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Fretting-corrosion between 316L SS and PMMA: influence of ionic strength, protein and electrochemical conditions on material wear. Application to orthopedic implants.

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Abstract

In biomedical field, fretting-corrosion between 316L SS femoral stem and bone cement is one of the significant causes of the hip prosthesis loosening. This article investigates wear by fretting-corrosion at the contact between 316L and PMMA . The influences of the ionic strength (NaCl solutions from 10⁻³ to 1 mol.L⁻¹), a model protein (albumin) and electrochemical conditions on contact behaviour are studied. At OCP (Open Circuit Potential) conditions, the chlorides concentration, i.e. the ionic strength, increases the 316L wear; and albumin, concentration of 1g.L⁻¹, does not play a significant role in total 316L wear. At cathodic applied potential E = -400 mV(SCE), a threshold concentration of $10^{-1} \text{ mol.L}^{-1}$ (NaCl solution), C_{th}, indicates two behaviours: a protective effect below C_{th}, and an additional anodic dissolution above C_{th}. One might suggest that, beyond C_{th}, the passive layer is not efficient for protecting against the corrosion. At this potential, albumin reduces wear due to corrosion and amplifies mechanical wear induced by corrosion. Albumin seems to act as an anodic inhibitor. To determine the mechanisms of synergism, a "more cathodic" potential is applied, E = -800 mV(SCE), during fretting-corrosion experiments. Consequently, the corrosive wear can be neglected and the mechanical wear can be only measured.

Keywords: Fretting-corrosion, 316L SS, PMMA, Ionic strength, Albumin.

1. Introduction

In Europe, each year, there are about 1 million hip replacements, and approximately the same number in USA. Nevertheless, after 10-15 years, approximately 10% of Total Hip Arthroplasties (THA) need a reintervention. Nowadays, the ageing of the population, injuries or diseases like arthrosis are public health issues. To avoid additional surgical reoperation and improve lifetime duration of implants, studies on orthopedic implants

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are needed. Three different metal alloys can be used for femoral stem: cobalt-chromium alloy, titanium alloy or austenitic stainless steel. We focus our attention on cemented hip prostheses. In 1960, in order to decrease the number of femoral stem debonding, Sir John Charnley decided to insert hip prostheses with poly(methyl methacrylate) called bone cement [1]. For a cemented prosthesis (Fig. 1), the austenitic stainless steel (316L SS alloy) is usually used for femoral stem thanks to its biocompatibility and high resistance against corrosion in biological environment without friction.

Because of the daily human gait and significant difference of mechanical properties (particularly, Young's moduli) between 316L SS and bone cement, stress shielding phenomenon appears. The metallic material and the polymer do not have the same strain properties, thereby leading to debonding between the femoral stem and the cement. It involves fretting (friction under small displacements, term precisely defined later) and friction. Finally, the upper part of femoral bone is unloaded and then becomes more susceptible to fracture; on the other hand, if bone around femoral stem tip is overloaded, it becomes stronger [2]. However, the bone cement acts as a "bumper" between femoral stem and bone, leveling peak forces acting around the prosthesis during the human gait.



Fig. 1 Total hip joint cemented prosthesis components

Femoral stem debonding and consequently friction between femoral stem and bone cement leads to cement degradation and cracking (Fig. 1). Debris from bone cement and femoral stem (metal oxides and ions) go through cement up to bone tissues [3]. Thus, the disturbance of osteoblasts cell activity and the tissue inflammation, respectively caused by bone cement particles and metallic oxides and ions, involve implant reoperation. It is the reason why fretting-corrosion between femoral stem and bone cement, particles generation, is one of the most

important causes of reintervention [4,5]. Consequently, fretting-corrosion experiments were carried out for understanding corrosion and mechanical influences, and improving the lifetime of implants.

Fretting means friction under small reciprocal displacements, and the amplitude of displacement is lower than contact width. Rubin *et al.* found out, by numerical modeling, the amplitude of displacement is about 50 μ m (depending on the location) between bone and stem under applied stresses during human gait [6]. Nonetheless, the displacement amplitude between bone cement and femoral stem is supposed identical [7].

Wear of a metallic material under the conjugated effects of a mechanical solicitation (friction or fretting in this case) and of corrosion by a medium (physiological liquid) is named tribocorrosion [8]. Thus, fretting-corrosion is fretting in a corrosive environment for a metal alloy. This phenomenon leads to passive layer destruction and debris generation, principally metal oxides (Fig. 2) and polymer particles [9-11].

The fretting-corrosion phenomenon between femoral stem and bone cement is modelized by frettingcorrosion between a 316L SS sample (the same material as that of femoral stem) and PMMA which has the same mechanical properties as bone cement. This polymer was chosen because of its optical transparency. In that way, a direct observation of the contact zone and the debris evolution between materials at contact are investigated.



Fig. 2 (a) Fretting-corrosion between femoral stem (Exeter V40TM) and bone cement [7]; (b) fretting-corrosion between a head (316L) and a femoral neck (Ti-6Al-4V)

Physiological liquid components are various, including numerous ions and proteins. Each component plays a different role in double layer constitution at material surface. To separate the contribution of each component, fretting-corrosion tests between 316L and PMMA were investigated in different solutions. The first one is Ringer solution. Its composition is close to physiological liquid. However, this solution is too complex: each ion may play a different role during a fretting-corrosion test. That's why this work focuses on 1:1 solutions, i.e. solutions constituted by only NaCl. In this way, the role of chlorides on the passivity behaviour of 316L can be determined

during a fretting-corrosion test. To compare corrosion and mechanical behaviours in these solutions, this study aims at investigating the influence of ionic strength I:

$$I = \frac{1}{2} \sum_{i} z_i^2 C_i \tag{1}$$

where z_i : ion charge; C_i : ion concentration. Indeed, the ionic strength can interfere in the double layer constitution by the solution content and in the "attraction-repulsion" phenomenon between the contact between two materials [12].

The role of proteins is also investigated. This work focuses on albumin, which is the main protein of blood plasma. Afonso *et al.* [13] have shown that bovine serum albumin, in NaCl 0.15 M at 37 °C, on UNS S31254 stainless steel, stops anodic dissolution of chromium and decreases iron dissolution. Valero Vidal *et al.* [14] worked with 316L and CoCrMo alloy in NaCl 0.14 M at 37 °C, with or without albumin (0.5 g.L⁻¹). They showed that for 316L, albumin does not modify open circuit potential (OCP) but it acts as an anodic inhibitor at applied potential, i.e. it increases cathodic reaction rate. Albumin also tends to decrease the polarization resistance, removing oxides from a protective layer and then exhibiting a material surface from corrosion. For CoCrMo alloy, albumin acts as a cathodic inhibitor and also increases anodic reaction rate, i.e. it accelerates corrosion. During a fretting-corrosion test between a Ti-6Al-4V alloy and alumina in pH buffered saline solutions at 37 °C, Hiromoto *et al.* [15] showed that albumin (1 g.L⁻¹ bovine serum albumin) mostly modifies cathodic behaviour and acts as a cathodic inhibitor, i.e. it reduces cathodic reaction rate. From a mechanical point of view, albumin does not affect friction coefficient µ and wear volume.

During a fretting-corrosion test, material wear is a combination of mechanical wear (fretting) and corrosion. However, the total material loss(W) is not only determined as the sum of two different material losses: (i) material loss due to mechanical wear obtained without the influence of $corrosion(W_m)$ and (ii) the material loss due to corrosion obtained without fretting $action(W_c)$. There is a synergistic effect between corrosion wear and mechanical wear. The synergy term(ΔW) means that corrosion increases wear due to fretting(ΔW_{cm}) and friction increases wear due to corrosion(ΔW_{mc}) [16,17].

$$W = W_m + W_c + \Delta W = W_m + W_c + (\Delta W_{cm} + \Delta W_{mc})$$
(2)

This work aims at understanding the role of mechanical action and corrosion on a synergy term, and also calculating each of these terms for a fretting-corrosion test. In a first place, we focus on the percentage of wear due to corrosion obtained from the wear volume due to corrosion V_{corr} (cm³), and the percentage can be found out with Faraday's law:

% corrosive wear =
$$\frac{V_{corr}}{W} * 100$$
 with $V_{corr} = \frac{1}{F} * \frac{M}{n\rho} * i * t$ (3)

where *F* is Faraday constant ($F = 96500 \text{ C.mol}^{-1}$), *M* is the molecular weight ($M = 56.39 \text{ g.mol}^{-1}$), *n* is the number of electrons involved in the anodic process (n = 2, hypothesis), *t* is the total duration of the fretting-corrosion test (t = 14,400 s), ρ is density ($\rho = 8 \text{ g.cm}^{-3}$) and *i* (A) is the difference between the average current during fretting and the current after fretting.

Three electrochemical conditions were investigated for this study. First, the Open Circuit Potential (OCP) condition allows being as close as possible to the *in vivo* conditions. However, no information on current is available at OCP. Thus, fretting-corrosion tests are also carried out at cathodic applied potentials. -400 mV(SCE) is a threshold potential for anodic and cathodic transition of current. To remove (or to neglect) corrosion and to only obtain mechanical wear (W_m), the cathodic potential of -800 mV(SCE) is chosen. For each electrochemical condition, the role and the influence of albumin are investigated.

2. Materials and methods

2.1 Materials

The 316L femoral stem is modeled by a plane 316L SS sample: 9 mm x 9 mm x 20 mm parallelepiped. One of the flat parts (9 mm x 20 mm) was polished with diamond paste down to 1 μ m. This frictional surface exhibits 3D roughness(S_a) of about 10 ± 2 nm (area of 1 x 2 mm, VeecoTM optic profilometer).

The bone cement, involved in THA, is modeled by a cylindrical PMMA, a polymer which has the same mechanical properties as those of bone cement. A sample maintains the length of 15 mm and the radius of curvature of 10 mm. The cylindrical face was polished with 3 μ m diamond solution, Altuglass Polish 1 and 2[®] and colloidal solution of silica (particles size of 0.06 μ m). The 3D roughness (S_a) is about 35 ± 5 nm.

The geometry of the contact, cylinder/plane, allows a better control of contact pressure and width. Before the fretting-corrosion test, 316L SS and PMMA samples were stored at least 24 hours in a desiccator. The 316L sample is electrically insulated with Zircalloy alloy sheets and some varnish.

The composition of the 316L austenitic stainless steel (X2CrNiMo17-12, improved cleanliness and very low carbon content) is in accordance with ISO standard 5832-1 (Table 1). Mechanical properties of both materials are shown in Table 2.

				•						
Elements	Cr	Ni	Mo	Mn	Si	Co	С	Cu	S	Fe
Composition (% w/w)	17.4	13.0	2.6	1.73	0.66	0.20	0.0165	0.097	< 0.001	Balance

Table 1 Chemical composition of 316 L stainless steel

Table 2 Mechanical properties of tested materials: 316L stainless steel and PMMA, *: obtained by ultrasonic measurements;

Mechanical properties	Young's modulus E (GPa)*	Poisson's ratio v*	Yield stress (MPa)**	Ultimate tensile strength (MPa)**
316L	197	0.30	280	635
PMMA	2.5	0.39	65	75

**: manufacturer data

2.2 Solutions

Two types of solution are used: Ringer solution (composition in Table 3) and NaCl solution, with concentrations of 10^{-3} , 10^{-2} , 10^{-1} and 1 mol.L⁻¹. Room temperature was of 22 ± 1 °C. To understand influence of proteins on fretting behaviour, albumin of 1 g.L⁻¹ was added in each saline solution.

Table 5 Kniger solution composition							
	NaCl	KCl	CaCl ₂	NaHCO ₃			
Concentration (g.L ⁻¹)	8.50	0.25	0.22	0.15			

Table 2 Dinger solution composition

2.3 Test device

To reproduce fretting-corrosion phenomenon, micro-displacements measured *in vivo* (about 50 μ m) and stresses applied *in vivo* (between 7 and 30 MPa) [6] must be replicated. A specific device (Fig. 3), conceived and developed in collaboration between ENSM-SE and Böse, was used for fretting-corrosion tests. An electromagnetic motor, first tried for fretting-corrosion, let us obtain the same results as those delivered by the device used previously [9-11]. The load cell is a piezoelectric transducer and the displacement measurer is a capacitive sensor.

The mechanical parameters of fretting-corrosion tests are realistic:

- the applied normal load (P) was 127.5 N, i.e. a pressure of about 22 MPa, which is spread from 7 to 30 MPa, range of pressure applied *in vivo* [6];
- the amplitude of displacement was $\pm 40 \,\mu$ m, close to the *in vivo* displacement [6]. Moreover, the displacement is sinusoidal and the frequency is 1 Hz, i.e. one cycle equals one second, corresponding to the frequency of the gait cycle.

For these mechanical values, the gross slip condition was checked.

To identify actual displacement δ between both materials, the accommodation of the device must be considered. Based on the hypothesis that there is an elastic behaviour with compliance C_s , the actual displacement δ is: $\delta = \delta_r - \delta_{acc} = \delta_r - C_s * Q$, where δ_r and δ_{acc} are respectively the monitored and accommodated displacements, and Q is the tangential load [18].

For electrochemical measurements (OCP), current density and Electrochemical Impedance Spectroscopy (EIS), a PARSTAT 2263 potentiostat with a three-electrode set up was used:

- the working electrode was the stainless steel sample,
- the counter electrode was a circular wire of Pt having the diameter of about 60 mm,
- the reference electrode was a Saturated Calomel Electrode (SCE, E = +246 mV(SHE) at T = 22 °C).



Fig. 3 Fretting-corrosion device: ENSM-SE and Böse prototype

EIS measurements were investigated from 10^5 Hz to 10^{-1} Hz at a rate of 10 measurement frequencies/decade, with an AC amplitude of 10 mV. The potential was equal to OCP determined before EIS measurements or to applied potential (-400 or -800 mV(SCE) in our case). During friction tests, because of the non-stationarity conditions, EIS measurements should not be performed for actual values, but only the evolution of electrical parameters was considered [19]. One may notice that regular EIS measurements, during the fretting phase, have no influence on OCP or on current values after the EIS measurements.

For each electrochemical condition, OCP or cathodic applied potential (-400 and -800 mV(SCE)), two experiments (steps defined in Table 4) were carried out for all solutions with or without albumin.

Table 4 Electrochemical measurements steps for each electrochemical conditions, OCP or cathodic applied potentials (E = -

400 or	-800	mV(SCI	E))
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Electrochemical conditions	OCP	Potential applied: -400 or -800 mV(SCE)			
Step 1	Cathodic polarization	n: -1 V(SCE), 5 min			
Step 2	without fretting: 1 h + EIS	without fretting: 10 min + EIS			
Step 3	with fretting: 4 h + 10 EIS regular distribution				
Step 4	without fretting: 14 h + 2 EIS after fretting stop and 1 EIS after 14 h	without fretting: 10 min + 2 EIS after fretting stop			
Step 5		without fretting: 14 h OCP measurement and 1 EIS after 14 h			

2.5 Wear profiles

Thanks to VeecoTM WYKO NT9100 optic profilometer, the measurements of the total wear volumes for 316L and PMMA were performed. Three measurements were carried out on 316L and PMMA surfaces. The total wear volume was thus extrapolated for the total wear track length (15 mm). This value was compared with the total wear made on the whole length of the track. The wear volume error is about 10 %. Raw data were filtered by an automatic cylindrical shape.

3. Results

3.1 Open Circuit Potential conditions

3.1.1 OCP evolution

After cathodic polarization of 5 min at -1 V(SCE), the open circuit potential (OCP) is recorded without fretting during 1 hour (steps described in Table 4, OCP evolution on Fig. 4). From the beginning of fretting, OCP is significantly decreased and then slightly stabilized. Instantly after the end of fretting, OCP nearly recovers the value before fretting. At regular intervals (every 20 minutes), the polarization resistance (R_p) is determined with Electrochemical Impedance Spectroscopy (EIS) in order to investigate the behaviour of the metal surface. R_p is extracted from the simple electrical circuit: $R_{sol}(CPE//R_p)$, R_{sol} is the solution resistance, CPE is a Constant Phase Element and R_p is the polarization resistance (resistances of the passive layer and the double layer at high frequencies).



Fig. 4 Detail of experiment: OCP measurement before, during and after fretting and R_p measurements from EIS. ΔR_p : difference between R_p before and R_p after 2400 cycles of fretting; ΔOCP : difference between OCP before fretting (E_{bf}) and OCP at the end of fretting (E_{ff}). Solution is constituted by NaCl 10⁻³ mol.L⁻¹, displacement amplitude is of ± 40 µm, 1 cycle is equal to 1 second

Fig. 4 exhibits OCP and R_p values in a NaCl solution of 10^{-3} mol.L⁻¹. It is worth noting OCP and R_p evolutions are the same as those of other solutions, only Δ OCP and ΔR_p vary in accordance with the different solutions (Fig. 5).

Fig. 5 exhibits the decrease of $\triangle OCP$ and $\triangle R_p$ according to the ionic strength. Both electrical values are consistent and they are related to the evolution of the contact submitted to fretting, i.e. less passive behaviour of the AISI 316L SS.

3.1.2 Mechanical behaviour

From a mechanical point of view, the dissipated energy (E_d) during one cycle, which corresponds to the mechanical work of the tangential force, was monitored. For the total duration of the test, the cumulated dissipated energy (E_{dc}) , i.e. the sum of E_d for each cycle (1 cycle = 1 second), was calculated. The cumulated dissipated energy during fretting increases with ionic strength (Fig. 6). One may notice that experiments were investigated at two compliances. This fact substantiates that, the dissipated energy variation is controlled by the composition of the solution, not the mechanical device.



Fig. 5 ΔR_p : difference between R_p before fretting and 2400 cycles after fretting start; and ΔOCP : difference between OCP before fretting (E_{bf}) and that at the end of fretting test (E_{ff}) versus ionic strength I



Fig. 6 Cumulated dissipated energy E_{dc} during the fretting-corrosion test in a function of ionic strength I. Two compliances were investigated, $\diamondsuit: C_s = 0.0707 \ \mu m.N^{-1}$ and $\diamondsuit: C_s = 0.2563 \ \mu m.N^{-1}$



Fig. 7 Influence of albumin (1g.L⁻¹) at OCP conditions on average OCP during fretting, the total wear volume of 316L SS and polarization resistance for NaCl 1 mol.L⁻¹

3.1.3 Albumin influence

In order to satisfy *in vivo* conditions and to understand the role of proteins, experiments were carried out in the same electrochemical conditions and the same solutions with the presence of a protein: albumin with a concentration of 1 g.L⁻¹. As shown on Fig. 7, albumin has no influence (ANOVA test, $\alpha = 0.05$) on the average value of OCP during fretting and the total material loss for NaCl 1 mol.L⁻¹. E_{dc}, Δ OCP and Δ R_p are not significantly different in cases with and without albumin in NaCl 1 mol.L⁻¹. The same conclusion is highlighted in other solutions. Nevertheless, albumin has influence on wear volume of PMMA: this one is slightly lower with albumin.

3.1.4 SEM images

Worn surfaces of 316L SS were observed with a Scanning Electron Microscope (SEM) 6400 JEOL combined with an Energy Dispersive X-Ray (EDX) probe. Fig. 8 presents surfaces of 316L samples submitted to fretting-corrosion tests. Corrosion crevices (Fig. 8 a) are mostly localized at the border of the wear track (Fig. 8 b). At 1 mol.L⁻¹, corrosion seems more localized than that at 10⁻³ mol.L⁻¹. With albumin, grooves are compacter and corrosion crevices follow friction direction (Fig. 8 c).



Fig. 8 SEM images of 316L SS in NaCl 1 mol.L⁻¹ (a), (b) without albumin (c) with albumin, arrows show the direction of sliding

3.2 Cathodic applied potential conditions (E = -400 mV(SCE))

At OCP conditions, although we are close to the *in vivo* conditions, no information on current is available. That's why the second part of this work focuses on cathodic applied potential, and more specifically E = -400 mV(SCE). This value was chosen because the influence of ionic strength on current during fretting can be highlighted.

3.2.1 Evolution of the current

Significant values, as the average current during the last 3 hours of fretting (I_{mf}), ΔI , the difference between the current at the end of fretting (I_{ff}) and the current after fretting (I_{af}), were investigated in accordance with ionic strength. As shown on Fig. 9, fretting involves a higher impact on current values for NaCl solution of 1 mol.L⁻¹ ($\Delta I \sim 11 \ \mu A$) than that for NaCl solution of $10^{-3} \ mol.L^{-1}$ ($\Delta I \sim 1 \ \mu A$): the applied potential is enough to protect a surface during friction phase for NaCl $10^{-3} \ mol.L^{-1}$. In the same way, I_{mf} is close to zero and is even positive, i.e. anodic for one experiment at NaCl 1 mol.L⁻¹ even though it is cathodic (~ -15 \ \mu A) for NaCl $10^{-3} \ mol.L^{-1}$.



Fig. 9 Average current during fretting-corrosion, I_{mf} , and difference between current at the end of fretting (I_{ff}) and that after fretting (I_{af}), ΔI versus ionic strength I

3.2.2 Wear volume evolution

As described in §2.5, wear volumes were measured on PMMA and 316L samples with the VeecoTM 3D optic profilometer (Fig. 10). Wear volume of PMMA is constant whatever the ionic strength (ANOVA test, $\alpha = 0.05$) at OCP and E = -400 mV(SCE) is, but the wear volume of PMMA is slightly higher at OCP conditions.



As illustrated on Fig. 11, for 316L samples, total material loss increases with ionic strength for both electrochemical conditions. A threshold concentration is then highlighted, about 10^{-1} mol.L⁻¹. For 10^{-3} and 10^{-2} mol.L⁻¹, wear volume measured in OCP conditions is higher than that for E = -400 mV(SCE). Moreover, for concentrations above 10^{-1} mol.L⁻¹, wear volume measured in OCP conditions is lower than that at E = -400 mV(SCE).



Fig. 11 Total wear volume of 316L versus ionic strength for OCP conditions and E = -400 mV(SCE)

3.2.3 Albumin influence

Contrary to OCP conditions, albumin plays a significant role at the cathodic applied potential of E = -400 mV(SCE). Mainly, two values are analyzed: the average current during fretting and the percentage of corrosive wear. In NaCl 1 mol.L⁻¹ (Fig. 12), albumin largely (ANOVA test, $\alpha = 0.05$) reduces the current during fretting (from ~ -0.5 µA without albumin to ~ -10 µA with albumin) and the percentage of corrosive wear (from ~ 80 % without albumin to ~ 50 % with albumin). One may suggest that albumin acts as an anodic inhibitor, as shown by Valero Vidal *et al.* [14].



Fig. 12 Influence of albumin (1g.L⁻¹) at E = -400 mV(SCE) on average current during fretting and the percentage of corrosive wear of 316L SS for NaCl solution of 1 mol.L⁻¹

3.2.4 SEM images

At E = -400 mV(SCE), the main difference between Fig. 13 a and Fig. 13 b, i.e. NaCl 1 mol.L⁻¹ with or without albumin, concerns the grooves inside crevices. Without albumin, grooves indicate that pulled out particles are enough hard to damage surfaces. With albumin, no groove is observed into crevices: particles are probably softer or are ejected more rapidly. At NaCl 10⁻³ mol.L⁻¹ (Fig. 13 c), the central zone of the contact is not damaged and localized corrosion is not observed.



Fig. 13 SEM images of 316L SS in NaCl 1 mol. L^{-1} (a) without albumin (b) with albumin (c) in NaCl 10⁻³ mol. L^{-1} without albumin, arrows show direction of sliding

3.3 Cathodic applied potential conditions (E = -800 mV(SCE))

To neglect wear due to corrosion and to obtain wear due to mechanical wear (W_m), the cathodic potential of -800 mV(SCE) was chosen. To validate this potential value, only extreme ionic strengths (10^{-3} and 1 mol.L⁻¹) were tested.

	$I_{mf}(\mu A)$		Albumin	W (10 ⁶	Albumin	
	without albumin	with albumin	influence?	without albumin	with albumin	influence?
10^{-3} mol.L ⁻¹	-17.60 ± 0.30	-14.93 ± 0.32	YES	0.25 ± 0.07	0.19 ± 0.03	NO
1 mol.L^{-1}	-27.96 ± 9.31	-45.57 ± 3.11	YES	0.38 ± 0.14	0.35 ± 0.08	NO
Ionic strength influence?	NO	YES		NO	NO	

Table 5 Average current during fretting I_{mf} and total wear volume W measured at E = -800 mV(SCE), with or without albumin. Influence of albumin and ionic strength is determined by an one-way ANOVA test ($\alpha = 0.05$)

As shown on Table 5, the average of current during fretting varies in a function of ionic strength and is modified by the presence or not of albumin. However, the total wear volume is same without regard to the condition (ionic strength and albumin). One may assume that the total wear volume equals the mechanical wear volume W_m , i.e. $W_m = 0.3 \pm 0.1 \ 10^6 \ \mu m^3$.

4. Discussion

4.1 Open Circuit Potential conditions

The evolution of OCP before, during and after fretting is shown on Fig. 4. One may notice that E_{bf} logarithmically decreases with the ionic strength in accordance with [20]. Chloride ions, Cl⁻, accelerate anodic dissolution with the formation of an unstable compound MCl_n (or M(H₂O)_yCl_x) [21]. At high concentration of chloride ions, one may suggest that metallic dissolution is higher and leads to OCP decrease.

The drastic drop of OCP occurs as soon as fretting starts (Fig. 4), expressing the surface degradation. The passive layer on 316L surface is partly destroyed and metal dissolution, i.e. anodic reaction, occurs. In addition, breakdown of the passive layer is highlighted by the polarization resistance (R_p) fall. R_p is indeed inversely proportional to corrosion rate (Stern-Geary equation): the R_p fall involves acceleration of corrosion.

One may notice that these falls, $\triangle OCP$ and $\triangle R_p$, decrease with the ionic strength (Fig. 5). According to Macdonald [22], when the concentration of Cl⁻ increases, the passive layer reconstruction becomes more and more difficult. The result highlighted on Fig. 5 translates the difficulty of passive layer reconstruction during fretting when chlorides concentration increases.

The OCP and R_p upturn, directly after cessation of fretting (Fig. 4), confirms corrosion activation during fretting test. However, the OCP value after fretting is lower than E_{bf} . The electrochemical behaviour of 316L surface can indeed be modified by the less protective worn zone and the PMMA and oxides debris deposed on the 316L surface.

From a mechanical point of view, it is worth noting that the cumulated dissipated energy (E_{dc}) during the fretting test increases with the ionic strength (Fig. 6). One may suggest that when the ionic strength increases, attraction between surfaces is reinforced [23].

As shown on Fig. 7, albumin has no influence on the average OCP during fretting and on total wear volume of 316L. Without friction, Valero Vidal *et al.* [14] showed that albumin does not modify OCP values. At this concentration, 1 g.L⁻¹, albumin does not modify electrochemical behaviour: it has no influence on OCP during fretting and R_p values; nor on total wear volume.

4.2 Cathodic applied potential conditions (E = -400 mV(SCE))

The applied potential E = -400 mV(SCE) is a cathodic potential, i.e. the current is negative, thus the surface is protected against corrosion. However, this protection is not effective for all solutions. For example, in NaCl 10⁻³ mol.L⁻¹, this protection is effective: $\Delta I \sim 1 \mu A$ while it is not effective in NaCl 10⁻¹ or NaCl 1 mol.L⁻¹, respectively $\Delta I \sim 7 \mu A$ and $\Delta I \sim 11 \mu A$ (Fig. 9). During the fretting-corrosion test, the current even becomes positive, i.e. anodic, for NaCl 1 mol.L⁻¹. In fact, a threshold concentration and two surface behaviours can be determined (Fig. 11). For concentrations below 10⁻¹ mol.L⁻¹, wear volumes at OCP condition are higher than those at E = -400 mV(SCE): the applied potential protects surface. For concentrations above 10⁻¹ mol.L⁻¹, wear volumes at OCP condition is higher and the passive layer is less protective against fretting.



Fig. 14 The total wear volume of 316L SS, W, as a function of wear due to corrosion, V_{corr} , in the logarithmic scale. The dashed straight line: case where wear is only due to corrosion $\rightarrow W = V_{corr}$. Ionic strength decreases from the right to the left

Fig. 14 allows underlining which of preponderant wear phenomena is highlighted: corrosion or mechanical wear. Experimental data close to the dashed straight line indicate preponderance of corrosive wear. Corrosion is also the preponderant wear phenomenon for high ionic strength: ~ 80 % in NaCl 1 mol.L⁻¹ and 10⁻¹ mol.L⁻¹ without albumin. One may notice that, in presence of albumin, the percentage of wear due to corrosion decreases: ~ 50 % in NaCl solutions of 1 mol.L⁻¹ and 10⁻¹ mol.L⁻¹, instead of 80 % without albumin. Especially, in the presence of albumin and for lower ionic strength, standard deviation values are large and the coefficient of variation is in the range from 10 to 100. The preponderant phenomenon is mechanical wear and in this case, the dispersion of corrosive and total wear values is larger. The main mechanical degradation, under albumin and NaCl solutions, should involve hazardous wrenching that should be related to the high standard deviation.

4.3 Cathodic applied potential conditions (E = -800 mV(SCE))

As explained in §1, the total wear volume of 316L SS is the sum of wear due to corrosion, W_c and mechanical wear, W_m . The total wear volume, W, follows the equation (2). At E = -800 mV(SCE), no anodic current is measured during a fretting-corrosion test. In this electrochemical condition, anodic dissolution, i.e. metallic dissolution, is neglected without regard to ionic strength and the presence of albumin. The mechanical wear W_m is also determined: $W_m = 0.3 \pm 0.1 \ 10^6 \ \mu m^3$. At E = -400 mV(SCE), thanks to Faraday's law, the wear due to corrosion V_{corr} can be calculated (equation (3)). In fact, in comparison with the total wear volume (equation

(2)), $V_{corr} = W_c + \Delta W_{mc}$. By deduction, the term ΔW_{cm} can be determined, as a function of ionic strength at E = -400 mV(SCE). To set apart the synergy term $\Delta W = \Delta W_{mc} + \Delta W_{cm}$, the term W_c must be measured. From polarization resistance values, the dissolution current of the material, *i*, can be calculated: $i = B/R_p$, with B = 24 mV (hypothesis coming from investigations in the COST 533 round robin). In that way, W_c is ~ 100 or 500 μ m³ depending on the ionic strength. W_c represents only 0.01% of V_{corr} and this can be neglected. Iwabuchi *et al.* found the same ratio: W_c/W of ~ 0.01% [25].

Fig. 15 a illustrates the contribution of ΔW_{mc} , i.e. the increase of corrosive wear due to fretting. Levels can be defined from NaCl solution of 10^{-1} mol.L⁻¹ without albumin: above this concentration, the contribution of corrosive wear is about 80%. Albumin at a concentration of 1 g.L⁻¹ shifts the transition and the percentage of corrosive wear: above 10^{-2} mol.L⁻¹, the contribution of corrosive wear is constant and about 50%. Thus, albumin decreases the percentage of corrosive wear. One may notice that material behaviour in Ringer solution is different from that in NaCl solutions. Because of the presence of other ions (Table 3), the contribution of corrosive wear decreases. For example, the role of carbonates could be determined in future work. Fig. 15 b illustrates the contribution of ΔW_{cm} , the increase of mechanical wear due to corrosion. No trend can be defined: mechanical wear is not a function of ionic strength, and this term matches with an arbitrary wrenching in the surface material.



Fig. 15 The contribution of the different synergy terms with albumin (empty symbols) or without albumin (full symbols) : (a) $\Delta W_{mc}/W$ is the increase of corrosive wear due to fretting and (b) $\Delta W_{cm}/W$ is the increase of mechanical wear due to corrosion

Diomidis *et al.* [26] worked with a new expression of synergy term. Synergism is in fact the acceleration of material deterioration by corrosion because of mechanical removal and re-growth of a passive layer. Continuous and intermittent sliding tests are completed to measure the material loss in the sliding track. The intermittent sliding tests allow the removal and re-growth of the passive layer. However, the sliding conditions in a fretting-corrosion test are not continuous and the passive layer destruction and re-growth must be different.

In future work, the influence of albumin concentration on metal behaviour will be studied. To be closer than *in vivo* conditions, experiments will be realized at 10 and 20 g.L⁻¹. For example, the ISO standard indicates a concentration of proteins equal to 17.5 g.L⁻¹ for improving the lifetime of hip implants [27,28].

As seen on Fig. 10 a, the characteristic shape of 316L wear track, 'W' wear shape as mentioned in [9,10], is the consequence of crevice corrosion. The evolution of local pH might lead to this particular shape. The influence of pH solution could give some information on the crevice corrosion effect. The crevice corrosion effect will be investigated in further investigations.

5. Conclusions

Under OCP conditions, the increase of chlorides concentration leads to higher corrosion and a more difficult passive layer reconstruction during the friction phase. From a mechanical point of view, the E_{dc} increases according to the ionic strength; consequently one might suggest attraction between surfaces is reinforced.

At E = -400 mV(SCE), a threshold concentration, around 10^{-1} mol.L⁻¹, highlights that a surface protective phenomenon occurs below this concentration, while more metal dissolution appears above this value.

At E = -800 mV(SCE), only the mechanical wear is determined: $W_m = 0.3 \pm 0.1 \ 10^6 \ \mu m^3$. This value is needed to calculate the term of synergy between corrosive and mechanical wear.

Albumin, at 1 g.L⁻¹, has no influence at OCP conditions. However, at E = -400 mV(SCE), the percentage of corrosive wear is lower in presence of albumin. Albumin seems to play the role of an anodic inhibitor.

When the chlorides concentration, i.e. the ionic strength in this study, increases, the 316L wear increases too. The most interesting result is: albumin promotes the mechanical degradation compared with the corrosive wear, under cathodic polarization. This assertion includes the pure mechanical degradation and the synergism mechanical wear assisted by the corrosion. One might suggest this synergistic term should be the key-point of the 316L wear, under cathodic polarization.

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