RSC Advances



PAPER

View Article Online

View Journal | View Issue



Cite this: RSC Adv., 2018, 8, 19208

Correlation between the corrosion rate and electrochemical noise energy of copper in chloride electrolyte

Chenxi Yi, 10 * Xiaoqing Du, Yumeng Yang, Benfeng Zhu and Zhao Zhang

An electrochemical noise technique has been applied to describe the corrosion process of copper. The results show that the sampling frequency clearly changes both the energy distribution plot and the power spectral density spectra, which should be taken into consideration strictly and logically before an electrochemical noise test. The corrosion energy, (E_c), deduced using the fast wavelet transform method showed a similar variation trend with corrosion rate. Hence, the proposed parameter E_c represents the corrosion rate or severity.

Received 7th February 2018 Accepted 29th April 2018

DOI: 10.1039/c8ra01203b

rsc.li/rsc-advances

Introduction

Electrochemical noise (EN) is the generic term given to the fluctuation of current and potential, and corrosion engineering is regarded as one of its important fields of application.¹⁻⁶ Fluctuations of electrode potential are connected to local anodic and cathodic reactions and are a consequence of events occurring at flaws (such as pitting nucleation, propagation, and re-passivation), which are strictly related to the corrosion process.⁷

From a physicochemical viewpoint, the electrode potential is defined as the change in Gibbs energy when a charged particle transfers from the infinite into an electrode, including both the electrical and chemical work done during the process, whereas, potential only consists of the electrical work in the above transfer process. Therefore, the variation in the electrode potential definitely comes from the energy exchange between the electrode system and the environment.8 Additionally, the energy is divided into potential energy and kinetic energy, and both of them can be converted into each other. Potential energy is "inert" and only reflects the stability of objects, meanwhile kinetic energy is "active" and directly depends on velocity. Similarly,9-11 the fluctuation of electrode potential always simultaneously consists of "slow DC drift" and a "fast random non-equilibrium fluctuation signal". The former is the traditional electrode potential, which indicates the thermodynamic stability; whilst the latter is designated as electrochemical noise, which represents the speed of the electrode reaction.

EN data are usually analyzed using FFT (fast Fourier transform) and MEM (maximum entropy method) techniques to obtain PSD (power spectral density) plots, 3,12 or using the FWT (fast wavelet transform) technique to obtain an EDP (energy

Department of Chemistry, Zhejiang University, Hangzhou, Zhejiang 310027, China. E-mail: yichenxi2013@163.com

$$(C_1^j, C_2^j) = (2^j \Delta t, 2^{j-1} \Delta t)$$
 (1)

The mainstream EN practices $^{17-20}$ establish the relationship between the EDP and corrosion morphology. However, the scale range is dependent on Δt ($\Delta t = 1/f$) and the features of the EDP may vary with different Δt even for the same corrosion morphology. Therefore, when obtaining an EDP, using a selection of adequate sampling frequencies (f) is logical.

While investigating the corrosion behaviour of mild steels in saturated calcium hydroxide with 20 g L⁻¹ CaCl₂, Searson and Dawson¹⁵ found that there existed a relationship between the corrosion rate (r_{corr}) (obtained from weight loss) and the standard deviation of the potential noise with a correlation of 10^{-5} , *i.e.*, standard deviation \times $10^{-5} = r_{corr}$ (mpy). Although the comparison of short-term EN (1024 s in their work) with long-term weight loss (an average of several days) seems inappropriate, their pioneering work undoubtedly demonstrates a correlation between EN features and the corrosion rate.

Encouraged by Searson and Dawson,¹⁵ and based on previous reports^{21,22} that pitting definitely occurred when copper was immersed into chloride solutions, this study was devoted to finding the influence of the adopted EN sampling

frequency on the characteristics of both EDP and PSD plots, which are obtained from MEM and FWT analyses of the same electrochemical potential noise, and especially to probing the correlation between the corrosion rate and the electrochemical noise energy.

Theory of EN analysis

EN can be analysed using FFT, MEM and FWT techniques. When compared with FFT or MEM techniques, FWT can remove the DC trend without any observed signal distortion. Therefore, the conclusions obtained using FWT are more rational. Moreover, the EN energy (E) of each wavelet crystal of a different time-scale can be obtained using the FWT technique. After removing the energy contribution of DC drift, the mainstream RP-EDP plot could be obtained from E_i^D .

$$E = \sum_{n=1}^{N} S_n^2 \quad (n = 1, 2, ..., N)$$
 (2)

$$E_{Dj} = \frac{1}{E} \sum_{k=1}^{N/2^j} D_{j,k}^2, \quad (j = 1, 2, ..., J)$$
 (3)

$$E_j^{\ D} = \sum_{k=1}^{N/2^j} D_{j,k}^2, \quad (j=1,2,...,J)$$
 (4)

where E_j^D is the energy value of the D_j crystal with units of V, and E_{Dj} is the value after unitization. Generally, the E_j^D of a specific timescale (or frequency range) can be related to a certain corrosion process. For example, the metastable pitting/nucleation process always takes place prior to the nuclei growth process, both of which are much faster than the diffusion process, which therefore results in the large E_j^D of low j values. He should be noted that the timescale range of D_j is strictly dependent on Δt (or frequency) (eqn (1)), therefore each D_j partition of the RP-EDP plot will change along with the variation of f.

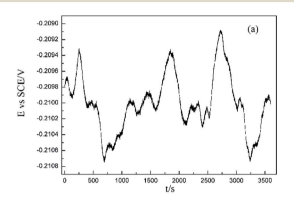
Experimental

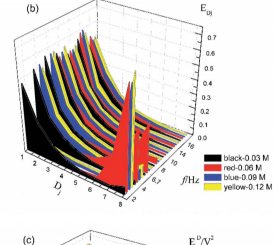
Corrosive electrolytes (NaCl at different concentrations, 0.03 M, 0.06 M, 0.09 M and 0.12 M, and 0.06 M HCl) were prepared from analytical grade NaCl or HCl and double distilled water. The working electrode (pure copper) was mechanically cut and embedded into Teflon, leaving an exposed area of 0.5 cm² as the working surface. Prior to each experiment, the working surface was successively abraded using sand paper from 400 to 1200 grit, polished to a mirror with 2.5 μ m diamond paste, rinsed with distilled water in an ultrasonic cleaner for about 3 min (KQ5200B, Youyi Instrument Co., Ltd., China), degreased with acetone and finally dried under a cool N₂ flow.

EN was monitored as a function of time between the working electrode and the reference electrode (SCE) for the $\mathbf{1}^{\text{st}}$ hour, using the Powerlab/4sp (made in Australia) electrochemical interface through a GP amplifier controlled by Chart 5 software for the Windows XP operating system. This instrument is equipped with analog/hardware filters including an AA filter

(anti-aliasing low-pass filters) to remove high frequency components before the signal is digitized, so that the acquisition of false data can be avoided.¹¹ The EN records were collected at different sampling intervals (sampling frequency, *f*).

The weight loss of copper was measured according to the standard ISO 8407:2009, IDT. Copper coupons with dimensions of 100 mm \times 80 mm \times 0.2 mm were sectioned, and a water bath was used to maintain a specific temperature. SEM (SIRION, FEI Company, made in Holland) was utilized to examine the corrosion morphologies of the specimens.





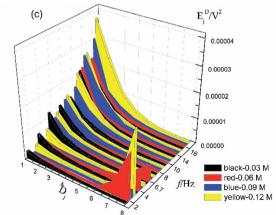


Fig. 1 (a) Typical E-t curve at a frequency of 2 Hz, (b) E_{Dj} map (with unitization) and (c) E_j^D map with varied sampling frequencies and $C_{\rm NaCl}$: black-0.03 M, red-0.06 M, blue-0.09 M and yellow-0.12 M.

RSC Advances Paper

Results and discussion

Influence of EN testing frequency

Fig. 1a shows the typical time-domain potential noise of the 1st hour for Cu corroding in 0.06 M NaCl neutral solutions at 20 °C, acquired at a commonly used data-sampling frequency (f) of 2 Hz. Fig. 1b and c show the RP-EDP plots of the noise generated during the same corrosion process at various concentrations of NaCl with different f(T=20 °C). The difference is that the data shown in Fig. 1b were obtained after unitization (eqn (3)) whereas the data shown in Fig. 1c were obtained without unitization (eqn (4)). For 0.06 M, 0.09 M and 0.12 M NaCl (in Fig. 1b), the RP-EDPs remain the same and the maximum (relative) energy is in D_1 – D_3 when $f \le 8$ Hz while it is in D_7 – D_8 when $f \le 4$ Hz. In 0.03 M NaCl, the maximum energy is in D_1 – D_3 at each testing frequency.

Fig. 2 shows the RP-EDP plots with and without unitization for Cu corroding in 0.06 M HCl solutions. The maximum energy of RP-EDP is in D_7 – D_8 when f < 4 Hz, while it is in D_1 – D_3 when f \geq 4 Hz. Considering Fig. 1 and 2, the characteristics of the RP-EDP plot are significantly influenced by sampling frequency. With increasing f, the maximum relative energy of the RP-EDP is in D_1 – D_3 regardless of the corrosion ionic strengths.

Fig. 3 shows the PSD analysed using the MEM technique in different concentrations of NaCl and Fig. 4 shows the PSD in 0.06 M HCl. It is evident that PSD parameters $(f_c, W, \text{ and } k)$ are

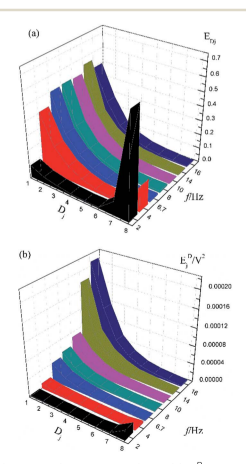
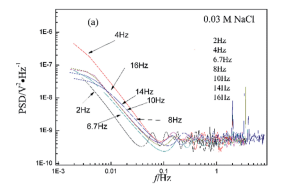
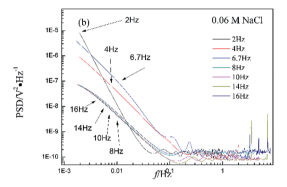
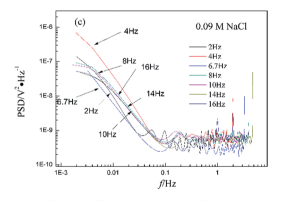


Fig. 2 (a) E_{Dj} map (with unitization) and (b) E_j^D map at different sampling frequencies in 0.06 M HCl solution.







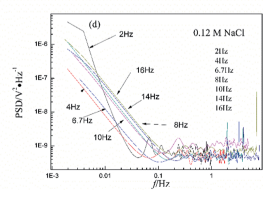
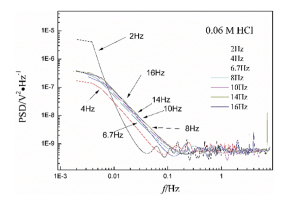


Fig. 3 Influence of frequency effects on the PSD spectra in different concentrations of NaCl.

also dependent on the adopted EN sampling frequency. In 0.06 M, 0.09 M and 0.12 M NaCl, these PSD parameters retain approximately equal values while $f \ge 8$ Hz, whereas others (<8

RSC Advances Paper



Influence of frequency effects on the PSD spectra in 0.06 M HCl.

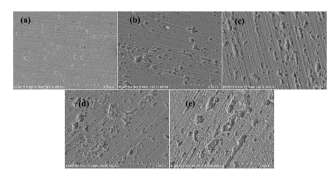


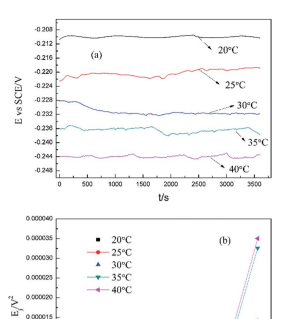
Fig. 5 SEM images of the Cu morphologies after corroding for 1 h in 0.06 M NaCl solutions at different temperatures: (a) 20 °C, (b) 25 °C, (c) 30 °C, (d) 35 °C and (e) 40 °C.

Hz) are distinct from each other. Meanwhile the turning frequency point is 6.7 Hz in 0.03 M NaCl and 0.06 M HCl. These results are in accordance with the RP-EDP plots (Fig. 1 and 2), and also demonstrate the significant influence of the sampling frequency on the EN results.

From a physicochemical viewpoint, the corrosion process should mainly depend on the natures of both the material (such as resistivity) and the environment (such as ambient temperature and erosive particles), but independent of the testing tools (such as the adopted EN sampling frequency). Additionally, the optimal or appropriate EN sampling frequency may be related to the investigated materials and their environment, and also possibly the research target of the researchers. Considering Fig. 1-4, the optimal testing frequency should be 6.7 Hz in 0.03 M NaCl and 0.06 M HCl and 8 Hz in 0.06 M, 0.09 M and 0.12 M NaCl. Previous reports14,15,29,30 claimed that an EN sampling frequency of 1-4 Hz ($\Delta t = 0.25$ -1 s) seems to be adequate, a reason for which may be that their efforts focused on the noise resistance deduced from PSD spectra of potential and current noise.

Relationship between EN features and corrosion rate

Based on the above analyses and inspired by Searson and Dawson,15 the corroded morphologies (Fig. 5), the weight loss



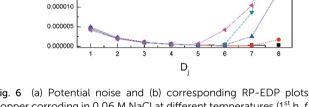


Fig. 6 (a) Potential noise and (b) corresponding RP-EDP plots for copper corroding in 0.06 M NaCl at different temperatures (1st h, f = 8

and the potential noise (Fig. 6a) of Cu corroding in 0.06 M NaCl neutral solutions at different temperatures were acquired, and the RP-EDPs without unitization were obtained (Fig. 6b). When performing EN tests, f = 8 Hz was adopted according to previous sections. Additionally, in order to ensure the comparability between the weight loss and the EN features, the same initial 1 h corroding period was selected as the timescale for both.

Fig. 5 clearly shows the pitting corrosion type, and the diameter and depth of the pits increase with temperature, which intuitively represents an enhancement of the corrosion rate. Since the passivation time of a metastable pit is typically

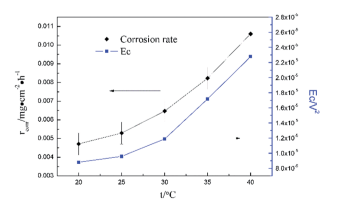


Fig. 7 Dependence of corrosion energy E_c and r_{corr} on temperature in 0.06 M NaCl.

no longer than 20 s,³¹ and D_8 mainly reflects the information of diffusion at f=8 Hz,³² the sum of the energy (Fig. 6b) deposited in crystals $D_1 \sim D_7$ (E_c), which represents the active pitting energy and therefore should reflect the severity of the corrosion of copper,^{8,33} is calculated and plotted *versus* the temperature (Fig. 7).

$$E_{c} = E_{1}^{D} + E_{2}^{D} + E_{3}^{D} + E_{4}^{D} + E_{5}^{D} + E_{6}^{D} + E_{7}^{D}$$
 (5)

 $r_{\rm corr}$ (units mg cm⁻² h⁻¹) obtained from the weight loss is also repeatedly plotted in Fig. 7, and both $r_{\rm corr}$ and $E_{\rm c}$ increase with the elevation of the temperature. In other words, a good parallel relationship between the $r_{\rm corr}$ and $E_{\rm c}$ can be confirmed during the investigated timescale.

Conclusions

The electrochemical noise technique has been applied to describe the corrosion process of copper. The results show that the sampling frequency clearly changes both the EDP and the PSD spectra, before establishing the relation between the EDP and surface morphology, or the PSD spectra and corrosion kinetics, the optimal testing frequency should be taken into consideration strictly and logically.

The corrosion energy, E_c , deduced from the FWT method, shows a similar variation trend with the corrosion rate. Hence, electrochemical noise offers a nondestructive on-line monitoring process, which can be easily carried out, and the proposed parameter E_c represents the corrosion rate or severity.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors wish to acknowledge financial support from the National Natural Science Foundation of China (Project 21273199 and 51131005).

Notes and references

- 1 D. H. Xia, S. Z. Song and Y. Behnamian, *Corros. Eng., Sci. Technol.*, 2016, 51, 527–544.
- 2 F. H. Cao, Z. Zhang, J. X. Su, Y. Y. Shi and J. Q. Zhang, *Electrochim. Acta*, 2006, **51**, 1359–1364.
- 3 T. Haruna, Y. Morikawa, S. Fujimoto and T. Shibata, *Corros. Sci.*, 2003, **45**, 2093–2104.
- 4 D. Xia, S. Song, J. Wang, J. Shi, H. Bi and Z. Gao, *Electrochem. Commun.*, 2012, **15**, 88–92.
- 5 A. Aballe, M. Bethencourt, F. J. Botana and M. Marcos, *Electrochem. Commun.*, 1999, 1, 266–270.
- 6 F. Zou and F. B. Cegla, *Electrochem. Commun.*, 2017, 82, 134–138.

- 7 C. Monticelli, G. Brunoro, A. Frignani and G. Trabanelli, *J. Electrochem. Soc.*, 1992, 139, 706–711.
- 8 Y. Chen, Z. N. Yang, Y. W. Liu, H. H. Zhang, J. Y. Yin, Y. Xie and Z. Zhang, *J. Taiwan Inst. Chem. Eng.*, 2017, **80**, 908–914.
- 9 A. Aballe, M. Bethencourt, F. J. Botana and M. Marcos, *Electrochim. Acta*, 1999, 44, 4805–4816.
- 10 A. M. Homborg, T. Tinga, X. Zhang, E. P. M. van Westing, P. J. Oonincx, G. M. Ferrari, J. H. W. de Wit and J. M. C. Mol, *Electrochim. Acta*, 2013, 104, 84–93.
- 11 E. García-Ochoa and F. Corvo, *Electrochem. Commun.*, 2010, 12, 826–830.
- 12 F. Mansfeld, L. T. Han, C. C. Lee, C. Chen, G. Zhang and H. Xiao, *Corros. Sci.*, 1997, 39, 255–279.
- 13 F. Mansfeld and H. Xiao, *J. Electrochem. Soc.*, 1993, **140**, 2205–2209.
- 14 T. Schauer, H. Greisiger and L. Dulog, *Electrochim. Acta*, 1998, **43**, 2423–2433.
- 15 P. C. Searson and J. L. Dawson, J. Electrochem. Soc., 1988, 135, 1908–1915.
- 16 X. Huang, Y. Chen, T. Fu, Z. Zhang and J. Zhang, *J. Electrochem. Soc.*, 2013, **160**, D530–D537.
- 17 L.-j. Zhang, X.-b. Zhu, Z. Zhang and J.-q. Zhang, *Trans. Nonferrous Met. Soc. China*, 2009, **19**, 496–503.
- 18 Z. Zhang, W. H. Leng, Q. Y. Cai, F. H. Cao and J. Q. Zhang, *J. Electroanal. Chem.*, 2005, **578**, 357–367.
- 19 B. Zhao, J.-H. Li, R.-G. Hu, R.-G. Du and C.-J. Lin, *Electrochim. Acta*, 2007, **52**, 3976–3984.
- 20 X. Liu, T. Zhang, Y. Shao, G. Meng and F. Wang, *Corros. Sci.*, 2010, 52, 892–900.
- 21 Y. Chen, D. Qi, H. Wang, Z. Xu, C. Yi and Z. Zhang, *Int. J. Electrochem. Sci.*, 2015, **10**, 9056–9072.
- 22 W. Qafsaoui, M. W. Kendig, S. Joiret, H. Perrot and H. Takenouti, *Corros. Sci.*, 2016, **106**, 96–107.
- 23 Z. Dong, X. Guo, J. Zheng and L. Xu, *Electrochem. Commun.*, 2001, 3, 561–565.
- 24 Z.-n. Yang, Z. Zhang, W.-h. Leng, K. Ling and J.-q. Zhang, Trans. Nonferrous Met. Soc. China, 2006, 16, 209–216.
- 25 J. Smulko, K. Darowicki and A. Zieliński, *Electrochem. Commun.*, 2002, 4, 388–391.
- 26 C. Cai, Z. Zhang, F. Cao, Z. Gao, J. Zhang and C. Cao, *J. Electroanal. Chem.*, 2005, **578**, 143–150.
- 27 S. M. Hoseinieh, A. M. Homborg, T. Shahrabi, J. M. C. Mol and B. Ramezanzadeh, *Electrochim. Acta*, 2016, 217, 226–241.
- 28 M. C. Lefebvre and B. E. Conway, *J. Electroanal. Chem.*, 2000, **480**, 46–58.
- 29 A. Nagiub and F. Mansfeld, Corros. Sci., 2001, 43, 2147-2171.
- 30 A. M. Homborg, T. Tinga, X. Zhang, E. P. M. van Westing, P. J. Oonincx, J. H. W. de Wit and J. M. C. Mol, *Electrochim. Acta*, 2012, 70, 199–209.
- 31 M. Shahidi, S. M. A. Hosseini and A. H. Jafari, *Electrochim. Acta*, 2011, **56**, 9986–9997.
- 32 Y. Y. Shi, Z. Zhang, F. H. Cao and J. Q. Zhang, *Electrochim. Acta*, 2008, **53**, 2688–2698.
- 33 C. Yi, X. Du, Y. Yang, Y. Chen, G. Wei, Z. Yang and Z. Zhang, *Int. J. Electrochem. Sci.*, 2017, **12**, 3597–3613.