

# Current state of electrochemical techniques and corrosion rate analysis for next-generation materials Junsoo Han, Oumaïma Gharbi

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Current state of electrochemical techniques and corrosion rate analysis for nextgeneration materials

Junsoo Han, Oumaïma Gharbi

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1	Current state of electrochemical techniques and corrosion rate analysis for next-generation		
2	materials		
3			
4	<sup>a</sup> Junsoo HAN, <sup>b</sup> Oumaïma GHARBI		
5	<sup>a</sup> Sorbonne Université, Laboratoire Interfaces et Systèmes Electrochimiques, LISE, F-75005		
6	Paris, France		
7	<sup>b</sup> Sorbonne Université, Laboratoire de Réactivité de Surface, LRS, F-75005 Paris, France		
8	Keywords: Corrosion rate; electrochemical techniques; new generation alloys		
9			
10	Abstract		
11	Current theoretical models and electrochemical techniques used to investigate corrosion mechanisms,		
12	including corrosion rates, have been tailored for conventional alloy systems. However, the application of		
13	conventional theories and techniques toward next-generation alloys such as multi-principal element alloys		
14	and additively manufactured alloys needs to be revisited due to the increased chemical complexity and		
15	refined microstructures of these alloys, which may yield different electrochemical properties from the		
16	conventional alloys. This review aims to discuss to which extent the current models and techniques used in		
17	corrosion science can be applied to these new alloy systems, and outline some of the challenges that need		

18 to be overcome to accurately describe their electrochemical reactivity.

### 19 Introduction

Since the Bronze Age, humanity has constantly developed new technological strategies and fabrication processes to produce alloys. The conventional alloy making procedure is based on adding minor alloying elements to one principal element to obtain the desired mechanical properties. In the early 2000s, the materials science community witnessed an important shift in the alloy making process with the emergence of two new alloy generations; namely multi-principal element alloys (MPEAs) [<sup>1</sup>, <sup>2</sup>, <sup>3</sup>] and additively manufactured (AM) alloys described in **Fig. 1**. [<sup>4</sup>, <sup>5</sup>].

In the light of recent technical advances, MPEAs are considered as promising materials that could be used in highly corrosive environments due to their superior aqueous corrosion resistance to conventional alloys such as stainless steel [<sup>6</sup>], as well as enhanced thermal resistance and irradiation-induced corrosion [<sup>7</sup>]. Conversely, AM alloys opened the door to complex net shape components production. However, the origins of their enhanced passivity, and the relationship between their unique microstructures (**Fig. 1**) and corrosion properties remain under investigation.

32 Theoretical models have been developed over the years to understand the dissolution/passivation 33 mechanisms of metals and alloys. These are often subsequently validated by experiments such as weight 34 loss measurements and electrochemical techniques. However, the concepts associated with passivity or 35 corrosion rate analysis were established based on "conventional" alloy systems (e.g., Al alloys, steels). Therefore, an important question is to which extent corrosion scientists may use such electrochemical tools 36 37 (*i.e.*, theories and experimental techniques) mostly built on the conventional alloys to the new emerging 38 class of alloys. Determination of a reliable corrosion rate of these alloys will be critical to control and 39 predict the sustainability of future engineered structures using these materials.

Herein, a brief review of the electrochemical theories and techniques used to determine the
corrosion rate of the conventional alloy systems is presented. The relevance of applying these current
electrochemical tools used on MPEAs and AM alloys by the corrosion science community is also discussed.
The objective of this review is to convey the current knowledge on the conventional alloys to newly
introduced complex materials.



**Fig. 1.** Schematic illustration of the microstructural characteristics of MPEAs and AM alloys. MPEAs, including high entropy alloys (HEAs), can be single phase multi-element solid solutions and are known to have complex oxide structures. AM alloys have been characterized to exhibit a refined grain size and microstructure with previously unreported phases such as quasicrystals, and grain directionality due to the high solidification rate of the manufacturing process and atypical nanometer sized particles.

#### 48 From passivity to corrosion rates on MPEAs and AM alloys: approximations and limitations

The electrochemical properties of the MPEAs and AM alloys are currently a subject of great interest to researchers due to their enhanced mechanical properties and corrosion resistance. This section firstly reviews the conventional theoretical models including passivation and corrosion rate determination, then discusses their application to the MPEAs and AM alloys. The range of spatial resolution of each alloy system and the reliability of the conventional theories which have been used to investigate these alloys are summarized in **Table 1**.

55

## 56 *Conventional metal oxidation and passivity models*

The passivity is often related to the formation of a protective film following the oxidation of a metal or an alloy. The passive film, generally assimilated to an oxide layer, is characterized by its composition, thickness, ionic and electronic properties which may regulate the overall corrosion of a system. Oxide growth models have been proposed and applied to understand the passivation kinetics of the metals and conventional alloys such as the Cabrera-Mott model [<sup>8</sup>], the Fehlner-Mott model [<sup>9</sup>], the place exchange model [<sup>10</sup>], the generalized growth model [<sup>11</sup>], and the point defect model (PDM) [<sup>12</sup>, <sup>13</sup>].

Kinetic models of passive film growth of MPEAs and AM alloys have been reported in the literature 63 based on theories built upon the conventional alloy systems. The question is whether the passive films of 64 65 the MPEAs and AM alloy systems exhibit unique properties or differ from the conventional systems. For 66 example, it has not been well understood whether these alloys show a critical threshold composition of the 67 principal passivating element as it is for the conventional alloys [2]. Another example is the PDM, originally 68 developed to describe the corrosion mechanism of austenitic alloys. The PDM was recently used to explain the transpassive dissolution mechanism of an AlTiVCr MPEA via conventional Mott-Schottky analysis 69 using EIS, coupled with element-resolved characterization techniques  $[1^{4}]$ . However, a general issue of 70 applying the PDM to the multi-principal element alloy systems is that the PDM does not account for the 71 72 non-constant electric field across the passive film, possible cation migration, and substrate composition. In addition, the PDM does not consider vet preferential segregation of alloving elements which may result to 73 the formation of multi-oxide structures as shown in recent studies of MPEAs [<sup>15</sup>, <sup>16</sup>, <sup>17</sup>]. Although a fourth 74 generation PDM is under development to account for these limitations and consider the preferential 75 76 segregation of alloying elements into the barrier or outer layer, the remaining question is how to validate 77 the PDM to be applicable to the newly developed compositionally complex alloys.

#### 79 Validity of corrosion rate determination: currently used techniques

80 Corrosion rate monitoring in aqueous environments is generally carried out by potentiodynamic 81 polarization experiments. Tafel extrapolation derived from the Butler-Volmer relation is subsequently used to experimentally determine the corrosion current (rate) and potential of a system. The "conventional" 82 corrosion rate has been measured using an experimentally determined polarization resistance ( $R_p$ ) by Stern-83 Geary equation assuming the system follows Tafel kinetics [<sup>18</sup>]. However, a number of other assumptions 84 85 have to be made to apply this equation to even a simple metal or alloy system, such as a stationarity during 86 the polarization, a uniform corrosion on the substrate (*i.e.*, no localized corrosion), and no changes in electrode potential due to additional reactions [<sup>19</sup>, <sup>20</sup>, <sup>21</sup>, <sup>22</sup>, <sup>23</sup>]. The application of these assumptions to a 87 non-uniform corrosion process (e.g., intergranular or pitting corrosion) is more challenging for MPEAs and 88 89 AM alloys due to their complex chemistry and microstructure.

90 An example illustrating the difficulty of determining a corrosion rate solely from Tafel 91 extrapolation is given in Fig. 2 in the case of an as-printed AM alloy AA2024 (AM2024) and a conventional AA2024-T3, exposed to a 0.01 M NaCl solution under anodic polarization [<sup>24</sup>]. Note that the AM2024 has 92 93 a similar elemental composition to AA2024-T3. The corrosion rate (icorr) determined from conventional 94 Tafel extrapolation (Fig. 2(a)) showed similar values for both alloys as provided in Fig. 2(b). The AM2024 95 alloy exhibited a slightly higher anodic current density than AA2024-T3 shown in Fig. 2(a). The elementalresolved atomic emission spectroelectrochemistry (AESEC) gives complementary information to the 96 97 conventional polarization curves as shown in Fig. 2(c) and 2(d). For both alloys, the total electrical current density measured from the potentiostat ( $i_e$ , Fig. 2(a)) was one order of magnitude higher than the equivalent 98 total current density ( $i_{tot} = i_{Me} + i_{Fe} + i_{Cu} + i_{Al}$ , as illustrated in Figs. 2(c) and 2(d)) measured by AESEC. This 99 100 result indicates the formation of a thicker passive film during anodic polarization on the AM2024 surface 101 (Fig. 2(d)) than AA2024-T3 (Fig. 2(c)) indicated by the larger  $i_{film}$  (=  $i_e$  -  $i_{tot}$ ) for the AM2024 (mass-charge 102 balance). This larger unaccounted charge (i<sub>film</sub>) was attributed to the refined microstructure of the AM2024, 103 limiting the onset and propagation of localized corrosion. Similar findings have been reported for other 104 MPEA systems highlighting how the conventional potentiodynamic polarization results can be even more 105 misleading, especially for MPEAs where the role of each alloying element in the anodic and cathodic reaction cannot be easily distinguished [<sup>25</sup>, <sup>26</sup>, <sup>27</sup>, <sup>28</sup>, <sup>29</sup>]. The AESEC technique, under this scenario, can be 106 107 useful to monitor the element-resolved dissolution kinetics including the elemental corrosion rate. 108 However, the current technical setup is limited to macroscale electrochemical analysis and does not provide information on nanometer level local events  $[^{30}, ^{31}]$ . 109

110 The recent literature reveals the growing need to use tools considering the nanostructural 111 complexity of these new alloys. The examples displayed herein highlight the inevitable upcoming

- 112 obsolescence of founding models and techniques used to characterize the corrosion susceptibility if no
- 113 forthcoming update is realized. Under this scenario, the collection of a valid experimental data-set would
- 114 be a first step to depict the origins of localized corrosion on these alloys, which ultimately means
- 115 downscaling local techniques to the nanometer level.



**Fig. 2**. (a) Potentiodynamic polarization curves of the conventional AA2024-T3 alloy and AM2024 alloy in 0.01 M NaCl with 1 mV/s scan rate. Tafel extrapolation indicates very similar corrosion rates for the two alloys. However, the AESEC anodic polarization curve (indicated by the pink rectangle) for the (c) AA2024-T3; and (d) AM2024 alloy reveals that in the case of AM2024 a substantial proportion of the anodic current forms of a thicker passive film (gray area). Adapted from [24].

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**Table 1**. Summary of the microstructural specificities of each alloy system and their potential impact onmechanisms interpretation.

Conventional alloys	MPEAs	AM alloys
<ul> <li>Well-reported micro- to nano-scales particles and precipitates.</li> <li>Micro-scale localized corrosion processes can be identified by current local techniques.</li> <li>Tafel extrapolation for corrosion rate determination can be applicable if occurring uniformly on the matrix.</li> <li>Passive film compositions are usually associated with one metal (matrix).</li> </ul>	<ul> <li>Tafel extrapolation and interpretation is difficult. Anodic current density can be associated with several elements from the matrix [25 - 29].</li> <li>Complex passive films (more than one metal present) with the presence of unoxidized species [14 - 17].</li> <li>Passive film formation mechanism different from conventional alloys, including stainless steels.</li> </ul>	<ul> <li>Formation of previously unreported non-equilibrium phases for new and well-known alloy systems due to high solidification rate [4, 5].</li> <li>Refined microstructures with nanometer sized particles and precipitates [4, 5, 24].</li> <li>Difficult to identify localized corrosion processes with the current local electrochemical techniques due to the limit of spatial resolution, and refined microstructure.</li> <li>Tafel extrapolation does not consider the proportion of current density associated with passive film formation [24]</li> </ul>

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## 127 Local electrochemical techniques for MPEA and AM alloys: to the origins of passivity and

#### 128 localized corrosion

If a system undergoes localized corrosion, the measured corrosion rate can significantly exceed that 129 of uniform corrosion  $[^{32}]$ . A passive film formed on an alloy surface can contribute to decreasing the 130 uniform corrosion rate, however, it can also accelerate the localized corrosion rate by pitting or crevice 131 132 corrosion associated with the local passive film breakdown. To this end, local electrochemical techniques 133 have been developed in the last few decades to address localized surface reactivity usually masked in 134 macroscale electrochemical measurements. The difference between global corrosion rate measurement and 135 the actual localized rate would be more significant for the AM alloys and some of the MPEAs where the 136 refined microstructural features have been reported. It is therefore essential to use high spatial resolution local techniques for these alloys, in particular to identify corrosion initiation sites. In this section, recent 137 138 studies using conventional and newly introduced local techniques to investigate the electrochemical 139 properties of a system including MPEAs and AM alloys are discussed. A better understanding of the local 140 reactivity of the surface could be obtained by combining conventional electrochemical and imaging techniques with novel high-resolution techniques. 141

#### 142 Local electrochemistry for MPEA and AM alloys: current limitations

Despite the technical advances, only limited studies on the MPEAs or AM alloys using local 143 144 electrochemical techniques have been reported to date. Recently, the scanning vibrating electrochemical technique (SVET) technique has been used to investigate a micro-galvanic corrosion between the two 145 eutectic phases of an AlCoCrFeNi MPEA with a 500 nm level step size scans [<sup>33</sup>]. The SVET technique 146 generally visualizes a real-time the local cathodic and anodic reactions by the potential difference in a 147 solution between a vibrating conductive probe and the sample of interest  $[^{34}, ^{35}, ^{36}]$ . Theoretically, it can 148 149 identify the location of half-reactions and quantify their associated corrosion rates. However, a few technical limitations, such as trade-off sample size  $[^{37}]$  and surface topography, could affect the potential 150 measurement leading to an error in quantitative analysis. In addition, real-time corrosion rate determination 151 152 using this technique for the MPEAs and AM alloys will be significantly influenced by the experimental protocol. For example, high vibrational amplitude may provide an artifact signal due to the accelerated  $O_2$ 153 reduction reaction which is often the rate-determining diffusion-controlled step, giving an incorrect overall 154 corrosion rate [<sup>38</sup>, <sup>39</sup>]. The SVET should be carried out more systematically to the MPEAs and AM alloys 155 considering these limitations to investigate the local corrosion processes. 156

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## 158 Downscaling electrochemical techniques for next-generation materials: towards the nanoscale?

159 Although nanoscale electrochemical measurement can be performed today using scanning 160 electrochemical microscopy (SECM), SECM measurement usually requires the use of a redox mediator that could alter the overall corrosion mechanism. Recently, a nanoscale electrochemical setup equipped 161 with nano-pipette imaging probe known as scanning electrochemical cell microscopy (SECCM) has been 162 used to monitor local electrochemical events  $\begin{bmatrix} 40 & 41 \end{bmatrix}$ . This concept resembles the electrochemical droplet 163 cell used to visualize anodic and cathodic reactions, in which a nanometer sized mobile meniscus (down to 164 ~ 30 nm  $[^{42}, ^{43}]$ ) is positioned on the alloy surface (Fig. 3.). The SECCM technique was applied to 165 investigate the reactivity of each crystallographic orientation on a polycrystalline alloy defining a structural 166 factor that governs the corrosion mechanism [44]. However, the SECCM technique applied to corrosion 167 168 studies is still limited to low carbon steel (using a 2 µm probe) and simple systems such as polycrystalline Zn to date  $\begin{bmatrix} 40, 41, 45, 46, 47 \end{bmatrix}$ . In addition, several technical challenges remain. For example, it has been shown 169 that the exposed surface is often non-reproducible and evolves with reaction time, e.g., by O<sub>2</sub> ingress into 170 the droplet if the O<sub>2</sub> reduction reaction is the dominant cathodic reaction, leading to a significant surface 171 alteration during measurement [<sup>48</sup>]. The stability of the probe can be affected by the formation of corrosion 172 173 products or the formation of gas bubbles ( $e.g., H_2$  reduction reaction) during the measurement. Moreover,

the tip-substrate distance (currently controlled by conductivity measurement) which can be influenced by the surface reactivity (and by extension, the surface composition), must be reproducible at each measurement.

177 Nonetheless, the nano-pipette probe could open the door to nanoscale electrochemical impedance spectroscopy measurements. Local electrochemical impedance spectroscopy (LEIS) has been utilized to 178 179 probe the surface reactivity giving a mechanistic information on a specific phase or location of the sample surface [49, 50, 51, 52]. LEIS has demonstrated to be a powerful technique to investigate local corrosion events, 180 however, the current experimental setup has a limited spatial resolution (down to 10 µm to date). Besides 181 182 the remaining technical challenges, the SECCM technique could be a pathway to nanoscale corrosion 183 studies and provide new insights into the corrosion mechanism of the next-generation alloys if the abovementioned experimental constraints are addressed in the future. 184

Through the miniaturization of electrochemical liquid cells designs, the development of 185 186 electrochemical chips (E-chips) operando transmission electron microscopy (TEM) has recently emerged  $[^{53}, ^{54}]$  to monitor real-time chemical/electrochemical reactions at the metal/oxide/electrolyte interface  $[^{55}]$ 187 188 Operando TEM shows nanometric morphological and compositional evolution during electrochemical 189 measurements as illustrated in Fig. 3 [<sup>56</sup>]. The liquid phase TEM (LP-TEM) consists of a 300 µm-thick E-190 chip containing a 5 µm width working, reference, and counter electrodes in a specially designed TEM sample holder [<sup>57</sup>]. Recent work investigated the dissolution of micro- and nanoscale MnS inclusions from 191 stainless steel lamellae during a real-time potentiodynamic polarization experiment. In situ monitoring of 192 193 surface evolution using TEM may give light to determine the local corrosion process (e.g., the effect of 194 nanostructure) of multi-oxide films or identify the reactivity of previously unreported phases recently found in the AM alloys [<sup>58</sup>, <sup>59</sup>]. However, the development of an optimized liquid cell is still at its premises and 195 196 requires further investigation. The quantitative measurement of the reaction rate using this technique is still 197 limitedly accessible and has not been applied to the new-generation alloys.



TEM techniques (LP-TEM), the TEM lamellae prepared by focused ion beam technique is placed in the E-chip and welded to the Pt working electrode. The E-chip is then integrated into a TEM sample holder consisting of two electrolyte inlets and one outlet. Such coupling allows a direct visualization of corrosion initiation mechanisms at the nanometer level.

### 201 **Perspectives and closing remarks**

Recent advances in alloy metallurgy require prompt action if we want to accurately estimate the lifetime of structural materials currently used and to be used in our society. Experimental discrepancies in the measurement of corrosion rate, for example, reveal the need to develop more elaborate theoretical models and appropriate experimental verification techniques.

206 First, the development of high spatial resolution nanoscale characterization and local in situ electrochemical techniques is necessary for a better mechanistic understanding of the corrosion mechanisms 207 208 of next-generation materials which are possibly more complex than conventional alloy systems. Promising 209 new imaging techniques coupled with electrochemistry using nano-pipettes have recently been introduced 210 to monitor the local reactivity of a system although they are still limited to the conventional alloys. The 211 novel local measurement techniques would allow a direct correlation between local corrosion sites and the nanostructure, particularly in the case of unreported or non-equilibrium phases which are difficult to isolate 212 213 from the matrix. Second, operando TEM technique will allow a direct visualization of corrosion and passivation mechanisms at the nanometer level. However, it is more experimentally challenging and has 214 215 not yet been applied to MPEAs or AM alloys.

Finally, the emergence of computational materials science and machine learning for thermodynamic simulations combined to an adequate experimental dataset, for the design of corrosion resistant alloys can be a path to future work. The adjustment of existing theoretical models with experimental justification is essential to rationalize their passivation and corrosion mechanisms and to accurately predict the lifetime of next-generation alloys.

#### 221 **Conflict of interest statement**

222 Nothing declared.

223

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- \* of special interest
- 231 \*\* of outstanding interest

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\* This paper classifies notions of the multi-principal element alloys with compositionally complex alloys (CCAs) and high-entropy alloys (HEAs). The up-to-date corrosion resistance researches on the MPEAs were reviewed and the perspectives of the future research are suggested.

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### **Declaration of interest statement**

Nothing declared.

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