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**Fretting-corrosion of cardiovascular stent materials: the role of electrochemical polarisation on debris generation mechanisms**

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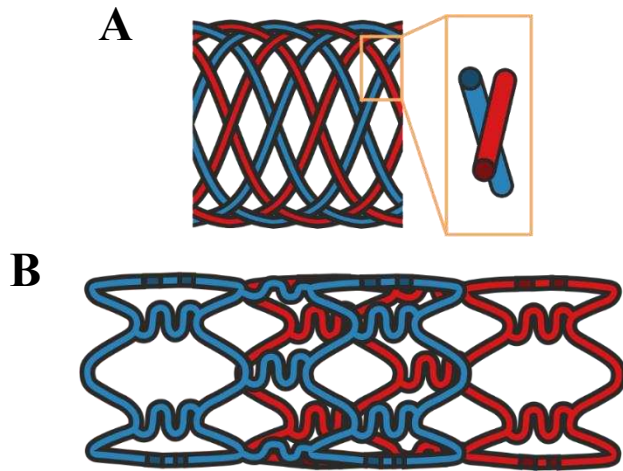
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## 1. Introduction

Balloon angioplasty and implantation of stents has been used as a successful intervention for atherosclerosis for upwards of 40 years, yet clinical limitations persist. In-stent restenosis (ISR) accounts for about 10% of coronary stent failures, increasing to 30% in peripheral stents, one year post-implantation in the modern stent era requiring further intervention (1, 2). In-stent restenosis (ISR) is characterised by a re-narrowing of the artery post-stent implantation to such an extent that blood flow is hindered to the downstream organ (3). The principle cellular component of the arterial wall are smooth muscle cells (SMCs) which migrate towards the lumen of the artery, proliferate and secrete extracellular matrix components during development of ISR (4). The advent of drug-eluting stents (DES) has gone some way to reducing the incidence of ISR but the issue remains in a non-negligible patient subset (5). In contrast with the clinical post-failure analysis conducted in orthopaedic implants, stents with ISR are typically not removed from the patient: re-stenting or bypass grafts are used (6, 7). Post-failure inspection and analysis of the devices is therefore hampered and the true occurrence of fretting and corrosion *in-vivo* is unknown.

The surfaces and interfaces established during the implantation of a vascular devices are receiving increased scrutiny due to the possible adverse soft tissue effects associated with implant derived debris. Surface analysis of retrieved vascular devices has demonstrated that interfaces are susceptible to fretting-corrosion; the combined action of mechanical wear and corrosion at small displacement amplitudes. Evidence of fretting-corrosion has been observed at the braided stent strut interface (7, 8) and *in vivo* at overlapping interfaces where the use of multiple stents may be necessary to treat long lesions and those arising from complex and dynamic biomechanics, particularly in peripheral stents (6, 7, 9). Such interfaces are shown in Figure 1. Within the vasculature, stenting undergoes large deformations through haemostatic forces (e.g. pulsatile flow, pressure) and daily-living activities (e.g., leg flexion) introducing micro-displacements at overlapping or braided interfaces. Previous studies (10, 11) have shown that overlapping and braided stents (Figure 1a) exhibit evidence of wear and corrosion after laboratory simulation. This is supported by *in-vivo* observations (12, 13) (Figure 1b), in which a tentative link between metallic degradation, inflammation and ISR has been hypothesised. Therefore it

46 is conceivable that the nature of micro-motion and contact conditions at the interface can be incredibly  
47 transient and activity dependant.



48

49 Figure 1. Schematic of fretting-corrosion interfaces in cardiovascular stents. A) Cross-wire interface in  
50 braided stents (shown with close up), more commonly used in peripheral arteries. B) Laser cut type  
51 stents in an overlapping configuration, often used for long atherosclerotic lesions or re-intervention due  
52 to ISR.

53

54 Degradation products (metal ions and nano-particles) arising from the fretting-corrosion interfaces will  
55 be released into the circulation and local tissues depending upon the degree of re-endothelialisation.  
56 Metallic debris arising from cardiovascular devices, in the form of metallic ions or bulk metal particles,  
57 has been shown to have a biological effect on cells and tissue and can mediate oxidative stress, release  
58 of proinflammatory mediators, cytotoxicity, DNA damage and functional cellular changes such as  
59 proliferation (14-17). These same biological effects are also implicated in the pathogenesis of in-stent  
60 restenosis (1, 4). To elucidate the links between device degradation and adverse biological processes,  
61 some studies have used sources such as metal chloride salts (15, 18, 19), passive corrosion through  
62 incubation under physiological conditions (20-22) or application of an electrical potential to simulate  
63 and accelerate the oxidising action of the physiological corrosive environment (16, 20). The role of  
64 tribocorrosion products (i.e. metal ions and particles) on vascular biology processes are yet to be fully  
65 considered. It is also well known that the characteristics (e.g. chemistry, thickness, stoichiometry,  
66 formation kinetics) of the passive layer on the surface of biomedical alloys, such as CoCrMo and NiTi,

67 is dependent upon the applied potential in corrosion systems (23, 24) and influences the synergistic  
68 electrochemical and mechanical degradation of a metallic alloy (25-28).

69 In order to accurately assess the biological responses to metallic debris, the pathways to their production  
70 must be fully considered, particular when qualifying materials and devices pre-clinically. In a  
71 comprehensive review Mischler found that the potentiostatic method, whereby the system is held at a  
72 nominal potential during fretting-corrosion, is the most commonly used tribo-electrochemical method  
73 (29). This method has been used variously to understand the mechanisms of fretting-corrosion which  
74 are representative of implanted metallic medical devices *in vivo* (23, 30-33). In reality, the potential  
75 under free corrosion conditions is dynamic and evolves over time (27, 34, 35). Conducting fretting-  
76 corrosion tests at the free potential of the system (or open circuit potential, OCP) has therefore more  
77 recently been used to simulate the physiological fretting-corrosion mechanisms of biomaterials (36, 37).  
78 The experimental methodology to understand bio-tribocorrosion mechanisms and released wear debris  
79 can fundamentally alter the results. Differences in wear debris chemistry and morphology can instigate  
80 differential biological effects (17, 38). Questions must be raised as to the suitability of imposing  
81 electrochemical potentials to simulate corrosion in tribological contacts, its relevance to the pre-clinical  
82 assessment, its implication on the nature of degradation and correlation between imposed  
83 electrochemical polarisation and free corrosion conditions.

84 The aim of this study was therefore to perform a detailed tribocorrosion study investigating the effects  
85 of polarisation on generated wear debris and material degradation during fretting in a simulated stented  
86 physiological environment. Two primary metallic biomaterials were investigated, cobalt-chromium-  
87 molybdenum (widely used biomaterial) and nickel-titanium (common vascular material) were  
88 investigated in a braided stent strut model simulating the geometric considerations. The need for further  
89 detailed investigation to understand the role of tribocorrosion and electrochemical potential/current  
90 distributions in the vascular environment has been emphasised by leading cardiologists (39). We have  
91 shown these degradation mechanisms to be detrimental to the performance of other metallic devices  
92 implanted into the biological environment (40).

93 **2. Materials and methods**

94 **Materials**

95 Superelastic nickel-titanium, NiTi, (Nitinol, NDC, USA) and low carbon cobalt-chromium-  
96 molybdenum alloy, CoCrMo, (ASTM F1537, Heymark Metals, UK) was machined into cylindrical  
97<sub>2.1.</sub> samples Ø6 mm to represent cylindrical braided stent struts. CoCrMo was for benchmarking due to our  
98 extensive experience with this alloy. Chemical composition of alloys used in this study are shown in  
99 Table 1 according to manufacturer. Prior to experimentation, all samples were polished to a surface  
100 roughness,  $R_a > 25$  nm using a polishing cloth (Buehler, Germany) and graded diamond polishing pastes  
101 (Kemet International Ltd, Kent, UK), cleaned with acetone and air-dried.

102 **Table 1.** Chemical composition of low carbon CoCrMo and superelastic NiTi used in this study as  
103 stated by the manufacturer.

Element (Wt%)	Cr	Mo	Mn	Si	Ni	Fe	N	C	Co
<b>CoCrMo</b>	26.0 – 30.3	5.0 – 7.0	0 – 1.0	0 – 1.0	0 – 1.0	0 – 0.75	0 – 0.25	0 – 0.14	Bal
Element (Wt%)	Ni	Ti							
<b>NiTi</b>	55.0	45.0							

104  
105 Serum-free culture media composed of Dulbecco’s Modified Eagle Medium supplemented with 1% v/v  
106 L-glutamine (necessary growth factor) and 1% v/v antibiotic-antimycotic (Gibco, Renfrewshire, UK).  
107 This formulation is widely used in cell culture to simulate the physiological biochemical environment  
108<sub>2.2.</sub> was used as the lubricant in this study (41, 42).

109 **Fretting tribometer**

110 Interactions between wear and corrosion were investigated using a custom-built reciprocating  
111 electromechanical fretting tribometer, previously developed, described and validated (43). A schematic  
112 for the setup is shown in Figure 2A. An electromechanical actuator provided reciprocating motion to  
113 the upper cylinder with an amplitude of 200  $\mu\text{m}$  ( $\delta_d$ ) and was maintained at a frequency of 1 Hz,  
114 simulating that of a physiological heartbeat.  $\delta_d = 200$   $\mu\text{m}$  was chosen to ensure we remain in the gross-  
115 slip regime for the duration of the test, to best represent the stent device observations (i.e. abrasion  
116 distances on the struts) (12) and FEA analysis (16). Reciprocating motion was measured by means of a  
117 fibre optic sensor fixed in the holder mounted to the base fixture of the fretting apparatus. The tangential

118 force ( $F_i$ ) was measured throughout the experiments using a cylindrical force transducer (Kistler, USA)  
 119 mounted axially to the actuator and the upper alloy cylinder. The initial normal load ( $F_n$ ) at the contact  
 120 between upper and lower samples was set using a force transducer (RDP Electronics, UK). A normal  
 121 load of 4 N was used for each experiment (Table 2), achieving contact pressures similar to values  
 122 reported by finite element analysis for overlapping cobalt-alloy stents (6). Analysis of the fretting-loop  
 123 data was done according to the criteria outlined by Fouvry *et al.*(44)

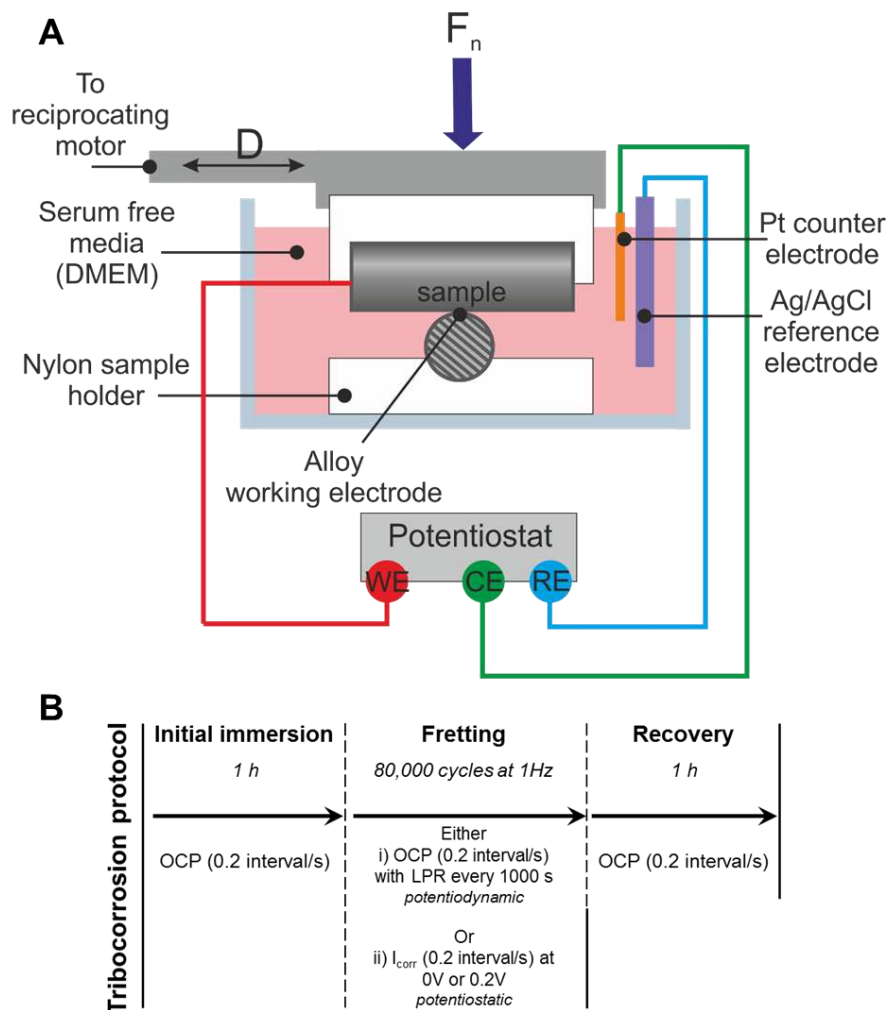
124 Table 2. Maximum pressure at cylinder contact as calculated by Hertzian analysis.

Alloy	Young's modulus (GPa)	Poisson's ratio	Maximum contact pressure ( $P_{max}$ ) (GPa)
CoCrMo	220	0.29	0.68
NiTi	34	0.33	0.21

125

126 **Fretting-corrosion arrangement**

127 <sup>2.3.</sup> A three electrode electrochemical cell was integrated into the fretting tribometer (Figure 2A). The  
 128 physical connection of the upper cylinder to the lower cylinder immersed in the serum-free culture  
 129 media acted as the net-working electrode (WE). A combined Ag/AgCl reference electrode (RE;  
 130 +0.196V vs. standard hydrogen electrode) and Pt counter electrode (CE) (Orion, ThermoFisher  
 131 Scientific, MA, US) was held in the serum-free culture media to complete the electrochemical cell. The  
 132 WE, RE and CE were connected to a computer controlled potentiostat (PGSTAT101, Metrohm,  
 133 Switzerland).



134

135 **Figure 2.** A) Experimental set-up of simulated braided stent strut fretting (perpendicular cylinder  
136 configuration). Hatching indicates cross section. B) Protocol for fretting-corrosion testing under either  
137 potentiodynamic or potentiostatic regimes.

### 138 2.3.1. Fretting-corrosion protocol

139 The protocol for the fretting-corrosion experiments is shown in Figure 2B. Briefly, the NiTi cylinders  
140 were polished, cleaned and dried before installation into the fretting rig in a perpendicular configuration.  
141 The serum-free culture media was then added and the upper and lower cylinders were immersed in  
142 contact with each other and left for 1 h. For each condition, the OCP was recorded in this interval.  
143 Subsequently, fretting occurred for 80,000 cycles at 1 Hz for all tests. After the fretting test, the alloy  
144 was left to recover for 1 h before the tests were dismantled and the cylinders and the serum-free culture  
145 media were collected for analysis. Serum-free culture media was frozen at  $-20^{\circ}\text{C}$  to prevent dissolution  
146 of particles.

147 The fretting-corrosion behaviour of the simulated braided strut interface was evaluated under three  
 148 conditions:  
 149 Open circuit potential (OCP) with intermittent linear polarization resistance (LPR) measurements at  
 150 which the alloy is under free corrosion conditions enabling characterisation of the interface at  
 151 equilibrium. This approach has been extensively used and published in the literature for the long term  
 152 assessment of tribocorrosion contacts (43, 45, 46). Measurement at free corrosion potential ( $E_{\text{corr}}$ ) with  
 153 LPR allows quantitative *in situ* dynamic corrosion measurements. LPR measurements were taken every  
 154 1000 s at an applied potential of  $\pm 0.02$  V versus OCP at a scan rate of  $0.25 \text{ mV}\cdot\text{s}^{-1}$ . It is widely accepted  
 155 that applying such a potential no permanent changes to the electrode surface are incurred (45). The scan  
 156 rates chosen in this study are sufficiently slow enough to eliminate any capacitive charging effects due  
 157 to polarisation and show no evidence of hysteresis (i.e. charging) on reversal of the potential (47). The  
 158 linear relationship between potential and current was used to calculate the polarisation resistance ( $R_p$ ).  
 159 This was then used in conjunction with the experimentally determined Tafel constants and the Stern-  
 160 Geary (SG) relationship (Equation 1) (48) to determine corrosion current ( $I_{\text{corr}}$ ) for the duration of the  
 161 experiment. Tafel constants for cross-cylinder contact for both CoCrMo and NiTi alloys in serum-free  
 162 culture media were experimentally determined under static conditions. Samples were immersed for 1 h  
 163 to allow the open circuit potential (OCP) to reach equilibrium before polarisation to  $\pm 0.6$  V vs RE at a  
 164 scan rate of  $0.25 \text{ mV}\cdot\text{s}^{-1}$  to capture the Tafel regions. For each alloy, the slope of the linear anodic and  
 165 cathodic regions of current in relation to overpotential ( $\beta_a$  and  $\beta_c$  respectively) were taken as the Tafel  
 166 constants.

$$I_{\text{corr}} = \text{SG} \left( \frac{1}{R_p} \right) \quad (1)$$

167 Where  $R_p$  = polarisation resistance, SG = Stern-Geary coefficient,  $I_{\text{corr}}$  = corrosion current density, and

$$\text{SG} = \frac{\beta_a \cdot \beta_c}{2.303 (\beta_a + \beta_c)} \quad (2)$$

168 Where  $\beta_a$  and  $\beta_c$  = anodic and cathodic Tafel constants respectively.

169 Potentiostatic polarization was conducted at either 0 V or 0.2 V vs RE for the duration of the experiment.

170 Tribocorrosion at increasing potentials within passive domain were selected to accelerate the anodic

171 processes. The current transient was recorded at 5 s intervals and both potentials were below the  
172 breakdown potential of each alloy in serum-free culture medium as experimentally determined during  
173 cyclic polarisation for determination of Tafel constants.

#### 174 **Optical microscopy (OM)**

175 Optical microscopy (DM6000M, Leica, Germany) was used to inspect the wear scar and observe  
176 surface changes on both the upper and lower cylinders after fretting corrosion. As the surfaces of the  
177 cylinders were curved, a 3D montage imaging technique (Z-stack) was used to ensure all parts of the  
178 images were in focus.

#### 179 **Vertical scanning interferometry (VSI)**

180<sup>2.5.</sup> The volume loss of material from the upper and lower cylinders was determined using vertical scanning  
181 interferometry (VSI) (NPFlex, Bruker, US) with a 10x objective. Prior to VSI analysis, the cylinders  
182 were lightly cleaned using acetone to remove superficially adhered debris. The measurement area was  
183 adjusted to accommodate the varying sizes of wear scar. The volume loss was defined as the negative  
184 volume of the wear scar from the mean zero plane of the surface. The total mass loss due to both wear  
185 and corrosion mechanisms was defined as the sum of the mass loss from the upper and lower cylinders  
186 and was extrapolated from the volume loss using appropriate material density (CoCrMo: 8.4 g/cm<sup>3</sup> ;  
187 NiTi: 6.5 g/cm<sup>3</sup>). VSI data was analysed using Vision 64 software (Bruker, MA, US). Cylindrical  
188<sup>2.6.</sup> curvature and tilt was removed and a median data filter of magnitude 3 was applied.

#### 189 **Scanning electron microscopy (SEM)**

190 Scanning electron microscopy with energy dispersive X-ray analysis (SEM-EDX) (Carl Zeiss EVO  
191 MA15, ZEISS, Germany) was used to determine changes in the chemical composition of the wear scar  
192<sup>2.7.</sup> at the fretting contact. Samples were imaged with a beam intensity of 20 kV and EDX was performed  
193 at a working distance of 8 mm.

#### 194 **Inductively coupled plasma mass spectroscopy (ICP-MS)**

195 Corrosion products in the media were analysed according the methods outlined by Simoes et al (49).  
196 Ion concentration and composition was measured with inductively coupled plasma mass spectroscopy

197 (ICP-MS). A small volume of the serum-free culture media from the fretting tests was taken and passed  
198 through a 2 kDa ultrafilter (Vivacon, Sartorius AG, Germany) at 14,000 x g for 1 h in a centrifuge to  
199 remove all particles. The ultrafiltered media was then diluted by 100x in 2% v/v nitric acid (Fluka,  
200 Honeywell, Romania) in ultrapure water and analysed for Co, Cr, Mo, Ni and Ti isotopes using ICP-  
201 MS standards for calibration for aforementioned elements (Sigma-Aldrich, USA). A matrix blank of  
202 serum-free media was also prepared to characterise background elemental signals from the media itself.

### 203 **Particle analysis**

204 A volume of 1 mL of the serum free culture media used as electrolyte during the fretting experiment  
205<sup>2,8</sup> was defrosted and centrifuged at 14,000 x g for 1.5 h. The supernatant was then carefully removed and  
206 the particles were resuspended in ultrapure water to remove traces of essential salts from the serum-free  
207 culture media. The centrifugation and resuspension process was repeated 5 times per sample when the  
208 supernatant was removed leaving a pellet of particles in 250 µl of ultrapure water. The particle pellets  
209 were then sonicated for 30 mins at 37°C. The particle suspension was then pipetted onto adhesive  
210 carbon SEM stubs and left to air dry. The particles were then imaged using SEM-EDX to analyse their  
211 morphology and chemical composition.

### 2.9.

### 212 **Statistical analysis**

213 A one-way ANOVA with post-hoc Tukey test was performed for each alloy (CoCrMo and NiTi) to  
214 assess the significance of impact of the polarisation regime (OCP, 0V, 0.2V) on total mass loss and  
215 composition of ions released to the electrolyte. A p-value of less than 0.05 was considered significant  
216<sup>3,1</sup> for all analyses. Error bars on graphs are presented as 95% confidence intervals.

## 217 **3. Results**

### 218 **Static cyclic polarisation**

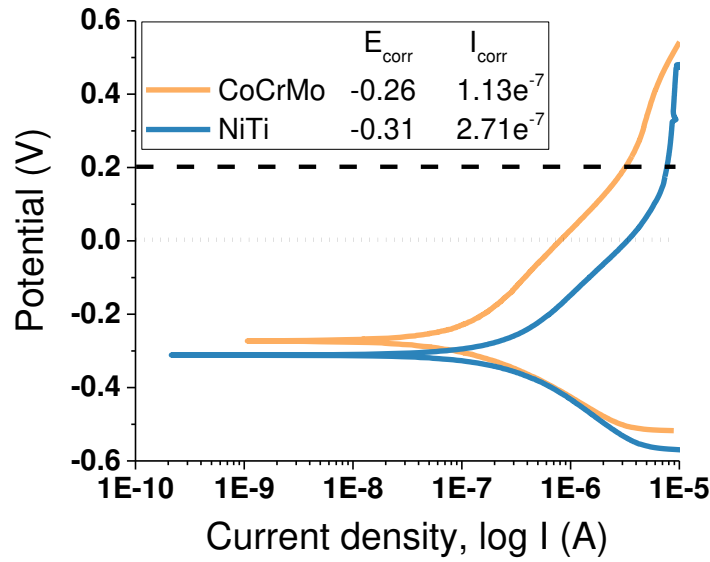
219 Figure 3 shows the Tafel plots obtained for CoCrMo and NiTi under static conditions. The corrosion  
220 current density ( $I_{corr}$ ) and free corrosion potential ( $E_{corr}$ ) determined through Tafel fitting are shown in  
221 the inset of Figure 3. Tafel constants and the associated Stern-Geary coefficient used later in the  
222 analysis of LPR data are tabulated in Table 3.

223

224 **Table 3.** Tafel constants of CoCrMo and NiTi alloy in serum-free culture media

Alloy	Anodic Tafel constant ( $\beta_{as}$ , V/dec)	Cathodic Tafel constant ( $\beta_{cs}$ , V/dec)	Stern-Geary coefficient
CoCrMo	0.31	0.17	0.048
NiTi	0.29	0.21	0.053

225



226

227 **Figure 3.** Polarisation curves of CoCrMo and NiTi alloy in serum-free media. Potential range -0.6 V to  
228 +0.6 V with scan rate of 0.167 mV.s<sup>-1</sup>. Polarisation potential for potentiostatic regime shown for 0V  
229 (dotted line) and 0.2V (dashed line).

230<sub>3.2.</sub>

### 231 Mechanical fretting results

232 Representative fretting loops for CoCrMo and NiTi alloy (n=3) under polarisation regimes at OCP and  
233 held at 0V and 0.2V are shown in Figure 4. For each condition, fretting loops from intermittent cycles  
234 are shown to demonstrate the evolution of the fretting regime over time. Cumulative dissipated energy  
235 over 80,000 cycles as characterised by the mathematical area of the fretting loop is shown in Figure 5A  
236 and B for CoCrMo and NiTi respectively. Representative optical microscopy and VSI images of the  
237 wear scar for CoCrMo and NiTi under all three polarisation conditions are shown in Figure 6.

### 238            **3.2.1. Fretting loops**

239 Under all polarisation regimes, CoCrMo alloy operated within a gross-slip regime characterised by a  
240 quasi-rectangular fretting loop with a sliding / displacement ratio,  $\frac{\delta_s}{\delta_a} = 0.93 - 0.95$ ; well within the  
241 gross-slip criteria given by Fouvry (50) . Under fretting-corrosion at OCP, CoCrMo alloy did not  
242 demonstrate any evolution in the fretting regimes over time and the loops over the course of 80,000  
243 cycles are relatively similar with an almost indistinguishable fretting loop area (Figure 4A). When the  
244 polarisation potential was increased to 0V, fretting remained in a gross-slip regime but the fretting loop  
245 area increased over time (Figure 4B). At 0.2V polarisation potential, CoCrMo again operated in a quasi-  
246 rectangular gross-slip regime with an increase in fretting loop area over time and was overall greater  
247 than at lower polarization potentials (Figure 4C). As CoCrMo existed in a gross-slip regime for OCP,  
248 0V and 0.2V, increasing the polarisation potential during fretting-corrosion did not therefore appear to  
249 alter the mechanism of the fretting regime. However, polarisation to both 0V and 0.2V resulted in  
250 changes to the fretting loop area over time which was not present at OCP.

251 The fretting conditions for superelastic NiTi alloy also exhibited a gross-slip fretting regime with  $\frac{\delta_s}{\delta_a}$   
252 varying between 0.83 and 0.95. However under such conditions the fretting loop adopts a quasi-  
253 elliptical shape with increasing tangential forces towards the end of each stroke (Figure 4D, E, F). The  
254 area of the fretting loops over the course of the fretting regime for NiTi are more consistent in  
255 comparison to those produced by CoCrMo alloy for all polarisation regimes. The trends in dissipated  
256 energy over-time were not consistent with those observed for CoCrMo and were not seen to increase  
257 with applied potential.

### 258            **4. Cumulative dissipated energy**

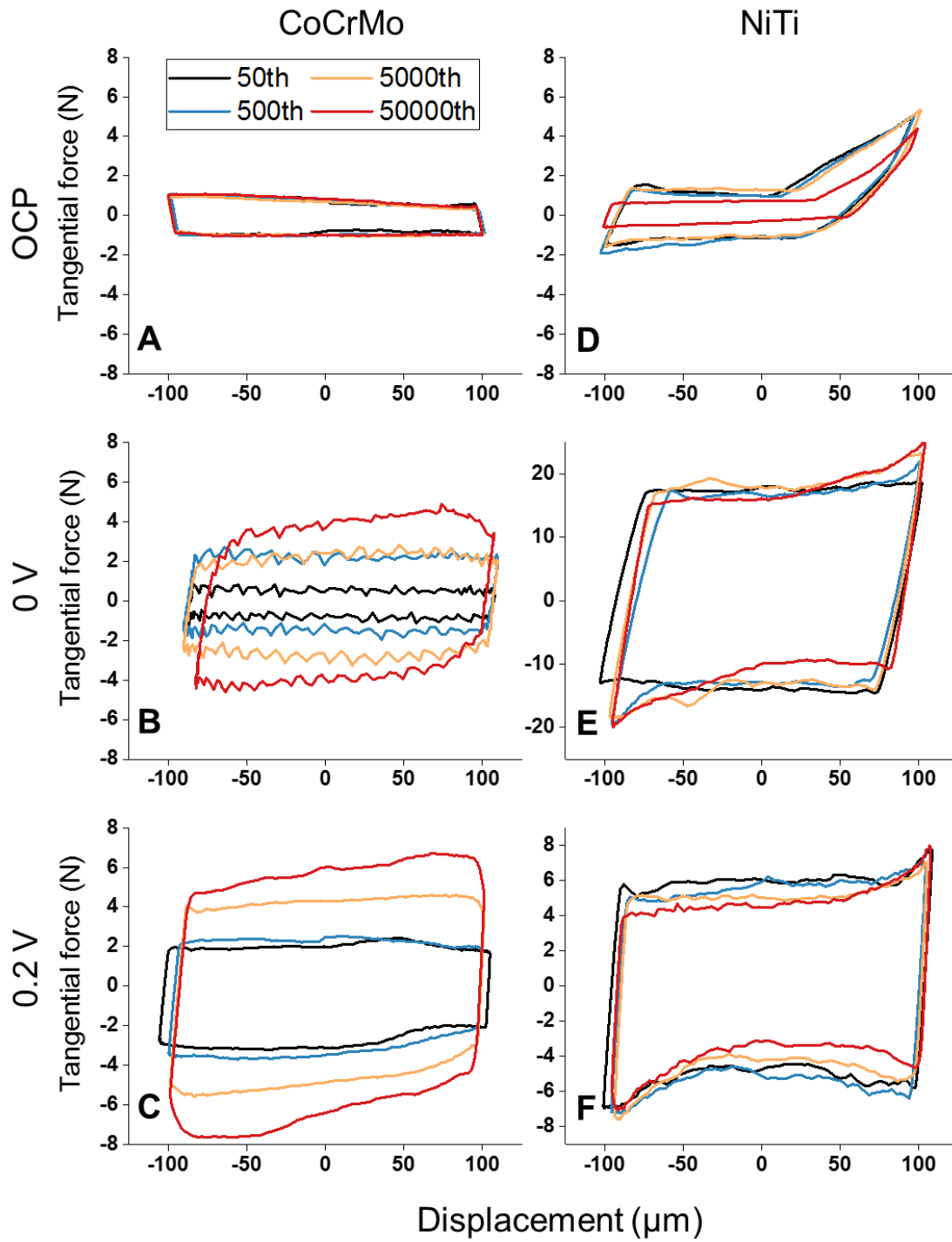
259 Figure 5A and B shows the cumulative energy dissipation as a function of cycles for CoCrMo and NiTi  
260 under different corrosion conditions for CoCrMo and NiTi, respectively. In general, an increase in the  
261 total energy dissipated during fretting increased with applied potential and in both material cases was  
262 non-linear with the number of cycles. When compared to CoCrMo, the energy dissipated during fretting  
263 for NiTi materials was greater.

264 The relationship between cumulative dissipated energy and total volume loss for CoCrMo and NiTi  
265 alloys at all polarisation potentials is shown in Figure 5C and D, respectively. For CoCrMo an  
266 increasing trend in total volume loss and total energy dissipated was observed. This relationship was  
267 not observed in NiTi alloy under all polarisation potentials (Figure 5C and D).

#### 268 **Wear scar morphology**

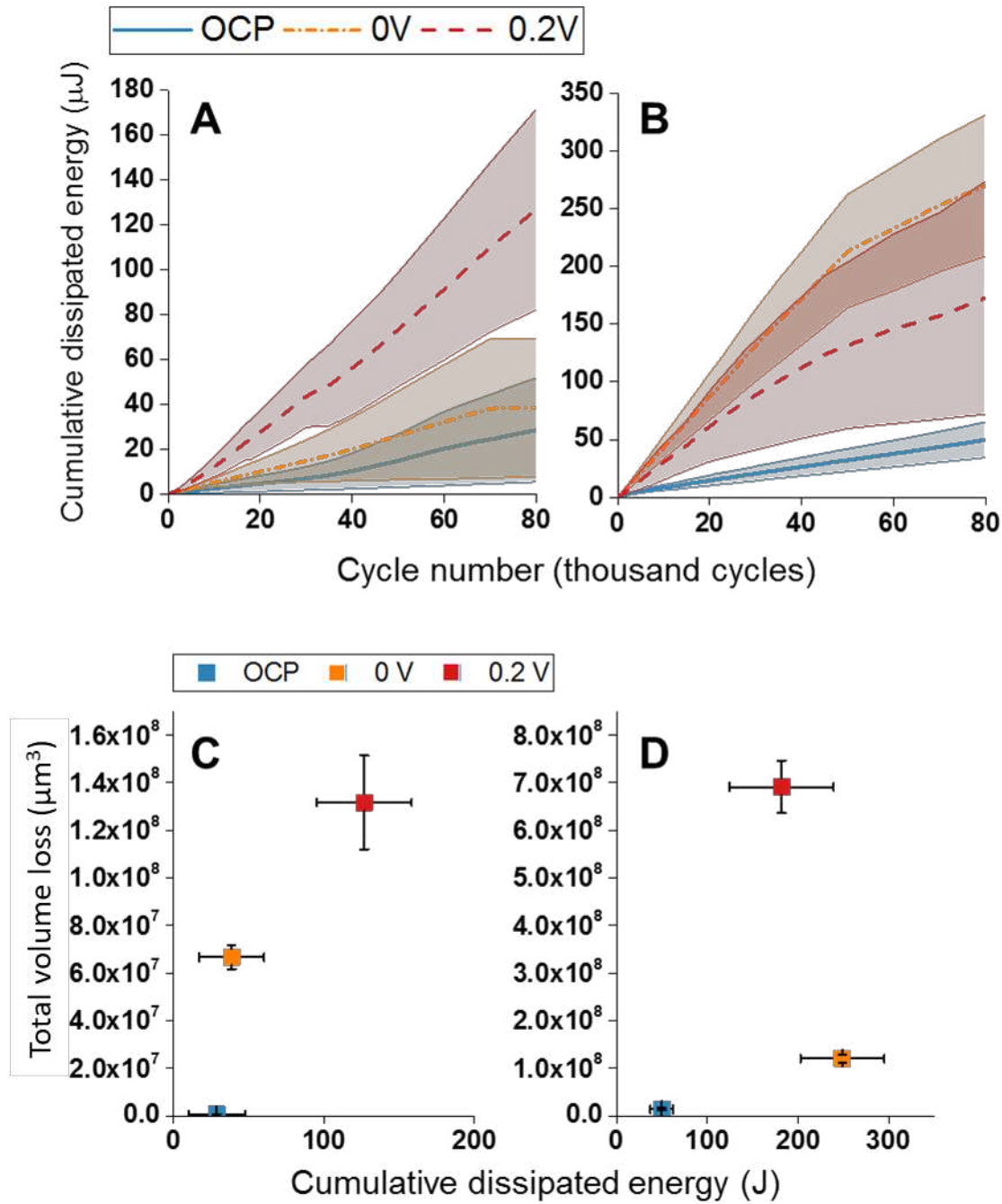
269 Representative wear scar morphology for both alloys under all polarisation conditions is shown with  
270 both OM and VSI images in Figure 6. Independent of the wear scar size, wear scar morphology is  
4.1. comparable over all polarisation conditions for CoCrMo alloy (Figure 6A, B, C). CoCrMo wear scar  
271 for all conditions is characteristic of a gross slip regime in agreement with fretting loop appearance  
272 (Figure 4A, B, C). The wear scar depth, as exhibited in the VSI images, appears to increase with  
273 polarisation potential. However for NiTi alloy, wear scar morphology appears to be affected by  
274 polarisation potential. At OCP, the wear scar directionality is relatively clear – parallel to the cylinder  
275 length for the top sample and perpendicular for the base (Figure 6D). As polarisation potential is  
276 increased, the wear scar surface became more heterogeneous and the directionality became  
277 indistinguishable at 0.2V (Figure 6E, F). CoCrMo alloy typically exhibited a classically abrasive wear  
278 scar morphology whilst NiTi showed an increasingly adhesive scar morphology with increasing  
279 potential. In summary, the inherent wear scar morphology was overtly altered in NiTi fretting by  
280 polarisation, but not for CoCrMo alloy.  
281

282



283

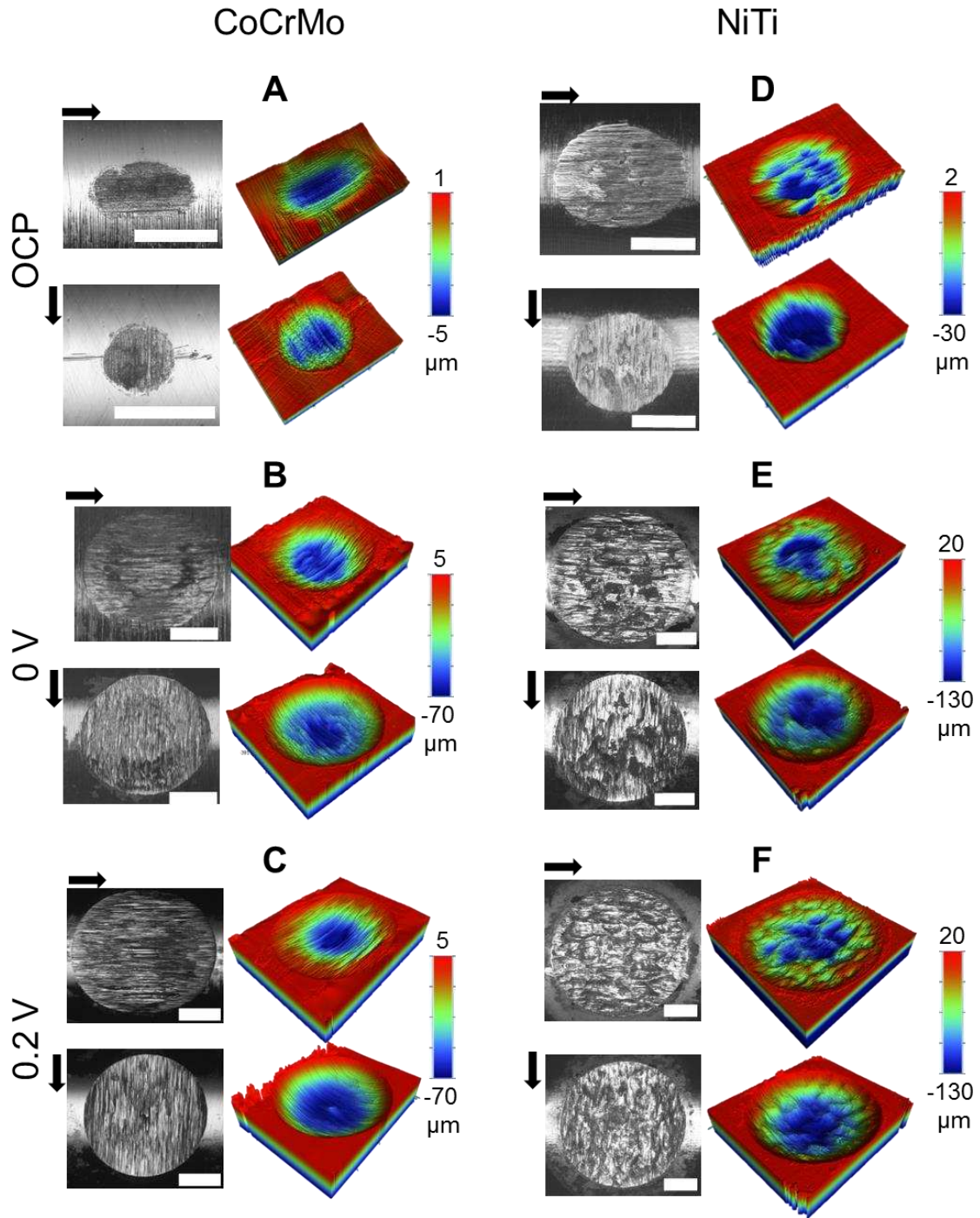
284 **Figure 4.** Representative fretting loops from the 50th and 50,000th cycle for CoCrMo and NiTi alloy  
 285 in polarisation regimes at OCP and held at 0V and 0.2V.



286

287 **Figure 5.** Cumulative dissipated friction energy over time for A) CoCrMo and B) NiTi alloy for fretting-  
 288 corrosion under OCP and held at 0V and 0.2V (n=3). Cumulative dissipated energy also shown as a  
 289 function of total wear volume for C) CoCrMo (red line = linear fit,  $\text{adj. } R^2 = 0.92$ ) and D) NiTi (red line  
 290 = exponential fit) alloy. Data shown is mean  $\pm$  95% confidence intervals (Panels A and B: shaded area).

291



292

293 **Figure 6.** Wear volume loss and wear scar morphology as shown by VSI and microscopy respectively  
 294 for CoCrMo and NiTi in all polarisation conditions. A) CoCrMo at OCP. B) CoCrMo at 0V  
 295 potentiostatic. C) CoCrMo at 0.2V potentiostatic. D) NiTi at OCP. E) NiTi at 0V potentiostatic. F) NiTi  
 296 at 0.2V potentiostatic. Scale bar for microscopy images = 500μm. Top and bottom images for each  
 297 panel represent top and bottom samples respectively. Sliding direction is horizontal for the top samples  
 298 and vertical for the bottom samples (arrows).

299

300

301

302 **Corrosion results**

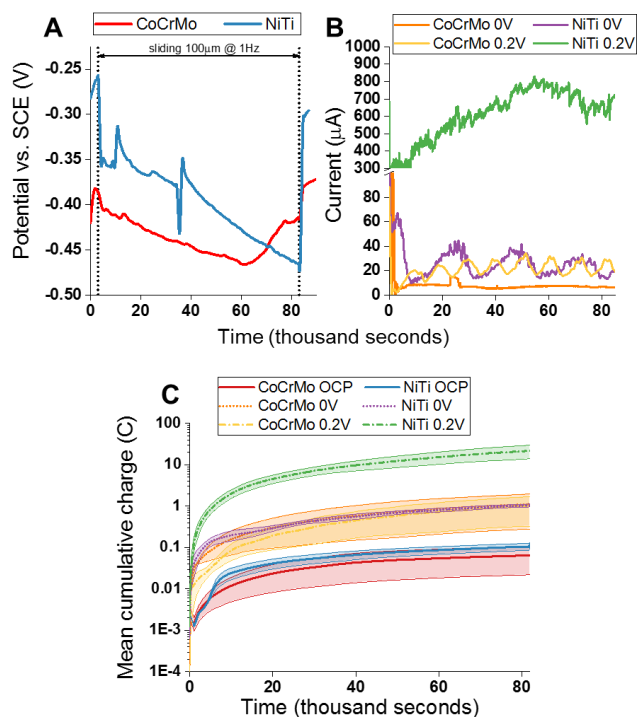
303 **4.2.1. Evolution of OCP**

304 Representative OCP readings for both CoCrMo and NiTi alloy are shown in Figure 7A. The OCP prior  
305 to initiation of sliding was more noble for CoCrMo when compared to NiTi. For both alloys, there is a  
306 sharp decrease in OCP upon initiation of sliding indicative of abrasion of the passive layer. There is a  
307 persistent decrease in OCP for CoCrMo and NiTi over time. Recovery of the OCP was observed when  
308 fretting was ceased although did not recover to values observed before the application of fretting  
309 suggesting changes in oxide chemistry and increased susceptibility to corrosion. (23, 29, 43, 51).

310 **4.2.2. Potentiostatic tests**

311 Representative current transients obtained under potentiostatic conditions (those held at 0V and 0.2V)  
312 are shown in Figure 7B. There was a consistent increase in current for both CoCrMo and NiTi at both  
313 0V and 0.2V upon initiation of sliding, indicating abrasion of the passive layer. For both alloys an  
314 increase in polarisation potential from 0V to 0.2V typically resulted in a consistently higher current  
315 although this increase was much starker in NiTi. Generally NiTi reached higher sustained currents than  
316 CoCrMo (Figure 7B). The cumulative charge for all experiments is given in Figure 7C. For CoCrMo,  
317 increasing the polarisation potential from 0V to 0.2V typically did not result in significant increase in  
318 cumulative charge. However, both polarised conditions have greater cumulative charge compared to  
319 experiments conducted at OCP. NiTi alloy was more sensitive to an increase in polarisation potential,  
320 resulting in a significant increase in both cases – particularly stark when increased to 0.2V.

321



322

323 **Figure 7. A)** Representative open circuit potential (OCP) readings for CoCrMo and NiTi cross cylinder  
 324 fretting corrosion. Samples were left to reach free potential for 1 h prior to initiation of sliding for  
 325 80,000 cycles at 1 Hz. Subsequently, samples were left for 1 h to recover. Linear polarization  
 326 measurements (OCP  $\pm$  0.02 mV) were conducted every 1000 s for the duration of the experiment. **B)**  
 327 Representative current transients for potentiostatic experiments held at 0V and 0.2V. **C)** Cumulative  
 328 charge for all experiments using Stern-Geary relationship with intermittent LPRs to give  $I_{\text{corr}}$  for OCP  
 329 experiments. Data shown is mean  $\pm$  SEM (shaded area).

330

4.3.

331

### Material loss

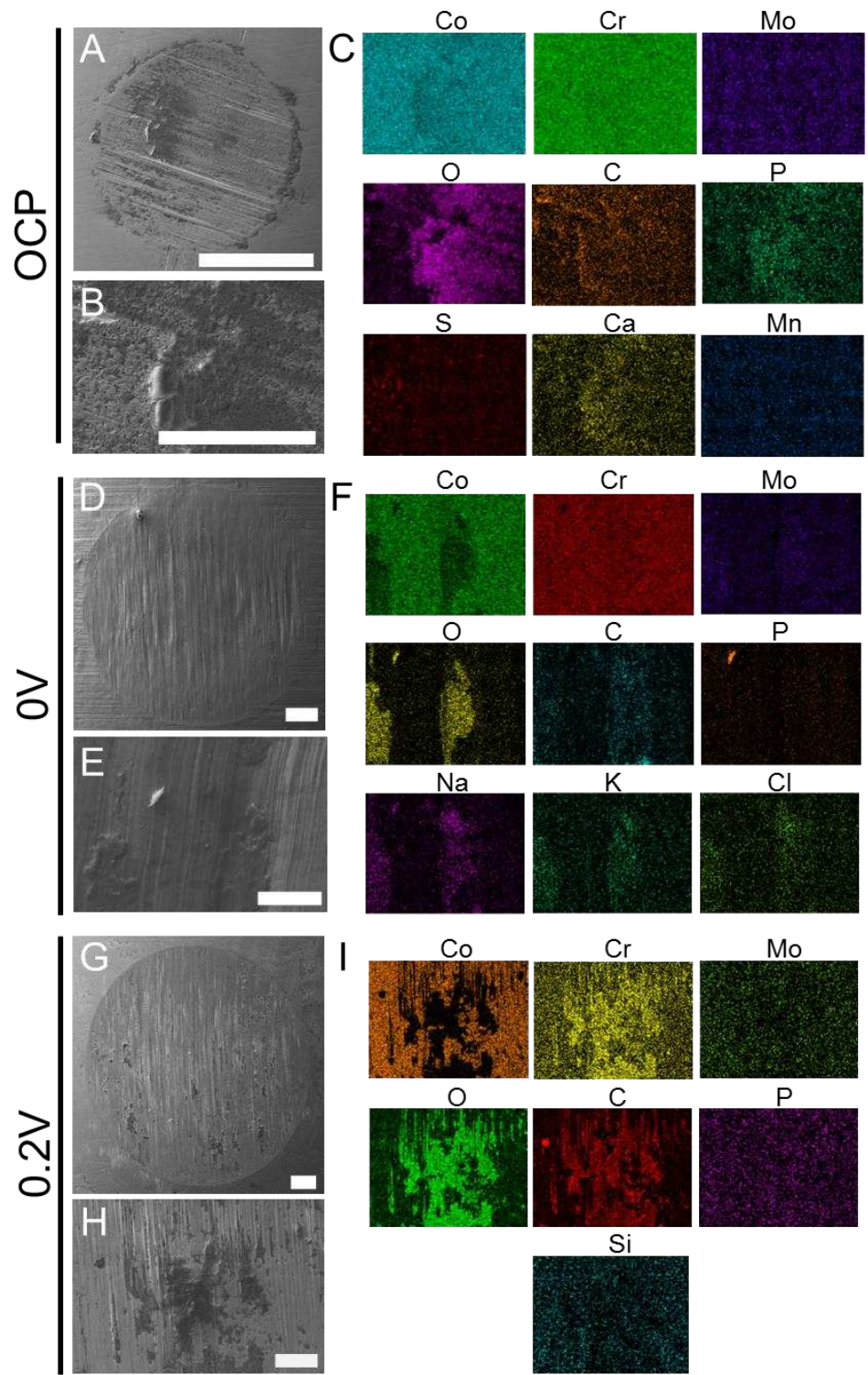
332 Total material loss ( $M_{\text{total}}$ ) was quantified using VSI and nominal alloy densities. The contributing  
 333 mechanisms to mass loss are presented later in this paper in Figure 13. There was no significant  
 334 difference in total mass loss between the potentiostatic regimes for CoCrMo alloy ( $0.73 \pm 0.42$  mg at  
 335 0V;  $1.21 \pm 0.49$  mg at 0.2V). Samples from both potentiostatic regimes had significantly greater total  
 336 mass loss vs. those which underwent fretting-corrosion at free corrosion potential ( $4.70 \pm 0.74$   $\mu$ g,  
 337  $p < 0.05$  vs. 0V and  $p < 0.001$  vs. 0.2V). NiTi alloy appeared to be particularly susceptible to an increase  
 338 in polarisation potential: total mass loss at 0.2 V ( $6.68 \pm 1.85$  mg) was significantly greater than that at  
 339 0V ( $0.79 \pm 0.06$  mg,  $p < 0.001$ ) and OCP ( $0.09 \pm 0.03$  mg,  $p < 0.001$ ). However, total mass loss at OCP  
 340 and 0V was not significantly changed. Total mass loss for both CoCrMo alloy and NiTi alloy were

341 relatively similar under both OCP and 0V regimes. However, at 0.2V NiTi had a mean 5.5-fold increase  
342 in total mass loss compared to CoCrMo alloy.

#### 343 **Wear scar chemistry**

344 Wear scar chemical composition and topography was analysed with SEM-EDX. Representative SEM  
345 images of the wear scar and SEM-EDX maps of the wear scar chemistry for each polarisation condition  
346<sup>4.4</sup> for CoCrMo are shown in Figure 8 and those for NiTi are shown in Figure 9. Elements such as  
347 phosphorus, potassium, calcium and chlorine were seen in the contact areas after fretting-corrosion. For  
348 CoCrMo alloy, increasing the polarisation potential from OCP up to 0.2V resulted in a stronger  
349 correlation of chromium with oxygen (Figure 8C, F, I). Molybdenum signal remained generalised over  
350 the whole wear scar under all polarisation conditions. At 0V only there was localised formation of  
351 phosphorous which was more generalised at both OCP and 0.2V. Under OCP conditions, a build-up  
352 formed at the edge of the wear scar which is noticeably absent under both 0V and 0.2V conditions.  
353 Interestingly, the element signals present differ between polarisation potential: calcium is only detected  
354 at OCP and potassium is solely detected at 0V.

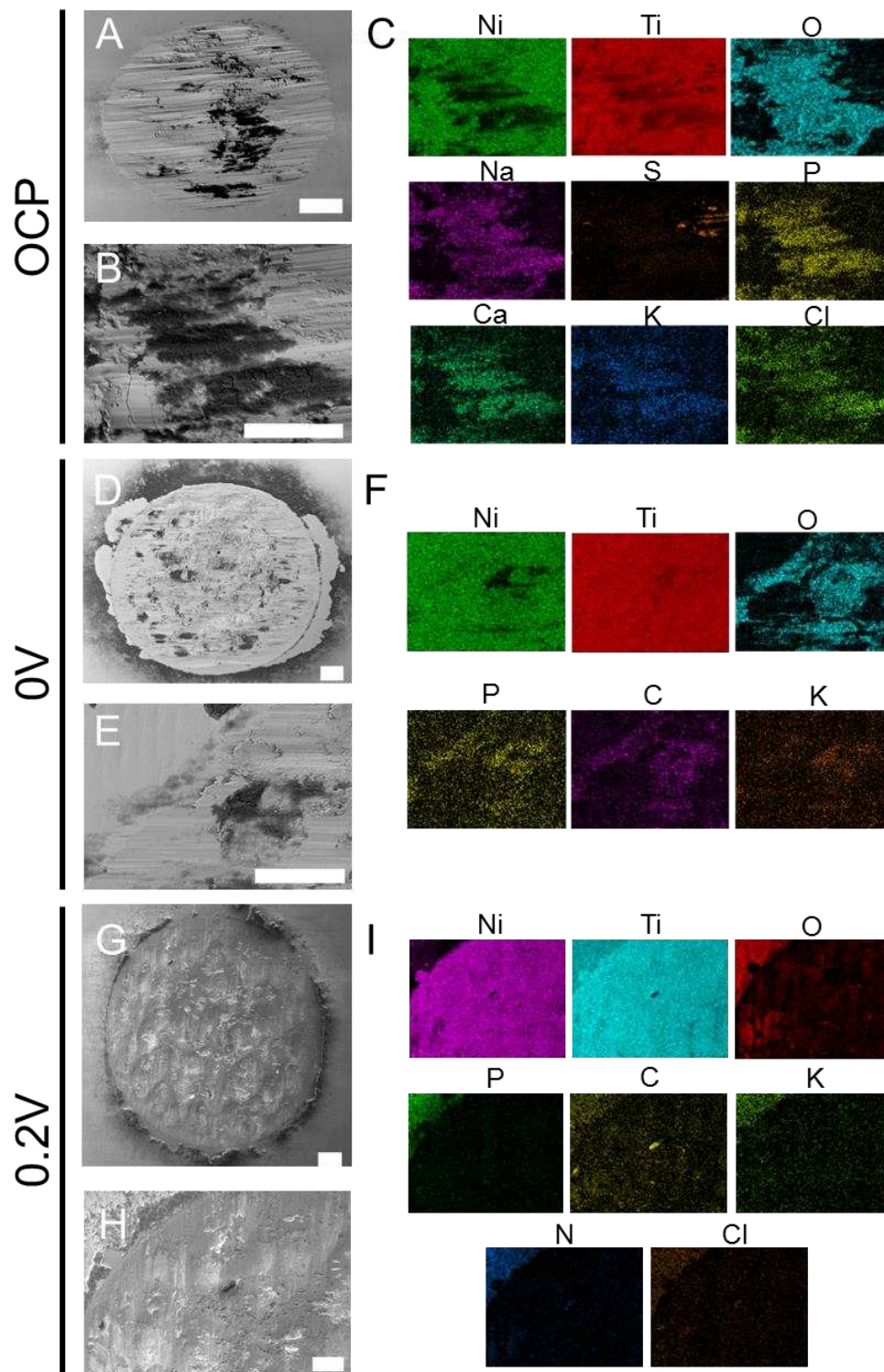
355 For NiTi alloy, in all polarisation conditions, the formation of complexes within the fretting contact  
356 were dominated by titanium (Figure 9; Panel C, F and I). Formation of titanium oxide film occurred  
357 generally over the wear scar, whereas sulphur and carbon complexes formed in specific locations at  
358 OCP and 0.2V respectively. Increasing the polarisation potential from instigates inherent changes in  
359 wear scar chemistry. At OCP, oxygen correlates with sodium, phosphorous, calcium, potassium and  
360 chlorine. However at 0V and 0.2V, carbon is detected which is not present at OCP. Conversely sodium,  
361 sulphur and calcium are only detected at OCP and are not present at 0V and 0.2V (Figure 9C, F, I).  
362 Contrary to CoCrMo, for NiTi a build-up of predominantly elements only contained within the serum-  
363 free media (O, C, P, K) rather than those contained within the bulk metal (Ni, Ti) can be observed at  
364 anodic polarisation conditions around the edge of the contact whereas this was absent at OCP (Figure  
365 9; Panels A, D and G).



366

367 **Figure 8.** Representative SEM images and SEM-EDX maps for CoCrMo alloy for all polarisation  
 368 conditions. A) SEM image of complete wear scar at OCP. B) SEM image of wear scar detail at OCP.  
 369 C) SEM-EDX map for wear scar chemistry at OCP. D) SEM complete wear scar 0V. E) SEM detail  
 370 wear scar 0V. F) SEM-EDX map 0V. G) SEM complete wear scar 0.2V. H) SEM detail wear scar 0.2V.  
 371 I) SEM-EDX map 0.2V. Panels A, D, G: scale bar = 200 μm. Panels B, E, H: scale bar = 100 μm.

372



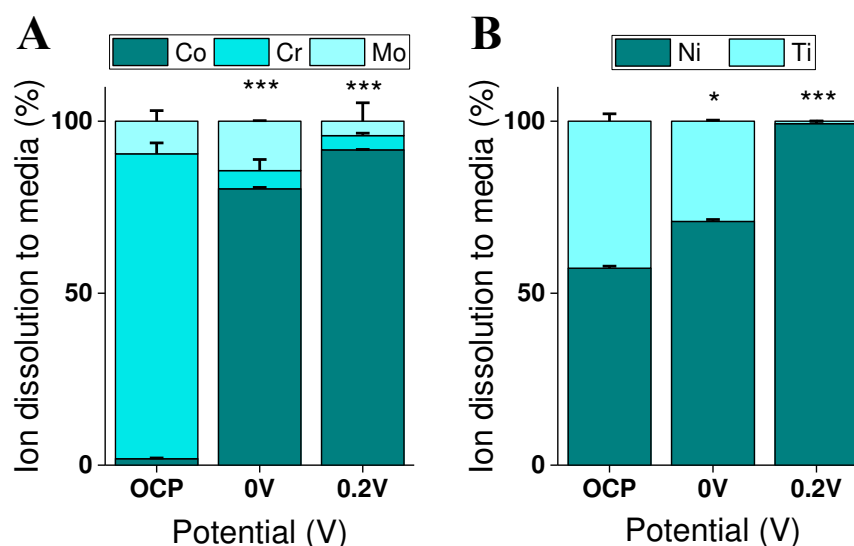
373

374 **Figure 9.** Representative SEM images and SEM-EDX maps for NiTi alloy for all polarisation  
 375 conditions. A) SEM image of complete wear scar at OCP. B) SEM image of wear scar detail at OCP.  
 376 C) SEM-EDX map for wear scar chemistry at OCP. D) SEM complete wear scar 0V. E) SEM detail  
 377 wear scar 0V. F) SEM-EDX map 0V. G) SEM complete wear scar 0.2V. H) SEM detail wear scar 0.2V.  
 378 I) SEM-EDX map 0.2V. Panels A, D, G: scale bar = 200μm. Panels B, E, H: scale bar = 100μm.

379

380 **Ion release into media**

381 Ion release to the electrolyte was determined through ICP-MS for CoCrMo and NiTi alloys at each  
382 polarisation condition (OCP, 0V and 0.2V) after centrifugation. The composition of ion release is shown  
383 in Figure 10. For CoCrMo, the total levels of ions released to the media was not significantly different  
384 across all polarisation regimes (11.1 ± 4.7 ppm, 2.9 ± 1.3 ppm and 10.5 ± 2.8 ppm for OCP, 0V and  
385 0.2V respectively). However the composition of the ion release was affected. Ion release for fretting-  
386 corrosion of CoCrMo was dominated by cobalt in both potentiostatic polarisation conditions; 80% at  
387 0V and 92% at 0.2V. Ion dissolution to media was dominated by chromium for CoCrMo alloy at OCP,  
388 at a level of 89% with Co dissolution dropping significantly to 2% (p<0.001).  
389 Total ion release for NiTi alloy was also not found to be different at OCP (0.22 ± 0.05 ppm) compared  
390 to 0V (1.9 ± 0.4 ppm). However, the total ion release at 0.2V (160 ± 79.6 ppm) was much higher and  
391 was significantly increased compared to both that at OCP and 0V (p<0.01). Ion release for NiTi alloy  
392 was approximately stoichiometric at OCP (57% Ni), but under electrochemical polarisation was  
393 dominated by Ni, increasing with polarisation potential; 71% at 0V and 99% at 0.2V. The proportion  
394 of Ni ions which were released into the media was significantly increased at both 0V (p<0.05) and 0.2V  
395 (p<0.001) versus OCP.



396

397 **Figure 10.** Proportion of ion release as measured through ICP-MS for A) CoCrMo alloy and B) NiTi  
398 alloy. \*p<0.05, \*\*\*p<0.001, one-way ANOVA with post-hoc Tukey test for A) Co ions and B) Ni ions.  
399 Data shown is mean ± 95% confidence intervals.

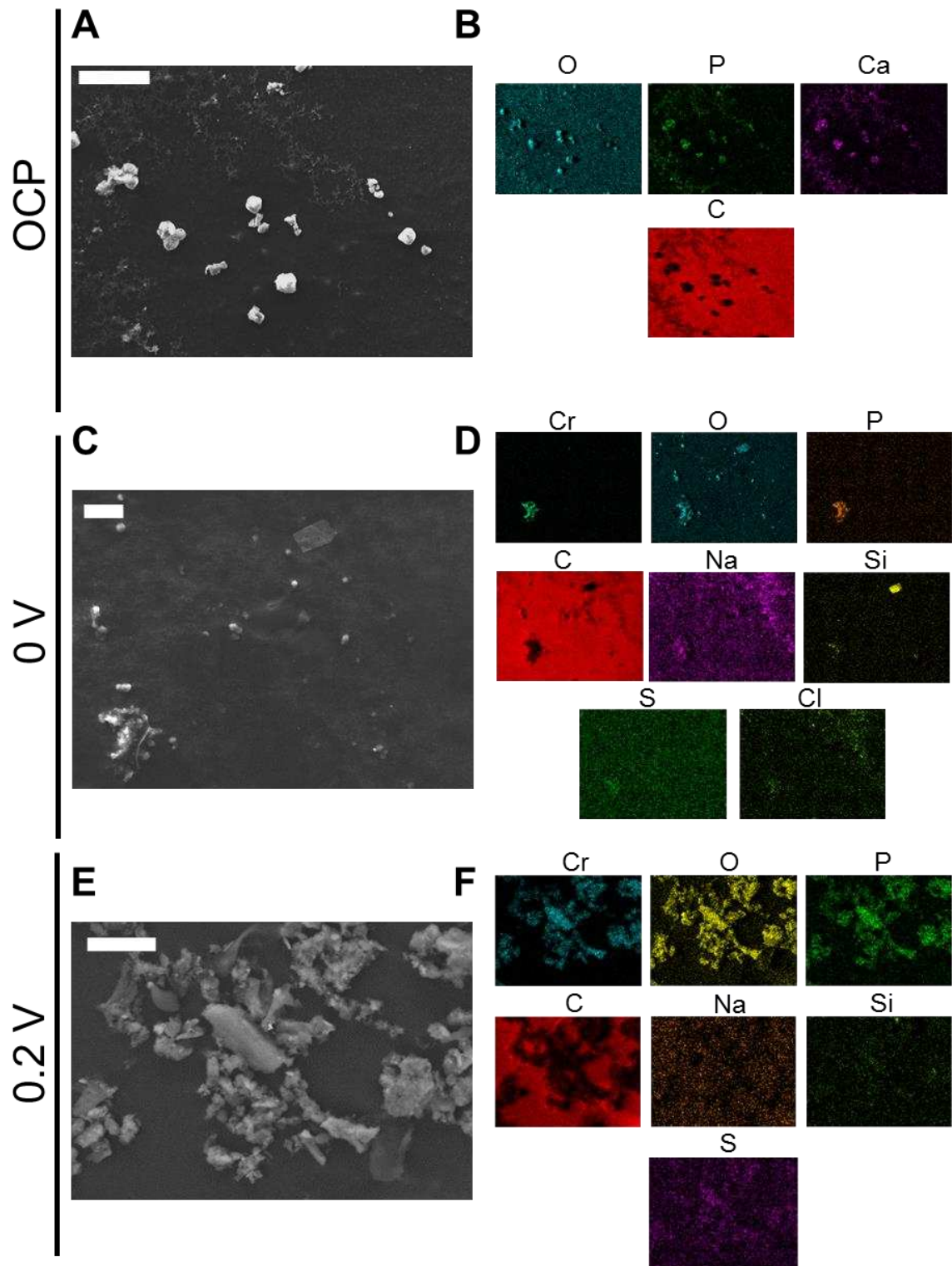
400

401        **Particle morphology and composition**

402 Particle morphology and composition was determined with SEM-EDX imaging. For CoCrMo alloy,  
403 increasing polarisation potential had a clear impact on particle morphology. At OCP, for all three tests  
404 which were performed, particles tended to be in the micron to sub-micron range and were typically  
405 smooth and relatively heterogeneous (Figure 11A). However, when analysed with SEM-EDX, it was  
406 not possible to identify any of the bulk metal elements (Co, Cr and Mo) in the particles after removal  
407 of soluble salts by washing with ultrapure water (Figure 11B). Instead, particles showed signals for  
408 oxygen, phosphorous and calcium which are all elements present in the serum-free culture medium  
409 electrolyte.

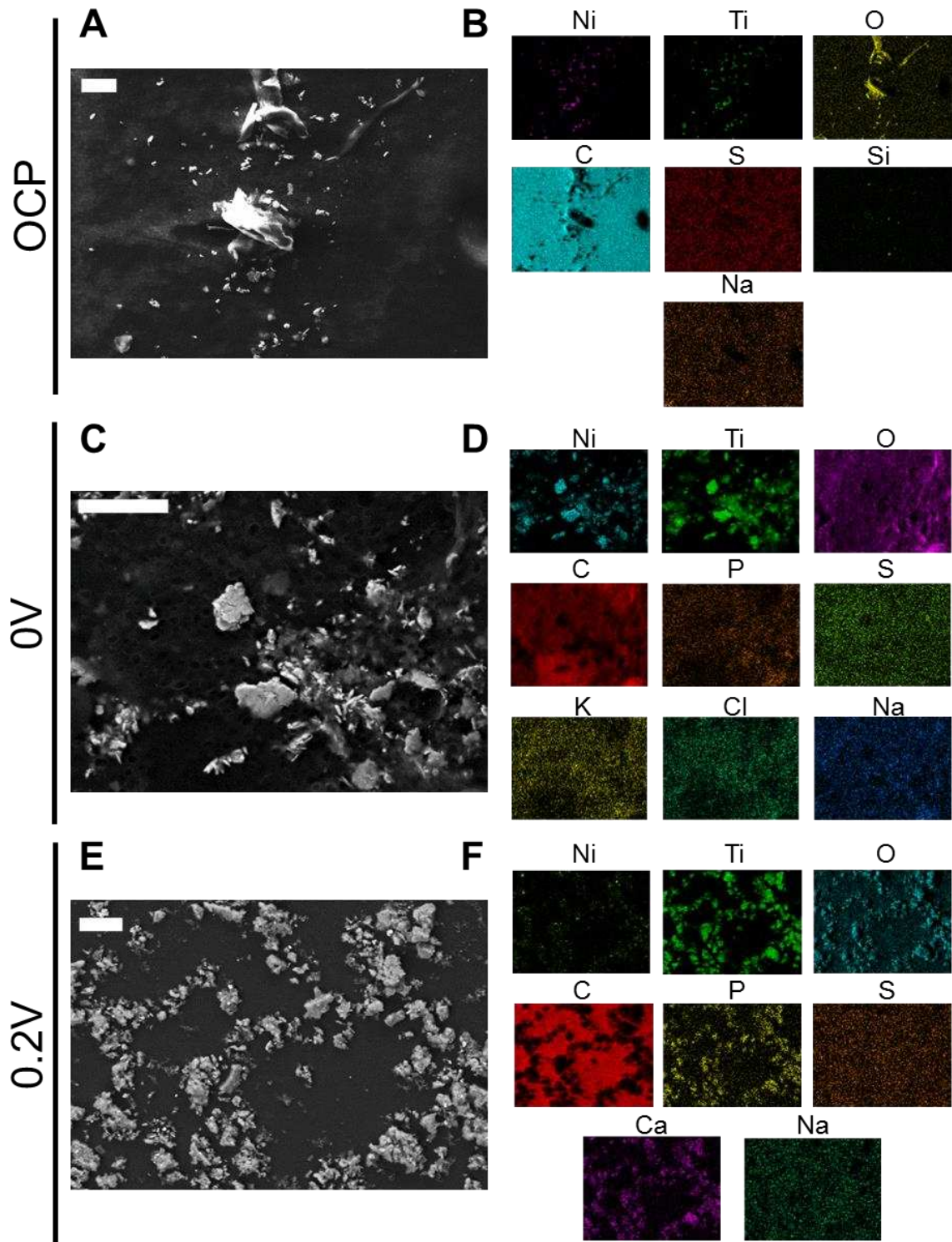
410 For the potentiostatic conditions, at 0V particles were typically in the micron to sub-micron range,  
411 although some larger ‘clusters’ were identified (Figure 11C). Chromium, oxygen and phosphorous rich  
412 particles were observed, with no evidence of cobalt being observed. Silicon was also detected which  
413 was not present in the wear scar interface of the bulk alloys (Figure 11D). At 0.2V, the particle  
414 morphology appeared to be less ‘particulate’ and was generally seen to have clusters of larger particles  
415 which were more irregular in shape (Figure 11E). The chemistry of the particles was relatively  
416 equivalent to that at 0V, with good correlation of chromium, oxygen and phosphorous and an absence  
417 of cobalt (Figure 11F).

418 For NiTi alloy, at OCP particles were typically smooth, uniform and in the single micron to nanoparticle  
419 range although this was occasionally interspersed with large multi-micron scale particles (Figure 12A).  
420 The smallest particles were comprised of bulk NiTi, demonstrating EDX signals for both nickel and  
421 titanium (Figure 12B). Larger particles showed oxide signals. At 0V, particle morphology was generally  
422 flakier; particles were rougher and flatter than in in OCP condition (Figure 12C). Particles ranged in  
423 nanometres to multi-micron in size. Composition at 0V polarisation was similar to that at OCP, where  
424 particles typically demonstrated nickel, titanium and oxygen signals (Figure 12D). Fretting tests at 0.2V  
425 polarisation were markedly different from the other conditions in that they tended to be larger and had  
426 a heterogeneous more ‘powdery’ appearance (Figure 12E). Nickel signals from EDX tended to be  
427 weaker than those from particles at other polarisation conditions and also showed complexing with  
428 calcium and phosphorus in addition to oxygen (Figure 12F).



429

430 Figure 11. Representative SEM images and SEM-EDX maps for wear particles from CoCrMo alloy  
 431 fretting experiments. A) SEM image of wear particles from CoCrMo fretting at OCP. B) SEM-EDX  
 432 map for particle chemical composition at OCP. C) SEM wear particles CoCrMo 0V. D) SEM-  
 433 EDX map wear particles CoCrMo 0V. E) SEM wear particles CoCrMo 0.2V. F) SEM-EDX map wear  
 434 particles CoCrMo 0.2V. Scale bar for all images = 25 $\mu$ m.



435

436 Figure 12. Representative SEM images and SEM-EDX maps for wear particles from NiTi alloy fretting  
 437 experiments. A) SEM image of wear particles from NiTi fretting at OCP. B) SEM-EDX map for particle  
 438 chemical composition at OCP. C) SEM wear particles NiTi 0V. D) SEM-EDX map wear particles NiTi  
 439 0V. E) SEM wear particles NiTi 0.2V. F) SEM-EDX map wear particles NiTi 0.2V. Scale bar for all  
 440 images = 25 $\mu$ m.

441

442

443 **5. Discussion**

444 This study has investigated the role of fretting -corrosion on the degradation of materials commonly  
445 used in biomedical applications. The resultant ionic and particulate debris has been quantified as a  
446 function of applied over-potential. Fretting-corrosion at the stent-strut interface has been observed  
447 clinically, with the roles of metal ion and particle release hypothesised to contribute to ISR processes  
448 (16). Whilst the role of metal ion release into the cardiovascular environment and prevalence in clinical  
449 complications is not fully understood, it is accepted that toxic and sub-toxic release of metal derived  
450 debris may induces some adverse clinical effects. Furthermore, pre-clinical assessment studies typically  
451 treat wear and corrosion in isolation (10, 14), only revealing part of the story concerning the degradation  
452 of metallic surfaces. Key findings from this study can be summarised as follows:

- 453 i) The synergistic interaction of wear and corrosion are inevitable at contacting metal-metal  
454 interfaces when subjected to micro-motion in a conductive biological environments. The  
455 presence of fretting increasing the rates of corrosion. This leads to the production of metallic  
456 ions and particulates; the nature of which are dependent on the tribological and corrosive  
457 conditions applied.
- 458 ii) The electrochemical parameters used to facilitate measurements of current transients  
459 associated with fretting-corrosion can significantly affect the mechanisms of degradation and  
460 the nature of the debris generated. This raises question around the suitability of imposing  
461 electrochemical potentials in the preclinical assessment of materials and how we interpret such  
462 5.1. data.

463 **The role of applied potential on degradation mechanisms**

464 The degradation mechanism in fretting-corrosion arises from a synergistic interaction between  
465 mechanical wear ( $M_{\text{mech}}$ ) and corrosion due to intermittent depassivation ( $M_{\text{chem}}$ )(29). Considering this,  
466 the relative contributions to total material loss in fretting-corrosion can be estimated using the Equations  
467 3 – 5.  $M_{\text{mech}}$  is subdivided into  $M_{\text{wear}}$ , a purely mechanical component, and  $M_{\text{cw}}$  representing the synergy  
468 between corrosion and wear (corrosion-enhanced wear). Equally  $M_{\text{chem}}$  is subdivided into  $M_{\text{corr}}$ , a purely  
469 chemical component, and  $M_{\text{wc}}$  representing the synergy between wear and corrosion (wear-enhanced

470 corrosion)(45). This approach is widely use in the area on tribocorrosion to estimate material mass loss  
 471 contributions. Bryant and Neville (45) have recently provided a detailed analysis of the approaches in  
 472 calculating material loss mechanisms in fretting-corrosion contacts and the limitations of this approach  
 473 (45, 52). Corrosion currents (Figure 7) were converted into corrosion mass losses where ‘m’ is the mass  
 474 liberated at the anode, ‘Q’ is the charge passing through the WE, ‘M’ is the molar mass of the element,  
 475 ‘z’ is the dissolution valence and ‘F’ is Faraday’s constant equal to 96,480 C mol<sup>-1</sup>. For each alloy, it  
 476 was assumed that there was stoichiometric dissolution of the component elements, giving molar mass  
 477 (M) as 58.0 g.mol<sup>-1</sup> for CoCrMo and 53.8 g.mol<sup>-1</sup> for NiTi. Equally, the dissolution valence (z) was  
 478 assumed to be 2.0 for CoCrMo and 2.9 for NiTi. Applying Faraday’s law to the electrochemical data  
 479 recorded throughout the duration of the fretting-corrosion experiments allowed conversion to mass loss  
 480 equal to chemical wear (M<sub>chem</sub>). The total mass loss (M<sub>total</sub>) was determined using VSI and known alloy  
 481 density and their difference represented mass loss due to mechanical wear (M<sub>mech</sub>).

$$M_{total} = M_{mech} + M_{chem} \quad (3)$$

$$M_{mech} = M_{wear} + M_{cw} \quad (4)$$

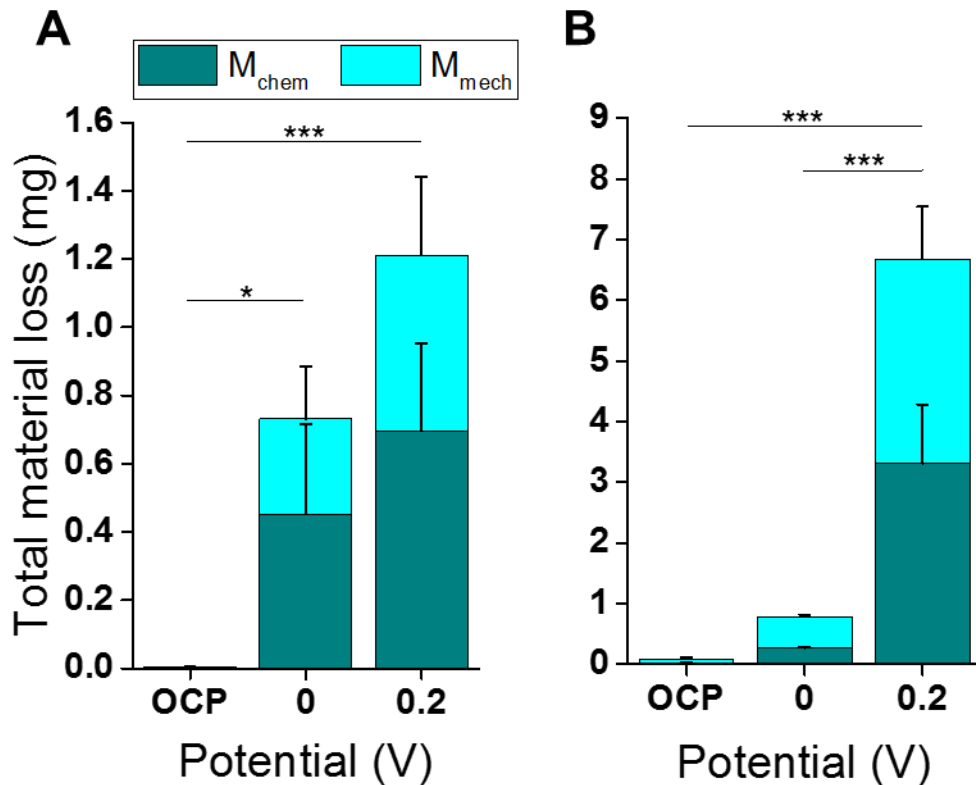
$$M_{chem} = M_{corr} + M_{wc} \quad (5)$$

$$m = \frac{Q \times M}{z \times F} \quad (6)$$

482  
 483 Figure 13 shows the total mass loss (M<sub>total</sub>) and their contributors using the approach described above  
 484 (Equation 6) for CoCrMo and NiTi alloys. At OCP, 0V and 0.2V, CoCrMo material loss mechanisms  
 485 were dominated by corrosion related processes. However with increasing applied polarisation potential  
 486 the role of mechanical contributors in the fretting-corrosion system increased (Figure 13A). As  
 487 mentioned above, M<sub>mech</sub> still contains a corrosive contributor. Corrosion enhanced wear processes,  
 488 whereby mechanical material losses are enhanced through corrosion processes (e.g. by increased  
 489 surface roughness), may contribute to this increase. With increased applied potential, higher currents  
 490 were observed which will in turn influence the nature of topography within the contact as evidenced in  
 491 Figure 6.

492 Conversely, NiTi operated in a regime where  $M_{\text{mech}}$  was dominant, with corrosive contributions  
493 increases with applied potential due to the increased over-potential driving the magnitude of current  
494 evolved during abrasion. An increase in polarisation potential appears to increase the susceptibility of  
495 NiTi to corrosive-dominated wear as demonstrated by the increase in corrosion current coupled with no  
496 change in dissipated frictional energy. Nevertheless, the wear degradation pathway tended towards a  
497 more wear-dominant regime with increasing polarisation potential as exhibited by the relative  
498 contributions to wear (Figure 13). However, the fretting-corrosion system exists as a synergy of wear  
499 and corrosion: although the corrosion currents exponentially increased with polarisation potential,  
500 mechanical depassivation is still required for this wear-enhanced corrosion. Lukina *et al.* demonstrated  
501 that under fretting corrosion, NiTi is particularly susceptible to very high corrosion currents compared  
502 to other alloys such as biomedical titanium and CoCrMo (53). Kosec *et al.* also concluded NiTi was  
503 susceptible to increased material degradation under wear-corrosion synergy conditions as opposed to  
504 static corrosion due to cyclic depassivation of the oxide film (54). In the case of overlapping NiTi stents,  
505 Trépanier *et al.* showed that the effect of wear on corrosion was comparable between stainless steel and  
506 NiTi (55). However, in this case, the wear-corrosion synergy was not considered and specimens  
507 underwent fretting wear damage and then were subsequently subjected to cyclic polarisation testing.  
508 NiTi is comparable to stainless steel in its ability to repassivate wear damage to the stent surface, but  
509 the present study shows that NiTi is more susceptible than CoCrMo to the wear-corrosion synergy under  
510 polarisation.

511



512

513 **Figure 13.** Relative contributions to total wear in terms of mechanical and chemical degradation  
 514 mechanisms for A) CoCrMo and B) NiTi alloy in three polarisation regimes (OCP, 0V and 0.2V) during  
 515 80,000 fretting cycles (n=3). Data shown is mean  $\pm$  95% confidence intervals. Total wear significance  
 516 shown with one-way ANOVA with post-hoc Tukey test, \* $p < 0.05$ , \*\*\* $p < 0.001$ .

517

## 518 5.2 The roles of applied potential on fretting mechanisms

519 The effect of applied over potential on the tribology at an interface has been well documented in sliding  
 520 contacts. Ponthiaux et al (56) showed the variation of the coefficient of friction,  $\mu$ , at different applied  
 521 potentials and hypothesised the coefficient of friction to be linked to lowest zero current potential  
 522 because ‘mechanical depassivation in the contact area is at maximum and leads to the largest total active  
 523 area’. They also hypothesised that higher coefficient of friction can be associated with passivity. Within  
 524 a biotribological context, Yan et al (51) demonstrated the roles between applied potential and the  
 525 coefficient of friction, attributing the complex behaviour to tribo-chemical processes occurring at the  
 526 metal – protein level.

527 A similar observation has been observed in this study. For CoCrMo alloy subjected to fretting-  
 528 corrosion, tangential forces and cumulative dissipated energy were seen to increase with increasing

529 applied potential. This is in line with other tribocorrosion studies on CoCrMo where the coefficient of  
530 friction correlates with applied over-potential (51, 57). In our case we have considered dissipated  
531 energy, which considers frictional forces and displacement at the interface, due to the difficulties in  
532 determining friction within a fretting contact (58-60). Interestingly, when CoCrMo was subjected to  
533 polarisation at 0 and 0.2V, the tangential forces increased with increasing number of cycles (Figure 4B  
534 and 4C). It is conceivable that frictional forces increase with increasing number of cycles due to the  
535 development of the contact area over the 80,000 cycles. The final nominal contact area is considerably  
536 larger than the initial Hertzian contact area calculated with an increased confirmation of the surfaces.  
537 It is evident that the development of the contact area is linked to the corrosion conditions at the contact  
538 and will be enhanced by the accelerated corrosion at the interface. In the literature, coefficients of  
539 friction ranging from  $\mu = 0.1$  to 1 have been quoted for CoCrMo in sliding interfaces (57). Furthermore  
540 the formation of and retention of oxides within the contact will influence the evolving tribology;  
541 although their roles in a fretting-corrosion contact past third body effects are not well understood and  
542 significantly different in nature to tribochemical reactions in sliding contacts.

543 In the case of NiTi alloy, the effect of applied potential from 0V and 0.2V was not as apparent as for  
544 CoCrMo alloy. All NiTi fretting loops showed a gross-slip regime, with some interesting observations  
545 between those at OCP and at an applied potential. A slight decrease in  $\frac{\delta_s}{\delta_d}$  was observed when compared  
546 to CoCrMo-CoCrMo interfaces likely due to the decreased reduced modulus of the NiTi-NiTi contact.  
547 As a result, a component of elastic deformation was introduced at the interface, although remained well  
548 within the gross-slip criteria as outlined by Fouvry (44). For all three instances of the experiment  
549 conducted at OCP the fretting loop shape exhibited a ‘tail-like’ region on the forward stroke. The  
550 reasons for this are not fully understood, but there are some links between the mechanical and  
551 electrochemical processes evident. At the wear scar interface there was very localised formation of  
552 sulphurous complexes solely at OCP, perhaps going some way towards explaining the unusual fretting-  
553 loop shape – as they are so localised, the tangential force peaks over the course of the fretting cycle as  
554 it interacts with them. Fretting loops at 0 and 0.2V presented a more conventional appearance with  
555 evidence of plastic pile up at the edges of the contact, causing the tangential force to increase rapidly

556 towards the end of the stroke (59). This is supported by Figure 9 which shows evidence of debris build-  
557 up at the edges of the contact. Due to the lack of fretting loop analysis for NiTi in physiological media,  
558 this phenomenon cannot be entirely explained. The root cause for this is unclear and further work to  
559 understand if this relates to a strain related transformation of the NiTi or localised build-up of debris  
560 within the contact is needed. No trend in the magnitude of tangential forces at the different polarisation  
561 conditions and with increasing number of cycles could be observed, in contrast to CoCrMo. Overtime,  
562 the rate of cumulative energy dissipation was seen to decrease suggesting a change in the degradation  
563 rate or mechanism at the interface. This may arise from the unique austenitic – martensitic strain-rate  
564 dependant transformation characteristics that NiTi alloys possess (61). It is conceivable that cyclic  
565 loading and fatigue at the interface will induce cumulative changes in the subsurface micro-structure  
566 effecting the evolution of mechanical and corrosive damage over time.

### 567 **5.3 The role of applied potential on ionic and particulate debris formation.**

568 In both metallic alloy cases, the debris and ion release was altered in terms of both chemistry and  
569 morphology with the application of electrochemical over-potential. The elements making up the bulk  
570 metal substrate (Co, Cr and Mo) were not detected when undertaking SEM-EDX analysis at OCP, but  
571 instead provided signals for O, P and Ca. These were also present on the wear scar surface, indicating  
572 that at OCP the removed wear debris consists of the passive layer formed continuously from abrasion  
573 and retained in the contact during fretting (62). With the application of over-potential, the elemental  
574 composition of the debris was seen to vary when compared to debris retrieved OCP. Cr, O and P rich  
575 debris was observed, similar to that retained within the contact. A similar observation was seen of the  
576 NiTi, with the application of over-potential influencing the nature and chemistry of the debris produced.  
577 At OCP, debris mainly consisting of Ni, Ti, and O was observed. The nature of the debris produced was  
578 seen to be altered with the application over potential during fretting-corrosion, with Ti, O, P and Ca  
579 rich particulates being observed at 0.2V. Without a doubt, the application of an applied electrochemical  
580 potential influences the nature of the debris generated at an interface, intrinsically linked to the changes  
581 tribological processes discussed above and the changes in passive film chemistry. It is well established  
582 that the application of over-potential influences the nature and quantity of preferentially absorbed

583 species on a surface, the thickness of the oxide film and thermodynamic driving forces for specie  
584 migration across the oxide layer. All of these factors will synergistically interact with the tribological  
585 processes, influencing the evolving degradation mechanism and nature of the debris produced. Whilst  
586 there is not a great deal of literature observing the nature and chemistry of debris derived from fretting-  
587 corrosion interfaces, the observations for CoCrMo correlate well with particle analysis retrieved from  
588 in-vivo orthopaedic implants. Cr and P rich particles have been observed in peri-implant soft tissues  
589 and hypothesised to arise from fretting interfaces which is consistent with this study (63). The nature  
590 NiTi debris is still yet to be fully understood.

591 Perhaps one of the most interesting observations made in this study is the variation in ion release  
592 stoichiometry with applied over-potential. For CoCrMo-CoCrMo fretting interfaces, the ion release  
593 stoichiometry changed from chromium-dominated to cobalt-dominated with the proportion of cobalt  
594 ions increasing with polarisation potential. Similarly for NiTi-NiTi interfaces, this went from a near  
595 stoichiometric release of Ni and Ti to a Ni dominated solution. This may be explained by the reliance  
596 of biomedical alloys to form a passive surface layer to protect against corrosion, the changes in the  
597 oxide film characteristics and nature of species absorbed at the interface as a result of polarisation (26).  
598 In the particular case of CoCrMo and NiTi, this is typically a chromium-oxide and titanium-oxide rich  
599 layer respectively (64, 65). Cyclic abrasion of the surfaces and retention of the debris within the  
600 interface is supported by both the wear scar chemistry and that of the wear particles. In both cases a  
601 correlation between the elemental composition of the debris and stoichiometry could be observed  
602 suggesting. For CoCrMo-CoCrMo contacts, where Cr rich debris was observed (Figure 11C and E),  
603 Cobalt dominated ion release was observed. This suggest that the mobility and of Co and Cr at the  
604 interfaces differs based on the electrochemical and tribological conditions at the interface. A similar  
605 observation was made for NiTi-NiTi interfaces. Ni dominated ion release was observed at 0.2V whereas  
606 the debris was Ti, O, P and Ca rich. In both cases, Cr and Ti are thermo-dynamically favourable  
607 reactions contributing the reformation of oxide layer once abraded (66). An initial release of Ni and Co  
608 will likely occur when the nascent metallic surface is exposed to the electrolyte until an effective oxide  
609 layer is formed, enabling the soluble Ni and Co to migrate from the interface. This will depend on the  
610 electrode potential and pH at the interface. This further points to the importance of considering the

611 nature of the test environment and experimental conditions on the debris generation mechanisms at an  
612 interface.

613 The enhanced metal ion release due to tribological processes has been reported mainly for sliding  
614 contacts, which significantly differ to fretting-corrosion interfaces. Espallargas *et al.* showed that ion  
615 release was dominated by cobalt under sliding-corrosion at OCP and cathodic and anodic polarisation  
616 (35). However, the interface in this study was a sliding (6 mm stroke length) CoCrMo-alumina interface  
617 as opposed to a fretting CoCrMo-CoCrMo contact. In a hip implant head/cup large sliding interface ion  
618 dissolution was shown to be stoichiometric rather than cobalt-dominated(67). Molybdenum in terms  
619 of ion release, wear scar chemistry and presence in wear particles also appeared to be relatively  
620 unaffected by polarisation. However it has previously been shown that molybdenum dissolution is  
621 significantly impacted by presence of proteins which were absent from this study (35, 49).

#### 622 5.4 Concluding remarks

623 Metallic biomaterials are commonly and widely used for cardiovascular stents and well as other areas  
624 of biomedical engineering. Although there is *in vivo* evidence of wear debris release at the strut interface  
625 of braided designs (12, 68), to the authors' knowledge there is no literature detailing the links between  
626 fretting-corrosion processes and the nature of the resultant debris. To investigate tribocorrosion  
627 mechanisms in biomaterials, studies have used potentiostatic polarisation to exacerbate material  
628 degradation during fretting. This study shows that the nature of the wear debris, from particle chemistry  
629 and morphology to ion release, is affected by the polarisation potential. In the case of CoCrMo, the  
630 wear debris was intrinsically altered during polarisation under fretting despite the maintenance of the  
631 tribological and electrochemical characteristics. However NiTi was particularly susceptible to wear-  
632 enhanced corrosion as polarisation potential was increased, affecting the tribocorrosion mechanism  
633 itself and the wear debris.

634 Gaining an appreciation of the stent in the physiological environment is imperative to elucidating the  
635 material degradation mechanisms occurring in stents *in vivo* and the biological impact of released  
636 debris. As proven in hip orthopaedic implants, wear debris and its morphology and composition may  
637 have unexpected and devastating consequences for patients (14, 17, 69). In the case of vascular stents,

638 it has been shown that fretting-corrosion and local release of wear debris to tissues occurs *in vivo* (12,  
639 68). This is hypothesised to be a compounding factor in the most common mechanism of stent failure,  
640 in-stent restenosis which is a biologically mediated phenomenon. Studies aiming to understand the  
641 biological effects of fretting-corrosion must therefore attempt to replicate the physiological scenario.  
642 This study shows that undertaking fretting-corrosion experiments under potentiostatic control as  
643 opposed to at the materials open circuit potential both CoCrMo and NiTi alloys affected the degradation  
644 mechanisms and released wear debris. It can be appreciated that useful triboelectrochemical techniques  
645 may inherently alter the system. Future work in this area will be focussed on the biological impact of  
646 wear debris under different polarisation regimes and the compounding factors of presence of serum  
647 proteins and inflammatory mediators.

## 648 **6. Conflicts of interest**

649 The authors have no conflicts of interest to disclose.

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652

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