indicate 'oversaturation' and a tendency for a protective CaCO₃ coating to form, whereas negative values indicate the tendency to dissolve an existing CaCO₃ coating

Higher CR values indicate higher corrosion rates in mils per year (mpy).

The Stiff-Davis (Stiff and Davis, 1952) method is one of the easiest ways to calculate calcium carbonate scaling tendencies (Calcite Saturation Index) in brines and it is valid

for temperatures from 0–90°C and ionic strengths from 0–4. This index does not take into account the pressure and carbon dioxide concentration. It requires that the pH is measured on a fresh sample to avoid inaccuracies. As ionic strength and/or the temperature increase, so the K value decreases, resulting in a higher SDI, indicating a higher calcium carbonate scaling tendency. Higher concentrations of calcium or alkalinity would also lead to higher SDI values, also resulting in increased scaling tendencies. Calculating the SDI requires a calculation of the ionic strength, knowing the temperature of the operation, and looking up the K value in a K *versus* ionic strength graph (Stiff and Davis, 1952).

The Oddo-Tomson (1982) method is an alternative index applicable to high ionic strength waters for predicting the formation of calcium carbonate and various sulphate scales. It is valid between temperatures of 0–200°C, ionic strengths of 0–4.0, and pressures of 1–1380 bar (0–20000 psig) (Equation [9]). The calculation was reported by Oddo and Tomson (1982) to be accurate at high and low temperatures and pressures. The calculation can be easily performed in the field and is said to work well when applied to geopressured wells.

The prediction of the corrosivity of underground minewaters towards mild steel was also explored by White and Higginson (1985). It was reported that the corrosion takes place under cathodic control, with the metal acting as a substrate for the cathodic reaction. Thus the corrosivity of minewaters is largely dependent upon on the oxygen concentration and the pH of the water.

Equipment and methodology

In determining the relationship between the calcium hardness and alkalinity and the corrosion rate of mild steel in brackish water at elevated temperatures (35–45°C), numerous laboratory tests were conducted with synthetic solutions.

The main aim of the laboratory evaluation was to determine the impacts of temperature (between 35° C and 45° C), calcium hardness (between 50 mg/l and 100 mg/l as Ca²⁺), and total alkalinity (between 55 and 220 mg/l as CaCO₃) on the corrosivity of brackish water towards mild steel. Thus, the limiting conditions for the applicability of this study are waters having a quality range defined in Table I.

C1010 (mild steel) corrosion coupons were subjected to synthetic test solutions (4000 ml) stirred at 100–110 revolutions per minute (r/min) for 72 hours. The coupons were then removed, cleaned with a water wash to finger-

Table I

Key parameters and their target concentrations and ranges

Variable	Target values (range)
рН	7.8 (6.0–8.1)
Calcium (mg/l as Ca2+)	50–100 (49–97)
Total alkalinity (mg/l as CaCO ₃)	55–220 (19–228)
Magnesium (mg/l as Mg ²⁺)	27.3 (22–30)
Chloride (mg/l as Cl ⁻)	750 (717–822)
Sulphate (mg/l as SO ₄ ²⁻)	1125 (1100–1400)
Fluoride (mg/l as F ⁻)	10 (8–10)

touch, followed by an ethanol wipe; and then oven-dried, weighed and the corrosion rates calculated based on their weight loss. The methods followed were based on ASTM methods:

- G31-72 (Reapproved 1999): Standard Practice for Laboratory Immersion Corrosion Testing of Metals
- G1-90 (Reapproved 1999: Standard Practice for Preparing, Cleaning, and Evaluating Corrosion Test Specimens.

A Corrater[®] (Rohrback Cosasco) was used to measure the general corrosion rate and imbalance (*i.e.* indicator of the tendency for localized corrosion). The test solutions were also tested for their total iron concentrations and compared against the coupon method and Corrater[®] readings. Each set of tests was performed in a batch of six tests in a 'Laboratory Scale and Corrosion Test Station', a Buckman proprietary corrosion testing device, over a period of three days (refer to Figures 1 and 2).

Each of the six stations has its dedicated 5-litre beaker and an overhead paddle stirrer, dedicated temperature probe, hot plate, three coupon holders, and a Corrater® probe

Results

Table II lists the target calcium and total alkalinity concentrations, the temperatures explored, and the total iron concentrations of the test solutions. Table III summarizes the corrosion coupon results and the Corrater® readings taken over the three days.

Figure 3 compares the mild steel coupon corrosion rates (mm/a) for the two positions, and Figure 4 provides a threedimensional view of the correlations between the three corrosion measurements: the average coupon corrosion rate, the average Corrater[®] general corrosion rate, and the total iron concentrations.

Figure 5 reflects the impact of a 10° C difference in temperature on the corrosion rates as well as the impact of higher and lower alkalinity. It includes the error bars based on the maximum standard deviation – that is, a value of 0.07 mm/a – for the entire set of coupon data recorded during the 30 test runs.

Statistical correlations were performed between the average coupon corrosion rate and the various parameters

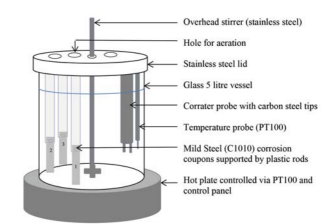
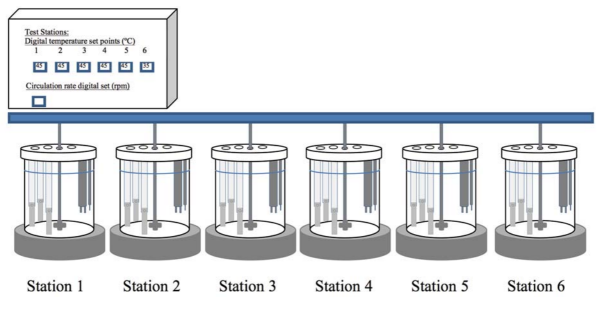


Figure 1 – Schematic diagram of Laboratory Scale and Corrosion Test Station



e 2 – Schematic of Laboratory Scale and Corrosion Test Station set-up

measured during the experiment. Statistically significant linear model correlations, with a 95% confidence level, were found between the average coupon corrosion rate and the following parameters: the initial and final calcium concentrations, the initial total alkalinity, the average Corrater® general corrosion rates, and the total iron levels. Inverse relationships were evident between the average coupon corrosion rate and the calcium and total alkalinity values,

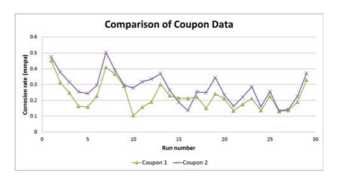


Figure 3 – Comparison of coupon data

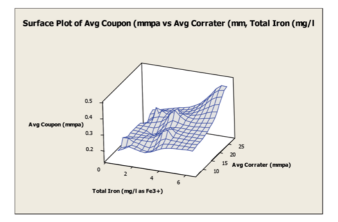


Figure 4 – Surface plot comparing the three corrosion measurement parameters: the average coupon rate (Avg Coupon mmpa) versus the average Corrater readings (mmpa) versus the total iron concentrations (mg/l as Fe²⁺)

Table II
Varying calcium and alkalinity: target values and

final total iron levels

Run						
	Calcium	Total alkalinity	Temperature			
	mg/l as Ca ²⁺	mg/l as CaCO ₃	°C	mg/l as Fe ³⁺ (total)		
1	50	55	45	6.4		
2	50	82.5	45	3.9		
3	50	110	45	4.3		
4	50	165	45	2.9		
5	50	220	45	2.7		
6	50	110	35	4.0		
7	62.5	55	45	6.1		
8	62.5	82.5	45	3.5		
9	62.5	110	45	3.0		
10	62.5	165	45	2.1		
11	62.5	220	45	2.0		
12	62.5	110	35	4.3		
13	75	55	45	1.7		
14	75	82.5	45	1.4		
15	75	110	45	1.2		
16	75	165	45	0.8		
17	75	220	45	1.4		
18	75	110	35	1.8		
19	87.5	55	45	3.3		
20	87.5	82.5	45	0.5		
21	87.5	110	45	2.9		
22	87.5	165	45	1.4		
23	87.5	220	45	0.7		
24	87.5	110	35	2.0		
25	100	55	45	2.8		
26	100	82.5	45	0.4		
27	100	110	45	2.0		
28	100	165	45	0.0		
29	100	220	45	0.0		
30	100	110	35	36.4		

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D	rrosion coupon and Corrater readings at varyious calcium and alkalinity levels										
Run	Corrosion coupon results			Corrater readings							
	Coupon 1 (mm/a)	Coupon 2 (mm/a)	Av. (mm/a)	Day 1 (mm/a)	Day 1 Imbal.	Day 2 (mm/a)	Day 2 Imbal.	Day 3 (mm/a)	Day 3 Imbal.	Day 4 (mm/a)	Day 4 Imbal
1	0.45	0.48	0.46	36.8	4.3	36.1	4.1	34	1.1	35.5	2.6
2	0.31	0.38	0.35	15.5	8.6	13.4	8.1	12.6	4.4	12.1	5.3
3	0.25	0.32	0.28	14	7.4	14.2	6.6	11.1	3.4	10.9	3.7
4	0.16	0.25	0.21	8.7	6.9	8.4	6.6	8.3	2.1	9.3	7.1
5	0.16	0.24	0.20	10.1	3.5	8.6	1.1	8.7	0.1	8.7	7.1
6	0.23	0.30	0.26	13	7.1	14.4	2.6	10.1	2.4	10.4	2.3
7	0.41	0.50	0.46	36.6	1.2	33.9	1.9	34.4	6.3	33.5	5.5
8	0.37	0.39	0.38	28.6	1.1	26.4	6.5	21.1	12.5	22.8	0.8
9	0.29	0.30	0.29	30.5	1.5	29.2	9.2	29.9	6.8	20.7	5.6
10	0.10	0.28	0.19	19.3	1.3	20.3	1.0	21.3	7.4	19.5	5.2
11	0.16	0.32	0.24	21.6	9.8	19.8	3.7	19.8	5.3	19.5	3.3
12	0.19	0.34	0.26	17.3	9.3	16.8	1.3	17.8	6.4	17.7	3.4
13	0.30	0.37	0.34	30.1	6.2	31.4	2.2	30.5	6.6	30.4	2.1
14	0.23	0.26	0.25	8.1	2.2	7.9	6.6	9.2	6.6	12.1	3.1
15	0.22	0.19	0.20	28.6	2.2	29.6	3.1	31.4	6.6	32.4	3.8
16	0.21	0.14	0.18	25.7	6.2	28.2	4.4	30.4	6.2	33.4	9.8
17	0.22	0.25	0.24	31.4	2.9	30.4	13.3	30.1	8.6	33.4	6.7
18	0.15	0.25	0.20	38.4	6.2	37.4	9.6	39.4	6.7	38.9	1.1
19	0.24	0.34	0.29	12.6	1.1	10.1	3.4	11.4	2.2	14.4	6.6
20	0.21	0.23	0.22	15.4	1.8	14.1	0.3	14.9	1.0	15.2	2.2
21	0.13	0.16	0.15	9.8	3.4	10.7	6.1	10.4	0.9	8.1	0.9
22	0.17	0.22	0.20	9.4	0.6	8.3	1.7	7.6	0.8	7.7	1.3
23	0.21	0.29	0.25	12.2	0.1	10.0	0.9	10.6	2.6	9.5	0.2
24	0.14	0.16	0.15	22.4	0.8	18.8	0.7	19.0	1.0	16.7	1.5
25	0.23	0.26	0.24	9.9	3.3	8.9	1.1	6.4	2.2	-	-
26	0.13	0.13	0.13	4.7	2.2	5.4	0.1	4.6	3.2	-	-
27	0.14	0.14	0.14	8.4	0.2	8.2	3.5	7.6	4.4	-	-
28	0.19	0.22	0.21	5.2	0.9	4.4	2.6	4.1	0.2	-	-
29	0.33	0.37	0.35	7.7	0.4	7.9	1.1	8.6	2.6	-	-
30	0.61	0.69	0.65	4.4	10.9	13.6	0.4	15.6	5.5	-	-

whereas direct moderate correlations were noted between the average coupon rates and both the average Corrater[®] general corrosion rates and the total iron concentrations. The average Corrater[®] general corrosion rates did not correlate well with the total iron values.

Contour plots (Figures 6 to 9) were drawn to reveal the combined effect of the calcium hardness and total alkalinity on the average coupon corrosion rate. The initial or final calcium concentrations and initial or final total alkalinity values were plotted against the average coupon rate.

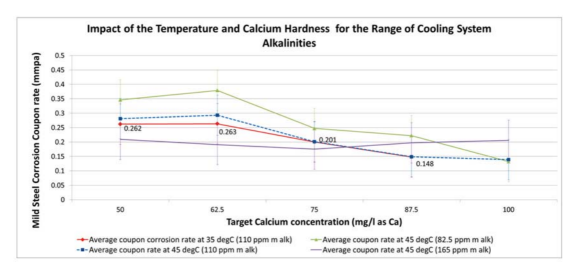


Figure 5 – Comparison of mild steel coupon data for the two temperatures (35°C and 45°C) at the same calcium hardness and alkalinity. Error bars: 0.07 mm/a (at max. std. deviation). Note: the tests containing high initial calcium concentration (above 75 mg/l as Ca) and high initial total alkalinity (165 mg/l as CaCO₃) produced visible quantities of white precipitate, confirmed by flame atomic absorption spectrometry and spot tests to be calcium carbonate. The test solutions also resulted in significantly reduced soluble calcium levels immediately after starting the tests

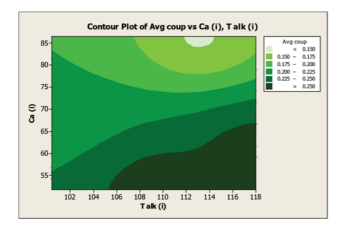


Figure 6 – Contour plot showing correlation of initial calcium hardness (Ca_i) and initial total alkalinity (T alki) on mild steel coupon corrosion (Avg coup) at 35° C

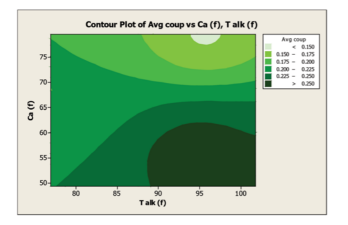


Figure 7 – Contour plot showing correlation of final calcium hardness (Ca_f) and final total alkalinity(T alk_f) on mild steel coupon corrosion at $35^{\circ}C$

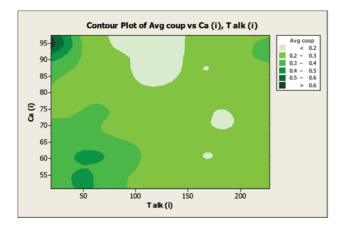


Figure 8 – Contour plot showing correlation of initial calcium hardness (Cai) and initial total alkalinity (T alk) on mild steel coupon corrosion at $45^{\circ}C$

A nonlinear multivariate regression analysis was conducted to predict the relationship between coupon corrosion rate and various predictors in order to determine

Contour Plot of Ava coup vs Ca (f) Talk (f)

Figure 9 – Contour plot showing correlation of final calcium hardness (Ca_t) and final total alkalinity (T alk_t) on mild steel coupon corrosion at 45° C

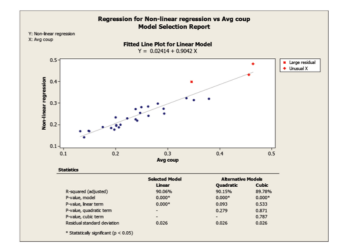


Figure 10 – Regression analysis for the derived nonlinear regression equation *versus* the average coupon rate (Avg coup)

how the response variable changes as the particular predictor variables change. This was done without taking into account the impact of the 10°C temperature difference (refer to Equation [10]):

Corrosion (mm/a) = 4.58E-6 Ca² - 8.85E-3 Ca + 4.64E-5 (Ca × M alk) - 9.30E-3 × M alk + 1.90E-5 M alk² + 1.26 [10]

where Ca = mg/l calcium as Ca and M alk = mg/l total alkalinity as CaCO₃.

Figure 10 serves to confirm the accuracy of the empirically derived nonlinear regression equation by comparing it to the laboratory coupon corrosion data. An R² adjusted value of 90.06% was obtained. The graph also indicates where large residuals and an unusual result occurred, explicitly at the upper section of curve.

The empirically derived equation was then compared statistically against the predictive indices discussed in the literature survey. This was performed by using the test solution target values and the comparison performed at both 35°C and 45°C. The results of the comparison are given in Table IV and shown by the scatter plot correlations in Figure 11. The statistical analysis indicated that the calculated rate

Table IV

Statistically significant relationships between the calculated rate and the other indices

	Calculated rate (mm/a)
Са	-0.408
	0.093
TOT Alk (SGO)	-0.344
	0.162
CR8 Corrosion In	-0.053
	0.833
Buffer Capacity	-0.344
	0.162
CR4 Corrosion In	0.529
	0.024
CCPP (mg/l)	-0.259
	0.300
ls (Oddo &Tomson, 1982)	-0.536
	0.022
onic Strength (M)	-0.531
	0.023
(Stiff and Davis,1952)	-0.570
	0.013
CaCO ₃	-0.180
	0.474
CaF ₂	0.176
	0.485
Langelier	-0.532
	0.023
RSI	0.526
	0.025
PSI	0.539
	0.021
Larson Skold	0.545
	0.019
CaCO ₃ FIME	-0.038
Note: the top number is the Pear	0.880

As a rule of thumb, r > 0.65 or R < 0.65 indicate correlation. The bottom number is the p-value. P-values ≤ 0.05 indicate correlation

at the 95% confidence level.

had the statistically significant moderately strong relationships at a 95% confidence level shown in Table V.

A contour plot (Figure 12) of the derived multivariate nonlinear regression equation [10]) was included to facilitate a visual comparison with the contour plots of the laboratory coupon corrosion data (Figures 6-9). Contour plots (Figures 12-19) of statistically significant indices taken from the literature, as per Table V, were also included for further comparisons.

Discussion

In order to first determine the correlation between the two coupon positions in the test vessel it was necessary to compare their results by means of a paired t-test. The paired t-test for the mean of coupon 1 versus the mean of coupon 2

Table V

Summary of the statistically significant correlations between the calculated corrosion rate and the established indices

Negative relationships	Positive relationships			
Is(Oddo), Oddo-Tomson	CR4, 4 variable model (Pisigan			
(1982) method	and Singley, 1984)			
Ionic strength	RSI, Ryznar Stability Index			
	(Ryznar, 1944)			
SDI, Stiff Davis Index	PSI, Puckorius or Practical			
(Stiff and Davis, 1952)	Scaling Index (Puckorius and			
	Brooke, 1990)			
LSI, Langelier Saturation Index	LR, Larson Skold Index, also			
(Langelier, 1936)	known as the Larson Ratio			
	(Larson and Skold, 1957, 1958).			

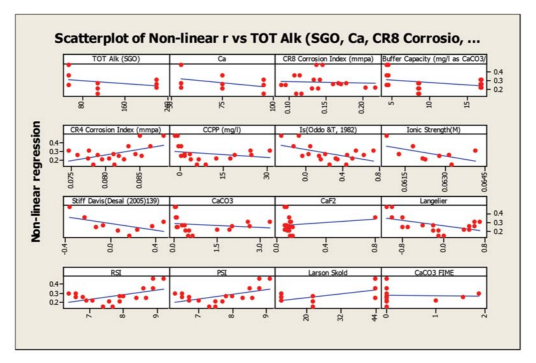


Figure 11 - Scatter plot correlations of the calculated corrosion rate based on the nonlinear regression equation versus the established indices

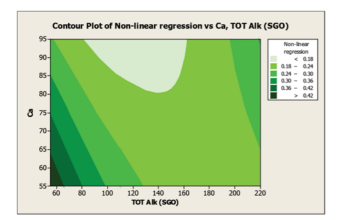


Figure 12 – Contour plot of the results of the multivariate nonlinear equation *versus* the targeted calcium (Ca) and total alkalinity concentrations (TOT Alk (SGO))

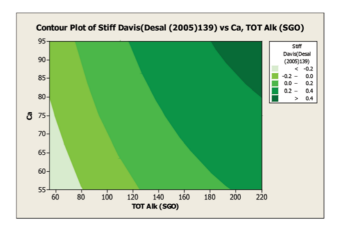


Figure 15 – Contour plot of the results of the Stiff and Davis Index (Stiff and Davis, 1952) *versus* the targeted calcium (Ca) and total alkalinity concentrations (TOT Alk (SGO))

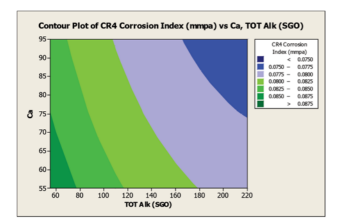


Figure 13 – Contour plot of the results of the C4 model (Pisigan and Singley, 1984) *versus* the targeted calcium (Ca) and total alkalinity concentrations (TOT Alk (SGO))

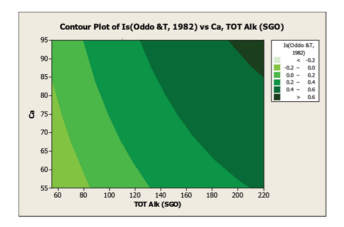


Figure 14 – Contour plot of the results of the Oddo and Tomson model (Oddo and Tomson, 1982) *versus* the targeted calcium (Ca) and total alkalinity concentrations (TOT Alk (SGO))

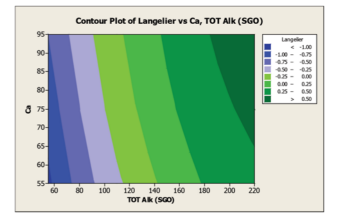


Figure 16 – Contour plot of the results of the Langelier Saturation Index (Langelier, 1936) *versus* the targeted calcium (Ca) and total alkalinity concentrations (TOT Alk (SGO))

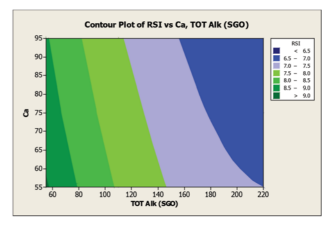


Figure 17 – Contour plot of the results of the Ryznar Stability Index (Ryznar, 1944) *versus* the targeted calcium (Ca) and total alkalinity concentrations (TOT Alk (SGO))

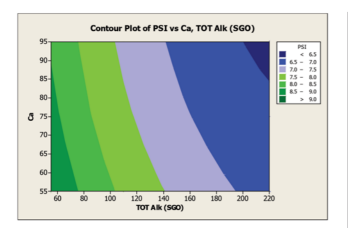


Figure 18 – Contour plot of the results of the Practical Scaling Index (Puckorius and Brooke, 1990) *versus* the targeted calcium (Ca) and total alkalinity concentrations (TOT Alk (SGO))

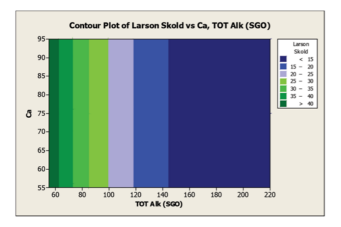


Figure 19 – Contour plot of the results of the Larson Skold Index ((Larson and Skold, 1957) *versus* the targeted calcium (Ca) and total alkalinity concentrations (TOT Alk (SGO))

showed that the two coupon positions were statistically different at a 95% confidence level. The two coupon positions did, however, demonstrate similar trends, as apparent in Figure 3; hence the average coupon rate was adopted as the result for each test.

A comparison of the plots of the average coupon rate against either the average Corrater® readings or the test solution total iron concentrations (Figure 4) demonstrated moderate direct correlations. The strength and direction of these correlations therefore supported the use of the average coupon rate for the continued corrosion studies. There was, however, a substantial discrepancy between the couponbased rates and the Corrater® readings, with the latter indicating rates that were approximately 54 times higher. The findings of this investigation were addressed by Van der Merwe and Palazzo (2015), and similar concerns over the lack of correlation between linear polarization resistance probe measurements and coupon immersion tests have also recently been investigated by Wu *et al.* (2015).

Figure 5 reflects the impact of a 10°C difference in temperature, where the higher temperature produced higher

corrosion rates at the lower targeted calcium hardness values of 50 and 62.5 mg/l (as Ca). At the higher calcium hardness values both temperatures resulted in similar corrosion rates. Figure 5 depicts the impact of different total alkalinities. It is apparent that higher alkalinity resulted in lower coupon corrosion rates, with the exception of the increased corrosion evident for the points corresponding to the combination of a high target calcium hardness (above 75 mg/l as Ca) and high target alkalinity (165 mg/l as CaCO₃).

The contour plots (Figures 6 and 7) for the average coupon rate *versus* calcium and total alkalinity for the corrosion tests performed at 35°C demonstrated:

- Lowest corrosion rates at high calcium (>70 mg/l as Ca) and high total alkalinity (>90 mg/l as CaCO₃)
- Highest corrosion rates at high total alkalinity (>90 mg/l as CaCO₃) and low calcium (<60 mg/l as Ca). The contour plots (Figures 8 and 9) for the average

coupon rate *versus* calcium and total alkalinity for the corrosion tests performed at 45°C demonstrated:

- ► Lowest corrosion rates at:
 - High calcium (80–100 mg/l as Ca) and moderately high total alkalinity (65–110 mg/l as CaCO₃)
 - Moderate calcium (60–80 mg/l as Ca) and moderate total alkalinity (150-200 mg/l as CaCO₃
- Highest corrosion rates at low total alkalinity (<50 mg/l as CaCO₃) and either low calcium (<50 mg/l as Ca) or high calcium (>85 mg/l as Ca).

The regions of either low or high corrosivity are similar for both the lower temperature of 35°C and higher temperature of 45°C.

The empirically derived nonlinear regression equation, based on only the solution's initial calcium and initial total alkalinity, was confirmed to account for 90.06% of the variations in the average corrosion coupon data.

Statistically significant moderately strong relationships, at a 95% confidence level, were evident between the derived nonlinear regression equation and the various established indices, namely: LSI, Langelier Saturation Index (Langelier, 1936), RSI, Ryznar Stability Index (Ryznar, 1944), LR, Larson Skold Index also known as the Larson Ratio (Larson and Skold, 1957, 1958), SDI, Stiff Davis Index (Stiff and Davis, 1952), I_s (Oddo), Oddo-Tomson (1982) method, ionic strength, CR4, four-variable model (Pisigan and Singley, 1984), and PSI, Puckorius, or Practical Scaling Index (Puckorius and Brooke, 1990).

A comparison of the contour plot of the new model *versus* those of other indices, for the set of target values considered revealed some subtle differences between them. One of the more obvious differences was the impact of the calcium concentration on the corrosivity, as evident by the relatively steeper diagonal contour lines for some of the indices, particularly in the new model, LSI, RSI, and PSI (Figures 12, 16, 17, and 18, respectively). As per the literature, it appears that a number of the authors included calcium in their empirically derived predictive models: LSI, RSI, SDI, and CCPP. The same can be said of pH for the LSI, RSI, PSI, SDI, and I_s indices. Linear regression analysis of the correlation between the average coupon corrosion rates and the various predictors confirmed the role of the initial pH and the initial calcium concentration.

Conclusion

Although the average coupon rate, the total iron concentrations, and the Corrater[®] general corrosion rate readings appeared closely related, it was possible to find statistically significant linear correlations, at a 95% confidence level, only between the average coupon rate and total iron concentration as well as the average coupon rate and the Corrater[®] general corrosion readings.

It was also confirmed that raising the calcium hardness and/or total alkalinity within the range of chemistries explored for the application of the brackish water in a cooling system does reduce mild steel corrosion. It did, however, become apparent that calcium carbonate saturation or supersaturation can lead to precipitation, resulting in reduced levels of calcium and alkalinity which in turn leads to increased corrosion rates.

The empirically derived equation was at best capable of only moderately strong correlations with any of the numerous calcium-carbonate-based models found in the literature. A comparison of the formulae of the statistically significant eight indices and a comparison to the remaining six non-statistically significant and weakly correlated indices revealed that the latter six were linearly related to either the calcium concentration or total alkalinity, whereas the former eight indices are a function of either the log or the inverse of either the calcium concentration and/or total alkalinity. The newly proposed empirically derived nonlinear regression model differes from both sets of indices in that it is based on a quadratic equation incorporating both the calcium concentration as Ca²⁺ and total alkalinity in mg/l as CaCO₃.

As stated in the introduction, it would be grossly inaccurate to base the calculation of the corrosivity of brackish water solely on its calcium concentration, total alkalinity, and temperature without giving any consideration to the many other factors or conditions prevailing in a typical industrial system. What is, however, apparent from the study, since it is based only on changes in the calcium hardness and total alkalinity, is that the empirically derived model may prove more accurate than most of the existing commonly found indices at estimating the corrosivity of brackish water, of similar characteristics to the chemistry explored, on mild steel between 35 and 45°C.

References

AWWARF and DVGW-TZW. 1996. Internal Corrosion of Water Distribution Systems. 2nd edn. American Water Works Association, Denver, CO. pp. 29–70.

Dye, J.F. 1958. Correlation of the two principle methods of calculating the three kinds of alkalinity. *Journal of the American Water Works Association*, vol. 50, no. 6. pp. 801–820.

- FEIGENBAUM, C., Gal-Or, L., and Yahalom, J. 1978. Scale protection criteria in natural waters. *Corrosion*, vol. 34, no. 4. pp. 133–137.
- IMRAN, S.A., DIETZ, J.D., MUTOTI, G., TAYLOR, J.S., and RANDALL, A.A. 2005a. Modified Larsons ratio incorporating temperature, water age, and electroneutrality effects on red water release. *Canadian Journal of Environmental Engineering*, vol. 131, no. 11. pp. 1514–1520.
- IMRAN, S.A., DIETZ, J.D., MUTOTI, G., TAYLOR, J.S., RANDALL, A.A., and COOPER, C.D. (2005b). Red water release in drinking water distribution systems. *Journal* of the American Water Works Association, vol. 97, no. 9. pp. 93–100.

- LANGELIER, W.F. 1936. The analytical control of anti-corrosion water treatment. *Journal of the American Water Works Association*, vol. 28, no. 10. pp. 1500–1521.
- LARSON, T.E. and SKOLD, R.V. 1957. Corrosion and tuberculation of cast iron. *Journal of the American Water Works Association*, vol. 49, no. 10. pp. 1294–1302.
- LARSON, T.E. and SKOLD, R.V. 1958. Laboratory studies relating mineral water quality of water to corrosion of steel and cast iron. *Corrosion*, vol. 14. pp. 285–288.
- LARSON, T.E. and SOLLO Jr., F.W. 1967. Loss in water main carrying capacity. *Journal of the American Water Works Association*, vol. 59, no. 12. pp. 1565–1572.
- MERRILL, D.T. and SANKS, R.L. 1978. Corrosion control by deposition of CaCO₃ films: Part 3, a practical approach for plant operators. *Journal of the American Water Works Association*, vol. 70, no. 1. pp. 12–18.
- MILLETTE, J.R., HAMMONDS, A.F., PANSING, F.J., HANSEN, C.E., and CLARK, P.J. 1980. Aggressive water: assessing the extent of the problem. *Journal of the American Water Works Association*, vol. 72, no. 5. pp. 262–266.
- ODDO, J.E. and TOMSON, M.B. 1982. Simplified calculation of CaCO₃ saturation at high temperatures and pressures in brine solutions. *Journal of Petroleum Technology*, July. pp. 1583–1590.
- PIRON, D.L., DESJARDINS, R., BRIERE, F., and ISMAEL, M. 1986. Corrosion rate of cast iron and copper pipe by drinking water. *Corrosion Monitoring in Industrial Plants Using Nondestructive Testing and Electrochemical Methods, ASTM STP 908.* Morgan, G.C. and Labine, P. (eds). American Society for Testing and Materials, Philadelphia.
- PISIGAN Jr., R.A. and SINGLEY, J.E. 1987. Influence of buffer capacity, chlorine residual, and flow rate on corrosion of mild steel and copper. *Journal of the American Water Works Association*, vol. 79, no. 2, pp. 62–70.
- PISIGAN, R.A. and SINGLEY, J. E. 1984. Evaluation of the corrosivity using the Langelier Index and relative corrosion rate models. *Material Performance*, vol. 24, no. 4. pp. 26–36.
- PUCKORIUS, P.R. and BROOKE, J.M. 1991. A new practical index for calcium carbonate scale prediction in cooling tower systems. *Corrosion*, vol. 47, no. 4. pp. 280–284.
- ROSSUM, J.R. and MERRILL, D.T. 1983. An evaluation of the calcium carbonate saturation indexes. *Journal of the American Water Works Association*, vol. 75, no. 2. pp. 95–100.
- RYZNAR, J.W. 1944. A new index for determining the amount of calcium carbonate scale formed by a water. *Journal of the American Water Works Association*, vol. 36, no. 4. pp. 472–475.
- SCHOCK, M.R. 1984. Temperature and ionic strength corrections to the Langelier Index revisited. *Journal of the American Water Works Association*, vol. 76, no. 8. pp. 72–76.
- SINGLEY, J.E. 1981. The search for a corrosion index. Journal of the American Water Works Association, vol. 73, no. 11. pp. 579–582.
- STIFF, Jr., H.A. and DAVIS, L.E. 1952. A method for predicting the tendency of oil field water to deposit calcium carbonate. *Petroleom Transactions of AIME*, vol. 195. p. 213.
- STUMM, W. 1960. Investigation of the corrosive behavior of waters. *Journal of the American Society of Civil Engineers, Sanitary Engineering Division*, vol. 86, no. SA6. pp. 27–45.
- VAN DER MERWE, J.W., and PALAZZO, A. 2015. Investigating the correlation between linear polarization resistance corrosion monitoring probe readings and immersion test results for typical cooling water conditions. *Journal of the Southern African Institute of Mining and Metallurgy*, vol. 115, no. 3. pp. 173–178.
- WHITE, R.T. and Higginson, A. 1985. Factors affecting the corrosivity of underground minewaters. *Proceedings of Mintek 50: International Conference on Mineral Science and Technology*, Sandton, South Africa.
- WU, J.W, BAI, D., BAKER, A.P., LI, Z.H., and LIU, X.B. 2015. Electrochemical techniques correlation study of on-line corrosion monitoring probes. *Material Corrosion*, vol. 66, no. 2. pp. 143–151.