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DEVELOPMENT OF A “2-D” TEST SYSTEM FOR VISUALIZING FRETting CORROsITION: A STUDY OF THE FRETting CORROsITION BEHAVIOR OF Co-Cr-Mo ALLOY

DongKai Zhu
Syracuse University

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ABSTRACT

Fretting corrosion, one of the most common forms of mechanically-assisted corrosion (MAC), has become a major concern in orthopedic medical devices. In order to a better understand of the mechanism of fretting corrosion in orthopedic alloys and the ways to prevent implants from corroding, custom test systems need to be developed to simulate fretting corrosion in vitro and to allow visualizing of the process. This study aimed to develop a new “two dimensional” fretting crevice corrosion test system with the capacity to visualize damage progression during systematic controlled fretting corrosion processes. Another goal of the study was to experimentally verify a tribocorrosion heredity integral approach to predict abrasion-current-impedance-voltage relationships by systematic variation of fretting frequency and area. The results of Open Circuit Potential (OCP) tests, fretting current tests and visualization of fretting showed this device could achieve the basic requirement for triboelectrochemical testing and also provided direct evidence of debris generation during fretting corrosion. The fretting region was surrounded by a halo of fretting debris after removal from the solution. One possible explanation for this phenomenon could be the redeposition of fretting product. It was also found that higher fretting frequency and smaller second electrode areas lead to larger and faster voltage drops during abrasion. This is because the smaller area of the second electrode resulted in less surface area for electrons to be reduced. Higher frequency abrasion also resulted in higher film currents and faster electron generation rate which would cause more electrons accumulating at the working electrode.
DEVELOPMENT OF A “2-D” TEST SYSTEM FOR VISUALIZING FRETTING CORROSION: A STUDY OF THE FRETting CORROSION BEHAVIOR OF Co-Cr-Mo ALLOY

By

Dongkai Zhu

B.E. Prosthetic and Orthotic Engineering, Capital Medical University, China

Thesis

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INTRODUCTION

Metallic orthopedic devices, such as joint replacements, spinal screws as well as bone plates, have been widely used over decades. Many biometallic materials like stainless steel (316LSS), titanium-based alloy (Ti6Al4V) and cobalt-chromium alloy (CoCrMo) have been documented as suitable implant materials. The main reason behind these popular materials is that they have excellent mechanical properties and biocompatibility. There are usually five contact combinations for total hip replacements: metal-on-body (MOB), metal-on-ceramic (MOC), ceramic-on-polymer (COP), metal-on-metal (MOM) and metal-on-polymer (MOP). The interface of each combination has its own unique characteristics. There are many designs for different material combinations that depend on the specific situation.

Implant material failures have been reported in vivo and many of them occur prior to the ideal lifetime of implant system [1][2][3][4]. Metal-on-metal (MOM) connections (e.g., modular taper junction) cannot avoid the generation of corrosion and wear debris that may result in biocompatibility concerns [5][6]. Degradation of implants is associated with loosening, local tissue reactions and other issues. Biological reactions, like local inflammatory reactions, stress shielding, pain in the joint and osteolysis, to name a few have been associated with metal degradation processes [7][8][9][10]. Typically, failed implant surgery results in revisions.

Many techniques and designs have been developed to help prevent implant corrosion. Surface coating is one approach to minimize interface corrosion [11][12][13].
For instance, Swaminathan et al., showed that chromium oxide-coated Ti6Al4V and CoCrMo alloy have suitable electrochemical properties as a hard bearing surface [14]. Two retrieval studies also showed that bearing surface materials and head size affect corrosion behavior at modular junctions for total hip replacement while other literature found that large femoral heads might increase the damage at the modular taper junction of MOP implants [15][16][17].

Implant materials modification is also a useful tool to enhance bearing surface corrosion resistance [18]. For instance, Kaushik et al., [19] developed a metallic glass thin film (TiCuNi) with good biocompatibility upon exposure to muscle cells. Low cost, viscoelastic behavior, high electrical conductivity, and the capability to coat various materials make this metallic glass thin film (TiCuNi) as an accretive material for biomedical application. However, failures still happens.

Many retrieved implants show evidence of corrosion and wear phenomena [20]. Mechanically assisted corrosion (MAC) is due to an interaction between mechanical and electrochemical factors leading to accelerated damage. MAC happens at the interface of two phases and may damage both surfaces. MAC consists of micro-motion, which is less than 100 µm, between two surfaces that disrupt the passive oxide film and accelerates corrosion. Asperities on both surfaces (if a metal-metal junction) will abrade the surface oxide film, when fretting motion in initiated, it will give rise to a series of redox reactions due to abrasion of the two surfaces. The redox reactions will shift the potential of the implant and generate currents called fretting corrosion current.

Oxide films are the most significant feature of the surface of the implant metal to limit corrosion (see Fig. 1). The potential of oxide-covered surfaces is relatively stable if
there is no fretting motion causing oxide disruption. The potential when no oxide disruption occurs is called the resting Open Circuit Potential (OCP) and it depends on the characteristics of the metal itself and the reduction reactions present. When the oxide film is disrupted, oxide repassivation and metal ion release into the solution will generate significant drops in electrode potential and will affect the local environment. For Ti6Al4V alloy, the oxide film formed is usually TiO₂. The main composition of oxide film for CoCrMo alloy and 316LSS is Cr₂O₃. The valance of Cr ion depends on the potential and may become Cr⁶⁺ at high potentials. These toxic species may have a negative effect on the human body [21][22]. Fretting motion can cause not only oxide film fracture but also debris generation. Debris may accumulate at the junction or be released into the periprosthetic tissue.

Figure 1. Schematic of the consequence of fretting corrosion. Note that green arrow stands for the direction of fretting motion.
The disruption and reformation of oxide films usually occurs in conjunction with ion release. Redox reactions, which happen at interface, will generate different ions, including metal ions (e.g. Ti$^{2+}$, Cr$^{3+}$) and other ions with positive or negative charge (e.g. H$^+$, OH$^-$). The ions will be released into the solution once the oxide film is disrupted and reformed. For phosphate buffered saline (PBS) solution, H$^+$ will attract Cl$^-$ due to the charge attraction force and form HCl. This environment can lead to a pH drop, thus creating a highly localized aggressive condition and will give rise to further corrosion known as crevice corrosion. Fretting corrosion is usually accompanied by crevice corrosion. Pitting corrosion is another reason for device failure and has been observed and documented in many studies [23][24][25][26]. Pitting corrosion results, in part, from depassivation of a very small region of the surface. Corrosion attack penetrates the oxide film and leaves a tiny hole in the surface. Many reasons may cause pitting corrosion, such as fretting, metal combination and surface geometry, etc. Recently, Gilbert et al, proposed that inflammatory cells may release chemical species that can cause corrosion on CoCrMo alloy based on retrieval analysis [27].

Although implant materials, including metals, have been studied for decades, there is more to be understood about fretting corrosion. There are several reasons for studying the fretting corrosion of CoCrMo alloys. Pearson et al., concluded that individual blood shows different responses to CoCrMo alloy debris and the responses can be an indicator of suitability for patient to receive MOM arthroplasty [28]. Liao et al., revealed that many studies point out that Cr ions show up in patient serum and urine and the concentration increases with time [29].
Many theories have been developed to better understand device failure and the study methods can be divide into several aspects: retrieved study, cell culture study, in vitro study and in vivo study. For example, Swaminathan and Gilbert developed a tribocorrosion equation to describe the interaction of mechanical and electrochemical factors: [30]

$$I_{\text{film}} = 2 \frac{\rho n F V_{\text{nom}} d\delta}{M_w \Delta} = 4 \frac{\rho n F V_{\text{nom}} \delta v}{M_w \Delta}$$

Eq. 1

The calculation of fretting current, denoted as $I_{\text{film}}$, is based on many parameters such as oxide density ($\rho$), effective valance of oxide ($n$), Faraday’s constant ($F$) and $M_w$ is the molecular weight of oxide. $V_{\text{nom}}$ is the nominal abrasion volume of oxide film and $\Delta$ is the average interasperity distance on the contact area. The time derivative of amplitude ($\delta$) can be defined as sliding speed and it equals the frequency ($v$) times two times ($\delta v$).

This equation shows that fretting currents are determined by a combination of mechanical factors and material properties. In addition, there are many other factors that affect the fretting currents such as the potential across the interface, the solution chemistry, the impedance of the whole system and the surface geometry, etc.

The potential drop across the electrode interface and the current. Though it can be measured using a three-electrode electrochemical cell. This system contains a working electrode (WE), which usually is the sample itself, a counter electrode (CE) and a reference electrode (RE). Carbon rod or platinum electrode are the most common counter electrodes and Ag/AgCl or saturated calomel electrode (SCE) is usually used as the reference electrode. As for the mechanical data, different sensors, such as linear variable
differential transformers (LVDTs) and differential variable reluctance transducer (DVRTs), are used depending on the situation. Electrochemical impedance spectroscopy (EIS) is another useful technique to study the electrochemical phenomenon of different materials and has been used to study corrosion behavior [31][32][33]. Haeri et al, developed a time-based equation to better describe the metallic biomaterials non-ideal impedance behavior of CoCrMo surface [34]. The basic principle for EIS is to apply an oscillating voltage, usually a sinusoidal wave, as input and to analyze the corresponding output of current. The results are then fit to one of several classic electrical analog models, like the Randles circuit, to obtain the resistance and capacitance of the test system.

Many factors affect EIS results based on published papers. Gilbert et al, showed that small increases in H₂O₂ concentration in phosphate buffered saline (PBS) solution result in large decreases in the low frequency (i.e., oxide) impedance of CoCrMo surface [27]. Ribeiro et al., revealed that 10% hydroxyapatite coating on CoCrMo (CoCrMo-10HAP) can decrease the low frequency impedance as well [35]. Time was another underlying factor that would lead to the change of impedance. Also, exposed electrode area is another factor known to affect impedance.

Electrode area affects current and potential behavior during fretting corrosion. Large electrode areas will cause lower voltage drops but higher fretting currents, and vice versa. The key factor of this phenomena is the electron reduction reaction rate. Electrons are generated from oxidation reactions and are accumulated at the metal surface and are removed by reduction reactions, usually at the counter electrode (CE). Greater electron generation and accumulation cause a larger potential drop due to the increased
accumulation of negative charges. Large areas provide more space for reduction reactions, which means more electrons from the fretting surface can be distributed on the surface to be reduced. This behavior will lead to less electron accumulation beneath fretting surface, which will thereby lead to lower voltage drops. It is important to study the relationship between electrode area and electrochemical parameters (potential and current) because both voltage and current affect fretting corrosion and electrode area is one of several parameters that can control their behavior.

For in-vitro testing, fretting currents are affected by normal load, fretting amplitude, surface roughness, exposure area, fretting frequency and solution chemistry. In order to control and capture these parameters simultaneously, fretting test systems have been established to help scientists understand fretting corrosion and wear behavior of different materials [36] [37][38][39][40][41]. Swaminathan and Gilbert, developed a pin-on-disk fretting testing system that can capture electrochemical data, like fretting current, and mechanical data, frictional force and relative displacement simultaneously. This device can also test different material combinations instead of only using fixed sample couples [30]. Billi et al., also presented a system to characterize modified Ti6Al4V alloy surfaces after exposure to fretting corrosion. The most attractive feature of this test system is that it can test three different sample combinations at same time [42]. Another set up from Royhman et al., can study the fretting corrosion of a single sample rod while it is in contact with two different materials. Note that all three alloys materials are in the same solution in same set up [43].

Even though the goal for each test is different, there are still many similarities for different fretting corrosion testing devices. They all need three elements, which include a
way to set enough load to make sample contact, a motor that can generate micro motion with appropriate amplitude and an electrochemical test device to adjust or record the corrosion behavior of whole system, to perform fretting corrosion tests. For each system, the ways to collect data depend on the goals that this system wants to achieve. But they all have something in common. For instance, a three electrode cell is widely used for most fretting corrosion test system. However, other corrosion test set ups may be designed to capture specific electrochemical data [44][45]. Different load and displacement sensors are used in fretting test systems to collect mechanical and kinematical data as well. It is essential to list all requirements before designing and building test system.

Even though numerous designs and theories help scientists have a better understanding of fretting corrosion, the process of how implants fail is still not clear, in part, due to a lack of ability to visualize the fretting corrosion process, especially dynamic video data. There are few case reports or papers involved in this aspect of the process. Geringer et al, designed a custom fretting corrosion test system to investigate the degradation between 316L steel and poly(methyl methacrylate), known as PMMA. In this study, a CCD camera was used to allow the continuous recording of contact conditions during fretting through transparent PMMA. Many pictures were acquired from that camera and showed a clear trend of increasing wear degradation with increasing load [46]. Other published literature from Kim et al, introduced a fretting test system which has similar features as the previous one, to study the coefficient of friction and fretting wear rate between 316L stainless steel and PMMA. This machine allows the operator to observe the fretting conditions or let the camera record the whole process [47]. However,
both systems focus on the mechanical aspects of fretting corrosion, such as friction force and energy loss, and pay little attention to the details of fretting corrosion, like debris generation, and changing of electrochemical parameters, (e.g. voltage and fretting current). It is useful to combine electrochemical and mechanical testing with visualization to better understand the fretting corrosion process.

Figure 2. Two fretting test systems with visualization capabilities build by Geringer et al, (up) and Kim et al, (down).
In this study, a new fretting corrosion testing system is developed to perform visualized testing and to collect electrochemical and mechanical data concurrently. Some of the advantages of this system are that it has less solution volume between two glass slides to capture debris, it interrogates small contact areas, and the transparent glass allows the operator to use a microscope to observe and record the details of fretting corrosion during the whole process. Another novel feature of this system relates to sample preparation. As a modified pin-on-disk fretting test system, the contact area is small and leads to relatively large contact stresses even with small normal forces applied to the surface. This combined set of capabilities should provide an ability to explore a range of research questions associated with the process of fretting corrosion.
GOALS AND HYPOTHESIS

Long-term goal

The long-term goal of this thesis is to develop an in vitro visualized fretting corrosion testing system to explore and understand the corrosion process of metallic biomaterials and to use this system to answer fundamental questions related to the mechanism of fretting corrosion. As the first step, this thesis has three specific aims and underlying hypotheses as follows:

Specific Aim 1

Design and develop a “two dimensional” visualizing fretting test system to study the interaction between two CoCrMo alloys engaged in fretting corrosion. The device should be able to control and monitor mechanical and electrochemical parameters during the fretting corrosion test and record dynamic data, like video, at the contact area, where the fretting corrosion is taking place during the entire experiment.

Specific Aim 2

To observe and record the phenomenon of debris generation that results from fretting corrosion interaction between two CoCrMo alloys.

Hypothesis
Metal and metal oxide debris is one of the main products resulting from fretting corrosion that may cause some local tissue process or even potentially affect device failure. Debris generation will occur during fretting and the quantities of debris will be affected by fretting duration. Longer fretting duration can generate more debris. Fretting corrosion will also leave obvious corrosion damage areas after fretting and this should be observed on the fretting surface. Fretting area may be surrounded by debris deposition due to the disruption of surface oxide film.

**Specific Aim 3**

To study the effect of fretting frequency and second electrode area on the current and open circuit potential drop during fretting corrosion of CoCrMo alloys.

**Hypothesis**

The voltage will become more negative during fretting due to the fracture and electrochemical repassivation of the oxide film. In addition, the fretting current will also be affected by the changes in OCP during fretting. These effects are linked together by the impedance character of the electrode surface which is, itself, affected by the area of the electrode. As more area of second electrode is exposed in the solution, there will be more surface for reduction reactions to occur, thus leading to less voltage drop. For the same area, higher frequency will give rise to higher fretting currents and a larger and more rapid voltage drop. These effects will be modeled based on the theory put forward by Gilbert and Swaminathan and Gilbert, and the tests of frequency and area will be assessed in this thesis.
INSTRUMENT DEVELOPMENT

Introduction

In order to have a better understanding of fretting corrosion, a new fretting test instrument was developed to visualize and record the dynamic process of fretting corrosion in vitro. From the visualization perspective, this system should contain transparent components to allow the operator to watch and record the whole process of fretting using a microscope or other observation devices. As for the mechanical aspect, a thin-section pin-on-disk model is designed to achieve this purpose. The pin-on-disk system has both mechanical and electrochemical aspects of testing. For corrosion, a three electrode method was used to control and monitor voltage and current. Like many traditional fretting test systems, this instrument can control the normal force, frequency, contact area, displacement, voltage and current by adjusting different components in the system. The operator can easily obtain many results, like coefficient of friction and the work per fretting cycle, based on the measurement outcomes. However, beyond that, this system can also allow the operator to observe the whole process of fretting corrosion, such as debris generation. This novel feature can lead to a better understanding of the in vitro fretting test and provide a visualized integration of the materials, mechanical and electrochemical field. The system was designed using SolidWorks2013® software and fabricated with the help of the department machine shop. It can be divided into three parts: mechanical part, electrochemical part and observational part. Each of them work independently. There are two systems for data collection, one uses a potentiostat to
measure and record impedance results and another one was designed with LabVIEW2013® software for mechanical control and measurement.

**Mechanical Elements**

The mechanical part includes three sub-components: load cell, movement system, and mechanical measurement system and each component has its own characteristics. The load cell was a double-cantilever design with an aluminum mount, two brass cantilevers, a displacement sensor (LVDT) with a core inside and a brass rod support. This custom load cell generated a spring-like system with lateral rigidity to achieve its function. There was a relative displacement between the stage mount and the sensor itself when two samples contacted with each other. The two cantilevers begin to bend and continuously deflect with increasing relative displacement. This process generated force that could be measured by the sensor. The more deflection of the cantilevers, the larger force that the load cell applied. The load was perpendicular to a stage that performed fretting motion.

The movement system (see Fig 3.) contained two parts, one was the fretting part and another was the vertical position adjustment. For the fretting part, fretting motion was performed by a piezoelectric actuator (burleigh®), which had a piezoelectric crystal inside, an amplifier (PZ-300M, buleigh®), and a function generator (FG-8002, EZ Digital®). The crystal could adjust its own length by receiving different magnitude of voltage. In order to have fretting motion, a triangle wave signal was generated by the function generator into the amplifier to change voltage magnitude from 0 V to 100 V.
The voltage allowed the piezoelectric actuator to provide up to 80 µm of micromotion. A stage was mounted to the actuator and moved along with it to perform fretting motion. Vertical position adjustment was accomplished by a linear stage (460A-X, Newport®). This stage could adjust the height of the disk stage relative to the pin for proper alignment.

In order to have complete measurement for the whole mechanical aspect of the system, three sensors were used to measure the normal load, friction load and the lateral displacement. Normal load could be measured through the relative displacement between the load cell stage and the cantilever sensor, which uses a linear variable differential transformers (LVDT) (DC 750-50, Macro Sensors®). The LVDT could measure the relative displacement between the coil and the core, which is connected with the rod in the load cell. Frictional force was obtained through a non-contact differential variable reluctance transducer (DVRT) (SN3327, micro-epsilon®). This sensor could measure the distance, which changed with frictional force, between the target and the sensors. The last parameter was relative displacement between the pin and disk samples and it was the difference between the stage movement, which could be measured by another LVDT connected with sample stage, and the deflection of the pin. The raw data captured by computer was in voltage and the data need to be converted to force based on the calibration of each sensor.
Figure 3. This is a schematic of the fretting corrosion test system from the SolidWorks program. Note that pin and disk are recognized by smaller rectangle and will be detailed in the electrochemical section below.

**Electrochemical Elements**

The detailed relationship between working, reference and counter electrodes are detailed in Fig. 14. The impedance of the electrode system was measured before and after a fretting test. A three electrode configuration was used with a 10 mV sinusoidal voltage.
applied to the working electrode (WE) over a range of frequencies from 20K Hz to 0.01 Hz. Solartron 1280C Potentiostat/Frequency Response Analyzer (Solartron Analytical Inc) and a software called ZPlot™).

As for studying the relationship between voltage drop and second electrode area, a revised three electrode method was established to meet the purpose. A custom built bipotentiostat was used to measure the OCP between working electrode (pin and disk) and reference electrode (Ag/AgCl) and a zero resistance ammeter (485, KEITHLEY®) was used to measure the current from the working electrode to a second electrode of known area (i.e. CoCrMo alloy slice). The surface of the pin, other than the contact surface, was covered with electrically isolating material (nail polish, Ciate). The disk sample was attached to a thin slice of PLA using super glue (Gorilla® super glue). The slice, made from polylactic acid (PLA), was printed from a 3-D printer into a rectangular shape (Replicator™2, Makerbot®). The combined sample was stuck to a glass slide (Fisherbrand®) using super glue. In order to ensure the alignment between the pin and disk, a holder, which mounted onto the stage was made for this purpose. A metal screw penetrated through the top of the holder and contacted with the disk sample, thus leading to an electrical connection between the disk sample and the outside circuit. Another glass slide was mounted to the top using double face tape (Scotch®) which left a small gap between the two glass slides to allow for free sliding motion. Considering the thickness of the tape, about 0.2 mm, there were two layers in the front area of the glass slide and three layers in the rear to keep the two glasses slides separated from each other. The reason for doing this was to keep the solution in the fretting region and prevent it leaking (see Fig. 4).
Figure 4. Top view of schematic of electrochemical part. Note that WE is the working electrode, RE stands for reference, CE stands for counter electrode and SE stands for second electrode.

Data Acquisition and Analysis Elements

All mechanical and electrochemical data were acquired using two National Instrument (NI) data acquisition boards (DAQs). A custom LabVIEW program (see Fig 5.) was developed to graph and save data from DAQ simultaneously. The program
include some parts that are specific to acquire data from two LVDTs and the DVRT. Electrochemical data, voltage and fretting current, was acquired from the biopotentiostat and ammeter and was transferred into a digital signal in order to be read by the DAQ. All data was saved as an Excel® file and was analyzed in Excel®.

Figure 5. Screenshot of LabVIEW program front panel that could graph and collect electrochemical and mechanical data simultaneously. Note that number 1 and number 2 waveform chart graphed voltage and current data while number 3 and number 4 displayed the frictional force and relative displacement at same time. Number 5 was a support waveform chart for the output voltage of DVRT.
EXPERIMENTAL METHODS & MATERIAL

The samples tested were CoCrMo alloy (ASTM 1537) slices and cone-shaped flat bottomed pins (interfacing surface area nominally: 0.002 cm² to 0.009 cm²). The disc slice was cut using a low speed diamond saw in rectangle shape (thickness nominally: 0.34 mm to 0.89 mm). Each disc test surface was polished by hand to 600 grit using emory paper until the scratches were aligned in a single direction and the whole process of polishing was done after the disk slice was attached to the glass slide. After polishing, the samples were cleaned in ethanol and DI water to remove any possible remaining particles from the sand paper. In order to have a better electrochemical environment, a relatively constant area, approximately 0.0045 cm² (nominally: 0.003 to 0.006 cm²), was exposed while the rest of the surface was covered by an electrically insulating material. Only one trial was used for initial simple demonstration but three trials were performed for each particular test where statistical analysis was performed. For basic operation experiments, like voltage vs time, current vs time and frictional force vs time etc., the results were to demonstrate that this system had the capability to meet the basic requirements for a fretting corrosion test system. One trial was sufficient to demonstrate system capabilities. As for fretting visualization experiments, three trials were needed to prove that debris showed up at each trials. In addition, three trials were also needed to explore the relationship between the potential drops and the other two factors: fretting frequency and electrode area.
In order to show that this fretting corrosion test system can achieve the expected goals, a series of tests were performed to demonstrate different functions of this system.

1. Basic Operation Experiment

The instrument’s basic operations were assessed in a series of preliminary experiments at room temperature (RT). After calibration of all motion and load sensors, the experimental set up, with phosphate buffered saline (PBS) solution, was assessed in its OCP vs time behavior with both pin and disk exposed to the solution. The three electrode electrochemical cell (WE, RE, CE) was used in this experiment. Test samples were the working electrode (WE), a custom-made Ag/AgCl electrode was the reference electrode (RE) and the gold foil was the counter electrode (CE). Then, impedance tests were performed until the OCP stabilizing (about 2200 s). Solartron 1280C and ZPlot® was used to record impedance data with the scanning frequency from 20K Hz to 0.01 Hz under 10 mV scanning potential. Fretting corrosion tests were performed at the following conditions: OCP condition, 0.5 N for normal load, 65 µm for sliding amplitude and 1.0 Hz for fretting frequency, to generate fretting corrosion current. The potential and fretting currents were measured by the custom biopotentiostat. The results were collected by DAQ and graphed by the custom LabVIEW program. Pin and disk were separated but still exposed in the solution to measure the impedance after fretting using the same software.

The next experiment was to assess the capability of load adjustment. Test samples were exposed in the PBS solution for the same time as the first experiment. The fretting amplitude was 65 µm and fretting frequency was 1.0 Hz. Eight different loads were
applied to the system from 0.08 N to 1.16 N at a fixed potential controlled by another 
potentiostat (Model 263, EG&G®), -100 mV vs Ag/AgCl, to generate fretting corrosion 
currents. Each test lasted for 5 second and under the same condition as the first 
experiment except for the load. There was a 30 s gap between each test. All parameters, 
including fretting current, normal load, frictional force and relative displacement, were 
still collected by the DAQ system and graphed by the custom LabVIEW program. For 
two experiments, one group of samples, including a single pin and a single disk, were 
used for each experiment.

2. Observation of Debris Generation

To show that metal debris was generated during fretting corrosion, the whole 
system was combined with a digital microscope (KH-870, Hirox®). The electrochemical 
cell used in this experiment was the same as the first. Both pin and disk was exposed in 
the PBS solution for 2200 s at room temperature (RT) before each test. Three trials were 
performed for this experiment. The potential, -100 mV, was controlled by a potentiostat 
(Model 263, EG&G®) during fretting. In order to have a better debris generation rate, 5 
Hz and 0.7 N was used with 65 µm sliding amplitude. 350x magnification was used to 
provide high resolution result during fretting. Video, which lasted for 5 seconds, was first 
recorded at 12 min after the test started and continually recorded after 20min, 30 min, 60 
min, and 120 min. The solution was removed after the test finished and images were 
captured for both fretting interfaces and observation surface (on the side).
3. Frequency and Electrode Area Affects on Voltage Drop During Fretting

Four pieces of CoCrMo alloy covered by electrical insulation material were exposed to solution with different areas: 0.018 cm$^2$, 0.213 cm$^2$, 0.352 cm$^2$, 0.927 cm$^2$. Both pin and disk was exposed in phosphate-buffered solution (PBS) for 2200 s at room temperature (RT) before test initiation and the impedance was measured using the basic three electrodes cell (WE, CE, RE). Then a modified three electrode electrochemical cell, where the counter electrode (CE) was replaced by the second electrode (SE), while the other electrodes stayed the same. The normal load for the whole experiments was 0.5 N and sliding amplitude was controlled at 65 µm. Three trials were needed for this experiment. There were five different frequency (0.2 Hz, 0.5 Hz, 1.0 Hz, 5.0 Hz, 10.0Hz) performed for each second electrode. For each test, the fretting duration lasted for 300 s and the next test frequency started until the potential recovered to the beginning because the potential would drop during fretting corrosion. Both mechanical and electrochemical data were collected by the DAQ system and graphed by the custom LabVIEW program. The impedance was also measured for each electrode after testing.
EXPERIMENTS RESULTS

1. Basic Operation Experiments

The OCP of both CoCrMo pins and disks stayed in a range from -400 mV to -150 mV (vs Ag/AgCl) in this experiment (see Fig 6. as an example). The OCP started around -12 mV and continued to fall exponentially with time, about -170 mV at 2000 s.

Figure 6. A representative plot of the OCP vs. time of both pin (CoCrMo) and disk (CoCrMo).

The impedance values for the different working electrode area varied with area (see Table 1.) $R_s$ stands for the solution resistance, $R_p$ is the oxide film resistance and CPE-T is the oxide film capacitance. There are several observations to discuss. First the
relationship between total area and $R_p$ follows the trend that is larger area relates to smaller $R_p$. The second concern is about the oxide film capacitance is increasing as the total exposed area increases. These trends with area are also present after fretting. The last one is about $R_s$, which is far smaller than $R_p$. It is clear that all $R_s$ values before fretting, in the table, is larger than the value after fretting.

<table>
<thead>
<tr>
<th>Second Electrode Area (cm²)</th>
<th>Before Fretting</th>
<th>After Fretting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit</td>
<td>0.02433</td>
<td>0.21883</td>
</tr>
<tr>
<td>$R_s$</td>
<td>Ω</td>
<td>311.1</td>
</tr>
<tr>
<td>CPE-T</td>
<td>F</td>
<td>1.25E-05</td>
</tr>
<tr>
<td>$R_p$</td>
<td>Ω</td>
<td>2.47E+06</td>
</tr>
</tbody>
</table>

Table 1. A representative impedance data before and after fretting. Note that the area in this chart is total area which includes both the surface area of the pin and disk and the exposed area of second electrode. The $R_p$ of 0.93283 cm² after fretting has extremely high error percentage, which is not reliable.

These is a specific and direct link between the mechanics of fretting motion and the electrochemical response (current and voltage). Potential keeps dropping during fretting corrosion while the corresponding fretting current rises at each time (Fig. 7). When the stage is moving, the pin first begins to bend which leads to the friction force changing and then sliding through the fretting surface, which makes the friction force stay relatively stable (Fig 8.). The coefficient of friction (COF) follows the same trend as friction force. The energy loop shows the relationship between relative displacement,
which is the difference between stage movement and lateral deflection of the pin, and the friction force (Fig 9.).

Figure 7. Potential vs time and fretting current vs time during fretting corrosion (5-7 s).

Note that potential drop when fretting current rise. The phenomenon indicate that the oxide film is disrupted and corrosion initiation.
Figure 8. Stage movement vs time, lateral deflection vs time and friction force vs time during fretting corrosion (5-7 s). Note that the pin first deflect (sticking) and then slide (sliding) through the fretting surface, thus causing fretting current. The friction force vs time plot follow the same trend.
Figure 9. COF vs time and energy loop during fretting corrosion (5-7 s). Note that coefficient of friction (COF) change a lot at sticking condition and stay relative stable at sliding condition. Energy loop shows the relationship between friction force and relative displacement.

In figure 10, the average fretting current vs. normal load is shown. It is clear that the average fretting current increases as normal load increases and reaches to a peak, which of almost 0.082 µA. After passing this point, the average fretting current begins to decrease as the load increases. The error bars in this plot is the RMS about the mean of
the fretting corrosion current. As the load increases, there will be more contact area for the interface causing more asperities to have a chance to abrade the oxide film. However, the pin and disk will stick together as the load increases and this will lead to less relative motion, which will decrease the average fretting current.

Figure 10. Average fretting current vs. normal load for CoCrMo pin and CoCrMo disk at -100 mV vs. Ag/AgCl (n=1). Note that the error bar stands for the root mean square (RMS) about the mean.

2. Observation of Fretting Debris Generation

Based on many retrieval studies and observational results, metal debris and/or oxide debris is generated through fretting corrosion which is thought to be associated with the optical image (see Fig 11) of the debris generation process. The interface-generated debris stated as early as 12 min and continued over 60 min. Debris first showed up at the fretting interface and grew with time. Newer debris generated during fretting
pushed the older debris to the peripheral area. This lead to a phenomena where the outer layer of debris was pushed faster away from the source.

Figure 11. Image (e) and the caption is on next page.
(e) 120 min

Figure 11. Digital microscope image of debris generation from CoCrMo pin and disk at different times. Note that the black part (left) is the pin covered by electrical insulation material and the light gray material was the mirror-polish surface of the CoCrMo disk. (a) 12 min after fretting (b) 20 min after fretting (c) 30 min after fretting (d) 60 min after fretting (e) 120 min after fretting

Two images (Fig. 12a and 12b) are captured after fretting by simply removing the solution between two glass slides and the disk sample was placed under the microscope until the observation surface dried. The left image (a) shows the region of the observation surface and the surrounding corrosion debris halo after fretting under 350x while the right one (b) is a high (700x) magnification of the same region. It is obvious that the contact point is corroded and a halo area is formed at the outer region of the contact area. Figure 12(c) is a side-view schematic comparing the ideal contact situation and real contact condition. Note the red circle stands for the different situation about the contact interface. The ideal condition is that pin and disk have full contact with each other, like the upper
condition in figure. 12(c), and the observation surface can reflect what happens at the interface. However, the fretting surface of the disc sample is too small to polish to a mirror-like finish and the observation surface needed to be mirror-like instead of 600 grit. The roughness of the fretting surface is 600 grit whereas the observation surface was polished to a mirror-like condition. This caused the edge of disk sample to become curved, which is demonstrated as the lower condition of figure. 12 (c).

The edge condition described in last paragraph (shown at Fig. 12) shows significant debris generated. In figure 13, image (a) and (b) are the fretting contact surface with different regions in focus. Red circle indicates the abrasion region, which can also be observed on the pin as well in figure 13(c), where the fretting direction is perpendicular to the original polished scratch direction. It is worth noting that the contact region is surrounded by a corrosion halo area and some debris, which is the dark parts at the interface.

Figure 12. Picture (c) and the caption is on next page.
(c) Comparison between real and ideal situation

Figure 12. Digital microscope image for fretting trace on the observation surface at different magnifications: (a) 350x and (b) 700x. Picture (c) is the schematic (side-view) for the comparison for ideal and real contact circumstance.

Figure 13. Image (c) and the caption is on next page.
Figure 13. Digital microscope images of both pin and disk. Image (a) and (b) are the fretting surface of the disk sample with different focus points due to the surface geometry. Image (c) is the fretting surface of the corresponding pin sample.

3. Frequency and Electrode Area Affection of Voltage Drop during Fretting

The voltage drop associated with the four different second electrode areas (Fig. 14) at five different frequencies show that higher frequencies cause faster voltage drops, which can be measured through the initial slope of each curve, and result in larger drops. For larger second electrode areas, smaller voltage drops (at a specific frequency) is also shown. A summary of area and frequency effects (Fig. 15) show that the voltage drop for 10.0 Hz and of 0.018 cm$^2$ second electrode area is -0.2 V and for 0.213 cm$^2$ is -0.17 V, for 0.352 cm$^2$ is -0.14 V and for 0.927 cm$^2$ is -0.11 V. Figure 15 shows that larger second electrode area results in smaller voltage drop.
Figure 14. Plot (c), (d) and the caption are on next page.
Figure 14. The voltage versus time for different second electrode area (0.018 cm$^2$, 0.213 cm$^2$, 0.352 cm$^2$, 0.927 cm$^2$) and frequencies (0.2 Hz to 10.0 Hz).
Figure 15. Voltage drop vs. frequency for different second electrode area. Note that the value of voltage drop equal to average value of the last second minus the average value of the first second. Mean and SD n=3.

Data are replotted (see Fig. 16) voltage drop versus area for different frequencies to emphasis the effect of area on voltage change. The plot shows that the voltage drop for 0.018 cm$^2$ and 0.2 Hz is -0.03 V, for 0.5 Hz is -0.06 V, for 1.0 Hz is -0.09 V, for 5.0 Hz is -0.15 V and for 10.0 Hz is -0.185 V. The plot also shows that higher fretting frequency results in larger voltage drop.
Figure 16. Voltage drop vs. second electrode area for different frequencies. Note the voltage drop in average potential just prior to fretting minus the average potential at the last second of fretting. Mean and SD n=3.

These data show that the voltage dropped exponentially with time and reached a plateau. However, for lower frequency tests (see Fig. 17), the voltage drops in an oscillating form and this behavior can be affected by different frequencies. The voltage drop for a single experimental sample during fretting at different frequencies shows this oscillation and how it is directly related to fretting frequency. Although frequencies (i.e. 10 Hz), the oscillation is eliminated whereas its process increase with lower frequencies.
Figure 17. Detail of voltage drop vs. time during fretting for a single sample at different frequencies. Note that the red circle and arrow (in the left plot) is enlarged and displayed at the right plot to demonstrate the effect.
DISCUSSION

Fretting corrosion and its mechanism have been studied for many years from different perspectives [48][49][50][51]. However, the details of the fretting corrosion processes are still not well understood. Many custom devices have been established to either focus on measuring the electrochemical parameters during fretting or to study the mechanical influence, like weight loss, of the fretting process. Few of the devices, mentioned in the introduction, have visualization capabilities but they still lack the capability to have high-resolution observations of the processes of fretting corrosion. In addition, it is hard for these devices to collect and study the fretting products due to the typically large solution volumes involved. Another objection for many fretting corrosion test systems is that they usually have a relative large contact areas compare with the sample contact condition in this study. This study was undertaken to develop a custom fretting crevice corrosion test system that combined electrochemical and mechanical features together and measured all parameters simultaneously. The whole device can be placed under a digital microscope to visualize the process of fretting corrosion directly. The sample chamber is made of two glass slides and regimes much less solution volume. This feature can be used to study the products of fretting corrosion. In addition, the small thickness of disk sample provides small contact areas and which is useful to study the fundamental theory of fretting corrosion.

The results show that debris, which is one of the products of fretting corrosion, is generated during fretting corrosion. The debris accumulated at the fretting region as time
increased. There is a halo that appeared on both fretting interfaces and the slide observation surface after the solution was removed. The fretting region was surrounded by the halo. The relationship between voltage drop and fretting frequency and second electrode area through the impedance of the surface, shows that higher frequency resulted in larger voltage drops at similar second electrode area. A similar phenomenon happens for smaller second electrode areas under different fretting frequencies. In addition, an oscillation phenomenon during the voltage drop at different frequencies was observed when enlarged plots of voltage versus time are generated. The oscillation was not obvious at lower frequency (i.e. 0.2 Hz) but became clear as the fretting frequency increased. This phenomenon was eliminated at higher fretting frequency (i.e. 10.0 Hz). These basic experiments show that the change of mechanical parameters (i.e. frictional force and relative displacement) have a direct connection with the electrochemical parameters (i.e. voltage and fretting current).

This study provides a new direction for the study of fretting corrosion. However, the test system still has some weakness. The first one is about the sample preparation. The sample edge was curved during polishing due to lack of a suitable holder. This circumstance influences the ability to visualize fretting corrosion. According to a published literature by Swaminathan and Gilbert [30], the change of mechanical parameters is different from this study. This is because the lateral stiffness of pin holder was not large enough and the pin was easier to be bend.

Even though this system has some weakness, many valuable insights were gained for future study. The visualization of debris generation during fretting corrosion can be used to collect debris. This system can also combined with many devices in the cell
culture field to study the effect of fretting products on different types of cells. There was one possible explanation for the halo which it is demonstrated in figure 17. The schematic shows some possible compounds based on the solution chemistry. Metal ions release into solution after the oxide film is ruptured. Some of them may combine with different anions in the solution to form some compounds. These possible compounds may gather together and adhere onto the oxide film or even the bare metal surface. It is likely that there is some form of deposition and/or redox reaction related to local impedance behavior. The compounds may reflect different color under light and cause the halo phenomenon.

![Figure 18](image)

Figure 18. Schematic of one possible mechanism of halo generation. Note that yellow triangles stand for surface asperity and orange rectangle means surface oxide film.

The relationship between voltage drop and the other two parameters (area and frequency) can be combined with the impedance and the current data together to develop a heredity integral approach of impedance-voltage-current model to better understand fretting corrosion. The theory is based on the assumption:
where the oxide film abrasion rate (left part of equation) equal to the oxide film repassivation rate (right part of equation). Two equations are used to predict the current and the potential behavior during fretting corrosion:

\[ I(t) = A^T V_{vol}(t = 0) + \int_0^t A^T(t - \lambda) \frac{dV_{vol}}{d\lambda} d\lambda \]  

**Eq.3**

\[ E(t) = Z(t)I(t = 0) + \int_0^t Z(t - \lambda) \frac{dI}{d\lambda} d\lambda \]  

**Eq.4**

where \( V_{vol} \) is oxide film volume. \( A^T \) is called the repassivation admittance and \( Z(t) \) is the time-dependent impedance of the Randle’s circuit. \( A^T \) can be calculated by following equation:

\[ A^T(t) = \frac{\varphi}{\tau} e^{-\frac{t}{\tau}} \]  

**Eq.5**

where \( \varphi \) is constant for each oxide as follow:

\[ \varphi = \frac{\rho n F}{M_w} \]  

**Eq.6**
where \( \rho \) is the oxide density, \( n \) is the effective valance of oxide, \( F \) is the Faraday’s constant and \( M_w \) is the molecular weight of oxide. \( Z(t) \) can be described using the following equation:

\[
Z(t) = R_s + R_{ox} (1 - e^{-\frac{t}{\tau}})
\]

Eq.7

The fretting current behavior during fretting corrosion is described by Eq. 3 and has close connection with the fretting frequency. Higher fretting frequency will lead to larger oxide generation rate, which is described as \( \frac{dV_{vol}}{d\lambda} \). The potential behavior during fretting corrosion is well-predicted by Eq. 4. \( R_s \) is relatively stable while \( R_{ox} \) decreases as electrode area increases (see Table.1). This is one of the reasons why larger second electrodes have smaller potential drops.

The electrochemical circuit has two parts (see Fig. 19). During fretting, electrons are going through the right circuit and generate current. The ammeter is used for measuring the current flow through the circuit (i.e., the combined pin-disk sample and the second electrode) and the voltmeter can measure the voltage between reference electrode and the sample.
Further investigations are needed to understand the details of fretting corrosion. One possible direction could be to use energy-dispersive spectroscopy (EDS) or XPS to examine the surface chemistry of the halo and find the majority chemical elements. Both the fretting surface and observational surface could be put into the scanning electron microscope (SEM) to observe the fretting edge condition and grain structure. The collection and analysis of debris is also a possible direction to study. Sensitive pH indicator can be add into the junction to see whether the change of pH because junction between two fretting surface could be drop very low due to the redox reaction and surface abrasion.
CONCLUSION

The primary conclusions of this study are as follows:

1. A 2-D fretting corrosion test system was designed, developed and evaluated. The system can contact and measure both mechanical and electrochemical properties, while also directly image microscopically the damage generation at the fretting surface.

2. The frequency of fretting corrosion motion is one factor that affects the potential drop of the electrode. Higher frequency leads to faster and larger potential drop under same second electrode area.

3. The area of the second electrode where reduction reactions take place is another factor that affects the potential drop of the fretting corrosion sample. Larger second electrode area causes lower and slower potential drops at the same fretting frequency.

4. The impedance characteristic of the fretting electrodes are area dependent and directly affect the fretting current-potential drop relationship.
SUMMARY

Fretting corrosion, as a major form of mechanically assisted corrosion, is one of the causes of implant material damage, especially for metal implants. This form of corrosion is caused by relative micromotion between two surfaces. Oxide films at each surface will be disrupted and repassivated after fretting and this will give rise to a series of redox reactions, which can cause the metal surface to corrode. Fretting corrosion can lead to many consequences, such as metal and oxide debris generation and is associated with local tissue reactions that may lead to device failure, etc.

Many studies have been performed to understand fretting corrosion from different aspects. Some of the factors are solution chemistry, applied potential, applied load, surface roughness, etc. This study focuses on developing a 2-D fretting corrosion test system that allow to observe the fretting process and collect mechanical and electrochemical data at same time. The disk samples were cut into very small thickness and polished to 600 grit to achieve appropriate roughness. Different frequency, from 0.2 Hz to 10.0 Hz, and second electrode area, from 0.018 cm$^2$ to 0.927 cm$^2$, were performed to test the potential behavior under OCP circumstance.

It was found that debris generated during fretting and accumulated as fretting continued. Fretting corrosion was found on both fretting surfaces of two samples and left a halo of corrosion products around the corrosion region, which could be the deposition of different compounds. The voltage drop of the fretting surface was affected by
frequency and second electrode area. Higher frequency with smaller area lead to the largest and fastest potential drop.

Although this system still needs to be refined including the depth of the holder, it is still a useful tool to use testing fretting corrosion in vitro. One of the possible topics to explore is the development of the halo and its effect on cell viability. Another topic for further study might be about solution chemistry effects (e.g., pH, etc.) and grain boundary behavior during fretting corrosion.
APPENDIX

The following figure is a screenshot of CorrWare® and ZPlot® working window with experiment parameters setup.

Figure 20. Screenshot of CorrWare (up) and ZPlot (down) working window with experiment setup. CorrWare is for OCP test and fretting current test at fixed potential, which is -0.1 V vs Ag/AgCl. ZPlot is for impedance measurement. Note that the scanning voltage amplitude is 10 mV vs. Open Circuit Potential.
REFERENCE


VITA

Name of Author: Dongkai Zhu

Date of Birth: Oct 06, 1990

Mobile Phone: (+1) 315-391-0385

Email Address: dozhu@syr.edu

Education Background:

2009.8-2013.6, School of Biochemical Engineering, Capital Medical University, China
Major in Prosthetics and Orthotics Engineering
GPA: 3.3/4.0, ranking: 10/29

2013.8-present, College of Engineering and Computer Science, Syracuse University, US
Major in Bioengineering, research advisor: Dr. Jeremy L Gilbert
GPA: 3.75/4.0

Software and Programming Skill:
Office Software: Word, Excel, PowerPoint
Engineering Software: SolidWorks, AutoCAD
Image Process Software: Insight Segmentation and Registration Toolkit (ITK), Visualization Toolkit (VTK)
Corrosion Test Software: CorrWare®, ZPlot®
Programming Language: C++, Visual Basic, LabVIEW, MATLAB

Extracurricular Activities:
2009.9-2010.5, Cadre of Sports Department of Students’ Union, School of Biomedical Engineering, China Capital Medical University.

2009-2010, the title of “Excellent Cadre of Youth League”


2010.05-2011.04, Secretary of Sports Department of Students’ Union, School of Biomedical Engineering, China Capital Medical University. Organizing sports activities for freshmen.

2013.5, Outstanding Graduate Student of Capital Medical University

**Internship:**

2012.9-2013.5, China Rehabilitation Research Center, response for making prosthesis and orthotics. Have experience working with doctor and other hospital faculties.