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CORROSION BEHAVIOR OF SURGICAL IMPLANT MATERIALS

by

ROBERT L. SUTHERBY

A THESIS

x ;

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH

OF MASTER OF SCIENCE

IN

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Aand a. Brady

Supervis

External Examiner

Date Feb 2, 1984

DEDICATION

I dedicate this thesis

to my parents

7

ABSTRACT

A study has been made of the corrosion and corrosion fatigue behavior of the surgical implant materials 316L stainless <u>steel and the cobalt alloy, Vitallium, in simulated</u> physiological solutions, both with and without organics. In solutions containing no organics 316L stainless steel was found to be highly susceptible to crevice and pitting corrosion. Under these same conditions, however, Vitallium remained completely passive.

In tests on Vitallium in solutions containing organics it was found that, while glucose had minimal effects on the polarization behavior, serum shifted the corrosion potential in the negative direction. Fortunately, the potential shift occurred wholly within the passive range, so that no increase in the corrosion rate resulted. These tests have also indicated that where Vitallium is involved a saline solution containing serum more closely approximates in vivo conditions, at least in the short term, than does an inorganic saline solution.

In the corrosion fatigue experiments it was determined, with a high degree of probability, that serum reduces the corrosion fatigue life of Vitallium. However, the means by which this reduction occurs, without changing the basic fracture mechanism, was not determined.

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1. INTRODUCTION

Implants, or prostheses, are devices surgically placed in the body to carry out either an active or a passive function which the body cannot naturally perform. It may be to replace a damaged or worn part as is the case for joint or heart valve replacements; to rectify the mode of operation of an organ, for instance a pacemaker; to correct a structural deformity, for instance, a spinal brace; or to simulate an absent or disfigured part of the anatomy, as are the cases in breast and facial reconstructions. Materials for these roles include metals, polymers, and recently, ceramics and modified transplanted materials, such as collagen and bone. Figure 1.1 shows several types of metallic orthopaedic implants, including total hip and knee replacements, a shoulder prosthesis and various bone nails and pins.

Regarding metallic surgical materials, Venable and Stuck wrote the following in 1947:

The effort to find strong materials attracted the interest of many different surgeons in the past and then, as now, they adopted the most immediately available metallic materials and modified them to their use.'

Before the electrochemical aspects of implant degradation were fully appreciated the above philosophy was responsible for the use of such ma - 1s as bronze and iron for external suture, lead shot to h (internal silver sutures, and magnesium as resorbable suture and bone plates.



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Plate 1.1 Assorted Orthopaedic Implants

According to Williams and Roaf², in 1906 John B. Murphy recommended the use of stove pipe wire and carpenter nails and screws for bone fixation.

In response to a flood of new alloys in the early years of this century, tests were conducted to determine their biocompatibilities. In 1926, Austenal Laboratories patented a CoCrMo alloy to be used for dental purposes; they named it 'Vitallium'. Seven years later, after conducting in vivo and clinical tests, Venable and Stuck reported that this material was "completely inert in body fluids". It appears that it was their recommendations based on these findings that initiated the long and extensive use of this alloy group in orthopaedic and other surgical applications.

Some time after 1926 Krupp "18-8 stainless steel" (roughly equivalent to AISI 302) was found to exhibit greater resistance to the body than the widely used vanadium steel, even though it was known to experience pitting corrosion. Later, molybdenum-modified "18-8 SMO" (an early version of AISI 316) with greater resistance to pitting displaced the earlier version.

Such an improvement were these new implant mæterials over their predecessors that, over forty years later, they continue to be the two most widely used. However, they have not been totally without failure for, in addition to infection and loosening of devices, excessive wear and fracture still claim a number of implants each year. For this reason, research and development into new materials is

ongoing.

1.1 Problem Statement

One of the major criteria for the acceptance of a prospective material is its resistance to corrosion in the human body. Since it is impractical or dangerous to make all the necessary tests in vivo (animal or human), simulated, in vitro, conditions are substituted. In the past, most researchers used solutions similar in content to the inorganic fraction of body fluids, based on the conventional wisdom that the extent of damage increases with the chloride ion concentration.

St 3

Recent findings, however, indicate that physiological organics can significantly affect the corrosion behavior of some materials. It is, therefore, the aim of this research to investigate the effects of organics on the corrosion and corrosion fatigue of one of the most often used implant materials, the cast cobalt alloy, Vitallium. Preliminary work will also involve the use of AISI 316L stainless steel.

2. SERVICE CONDITIONS .

Surrounding a prosthesis is a dynamic system of flowing electrolytes, together with living and moving muscle and bone. The following is a brief description of the in vivo conditions in which orthopaedic implants must perform³.

2.1 Fluids

About 56% of the body weight is made up of fluids. Of this, 38% exists outside the cells as extracellular fluids which include plasma, cerebrospinal, gastrointestinal, synovial, and interstitial fluids. It is primarily with the latter two of these that prostheses interact.

Synovial fluid, contained within the joints, is basically a dialysate of blood plasma. With the addition of hyaluronic acid, the fluid is responsible for the lubrication and nutrition of the adjacent articular cartilage joint surfaces. When hips, elbows, or knee joints are involved this fluid is in contact with the prosthesis, Figure 2.1.

Interstitial fluid, or physiological saline solution, lies in the spaces between the cells of both soft tissue and bone. Its major physiological roles are in the maintenance of constant conditions in the internal environment, and as a medium for the transport of nutrients, oxygen, and carbon dioxide between the cells and the capillaries. As with all extracellular fluids, the ionic and nonionic components are constantly being mixed throughout the body by various



transport mechanisms. The normal static concentrations of the fluid components' are given in Table 2.1.

•	
Component	Interstitial (mg/L of water)
Na⁺	3150
К *	184
Catt	96
Mg * *	34
C1-	4000
HCO ₃	1726
HPO4 ⁻ ,H ₂ PO4	194
SO 4	48
Amino Acids	0.3
Creatine	26
Lactate	108
Glucose	1000
Protein	20
Urea	240
· · · · · · · · · · · · · · · · · · ·	

Table 2.1 Static Composition of Interstitial Fluid

Arterial blood is regulated by several buffer and transport systems to within a few percentiles of pH 7.4. Due to a higher carbon dioxide concentration gradient interstitial fluid is maintained closer to pH 7.35.

The temperature of the body is rigorously maintained at 37°C.

2.2 Structural Members

The structural members of the body consist mainly of muscle and bone; muscle developing and supporting the tensile loads, and bones supporting the tensile and compressive loads.

Bone can be described as a biological tissue consisting of long and highly oriented fibres of collagen, densely packed with small crystallites of hydroxyapatite. When reacting to short term loads, bone behaves in a viscoelastic manner, and in response to long term static and dynamic stresses the bone fibres become reoriented to facilitate support of the load.

2.2.1 Loading

Loading on the bones and joints varies between that produced by the minor postural changes during sleep, to that developed during the more strenuous activities of walking, running, and jumping. Rydell⁴ showed that loads on the hip can be as high as two to three times the body weight while walking, and, according to Paul⁵, can reach seven times body weight for young people. Figure 2.2 shows a typical load profile developed on the hip during one walking cycle.

As walking produces between 1 and 2.5 million cycles per year, and because the average age of implant recipients is falling, there is a growing need for prosthesis materials with longer fatigue lives.

: 8



Figure 2.2 Hip Joint Load Profile During Walking

3. THEORY

This section will deal with the theory of corrosion commencing with the thermodynamics and kinetics. With the

aid of the general rate equation, potential-current plots, or Evans' Diagrams, will be presented and used to discuss mixed potential theory.

3.1 Thermodynamics - The Driving Force

In electrochemistry, the free energy of a reactant or product is given by the addition of its chemical free energy and an electrical energy term, as follows:

where

 G_{ec} = the electrochemical free energy, G_{c} = the chemical free energy,

 $q\Phi$ = the electrical energy given by the product

of the charge and the electric potential

at that point.

When molar quantities are in use, the electrochemical free energy of a substance is given by:

$$G_{ec} = G_c + zF\Phi$$
(2)

where

zF = the charge transferred in the reaction. According to the second law of thermodynamics, a reaction will be spontaneous if there is a net loss of free energy as the reactants form products. Thus, in terms of the above expressions, the condition for spontaneity is given

 ΔG_{ec} + zFE < 0

where

by:

E = the potential difference, or voltage. From this last expression it is evident that a corollary to the second law is that at electrochemical equilibrium, the magnitudes of the chemical free energy and the electrical energy are equal:

$$\Delta G_{ec} = -zF$$

When conditions are other than standard, and when the reaction is written in the reduction form, the change in free energy varies with the chemical activities or concentrations according to the van't Hoff isotherm:

 $\Delta G = \Delta G^{\circ} + RT \ln ([REDUCED]/[OXIDIZED]) \dots (5)$ where

ΔG° = the chemical free energy at standard conditions.

By inserting equation (4) into (5) the equilibrium conditions can be expressed in terms of the readily

measurable electric potential difference. Thus:

 $E_r = E^\circ - (RT/zF) \ln ([Re]/[Ox]) \dots (6)$ = $E^\circ - (0.0591/z) \log ([Re]/[Ox]) \text{ at } 25^\circ C$

where

E° = the equilibrium redox potential at standard conditions.

This relationship, known as the Nernst Equation, defines the equilibrium redox potential at a given .(3)

.(4)

temperature and concentrations of reactants and products.

Consider, for example, the following cathodic reaction:

 $O_2 + 2H_2O + \cdot 4e^- \rightarrow 4OH^-$

According to equation (6) the equilibrium potential at 25°C is:

 $E_r = 0.40 + (0.0591/4) \log(pO_2) - 0.0591 \log[OH^-]$ = 1.227 + 0.0148 log(pO_2) - 0.0591pH

If potential, E, is plotted against pH, this equation would describe a series of parallel lines of slope -0.0591, and position depending on the oxygen partial pressure.

More generally, if the reactions involving a metal and an aqueous solution are plotted, a potential-pH, or Pourbaix, diagram could be constructed showing the domains of stability of all reactants and products, and would, therefore, define the conditions under which/the metal should thermodynamically be either free from corrosion, or should corrode.

3.2 Corrosion Kinetics

Although a spontaneous reaction will, by definition, reduce the energy of the system, some minimum activation energy is required before the reaction can proceed. This notion of activation energy is illustrated in Figure 3.1 for the following hypothetical reaction:

 $A \rightarrow A^{z+} + ze^{-}$

Although the reaction involves a reduction in energy, a quantity of energy at least equal to ΔG^{*} must be supplied to



overcome the energy barrier. Where corrosion reactions are involved this 'activation energy' results from the inherent, random, thermal vibration among the atoms.

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.(7)

.(8)

3.2.1 The General Rate Equation

According to Maxwell's Distribution Law the number of particles, N, in a total population, M, having an energy at least equal to a value, E, is given by:

 $N = M \exp(-E/kT)$

where

k = Boltzmann's constant,

T = absolute temperature.

In corrosion, N is proportional to the rate of the reaction, and the equation, on a molar basis, takes on the following form:

Rate = K [A] $exp(-\Delta G^{*}/RT)$

where

[A] = the concentration of reactant A, R = the universal gas constant, K = a constant of proportionality which includes the rate of thermal vibration, a measure of the reversibility of the reaction, etc.

If the rate of the reaction is expressed in moles/second, the rate of the corrosion process can be expressed in terms of electrical current using Faraday's Law:

rate =
$$i/(zF)$$

where

i is the current density, or the electrical

3.2.2 Redox Reactions at Equilibrium Conditions

current-involved-per-unit-area.

When a metal is placed in a solution of its ions the activation energy changes as a charge separation occurs across the interface. Because this electrified interface has an associated electric field any particle crossing the region will be acted upon by the field, and the height of the activation energy barrier will be altered. Substituting the new activation energy term into the basic anodic rate equation gives the following expression for the equilibrium corrosion rate:

> $i_a = KzF exp[-(\Delta G^{\times} - azFE_r)/RT]$ = K' zF exp[azFE_r/RT] ...

The cathodic reaction has a similar form:

 $i_{c} = K" zF exp[-(1-a)zFE_{r}/RT]$

where

 a is a measure of the symmetry of the energy barrier with respect to the peak.
 i_a, i_c are the anodic and cathodic

current densities, respectively.

As the magnitudes of the anodic and the cathodic reactions are equal at equilibrium the net current flow is

.(9)

..(10)

....(11)

zero. The magnitude of the current flowing in either direction is referred to as the exchange current density, i_0 .

3.2.3 Non-Equilibrium Conditions

If a potential is supplied from an external power source the electrode reactions will deviate from the equilibrium state. If, for example, the metal is made more positive (noble), the activation energy for dissolution will be reduced while that for oxidation will be increased.

Substituting the new value of the activation energy gives the following equations:

Anodic $i_a = i_0 \exp[azF\eta/RT]$ (12) Cathodic $i_c = i_0 \exp[-(1-a)zF\eta/RT]$ (13) and a net rate (in this case anodic) of:

 $i_{net} = i_a - i_c$

= i_o [exp(azF η/RT)-exp(-(1-a)zF η/RT)](14)

where

 η is the overpotential, or difference between the electrode potential and

the corrosion potential, thus $E - E_r$.

A visual summary of the corrosion kinetics can be obtained by plotting this latter equation, in terms of potential, E, versus the logarithm of the reaction current density, i, as shown in Figure 3.2 for two redox reactions involving species 'A' and 'B'. This is sometimes referred to as an 'Evans Diagram'.





Reactions

Considering the reactions involving 'A' and 'B', it is noted that as the electrodes are polarized in the positive (noble) direction the cathodic currents become zero, and the net_currents_(equation_(14))_are_equal_to_the_anodic______ currents as given by equation (12). However, in terms of the semi-log plot, the relationship is expressed in the so-called Tafel Equation form:

 $\eta_a = 2.303(RT/azF) \log i_a - 2.303(RT/azF) \ln^3 i_0 \dots (15)$

As the figure implies, and as this equation states the overpotential has a linear relationship with the logarithm of the current density; this slope is referred to as the anodic Tafel slope and is written:

$$b_a = 2.303 \text{ RT}/(azF)$$
(16)

A similar argument applies when the electrodes are polarized in the negative (active) direction, which results in a cathodic Tafel Equation and slope, as follows:

 $\eta_{c} = -2.303(RT/(1-a)zF) \log i_{c} - 2.303(RT/(1-a)zF) \log i_{o}$(17)

 $b_{c} = -2.303 \text{ RT}/((1-a)zF)$ (18) #

As would be expected, the intersection of the extensions of the linear anodic and cathodic Tafel regions occurs at the point given by $(E_r, \log i_0)$ for each reaction.

3.2.4 Concentration Polarization

As the cathodic overpotential is increased the rate of the reduction reaction approaches the diffusion rate of reducible species to the electrode, and the cathodic current line deviates from the linear Tafel slope. When the diffusion rate becomes the rate determining step, that is when the reduction reaction consumes the reducible species as fast as it can arrive, the current reaches a limiting value, i_T .

3.3 Mixed Potential Theory

icorr

When the sites of the two redox reactions are connected through electric and ionic conductors, the redox reactions may proceed concurrently and the total current at any given potential will be the algebraic sum of the currents for the individual reactions. Thus at very negative potentials, in Figure 3.2, the cathodic current will be primarily the cathodic current resulting from the reduction reaction involving 'B'. However, as the intersection of the cathodic curves is approached the contribution from the other reduction reaction will become significant, shown by the dashed line, and finally dominant. A similar situation exists where the total anodic curve crosses the cathodic curve; the total currents become equal at the equilibrium corrosion potential,

 ${\rm E}_{\rm corr}$. The intersection of the two partial curves defines the corrosion current density,

Typical cathodic reactions that interact with anodic processes in neutral aerated solutions are as follows:

 $O_2 + 2H_2O + 4e^- \rightarrow 4OH^-$
$2H_2O + 2e^- \rightarrow H_2 + 2OH^-$

3.3.1 Passivity

In terms of oxidation behavior beyond the linear Tafel region, metals are divided into two groups: those abnormal metals, such as chromium, nickel, iron, aluminum, and titanium, whose corrosion rates fall significantly with further increase in anodic overpotential, and those normal metals whose corrosion rates continue to increase with overpotential. The idealized form of the Evans Diagram for the former of these is shown in Figure 3.2 for material 'B'.

While it is appreciated that the passive metal owes its low corrosion rates to a protective film on the surface, the exact nature of the film is still a subject of controversy. Two theories, however, have wide acceptance today: the adsorption theory and the oxide film theory.

The adsorption theory credits the low corrosion rates with the presence of an array of an adsorbed species, in some cases less than a monolayer thick, on the surface. The presence of such a layer has been demonstrated by Langmuir' and Frankenthal', among others, and its passivating effect has been reported by Agius and Siejka'.

The oxide film theory credits a three dimensional metal oxide layer with reducing the further dissolution of the metal. DeGr moboy and Shreir', in anodic polarization studies on mick 1 in sulphuric acid, found current arrests at potentials which corresponded with several metal oxide/metal reversible potentials. Verink'^o has also demonstrated agreement of polarization data, at several pH

values, with the respective thermodynamic E-pH diagrams.

3.3.2 Breakdown of Passivity

At the noble end of the passive range a sharp increase in current is usually observed. This current rise is associated with a disruption or breakdown of the passive film. Although several mechanisms have been proposed and controversy continues, all the theories involve some damaging species, such as the chloride ion. Whatever the mechanism, it is agreed that surface imperfections, such as inclusions, lattice defects, and micro-pores in the film become initial sites for breakdown and localized attack in the form of pitting corrosion or grain boundary attack.

Polarization studies also reveal a phenomenon which mimics breakdown, however, at potentials below the breakdown potential. Crevice corrosion is a form of localized attack which is limited to regions which have become depleted of the primary oxidizing agent, such as oxygen, and continue to corrode by means of another cathodic reaction, such as hydrogen ion reduction, which depolarizes the anode to a high anodic current regime. The new potential difference, or driving force, can be determined by applying the Nernst Equation to the conditions of the anodic and cathodic half cells. Hydrolysis within the crevice causes the pH to fall and the reaction to become self propagating, with severe localized attack.

The sensitivity of a material to suffer localized attack can be determined qualitatively by reversing the

potential scan after film breakdown has occurred. The form of the polarization curve thus produced can have one of the two forms shown in Figure 3.3. In Figure 3.3(A) the current is seen to fall to values lower than those reached on the forward scan. This indicates that the surface film has become even more protective, a very desirable situation. On the other hand, the current may remain high and exhibit a hysteresis effect. According to Wilde'' the size of the hysteresis is a qualitative measure of the susceptibility of the material to locally corrode. It has been demonstrated¹² that when potentials are held above the crossover, or repassivation potential, E_{rp}, localized attack will continue at any pre-existing pits or crevices. Below the repassivation potential all pitting and crevice attack should cease. A more quantitative assessment of the susceptibility to localized attack can, therefore, be made by_comparing the corrosion, breakdown, and repassivation potentials.

3.3.3 Corrosion Inhibition

In order for corrosion to occur, three conditions must be met:



1. There must be a potential difference, given by the Nernst Equation,

 There must exist mechanisms for the transfer of charge between the electronic and electrolytic conductors, and
There must exist a continuous electronic conductor between the anode and cathode.

Corrosion inhibitors act on the first two of these requirements. The former involves neutralization of the corrodent, while the latter involves the action of a barrier layer former.

Neutralization of the corrodent is achieved by adding a reducing agent to react with the oxidizer which is responsible for the high corrosion rates. An example is the use of hydrazine (N_2H_4) against oxygen in acid and neutral solutions.

Two of the most common types of barrier layer formers are oxidizing inhibitors and surface-active inhibitors. The former of these are used with passive metals when the addition of the inhibitor polarizes the metal into its passive region, for example, the nitrite inhibition of steel, as described by Dean et al.'³.

Surface-active inhibitors adsorb on the metal surface and interfere with the anodic reaction, cathodic reaction, or both. The mechanism of the inhibitor may be to physically block a site, preventing diffusion of reactants, to alter the electrified interface thus changing the kinetics, or to block specific high energy sites, such as surface defects and grain boundaries where dissolution may be concentrated.

Organics make up a large proportion of adsorbed layer \cdot

formers, and have found uses in a wide variety of environmental conditions, as the following examples show:

Table 3.1 Examples of Organic Corrosion Inhibitors

Metal	Reagent	Inhibitor
Fe	H ₂ SO ₄	Amines,
		thiouręas
Al, Zn	NaOH	Glucose

Depending on the action of the inhibitor it may be classified as either an anodic or a cathodic inhibitor, or both. An anodic inhibitor, as the name implies, inhibits the anodic reaction and reduces the corrosion rate as illustrated in Figure 3.4.

Similarly a cathodic inhibitor reduces the rate of corrosion, Figure 3.5, by interfering with the cathodic reaction.

3.3.4 Potentiostatic Polarization

The E-log I plots, discussed above, are generated with a potentiostat, an instrument which maintains a known potential between a working electrode (the metal sample) and a reference electrode, and monitors the corrosion rate in terms of electrical current. By changing the potential, or



Figure 3.5 Cathodic Inhibition

. strictly speaking, the overpotential, on the working electrode in a stepwise manner (potentiostatic polarization), or scanning at a preset rate (potentiodynamic polarization), the corrosion behavior of a material can be evaluated over a wide range of potentials. The former of these methods will be used to examine the implant materials in the subsequent tests.

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3.4 Corrosion Fatigue

Corrosion fatigue, or more generally, environmentally assisted fatigue can be defined as the cracking of a material in response to combined cyclic loading and chemical attack. Significant reductions below expected life can be caused by cyclic loads well below the ultimate tensile stress and, in many cases, below the average yield stress. Often the chemical environment is not excessively aggressive toward the material in the absence of the loading; however, in the presence of a cyclic stress service life may be significantly reduced. It has been demonstrated for an aluminum alloy that the crack growth rate increases as the relative humidity of the air is increased'4. For reasons such as this, some consider that any fatigue testing performed in other than a high vacuum is actually environmentally-assisted-fatigue testing.

To state that corrosion fatigue is simply a superposition of corrosion and fatigue is a gross over-simplification. However, some insight into the phenomenon can be obtained by considering the two components separately, but in the context of the overall problem.

3.4.1 The Fatigue Aspect

The fatigue process has been divided into three stages. Stage I involves crack initiation and the subsequent propagation of the crack through the two to five surface grains. Initiation is considered to have occurred when dislocations accumulate at the metal surface in such a configuration as to be equivalent to irreversible extrusions and intrusions of layers of metal. If such deformation continues, the intrusions, particularly, will become initiation sites for cracks.

In response to the applied stresses cracks will propagate into the metal following slip planes in the surface grains which are under the highest critical resolved shear stress. Because the surface grains may have a random orientation in the metal, Stage I fracture surfaces will tend to be at 45° to the direction of the applied stress.

According to Laird'⁵ Stage II fracture occurs by a plastic blunting process; Figure 3.6 illustrates the steps involved. As the tensile stress is increased toward the ultimate, slip becomes concentrated at the crack tip, in the lobes of the stress field, resulting in irreversible plastic deformation and finally in an incremental extension of the crack. When the load is reversed the crack faces are forced together where the blunt tip partly folds to form a resharpened tip. Thereafter, the cycle is repeated. As illustrated, cracking is usually straight, transgranular and perpendicular to the axis of the applied stress.'

Another theory for Stage II crack growth, called crack closure, has been proposed by Elber'. While this mechanism does not offer a radical alternative for the extension mechanism occurring at the tip of a propagating fatigue crack, it argues that residual tensile stresses, remaining



Figure 3.6 Plastic Blunting Fatigue Crack Propagation Mechanism

after tensile deformation, play a significant role in the kinetics of propagation. "As a consequence of the permanent tensile_plastic_deformation_left_in_the_wake_of_a_fatiquecrack, one should expect partial crack closure after unloading." The crack should also be expected to remain closed for at least a portion of the tensile part of the load cycle. This mechanism means, therefore, that rather than the crack propagation rate being a function of the total amplitude of the stress waveform, or the 'R' value, only that portion of the cycle during which the crack is open actually contributes to propagation. In addition to its use in situations where loading is known to be regular (constant amplitude and frequency), this theory also gives insight into crack retardation and acceleration following low and high loading blocks, and in the propagation rate during random loading' 7. 18, 19.

Finally, Stage III involves the final rupture when the crack length exceeds a critical value and the crack growth rate becomes exceedingly rapid. The advancing crack becomes highly dependent upon inherent weaknesses in the remaining ligament, and is typically characterized, therefore, by a rough surface composed of regions of brittle and/or ductile fracture.

3.4.2 The Corrosion Aspect

In the simplest case, preferential anodic dissolution at the highly stressed crack tip may be the only contribution made by corrosion to the overall mechanism. However, the contribution may be more subtle and <u>devastating. For example, the anodic dissolution reaction</u> may produce a chemical species which embrittles the material by either absorption and accumulation within the metal lattice, or it may adsorb at the high energy crack tip, presumably reducing the surface energy, and leading to decohesion of the metal.

On the other hand, if the species is an effective corrosion inhibitor, the effect of the environment may be significantly reduced, possibly returning the situation to one resembling pure fatigue.

3.4.3 Variables Affecting Corrosion Fatigue Cracking Rates Wei and Speidel² have outlined a large number of variables which can affect the corrosion fatigue cracking rate. These have been divided into three groups, as follows.

Mechanical Variables

Maximum stress or stress ratio, R. Cyclic loading frequency. Cyclic load waveform (e.g. square or sinusoidal waves). Stress state (e.g. bi- or triaxial). Residual stress. Notch or crack size. Specimen geometry.

Metallurgical Variables

Alloy composition.

Distribution of alloying elements.

Crystal structure and microstructure. Prior heat treatment.

Mechanical properties.

Environmental Variables.

Temperature.

Concentration of damaging species.

Electrochemical potential.

Coatings, inhibitors, etc.

Fundamental to all theories of fatigue and corrosion fatigue crack propagation is the involvement of slip in plastic deformation and work hardening. Variables, therefore, which limit the number and movement of the dislocations will have a major effect on the cracking mechanism.

3.4.3.1 Mechanical Variables

Slip occurs in response to shearing stresses. Load configurations, therefore, which tend to produce low shear stress zones, such as under triaxial stress conditions, will result in the accumulation of dislocations and possibly in the formation of a crack. This sequence of events can occur in thick materials in response to external loads, or at the roots of cracks or notches. Just as the applied stress is a major concern, the presence of residual stresses from prior forming and heat treating practice can also have a significant effect.

Of particular importance in corrosion fatigue is the cyclic loading frequency and waveform. Barsom² ¹ has demonstrated that the crack propagation rate often increases as the loading frequency falls. Furthermore, the loading waveform is also seen to be a controlling factor². Crack rates are highest when the rise-time of the tensile waveform is longest; a sinusoidal loading curve is, therefore, more destructive than a square wave.

These phenomena, unique to corrosion fatigue, appear to be the results of synergistic combinations of the fatigue and corrosion mechanisms.

3.4.3.2 Metallurgical Variables

As the generation and movement of dislocations has a fundamental dependence on the particular material, the metallurgical condition has a profound effect on crack initiation and growth. In addition to its crystal structure, one of the most important fundamental properties of a material is the relative magnitude of its stacking fault energy (SFE). This energy term is simply the surface energy of a plane of atoms in the solid which is stacked in a pattern out of sequence from the ideal. As the energy term falls, the length of the stacking fault increases as does the force required to make it cross slip. With regard to low cycle fatigue crack propagation behavior, face centre cubic (FCC) alloys can be grouped according to relative magnitudes of SFE, which bear directly on the dislocation structures generated in those materials in response to cyclic loading. Low SFE alloys, such as austenitic stainless steels and cobalt-chromium alloys, having high resistances to cross slip form fine planar dislocation structures; high SFE materials, for example copper and aluminum, form cell-like dislocation structures owing to their low resistance to cross slip.

A consequence of the higher resistance to cross slip, in low SFE materials, is the formation of fine persistent slip bands, which cause the material to reach its saturation hardness at a higher number of cycles than the high SFE materials. This capacity to retain the reversible nature of slip prolongs the time required for crack initiation. Furthermore, fine slip more evenly distributes plastic deformation at the crack tip, which leads to lower stress concentrations and slower crack propagation rates than for high SFE alloys.

Two other factors which are considered to have significance in fatigue crack propagation are grain size and ductility. Whereas in high cycle fatigue, where life increases with decreasing grain size, the same is not considered to be the case for low cycle fatigue. According to Feltner and Laird²³ the grain size has no effect on the rate of crack growth in high SFE alloys, presumably as a consequence of the dislocation cell structures. Concerning low SFE materials, on the other hand, it has been hypothesized by Laird²⁴ that fatigue cracks propagating by

plastic blunting should be slowed with decreasing grain size.

The effects of ductility vary according to whether the fatigue is strain-, or stress-controlled. In strain-controlled, low cýcle fatigue, the greater the ductility the greater the capacity for deformation and blunting²⁵, therefore, the lower the propagation rate.

In stress-controlled, low cycle fatigue, high ductility is especially important for cold-worked materials, which fatigue soften². If the material posesses insufficient ductility the initiation, time will be short, and propagation rates will be high. For annealed alloys the importance of ductility will increase with the level of the stress.

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4. LITERATURE REVIEW

This chapter reviews the relevant clinical and scientific literature on the corrosion and corrosion fatigue behavior of 316L stainless steel and cobalt alloys. A review is also made of the literature dealing with the effects of organics on these and other materials.

4.1 Stainless Steel

4.1.1 Clinical Findings

Clinical reports^{2,7-3,6} indicate that the fracture of 316L stainless steel implants is usually accompanied by metallurgical defects in the form of porosity, manufacturing defects such as grinding marks, and various design problems. However, the actual fracture mechanisms generally fall into three groups: fatigue, corrosion fatigue, and stress corrosion cracking. While it is sometimes equivocal whether the responsible mechanism is actually fatigue or corrosion fatigue, some cases have been reported where two mechanisms were clearly active during the life of a single component^{3,6,432}.

Some form of corrosion is usually associated with used stainless steel implants. In one study by Colangelo and Greene³⁵ corrosion was observed on 91% of all retrieved components, and crevice corrosion was present at 42% of the possible sites. In a number of cases, reported by Cahoon and Paxton²⁸, crevice corrosion was credited with causing such

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severe tissue irritation that implant removal was required.

4.1.2 Corrosion Studies

Considerable research has been conducted on the corrosion behavior of austenitic stainless steels under saline conditions. While some reports have dealt with stainless steel as an implant material, much of the published results deal with material selection for use in seawater, chemical process lines, and recently, geothermal power plants. Fortunately, these latter service conditions are similar enough to the inorganic fraction of body fluids to make, at least, a first approximation to the behavior of these materials in vivo.

It has been known since the earliest applications of the material that "it suffers from localized attack. In 1943 Venable and Stuck'' reported that '18-8' stainless steel experienced "pit corrosion" in body fluids, but the addition of molybdenum reduced this tendency.

Recent tests in various other media have indicated similar problems. Streicher' studied the corrosion of twenty-five different alloys in seawater and found that the high Cr-Mo alloys, such as Hastelloy C-276 and Inconel 625, had the greatest resistance to localized attack, while AISI 316 had one of the worst. In similar tests Hack' found that crevice corrosion on 316 was variable.

Susceptibility to localized attack has also been demonstrated by electrochemical means. Bennett et al. "

using cyclic anodic polarization compared the relative resistances of AISI 304, 316, and 317 stainless steels in pulp mill white water; 316 was rated between the other two.

Bandy and Cahoon' studied the effect of alloy composition on the resistance of austenitic stainless steels to localized attack in Ringer's Solution. In the case of 316L the repassivation potential was below the corrosion potential; increasing the molybdenum content; however, produced a higher breakdown potential, and a very small hysteresis loop.

While polarization tests usually give corrosion .potentials in the vicinity of -0.2 V (Mueller and Greener**) to 0.065 V (Bandy and Cahoon*'), long term tests indicate that the corrosion potential generally drifts in the positive direction to an average value, after twenty days, of approximately 0.14 V (Levine and Staehle*'). Brettle and Hughes** also show a similar trend; however, the potential after 100 days, in their test, was only about 0.014 V. Because they did not report if the solution had been oxygenated, this potential value may reflect deoxygenated conditions. Regardless of the source, however, all potential-time plots show frequent potential fluctuations, similar to the results of Levine and Staehle*', which Hoar and Mears*' suggest are caused by passive film breakdown and pitting.

Hoar and Mears's, who conducted both potentiostatic polarization and potential monitoring tests, report that after 480 hours a corrosion potential of 0.26 V was reached. Because they also recorded breakdown potentials of 0.116 V to 0.24 V, they concluded that spontaneous film breakdown is

possible if the stainless steel remains in aerated saline for long periods.

Some confusion appears to be associated with the value of the breakdown potential. Anodic polarization curves generated by Mueller and Greener⁴³ and Bandy and Cahoon⁴⁴ for annealed 316 in aerated Ringer's Solution show breakdown in the vicinities of 0.35 V and 0.55 V. Mueller and Greener⁴⁴ suggest that the difference is due to a variation in oxygen partial pressure, an explanation which is consistent with another of their findings, that breakdown under deaerated conditions occurs at 0.16 V. In similar tests using cold worked 316L, however, Syrett⁴⁴ found that the breakdown potential was independent of oxygen content.

Concerning Bandy and Cahoon's'' results, no mention is made of varying oxygen concentrations. However, it is reported that in the early tests, possibly referring to those giving lower breakdown potentials, crevice attack occurred at the gasket-sample interface. As the breakdown potential for crevice corrosion is lower than that for pitting, it is possible that this phenomenon was responsible for the lower values in the cases of both the annealed and cold worked materials.

A few papers have been published recently dealing with. the effects of physiological organics on the corrosion behavior of 316 stainless steel; however, the results appear to be contradictory.

Samitz and Katz⁴⁷ compared nickel dissolution from 316L in various solutions including isotonic saline (0.17M NaCl), pooled sweat, whole blood, and plasma. After one week, they found nickel concentrations to be greater, for the most part, in the physiological solutions than in the isotonic saline.

In 1980, Brown and Merritt'' reported results which supported those of Samitz and Katz''. 316L coupons were immersed in solutions of saline, saline plus 1% serum, and saline plus 10% serum. After maintaining an anodic potential of 5 V for thirty minutes, the samples were compared. Samples in solutions containing serum were attacked more than those in saline, and the extent of damage increased with serum content. Anodic polarization curves in the same solutions showed that the critical current for passivation was increased in saline containing serum.

Contrary to the results of the two reports above, however, another series of tests, also reported by Brown and Merritt'', on fretting corrosion of 316L, indicated that the serum may have actually been beneficial; weight losses were over seven times greater in isotonic saline than in 10%

serum. As a side note, their results showed a more negative corrosion potential in serum than in saline.

4.2 Cobalt Alloys

4.2.1 Clinical Findings

In a survey of ten reports'''' dealing with both stainless steel and CoCrMo implants, the only failure mechanism reported for the latter was fatigue. In one paper, by Cahoon and Paxton'', however, this mechanism was assumed simply on the basis that the implant failed in a brittle manner and because it had been cyclically loaded. This case emphasizes a caution stated by White et al.'' on the identification of fracture mechanisms; while fracture surfaces are often obliterated subsequent to failure, they note, it is important to consider that the necessary conditions for corrosion fatigue and stress corrosion cracking, and other mechanisms also exist, and that the identification of all brittle failures as fatigue will only "lead to complete ignorance of the root cause of the problem."

As was the case with the stainless steel implants, all the CoCrMo fractures had associated defects. However, unlike the stainless steels, no corrosion of any type was reported.

4.2.2 Corrosion Studies

Compared to the volume of literature on the corrosion of stainless steels, relatively little is available on the ambient temperature corrosion of CoCrMo alloys. Most of what is available, however, deals with the use of the material in surgical implants.

The/anodic polarization of cast CoCrMo has been studied by several teams in solutions simulating the physiological milieu. Cahoon et al. " performed potentiodynamic polarization on a cast CoCrMo in Ringer's Solution containing physiological pressures of oxygen and carbon dioxide. The anodic curves, thus produced, were similar to those for the stainless steel in that the material self-passivated with a current of approximately 10^{-6} A/cm². On reversing the potential scan, however, unlike the 316L, the material repassivated and exhibited only a small hysteresis. This behavior agrees with the results of Sury⁵¹ who studied Vitallium in 10% HCl, and Uhlig and Asphahanis', who examined HS-21 in 4% NaCl at 70°C, all without experiencing localized attack. In fact, this material is so resistant to localized attack that it took Pugh et al. 53 74 days to detect crevice corrosion on Vitallium in 10% HCl plus 1% FeCl₃, and of 40 samples tested only two showed evidence of the attack.

While Hoar and Mears's did not study reverse polarization behavior, they found forward anodic curve forms similar to those reported by the authors mentioned above in Hank's Solution, human blood, and in NaCl solutions.

Depending on the source, breakdown potentials are reported from approximately 0.3 V, in the case of Mueller and Greener's, who used aerated Ringer's, to 0.87 V for Brettle and Hughes'' where 0.17 M NaCl was the test solution. While there are several cases when high breakdown potentials arise (such as high potentiodynamic scanning rates), Hoar and Mears's report that the potentials in 0.17 M NaCl are always higher than in balanced saline solutions, such as Hank's Solution.

As was the case with 316L, corrosion potentials of CoCrMo rise with time. Lucas et al.⁵⁴ reported a corrosion potential after a number of hours in 0.9% saline to be -0.24 V, while Hoar and Mears⁴⁵ reported an average value of 0.26 V, between 20 and 200 days in 0.17 M NaCl, and 0.07 V, after 71 days in vivo. Because these long term values, and particularly that for the in vivo situation, are lower than the breakdown potential, Hoar and Mears⁴⁵ consider spontaneous film breakdown to be highly unlikely.

Regarding the interactions between implant materials and physiological organics, research has continued, to varying degrees, from two standpoints: the effects of body organics on the implant metal, and the effects of the dissolved metal ions on the body. Work on the first of these has been relatively sparse.

Samitz and Katz^{*}, in addition to working with stainless steel, also studied Vitallium and found the concentrations of nickel to be greatest in physiological solutions, as was the case for the steel.

Hoar and Mears⁴⁵ monitored corrosion potentials in vivo and in vitro and found the former to be up to 200 mV more negative after 71 days. In a test reported by Clark and Williams^{5,5}, a CoCrMo alloy was immersed in a saline solution containing serum albumin. They found that the cobalt and molybdenum were being selectively removed from the surface, possibly by the formation of stable complexes with the surrounding organics.

Perhaps the first unifying theory relating the role of physiological organics to the corrosion of an implant material was made recently by Woodman et al.⁵. In tests on Vitallium total finger prostheses implanted into cats they learned that the concentrations of dissolved cobalt and nickel increased exponentially over the one year test period. Furthermore, both ions were only detected bound to serum proteins. It was hypothesized that the continued increase in the metal ion levels was due to an autocatalytic mechanism directly involving the serum proteins: the presence of dissolved metal ions stimulates the immune system which results in a supply of proteins; metal ions complex with the proteins, diminishing the ion concentration adjacent to the metal, which leads to continued dissolution.

In comparison, a greater number of studies have considered the interactions between metal ions and physiological organics. Among the information gathered to date are the following points:

Cr, Co, and Ni tend to circulate in the body and

become stored in various organs, 57

at least Co forms complexes with serum proteins, and possibly other organics, ⁵ *

Cr levels are highest near the implant, "" the metal-protein bond may grow stronger with time."

4.3 Effects of Organics on Other Alloy Systems

In addition to the research performed on the effects of organics on stainless steel and CoCrMo, work has also been made on other alloy groups.

While testing candidate materials for implantable electrodes, Niazy et al.¹ studied the effects of various organics on the anodic polarization behavior of aluminum and zinc. While none of the test solutions had any visible effect on the passivation of aluminum, solutions containing bovine plasma protein, and a mixture of chondroitin sulphate and arginine produced passivity on zinc where none existed in isotonic saline.

Similarly, in anodic polarization experiments on copper, nickel, cobalt, and titanium in Ringer's Solution, Svare et al.² found that solutions containing the amino acid cystine enhanced the passivity of copper, while for nickel passivity was inhibited.

The effects of amino acids were also investigated by Solar et al. ' on Ti-6Al-4V. In anodic polarization tests arginine and methionine shifted the reversible potential in the negative direction, while proline, tyrosine, glysine, and cystine shifted it in the positive sense. No effects were noted in the passive regions. Ti-6Al-4V was also the material examined by Aragon and Hulbert'. In tests similar to those conducted by Solar et al.', lower current densities were attained using plasma than with isotonic saline. Furthermore, while lactated Ringer's Solution shifted the corrosion potential in the negative direction by over 300 mV, the corrosion current increased by approximately one order of magnitude over the value in isotonic saline.

4.4 Corrosion Fatigue of Cast CoCrMo

As was indicated above no corrosion related fracture mechanisms have been reported in the clinical literature for CoCrMo. However, this may not reflect the actual mechanisms, for when faced with equivocal data, one may choose to identify the fracture mode as fatigue, based on previous experience with the material, rather than the more debatable mode of corrosion fatigue.

A corrosion fatigue study was conducted by Luckey." using deaerated lactated Ringer's Solution. Samples were sinusoidally loaded in reversed bending at a rate of 30 Hz. The endurance limit at 10⁷ cycles was reported to be 419 MPa (60.8 ksi); however, he remarked that the endurance curve was still decreasing at this point. While he did not discuss his findings in detail, he concluded that his work "... has shown potential environmental effects on the high cycle corrosion fatigue behavior of cast Co-Cr-Mo ...". Corrosion fatigue tests were also conducted by Miller et al. ". Cast CoCrMo samples were sinusoidally loaded in an aerated tissue-culture medium at a rate of 2500 Hz. They justified their use of this high loading frequency by

stating that "... the work of Bowers and Staehle and Devine and Wulff indicate that corrosion fatigue has a relatively small effect on this alloy." (the works referred to are unpublished). From their results, however, they calculate an allowable design stress of 321 MPa (46.5 ksi), such that there is a 95% confidence of less than one failure in one thousand after 5 x 10^7 cycles, assuming no manufacturing defects.

Fatigue testing in air was performed by Ducheyne et al.''. In a four point bending jig, intact hip prostheses were loaded at 30 Hz to various stress levels. They reported a fatigue limit of 424 MPa (61.5 ksi) at 10⁷ cycles.

Gilbertson^{••}, also using intact hip prostheses, studied the relationship between fatigue strength and the loading orientation. He found that the endurance limit at 10^7 cycles increased from 1054 to 2122 N (237 to 477 lbs), or approximately 1 to 3 body weights, as the loading axis approached the long axis of the device.

In a review of the fatigue strengths of various cobalt-based implant alloys manufactured by Sulzer Brothers Ltd., Lorenz et al.'' discuss the microstructures, the fatigue resistance, and the fatigue fracture morphologies of cast CoCrMo tested in air. From the basic fatigue strength of 190 - 280 MPa (27.6 - 40.6 ksi) in the as-cast condition, the strength increases to 280 - 350 MPa (40.6 - 50.8 ksi) following a homogenizing anneal. Solution annealing, however, while increasing the impact and tensrle strengths, produces fatigue strengths of only 220 - 280 MPa (31.9 -40.6 ksi). This lower fatigue strength is blamed on the presence of voids, termed "Kirkendall holes"'', which remain after the carbides are dissolved.

The relative effects of notching and cyclic loading in a corrosive environment were investigated by Hughes et al.''. Cast CoCrMo samples were tested as both smooth and notched specimens in air and in saline, and the endurances at 10⁸ cycles were compared. Their results are listed below:

Smooth S			Specimens
Tested in <u>Air</u>	Tested in <u>Saline</u>	Tested i <u>Air</u>	n Tested in <u>Saline</u>
248	234	138	48
(36.0)	(33.9)	(20.0)	(7.0)

Endurance Limit at 10⁸ Cycles, MPa (ksi)

Table 4.1 Results Reported by Hughes et al.

Based on these findings, they calculated that the total effect of notching and testing in saline was to reduce the endurance limit by 81%. Of this the presence of the notch had the greatest effect accounting for 55% of the total reduction, while the chemical effect only accounted for 7%. Finally, an interaction effect, defined as the difference between the total effect and the individual contributions of the notch and the environment, was calculated to contribute over 30% to the total. Therefore, although the individual effect of the saline was small, the combination of it with

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the notch was very significant, and may have even been greater if the loading frequency were less than 100 Hz.

Regarding fractographic features, most sources, including the clinical reports, agree that fatigue crack propagation in cast CoCrMo is charaterized by cleavage fracture resembling Stage I crack growth. Both Gilbertson'' and Lorenz et al.'' suggest that this is related to the low stacking fault energy, which makes cross slip difficult. Ducheyne et al.'', on the other hand, suggest that a deformation-induced martensite phase transformation from FCC. to HCP may be responsible.

5. MATERIALS AND PROCEDURES

The research was divided into four sections according to the objectives. The first section was the development of an experimental procedure which could be used to generate

reliable potentiostatic polarization curves. The second and third parts dealt mainly with the polarization behavior of a cast cobalt alloy in solutions containing combinations of glucose and other physiological organics, in the form of calf blood serum. The fourth, and last part, was a study of the effect of physiological organics on the corrosion fatigue crack propagation rate and fracture morphology in a cast cobalt alloy.

The following section will review the chemical compositions, metallurgical conditions, and mechanical properties of the three materials studied, and will describe the equipment used and the procedures followed.

5.1 Test Materials

In addition to the 430 stainless steel used in the procedure development runs, the three materials involved in the tests were: 316L stainless steel and cast Vitallium both machined from used total hip implants, and Vitallium cast into the form of round tensile specimens. Plate 5.1 shows that the stainless steel implant, used as the source of material, had fractured in service. The Vitallium implant, similar to the one shown in Plate 5.2, was, on the other hand, intact. It is assumed that it was removed because of

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Plate 5.1 Fractured Stainless Steel Implant

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Plate 5.2 Intact Vitallium Implant

infection or loosening.

The Vitallium fatigue specimens, produced and supplied by Howmedica Inc. of Rutherford, New Jersey, were purported to be in the same metallurgical condition as the "Vitallium" prostheses that they manufacture.

Analysis of the test materials consisted of determining their chemical compositions, hardnesses, microstructures, and in the case of the fatigue specimens, tensile.

productives. Chemical analysis was performed by the Alberta Research Council (ARC) dsing their Induction Coupled Plasma (ICP) spectroscopy equipment, and in the Department of Mineral Engineering using the Leco carbon analyser. Among the results reported by ARC the compositions given for manganese, phosphorus, and sulphur were all high, in most cases, several times the maximum specified limits. In the case of the stainless steel the concentrations reported were much higher than the levels readily obtainable in even the lowest steel grades. It was, therefore, considered that the values for these elements did not reflect actual

concentrations, but, rather, erroneous analyses on the part of the laboratory. While these results are not included in the following summary of results, they are attached as Appendix B.

All other analyses were carried out in the Department of Mineral Engineering.

5.1.1 Stainless Steel

One grade of stainless steel currently used in the manufacture of implants is 316L. In North America this material is often required to conform to the minimum requirements of ASTM F55-82. The compositional requirements of this standard and the assay of the test material are given below.

Table 5.1 Chemical Composition of Stainless Steel Test Material

Element	ASTM F55-82 Grade 2	Test Material
C 🎢	0.030 [%] (max)	°; <0.01%
Cr	17.0-19.0%	18.0%
Ni	12.0-14.0%	14.0%
Мо	2.0-3.0%	2.55%
Cu	0.50%(max)	0.21%
N	.0.10%(max)	Not Analysed

These results indicate that the material conformed to the chemical requirements of ASTM F55-82

5.1.1.1 Metallographic Examination

Longitudinal and transverse sections were cut from the stem of the hip prosthesis. Plate 5.3 shows a transverse cross section to have an equiaxed grath structure with an ASTM size of approximately #5. These features together with the absence of any deformed twins indicate that this austenitic material was in the annealed condition.



Plate 5.3 Typical Microstructure of the 316L Stainless Steel (100X)

5.1.1.2 Hardness Results

Rockwell hardness and Vickers microhardness measurements were made on a transverse section of the implant with these results:
naraness vi	Ofgintess Steet	iest Material
· · · · · · · · · · · · · · · · · · ·	Aler.	4
	Rockwell Hardness	Vickers Microhardness
	алан айтан алан алан алан айтан а Айтан айтан айта Айтан айтан айт	
Test Sample	HRB 75 (average of 5 readings)	HV 162 (average of 5 readings)
Typical		
Values ⁷	HRB 85 - 95	Not Given
		3
Dead Soft''	• HRB 80	Not Given

Table 5.2Hardness of Stainless Steel Test Material

Although the hardness specification was not obtained from the manufacturer, the literature values would indicate that the material was in the fully annealed condition.

Although a failure analysis was not performed on the implant, it is probable that the mechanism of fracture was aggravated by the low hardness and, therefore, a low tensile strength. However, based on the chemical composition, it is considered that the corrosion properties would have been similar to material in a harder condition.

5.1.2 Cast Vitallium

Of the several cast CoerMs prostheses on the market the one used in this research was manufactured by Howmedica Inc. and is referred to by them as 'Vitallium'. Similar, if not identical alloys are also produced by other manufacturers under other proprietary names, such as: Haynes 21 (Cabot Co.), Zimaloy (Zimmer Co.), Stellite (Union Carbide), and Protasul-2 (Sulzer-Brothers-Ltd.).

In North America the materials for CoCrMo implants are usually produced in conformance with the minimum requirements of ASTM F75-76. The chemical composition required for that specification, and the assays for the two alloys studied are as follows:

Table 5.3 Chemical Composition of Vitallium Test Materials

Elements	ASTM F75-76	Implant Sample	Fatigue Sample
ć	0.35%(max)	0.21%	0.03%
Si	1.00%(max)	0.82%	0.99%
Cr	27.0-30.0%	27.1%	24.0%
Ni	2.5%(max)	2.10%	0.16%
Мо	5.0-7.0%	6.05%	5.35%
Fe	0.75%(max)	0.18%	0.17%
Cu	Not Specified	0.05%	0.05%

Of these two alloys only the implant material conforms to the ASTM specification; the fatigue sample does not conform as a result of the low chromium content. While this discrepancy means that the average corrosion properties of the fatigue sample material would be inferior to that of the

implant material, it cannot be said that the alloy is unacceptable for use in the body as it is not known if conformance_to_the_quoted_specification_is_required_in_this____ instance.

5.1.2.1 Metallographic Examination

Transverse sections were cut from the neck of the prosthesis and from the gage length of a fatigue specimen. In Plates 5.4 and 5.5 the microstructures of the two samples appear to be different; the prosthesis material contains more interdendritic carbide than the fatigue test material. It is believed that the only cause for this was the lower carbon content in the latter.

Common to both materials was the large dendritic grain structure (ASTM size 0), and only a single layer of fine surface grains. These features are typical of castings which were poured into heated molds and slowly cooled.

Both materials also contained porosity. In the case of the prosthesis material it was predominantly present at the grain boundaries, while in the fatigue specimen the porosity was located both within the grains, usually associated with carbides, and at the grain boundaries.

Based on these observations, in addition to the lack of extensive coring it is likely that both materials were in the cast and homogenized conditions. In the case of the fatigue specimen, the annealing temperature may have been somewhat high and resulted in a partial dissolution of the



Plate 5.4 Typical Microstructure of Vitallium Hip Implant

(100X)



Plate 5.5 Typical Microstructure of Cast Vitallium Fatigue Specimen (100X)

interdendritic carbides and the formation of porosity.

5.1.2.2 Hardness Results

Rockwell 'C' hardness tests and Vickers microhardness tests were made on both materials with the following results.

Table 5.4Hardness of Vitallium Test Materials

	ASTM F75-76	Implant Sample	Fatigue Sample
Rockwell ^(C)			
Hardness	25-34	27	25
Vickers Microhardness	Not Specified	L 295	322

Although the hardnesses are near the lower limit of the specified hardness range, both material's conform to the specification.

The discrepancy between the microhardnesses and the Rockwell hardnesses of the two samples is likely a result of a variation in hardnesses among various heats.

5.1.2.3 Tensile Results

Three of the fatigue samples were tensile tested with the following results:

Table 5.5 Tensile Properties of Vitallium Fatigue Specimens

	ASTM F75-76	<u> </u>	Samples	
Tensile Strength	655 MPa (min) (95 ksi)	630 MPa (91.4 ksi)	663 MPa (96.2 ksi)	727 MPa) (105.5 ksi)
Yield Strength (0.2% Offse	450 MPa (min) (65 ksi) et)	418 MPa (60.6 ksi)	397 MPa (57.6 ksi)	272 MPa (39.4 ksi)

Elongation 8% (min) 8.6%

These results show the tensile strengths of the fatigue samples to be highly variable (from -3% to 11% based on the minimum specified) as well as only marginally acceptable. Furthermore, the yield stresses are even more variable, and below the specified minimum values. While conformance to this specification may not be required, it is almost certain that the variability is unacceptable.

Although the corrosion behavior of the fatigue specimens would be almost identical to the prosthesis material, the endurance limit may be somewhat low as a result of the marginally acceptable tensile strength, and low, but acceptable, dectilit

5.1.3 Water

Common to all tests was the water used to prepare the test solutions. It was assumed that the building's purified

water would be suitable. However, analysis showed the resistivity to be well below the value given in the system specifications. Since maintenance was scheduled on the order of weeks per interval, a fluctuation in water quality was expected. This source was, therefore, judged not to be sufficiently consistent, or dependable for the research. This judgement was reinforced when, during early experimentation, the deionizer resin was changed and, for several days thereafter, the water had a strong organic odor and a surfactant quality.

To ensure consistent quality, water from the building's system was first passed through a series of ion exchange columns. Any organics present were then oxidized during a single distillation from a potassium permanganate solution. Water produced by this means had an average resistivity of 10K ohms-cm.

5.1.4 Simulated Physiological Solutions

The ingredients of the basic simulated physiological solution were as follows:

		Table 5.6	
Composition	of	Simulated	Physiological
17		Solution	

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Component	Weight (g/l)
NaCl	6.30
КСІ	0.374
NaHCO3	2.76
Na 2HPO 4	0.071
NaH ₂ PO ₄ .H ₂ O	0.069
Glucose	1.000
MgCl ₂ .6H ₂ O	0.203
MgSO	0.060

The dry ingredients for simulated physiological solutions (SPS) were prepared in batch quantities of up to thirty times the nominal composition. To ensure homogeneity, all ingredients were first pulverized with an agate mortar and pestle, and to avoid precipitation of magnesium salts during dissolution, all components bearing magnesium were segregated into a separate vial.

Besides the glucose, organics were also added in the form of calf serum, prepared by Flow Laboratories. Primarily concentrations of 0.01%, 0.1%, 5%, and 15% were used; however, in one test 0.001% serum was used.

Solutions were prepared in a one litre volumetric $\frac{1}{4}$ flask. First the serum was diluted, then the

non-magnesium-bearing salts were dissolved, followed by those containing magnesium. The solutions were mixed with a magnetic stirrer.

5.1.5 Gases

The gases used in the testing of the cobalt alloys were. prepared by Matheson of Canada Ltd. and hac the following compositions:

Table 5.7 Gas Compositions

(0.056 atm) CO ₂ 45 mm	Interstitial ³	02	40 mm (Hg)
그는 그 같은 것 같은			(0.056 atm)
(0, 0, 0, 2, -+-)		CO 2	45 mm
(U.U63 atm)			(0.063 atm)

0 2

N₂

6.65 molar percent D_2 CO₂ 6.11 Ar Balance

Bottle 2

Bottle

6.47 molar percent CO₂ 6.11 Balance

5.2 Polarization Studies.

5.2.1 Development of Experimental Procedure

Before potentiostatic polarization testing could begin it was necessary to develop a standardized experimental procedure that would yield reproducible results. For this purpose ASTM designation G5-78 was used. This practice involves the generation of anodic polarization curves for

430 stainless steel in deaerated $1N H_2SO_4$. It provides standard curves with which to compare results, and discusses experimental problems that may be encountered.

Instructions for solution and sample preparation were generally adhered to, with the following exceptions:

- 1. It was recommended that the working electrode be
 - degreased in boiling benzene; this was not done. Rather, samples were ultrasonically rinsed in acetone.
- 2. Argon replaced hydrogen as the purging gas. According to Chance et al.^{1,2}, when all other factors remain the same, this does not affect the results.
- The platinum counter electrodes were not platinized.
 The starting potential was below the corrosion potential.

5.2.1.1 Test Apparatus

Figure 5.1 gives an illustration of the equipment used in this initial work. The system has four main units: the polarization cell, the gas purification and monitoring system, the potentiostat, and the current measuring and recording system:

The Cell

The cell and the glassware for the five electrodes were similar in design to those described by Greene''. The temperature of the test solution was maintained at 37°C by partial immersion in an oil bath.

The Gas System

As mentioned, argon was used as the purging gas. Because it was found to contain oil, which accumulated in



the Tygon tubing, an effort was made to remove the dil by passing the gas through calcium sulphate chips at a rate of 20 mL/min. Any residual oxygen was removed by passing the gas over copper turnings or titanium powder at 600°C.

The Potentiostat and Current Measuring Systems

During the course of the experimentation two potentiostars and current measuring systems were utilized. Initially, an AMEL 549 potentiostat was used. Current was determined by measuring the voltage drop actops a decade resistance box, placed in series between the counter, electrode and the potentiostat. A strip recorder and

voltmeter were isolabed from the Cell by an operational amplifier wired as a voltage follower with a gain of 10:1. The strip recorder monitored corrosion current with time. This system had a number of problems and limit tions. As Figure 5.2 shows, the characteristic curve for the

amplifier deviates from linearity below 0:03 volts output, and gives negative output for input potentials below 2 x 10⁻⁴ V. These features of the amplifier became important as currents in this range were often encountered as the corrosion potential was approached.

One other limitation resulted from the fact that resistors of greater value than 10KΩ were found to affect child reactions. The decade resis box, therefore, could only contain five ranges; the resistor values are given

below.



Table 5.8 Decade Resistor Values

Nominal-Value (Ohms)	Measured Val (Ohms)	<u>ับ</u> ย
1	1.0	
10 1	. 10.0	•
100-	99.6	
e # 1K	980.3	G
10K .	8826 0	

The Second system consisted of an AMEL 551 potentiostat with an AMEL 560 log interface. As before, the trip recorder monitored current with time and an electrometer was used to digitize the analog durgent values. The experimental arrangement for this system is illustrated in Plate 5.6 and in Figure 5.3.

5.2.1.2 Procedures

Samples were prepared by wet abrasion down to 600 grit, with SiC paper. They were stored in acetone during transport from room to room. Prior to immersion they were rinsed in the building's purified water, then dipped in test solution. Before the test began the solution was purged of oxygen for at least one-half hour prior to the introduction of the sample, then the system was allowed to equilibrate for one-half to one hour.





At startup, a potential below the corrosion potential was set on the potentiostat, and a maximum level and stepping rate were set on the function generator. Finally, the cell was switched into the potentiostatic circuit, and the strip recorder turned on.

5.2.2 Polarization Studies in Simulated Physiological Solution

When it was found that the experimental methods could be used to generate reproducible results, work began on testing the stainless steel and cobalt alloy in simulated physiological solution.

5.2.2.1 Procedures

The solution was prepared by first diluting the serum, where it was used, followed by the dissolution of the magnesium-free salts, and the magnesium salts.

As it was found that aeration of solutions containing serum, by bubbling, produced large quantities of stable foam, another method was required. This new method involved producing a vortex with a magnetic stirrer such that a swirling mass of bubbles and solution was formed. Any foam produced by this method was immediately broken by the turbulence. This process lasted approximately twenty minutes prior to the introduction of the sample.

Concurrent with solution aeration, the samples were prepared. They were first wet abraded to a 600 grit tinish, then immersed in acetone while being transported, washed with a mild detergent, rinsed in tap water, and, finally, ultrasonically degreated, again in acetone. Before the samples were placed in the cell, they were rinsed in distilled water and dipped in test solution.

With the stirrer rotating at about 40 rpm, the system was allowed to equilibrate for one hour. After this period, the corrosion potential was measured, the starting potential set, and the cell switched into the potentiostatic circuit. Where applicable, the reverse scan was made at the same scan rate. The value of the current just before the next potential step was recorded.

5.3 Corrosion Fatigue

5.3.1 Apparatus

These tests used a system composed of three units: the load frame, the cell, and the solution conditioning and circulation unit. The experimental arrangement is illustrated in Plate 5.7.

5.3.1.1 The Load Frame

All fatigue tests were performed on the MTS 810 Load Frame. This is a versatile closed-loop hydraulic system with adjustable cycle frequencies and waveforms, and a maximum load capacity of twenty-five metric tons.



5.3.1.2 The Cell

The cell, constructed of acrylic, Plate 5.8, was designed to allow axial loading of a threaded tensile specimen in a liquid environment. The assembly consisted of a cylinder with endplates, one of which was removable. To accommodate grip extensions, an opening was made in each end with rubber 'O' rings to provide leak-free operation. Two ports in the side of the cylindrical section served as inlet and outlet for the circularing solution, while a third hole was made available for an optional reference electrode.

5.3.1.3 Solution Conditioning And Circulation

Constant test conditions were maintained at all times during a test. In a reservoir outside the cell the solution was aerated by the previously described 'vortex' method and the gas pressure above the solution was set to approximately 1 Pa by a water bubbler. Solution temperature was regulated by immersion in an oil bath.

A peristaltic pump circulated the solution to the cell at a rate of 1 mL per second; return flow was by gravity. The sample potential was not monitored.

5.3.2 Samples'

All the specimens were found to have some degree of bend; some more than others. In one case the testing machine grips could not accommodate the bend and the test could not be performed.



The specimens were notched with a low speed diamond wheel to a nominal depth of 0.17 of the diameter. In cases where the bend was noticeable, the notch-was made on the concave side. Plate 5.9 shows the typical location of the

notches and Plate 5.10 shows a cross section of a typical notch to have a tip radius of approximately 0.25 mm (note porosity).

.3.3 Test Solutions

Solutions consisted only of zero and five percent serum mixed with the other ingredients in the same manner as in the previous tests.

5.3.4 Procedures

The specimens were prepared by rinsing them in acetone to remove felt pen markings, followed by washing, rinsing, and drying.

The sample was then threaded into the grip extensions and the crevices at each end covered with Unichrome 'Quick Dry' Stop-Off Lacquer 323 (a fast drying vinyl lacquer produced by M&T Chemicals Inc. for use in the electroplating industry). After the lacquer had dried (one-half hour), the cell was sealed shut, and the solution pumped in.

In order to simple loading on the hip, fatigue testing involved the use of a sinusoidal load waveform at a frequency of 1 Hz. As the tests were aimed at examining the corrosion fatigue propagation rates and fracture



Plate 5.9 Fatigue Specimens showing the Typical Location of the Notch



Plate 5.10 Notch Geometry (50X)

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morphologies, and not the total fatigue lives, loads were increased well above the endurance limit at 10⁷ cycles. Selection of the loads wa d on the test stress and on the area of the ligament or after notching. Minimum

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and maximum stresses were it 48.3 and 448.2 MPa (7.0 and 65.0 ksi), for a load ratio of approximately 0.007. The number of cycles to failure was recorded by a counter panel on the 810 system.

After the samples had failed, the total number of load cycles was recorded and the fracture surfaces were examined by means of a stereo optical microscope and a scanning lectron microscope.

4[°] Conventi<u>ns</u>

Throughout this thesis all electrical potential differences will be referenced to the Saturated Calomel Electrode (SCE), which is related to the Standard Hydrogen Electrode (SHE) by the following relation:

 $E(SCE) = 0.244 V(SHE) - 0.00065 V/^{\circ}C$

Whenever a potential is calculated by the Nernst Equation the free energy value used will be the reported value at 25°C, but the prevature will be the test value, 37°C. Thus:

2.303 RT/F = 0.0615 V

It is considered that the error resulting from this procedure will be insignificant from a practical viewpoint.

6. RESULTS AND DISCUSSION

6.1 Technique Development

The results of the procedure standardization tests are

given in Figure 6.1. Figure 6.2 gives the reference plot taken from ASTM G5-78. Comparison between these curves indicates that the procedure can be used to obtain satisfactory and reproducible results.

316L.Stainless Steel and Cast Vitallium in Serum-Free and Glycose-free Solution

In order to obtain a perspective on the corrosion behavior of the cast Vitallium, several tests were also made a on 316L stainless steel in glucose and serum-free simulated physiological solution. The results of these potentiostatic polarization tests are presented and discussed below.

6.2.1 Stainless Steel

Three tests were performed on the 316L stainless steel to determine its corrosion behavior in a simulated physiological solution. Parameters of particular interest were the corrosion and passive current densities, and the corrosion, breakdown, and repassivation potentials: In each case the start potentials were within 25 mV of -0.50 V (SCE), and the potential was increased by 50 mV every five minutes.





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The forward anodic scans of all three tests (Figure 6.3) had essentially the same features; the reversible potentials all lay within a narrow range, the corrosion currents dere all similar, as were the positions of the cathodic and anodic curves. The average values of the polarization parameters are given in Table 6.1.

Polarization	Parameters for Solution	316L in Saline
	Current (A/cm²)	Potential (V)
Corrosion	1.6 x 10 ⁻⁷	-0.26
Passive	4.0 x 10 ⁻⁷	
Breakdown		: 0.15

-0.14

Repassivation

Table 6.1

In the case of scan 1, film breakdown had not occurred at the potential of reversal and the surface passivated to a greater degree than on the forward scan. In the cases of the other two scans breakdown occurred within the range of 0.1 to 0.2 V, whereupon the current increased toward an apparent transpassive region. Following reversal, the currents remained high forming a hysteresis; repassivation was achieved between -0.13 and -0.15 V.

After the latter two of these tests the area beneath the PTFE gasket was corroded. This localized attack was interpreted to be responsible for the hysteresis effect.



Serum-Free Simulated Physiological Solution

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In order to determine whether or not a higher breakdown potential, due to pitting, existed in the absence of a crevice a further test was performed with the sample only partially submerged; the gasket-sample interface remained

above the solution surface. Table 6.2 summarizes the parameters for this test.

Corrosion	Table 6 Parameters f Crevic	or	316L without	a	
		-	•		

	Current (A/cm ²)	Potential (V)
Corrosion	1.6 x 10 ⁻⁷	-0.28
Passive	4.4 x 10 ⁻⁷	
Breakdown		0.60

Figure 6.4 shows that the position of the cathodic curve, the reversible potential, and the passive region for this crevice-free situation to be similar to those of the previous results. However, a higher breakdown potential now exists at 0.5 to 0.6 V, 300 to 400 mV above the previous breakdown position.

According to the polarization curve and the current-time trace (not included), at potentials of approximately 0.1 V, current peaks were experienced, but³ when 0.2 V was reached the current trace was steady once again. As breakdown previously occurred in this range, the disturbance is considered to have been due to an unstable crevice-like situation, perhaps at the meniscus.



Figure 6.4 Polarization Curve for 316L in Semi-Submerged Position

Although the reverse scan was not extended back to the repassivation potential, enough has been completed to indicate the presence of the hysteresis.

6.2.2 Cast Vitallium

In order to evaluate the corrosion behavior of the Vitallium and to compare it with that of the stainless steel, several tests were conducted in identical serum-free and glucose-free solutions as the previous tests. As before the potential was increased in 50 mV steps every five minutes from the starting potentials of -0.40 and -0.60 V. No precautions were taken to avoid the formation of a crevice between the PTFE gasket and the sample.

Samples exhibited self-passivation in every case. The corrosion parameters corresponding to Figure 6.5 are listed below.

• · · · · · · · · · · · · · · · · · · ·	A A	•
: L	Current (A/cm ²)	Potential (V)
Corrosion	2.7 x 10-7	-0.38
Passive	4.0×10^{-7}	
Breakdown		0.38
Repassivation		0.35

Table 6.3Polarization Parameters for Vitallium in
Glucose-Free Saline Solution

The material displayed a very small hysteresis which increased slightly as the reverse potential was taken to





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more positive values, in agreement with the results of Wilde''. No crevice attack or pitting was observed on the sample at the end of any test.

It is also interesting to compare the current-time traces for the stainless steel and the Vitallium, Figure 6.6. In contrast to the instability shown by the stainless steel prior to breakdown the cobalt alloy exhibited a stable low current until the sharp rise occurred at the breakdown potential. This stability would suggest that in the short time it took to perform this test (less than six hours) passive film breakdown and repair did not occur.

6.2.3 316L vs. Vitallium

Although both materials exhibit self-passivation, and the corrosion and passive currents were approximately the same, the relative sizes and positions of the hystereses give a more significant measure of the relative long term corrosion behaviors. Based on the values of the repassivation potentials with respect to the breakdown potentials, it is, therefore, concluded that the cobalt alloy should be more resistant to localized attack than the stainless steel. Quantitatively, this resistance can be expressed in terms of the ratio of the differences between the breakdown and the repassivation potentials and between the breakdown and the reversible potentials. These results indicate that the likelihood of localized attack initiating spontaneously on Vitallium is much less than on 316L. These



316L Stainless Steel



Vitallium

Figure 6.6 Current-Time Traces for 316L and Vitallium

findings are in agreement with those of others including Hoar and Mears⁴⁵.

The relevant parameters of both materials are given below.

Table 6.4Anodic Polarization Parameters

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Paramete	er	5	316I Stainless		Cast Vitall	
E _{corr}			-0.26	v/	-0.38	v
E _b .			0.15	v	0.38	V
(E _b - E _{corr})	/(E _b - E _{rp}	,)	1.4		13.8	· .

The implication of these results is that, from a corrosion point of view, Vitallium is more suitable as an implant material owing to its higher resistance to localized attack, and to its broad passive potential range. Although the stainless steel shows some resistance to localized attack in the absence of crevices, this situation cannot be guaranteed, and even if it could, the current instability, noted in the crevice breakdown potential range, may indicate that the resistance would only have short term value.
6.3 The Effects of Organics on the Corrosion of Cast

Vitallium

The next series of experiments was conducted to evaluate the effects of combinations of glucose and serum on

the corrosion behavior of the cobalt alloy. These tests consisted of two sets: the first was designed to determine whether or not the organics had any major effects on the polarization curves. Tests were made using serum at concentrations of 0%, 0.01%, 0.1%, 5%, and 15%, and at a stepping rate of 50 mV every five minutes in the manner described in the previous chapter. During all of these tests solution samples were drawn at intervals for pH determination.

One test was also performed to monitor the change in corrosion potential with time. A Vitallium sample was immersed in oxygenated saline solution containing glucose but no serum, and the potential was recorded approximately every five minutes over twenty-four hours using the Hewlett-Packard 9825A desktop computer.

When it was found that the organics only affected the position of the cathodic curve and the value of the corrosion potential, the second set of tests was conducted

to examine these changes in more detail. Cathodic potentiostatic polarization tests were made with the following glucose-serum combinations:

No	Glucose	ľ	No Serum	NG/NS
	Glucose	/	No Serum	G/NS
	Glucose	/	Serum	G/S
No	Glucose		Serum	NG/S

In contrast to the first set of tests the potential stepping rate was decreased, in most cases, from 50 mV to 10 mV every five minutes.

In the following section, the results of each set of tests will be presented and discussed. At the end, the overall effects of the organics on the potentiostatic polarization behavior of the alloy will be discussed.

6.3.1 Preliminary Tests on the Effects of Organics

6.3.1.1 The Effects of Glucose in the Absence of Serum

The results of the preliminary polarization tests in solutions containing glucose but no serum are given in Figure 6.7 and Table 6.5. 93

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Table 6.5 Corrosion Parameters in Solution Contains Glucose-but-no-Serum				
	Current (A/cm²)	Potential (V)		
Corrosion	2.7 x 10 ⁻⁷ .	-0.35		
Passive	4 x 10 ⁻⁷			
Breakdown		0.38		
Repassivation		0.35		

Comparing the results for a solution without glucose (Figure 6.5) and a solution with glucose (Figure 6.7) it is evident that they both have essentially the same

characteristics: the values of the corrosion and breakdown potentials, as well as the corrosion and passive currents are not significantly different, and the hystereses are identical. These similarities indicate that, in the absence of serum, glucose does not hinder the formation or short term stability of the passive surface film.

The Effects of Serum in the Presence of Glucose Figure 6.7 gives the results of the preliminary tests using glucose and serum. Table 6.6 summarizes the corrosion parameters.

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Figure 6.7 Preliminary Polarization Curves for Vitallium in Solution Containing Glucose but no Serum



Solution Containing Glucose and Serum

Preliminary Polarization Curves for Vitallium-in-Solution-Containing-Glucose-and				
	Current (A/cm²)	Potential (A)		
Corrosion	3 x 10 ⁻⁷	Varies		
Pássive	4.5 x 10 ⁻⁷			
Breakdown	`	0.38		
Repassivation	· · · · · · · · · · · · · · · · · · ·	0.35		

Table 6.6

Considering the scans given in Figures 6.7 and 6.8, (G/NS and G/S), it is evident that they all have the same form of anodic polarization curves. The only differences between the scans are in the value of the corrosion potential and, possibly, the position of the cathodic curve. Although only one test was performed at each serum concentration, it appears that as the serum content is increased the corrosion potential is shifted to more negative values. However, while the change in corrosion potential was on the order of 100 mV, the almost vertical slope of the passive region resulted in a negligible decrease in the corrosion current.

6.3.1.2 Solution pH

During these preliminary tests samples of solution were intermittently drawn for pH determination. The results, given below, show no significant differences or trends,

neither from one test to another, nor among the serum-free

and serum-containing tests. The overall average pH was

<u>7.7±0.1</u>.

Table 6.7 pH Readings			·			
	erum ontent	Potential (V)	рН	Serum Content	Potential (V)	рН
	0%	-0.40 0.05 0.45	7.8 7.6 7.6	0.01%	-0.60 0.20	7.8
X	•	0.80 0.85 0.15 -0.40	7.6 7.6 7.6 7.6	0.1%	0.60 0.15 0.65 0.15	7.8 7.8 7.7 7.7
	0 %	-0.60	7.7 7.6	•	-0.45	7.7 7.7
		0.40 -0.25 -0.60	7.6 7.6 7.6	5%	-0.60 0.10 0.55 0.00	7.7 7.7 7.7 7.6
	0%	-0.60 -0.20 0.50 -0.60	7.6 7.6 7.7 7.7	15%	-0.35 0.20 0.65	7.7 7.7 7.8

Long Term Corrosion Potential

The variation of corrosion potential with time is shown in Figure 6.9. As indicated, the potential increased over the first eighteen hours from approximately -0.41 V, to a maximum value of -0.15 V, then, after eighteen hours fell to approximately -0.25 V. Both these peak and minimum values are consistent with the potentials measured by Brettle and Hughes⁴⁴ within the first ten days of their ential-time tests. Furthermore, as no localized attack noted at the



end of the test, it would seem that these results corroborate the speculation by Hoar and Mears'⁵ that -spontaneous-film-breakdown-is-unlikely-for-this-materialunder these conditions, at least in the short term.

The results of the foregoing tests have indicated that neither the glucose, in the absence of serum, nor the serum, in the presence of glucose, affect the anodic polarization. behavior. It is also apparent that the serum, in the presence of glucose, may decrease the corrosion potential. thus making it a possible cathodic corrosion inhibitor.

The following series of tests will concentrate on the effects of glucose and serum on the cathodic polarization behavior of the alloy.

6.3.2 The Effects of Glucose and Serum on the Cathodic

6.3.2.1 No Glucose and No Serum - NG/NS

The first set of experiments in this series was conducted with no glucose and no serum and gave the results shown in Figure 6.10. As shown the curves have three relative positions, which only in part, reflect variations in procedure. The curves at the left and centre were obtained by the procedure described in the earlier chapter, the only major difference being that the central curve equilibrated for two and one-half hours before starting, whereas those at the left equilibrated for only one hour.



This difference is considered to explain the higher corrosion potential in the case of the central curve. The procedure leading to the curve at the right varied significantly from the others. Rather than equilibrating at the rest potential, the sample was cathodically charged at -1 V for at least one hour prior to commencing the scan. The higher currents at all potentials, therefore, are a result of the reduced activation energy for the cathodic reaction on a surface whose anodic film has been either partially or completely removed by cathodic charging. Why the corrosion potential of the clean electrode (the curve at the right). should be higher than the filmed electrode (curves at the left) is not known for certain, but may be a result of either a longer equilibration time, or the longer time required to reach a given potential as a result of the lower (start potential.

If the film reduction mechanism is also active, at least intermittently, in the region -0.65 to -0.70 V, it might also explain the higher currents in the case of the central curve.

In addition to the effects of variations in procedure these curves exhibit several other interesting features: the central curve exhibits a 'hook' or positive slope near the starting end of the curve, and the curve at the right appears to be formed by the addition of two cathodic curves. The latter of these observations is discussed now, while the former is dealt with later.

Cathodic Reactions

The curve at the right, in Figure 6.10, consists of two distinct regions, which are interpreted to indicate the presence of two separate cathodic reactions. The first, which dominates from the corrosion potential to the limiting current at approximately -0.7 V, is likely due to the oxygen reduction reaction which can be written as:

$$O_2 + 2H_2O + 4e^- \rightarrow 4OH^-$$

and which has, under these conditions, a redox potential given by:

 $E = E^{\circ} + 0.861 - 0.062 \text{ pH} + 0.015 \log pO_2$ = 0.77 V (SHE), or $E^{\circ} = 0.53 \text{ V (SCE)}$

At potentials below -0.70 V the curve deviates from the limiting current toward higher current values. As the test solutions were neutral, the predominant reaction is expected to be the reduction of water according to the following:

 $2H_2O + 2e^- \rightarrow H_2 + 2OH^-$

which has a redox potential of:

E = -0.062 pH= -0.48 V (SHE), or E = -0.71 V (SCE)

As this potential is within the vicinity of the point of deviation of the second slope, and because the overpotential of hydrogen evolution is small, this water reduction reaction is most likely responsible for the curve form below

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-0.7 V.

6.3.2.2 Glucose With and Without Serum - G/S and G/NS

Figures 6.11 to 6.15 give the results of the potentiostatic cathodic polarization tests using no serum, and serum at various concentrations in solutions containing glucose. Several features are immediately apparent: the curves for 0%, 0.01%, and 0.1% serum are relatively closely grouped, while those for 5% and 15% have wider variations, and the curves obtained at 0.01% serum consist of two groups, those with and without positive slopes on the cathodic curves. These results will now be considered in more detail.



Figure 6.11 Cathodic Polarization of Vitallium in Serum-Free Solution





Figure 6.13 Polarization Behavior of Vitallium in Solution Containing 0.1% Serum



Figure 6.14 Cathodic Polarization Behavior of Vitallium in Solution Containing 5% Serum



Figure 6.15 Polarization Behavior of Vitallium in Solution Containing 15% Serum

Effect of Serum on the Corrosion Potential

As was suggested in the preliminary results, the corrosion potential was found to fall with increasing serum contents, Figure 6.16 (the left-most error bar refers to zero serum conditions). As can be seen, the corrosion potential falls perceptibly after 0.01% serum, and may be continuing to fall, albeit at a lower rate, after 15%. Although the mechanism behind the potential change is unknown, it would appear that these results could be grouped with those of Solar et al.^{4,3}, who found that certain organics shifted the corrosion potential of Ti-6Al-4V in the negative (active) direction. Furthermore, these results corroborate the findings of Brown and Merritt^{4,4} who found that the corrosion potential of 316L, a material also protected by a chromium oxide film, was shifted in the same direction by a physiological solution.

The difference in the corrosion potentials between the serum-free and 15% serum situations also resembles the difference reported by Hoar and Mears⁴⁵ between in vitro tests in saline and in vivo tests using human subjects. Although the actual potential values depend on the procedures used in each case, the similarity in the differences suggest that the conditions used in these tests simulate the in vivo conditions more accurately than inorganic saline solutions.

In terms of the mechanism involved in shifting the corrosion potential, the results are interpreted to indicate



that the serum hinders the diffusion of oxygen to the metal surface.

Effect of Serum on the Corrosion Current

Because the corrosion potentials were remote from the reversible potentials for the anodic and cathodic reactions, no Tafel lines existed in the vicinity, and, therefore, another method was used to estimate the corrosion currents. The preliminary tests showed that the slope of the passive curve was reproducible and independent of the serum content. In tests, therefore, where the scans had been taken into the anodic range (0.01%, 0.1%, and 15% serum), the best fit line describing the passive ranges of the preliminary data was applied to the anodic current values, and the intersection of these lines with the corrosion potentials were taken to be the corrosion currents. Where the scans had been terminated at the corrosion potentials, the corrosioncurrents were estimated on the basis of the positions of the cathodic curves with respect to the former data. In all cases the estimates of the corrosion currents were essentially equal at 1 x 10^{-7} A/cm².

Positive Cathodic Slopes

During some of the previous tests and these latter tests, regions of positive slope were encountered on the cathodic curves. These include one curve obtained with glucose and serum-free solution, one with serum-free solution, one partially completed curve at 0.001% serum,

Figure 6.17, four with 0.01% serum, and one at 5% serum





where the curve seemed to alternate between two courses. No such positive slopes were observed at 15% serum.

Common to all these curves is the location of the region of positive slope, beginning near the start potential, -0.65 or -0.70 V, and increasing to maximum current values between -0.63 and -0.67 V. The maximum increase in current is only 22%.

Regarding corrosion potentials, values were either higher or the same as in the curves without positive slopes. In the case of the 0.01% serum results, those curves with positive slopes had an average corrosion potential 40 mV higher than the other curves in the same group, and at 5% serum the difference was over 100 mV. In addition, the curve obtained with glucose and serum-free solution was up to 125 mV higher. Perhaps the two most significant observations are that serum and glucose are not necessary for a positive slope to occur, and no positive slope was encountered at 15% serum.

In general several situations can result in positive cathodic slopes: a cathodic curve which closely approaches an anodic critical current peak, a simultaneous oxidation reaction which exhausts its reactant supply, or a second cathodic reaction whose rate increases with time.

The first of these possibilities is not responsible as it has been reported by Lucas et al.⁵⁴ that Vitallium exhibits self-passivation in deoxygenated conditions. Similarly, the simultaneous anodic reaction mechanism is considered unlikely as no such reaction is known to be active under glucose-free and serum-free conditions.

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The third situation, however, may be closer to the actual mechanism. Since it is thermodynamically impossible for the kinetics of a reaction to increase as the driving force is reduced, the only explanation for such an observation is that time is the controlling factor. It is possible, therefore, that the increasing current is a result of a partial reduction of the anodic film (a mechanism discussed in the glucose-free and serum-free results, above) whose rate increases with time until the overpotential becomes too low, at approximately -0.65 V, for the reaction to proceed. In addition, the decrease in the frequency of positive slopes as the serum content is increased may indicate that the serum has a stabilizing effect on the film.

6.3.2.3 Glucose-Free Solution Containing Serum - NG/S Two tests were conducted to determine the polarization behavior of the alloy in glucose-free solutions containing 15% serum.

As Figure 6.18 shows, the two curves were not entirely consistent either with one another or with the previous results. One curve reached its corrosion potential at -0.63 V, while the other reached it at -0.69 V, but without the asymtotic decrease in current normally observed as the corrosion potential is approached. The form and corrosion



potential of the latter of the above mentioned curves would suggest that it was obtained under deaerated test conditions; however this is not considered to have been the case. The following discussion will include a possible explanation for these findings.

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6.3.3 The Effects of Organic Additions on the Polarization Behavior of Vitallium

As the results show, the most significant effect that the glucose and serum have on the corrosion behavior of Vitallium is to alter the value of the corrosion potential. For this reason the following table summarizes the results of the four glucose/serum combinations in terms of the corrosion potentials:

Table 6.8 Corrosion Potentials at the Four Glucose-Serum Combinations

	No Glucose	Glucose
No Serum	-0.32	-0.35
Serum	-0.63	-0.46

According to this summary glucose has little, if any, significant effect on the corrosion potential when serum is absent. However, when serum is present, the corrosion potential is pushed more negative values, until when glucose is absent the corrosion potentials approach the redox potential for hydrogen reduction. The mechanisms responsible for these potential shifts almost certainly involve the relative adsorption of the glucose and other serum components, and appear to be dependent on the relative concentrations of the various components.

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It is believed that, while the glucose competes with the serum for adsorption, its 'adsorbability' is weaker than that of the serum. The fact that the corrosion potential is higher when glucose is present with serum indicates that the glucose is competing for adsorption, but does not hinder the diffusion of oxygen/to the cathodic sites. However, when only serum is added the lower glucose content (0.15 g maximum) is not sufficient to produce a significant effect and, as a result, other serum components adsorb and hinder, if not completely block the diffusion of oxygen.

6.4 Corrosion Fatigue

The results of the corrosion fatigue tests in

serum-free and 5% serum are summarized in Table 6.9.

Table 6.9 Corrosion Fatigue Lives

Teșt	0% Serum Cyclés	5% Serum Cycles
1	44000	25892
2	29613	17410
3	. 36660	28399
4	30710	177000†
5	38852	
6	26621	

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† - Rejected on a statistical basis; refer

to Appendix C.

According to ASTM committee E-9 fatigue life data can be considered to be log-normally distributed; that is, the data, itself, is significantly skewed, while the natural logarithm of the data is normally distributed. The above data was analysed on the basis of this assumption (Appendix C).

Table 6.10 summarizes the means and standard deviations of the two data sets in terms of the the log-normal distribution, as well as the 80% confidence limits and other statistical parameters for the two data sets in terms of the arithmetic values.

Table 6.10 Results of Statistical Analyses of Corrosion Fatigue Life Data

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N	<u>0% Serum</u>	<u>5% Serum</u>			
Mean (log-normal)	10.517	10.060			
Standard Deviation (log-normal)	Q.286	0.260			
At 80% Confidence	Interval				
Lower Limit	10.361 (31603 cycles)	9.777 • (17622 cyćles)			
Upper Limit	10.673 (43160 cycles)	10.343 (31042 cycles)			
Arithmetic Statistical Values					
Mean	38432 cycles	29194 cycles			
Median	36897 cycles	23388 cycles			
Mode	34009 cycles	21857 cycles			

According to the statistical analysis, there is a high probability that the two data sets are from different populations' (based on an 80% confidence interval). The fatigue life of Vitallium, therefore, is likely to be less when tested in a 5% serum solution, as opposed to when tested in serum-free solution.

While the standard deviations for the two data sets are very similar, which may reflect similar basic fracture mechanisms, no solid fatigue life data indicates a possible cause for the detrimental effect of the serum.

6.4.1 Fractographic Results

Several of the fatigued samples were examined in the scanning electron microscope to determine if the fracture morphologies changed with the introduction of serum. Photomicrograph composites of the fracture surfaces of one specimen from each group are enclosed in an envelope at the back. As these composites illustrate, the fracture surfaces typically consist of two regions: a flat transgramular region adjacent to the notch, interpreted to be the extent of corrosion fatigue crack propagation, and a second mixed

trans- and intergranular, likely final, fracture over the remaining ligaments. Both these regions were examined at higher magnifications and are dealt with as follows.

6.4.1.1 Flat Fracture

As the composites illustrate the flat fracture regions typically consisted of cleavage steps, and porosity and/or

incoherent inclusions. Plate 6.1 from the serum-free group shows the surface adjacent the notch to be generally flat with cleavage steps beginning at the notch. This micrograph

shows both porosity, labelled 'P', and holes from where inclusions had been pulled out, labelled 'I'.

Plate 6.2 illustrates the change in pattern of the cleavage steps which resulted when the crack propagated across a grain boundary.

Much of the flat fractures consisted of the type of cleavage steps shown in Plate 6.3. In all cases, either subsidiary cleavage cracks or dissolved slip planes'⁴ intersected the surface, while inclusions pock-marked the surface. A qualitative X-ray analysis of one such precipitate revealed it to be high in niobium.



Plate 6.1 Fracture Surface Adjacent to the Notch on a Specimen Tested in Serum-Free Solution (1000X)







Plate 6.2 Cleavage in Two Adjacent Grains (1200X)



Plate 6.3 Typical Cleavage Fracture near the Final Fracture Zone (900X)

6.4.1.2 Mixed-Mode Fracture

This second fracture zone was typically brittle, showing very little plastic deformation. The fact that the path of least resistance included interdendritic and grain boundary porosity indicates that the fracture involved the expenditure of less energy. This would suggest, therefore, that the material would have a greater fatigue life if such defects were not present.

6.4.2 The Effects of Organics on the Corrosion Fatigue Behavior of Vitallium

At a confidence interval of 80% the corrosion fatigue life of the Vitallium is significantly less in saline solution containing serum, than in serum-free saline. However, the fracture modes encountered in both sets of tests were identical. While the results indicate no cause for the reduction in fatigue life with the introduction of serum, the mechanism of propagation does not change. Therefore, the serum must alter, by degree only, some rate controlling step or property within the material.

According to Tyson and Alfred's, "Propagation of a truly brittle cleavage crack through a crystal is possible only if the stress required to rupture atomic bonds at the crack tip is reached before the shear strength of the lattice is exceeded." Therefore, if adsorption of some serum component reduces the cohesion between strained crystal . planes at the tip, the fatigue life would be reduced. Another possibility is that the serum affects the dissolution rate at the tip. If, as Woodson et al.⁵ ' suggest (that organics can lead to an autocatalytic corrosion

mechanism), an increased dissolution rate may be responsible for the reduction in life.

7. CONCLUSIONS

The effects of serum and glucose additions on the corrosion and corrosion fatigue behavior of a cast CoCrMo alloy have

been investigated with the following conclusions:

7.1 Potentiostatic Polarization Tests

- Both the 316L stainless steel and the cobalt alloy, machined from the prostheses, exhibited self-passivation with similar values of corrosion and passive currents in the oxygenated, neutral saline solution.
- 316L stainless steel is highly susceptible to spontaneous passive film breakdown and crevice corrosion in these chloride solutions.
- The cobalt alloy tested here is highly resistant to passive film breakdown.
- Glucose and other serum components compete for adsorption on the cobalt alloy in simulated physiological solutions.
- 5. Neither glucose nor serum have any effect on the anodic polarization behavior of the Vitallium.

While glucose does not appear to alter the corrosion potential, serum tends to shift it in the negative (active) direction. It is possible that this shift is a result of lower oxygen reduction rates caused by low diffusion rates through an adsorbed serum film. By this argument, it would appear that the glucose does not hinder oxygen diffusion.

7.2 Corrosion Fatigue Tests

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- Fatigue cracks in the cobalt alloy, from which the fatigue samples were cast, propagate by a cleavage mechanism.
- According to the fatigue life data and a confidence interval of 80%, the corrosion fatigue life of cast Vitallium is less when tested in 5% serum than when tested in serum-free solution. However, the means by which this reduction occurs, without changing the basic fracture mechanism, is not known.
- 9. Casting porosity is believed to have been, at least, partially responsible for reducing the fatigue lives of all samples.
8. RECOMMENDATIONS

Tests have been conducted to evaluate the effects of serum and glucose on the corrosion of a cast CoCrMo alloy in

solutions approximating the physiological conditions of healthy tissue. Results have indicated that while glucose and serum compete for adsorption on the metal surface, only the serum has any significant effect on the corrosion process. It has, furthermore, been demonstrated that the addition of serum to inorganic simulated physiological solution results in test conditions which approximate more accurately the in vivo conditions than do many of the traditional inorganic test media.

Several important questions concerning the effects of organics on the corrosion and corrosion fatigue, however, have not been addressed. The following questions are suggested as topics of further research:

- Are any physiological organics involved in electrochemical reactions?
- What specific organics are responsible for the effects noted above, and is the adsorption potential dependent?
- What is the effect of physiological organics in healing tissue where the conditions are significantly different from those used in this study?
- What is the effect of the organics in crevice situations?

Potentiostatic polarization is an accelerated corrosion testing technique the results of which only indirectly forecast the long term behavior of a system. It is

suggested, therefore, that long term tests be conducted to examine the effects of physiological organics. In addition to studying the variation of the corrosion potential and current with time, the following questions might also be addressed:

> What is the nature of the adsorbed and anodic films, and how do they change with time? Are specific constituents of the alloy selectively leached and, if so, what is the role of the organics?

Regarding the corrosion fatigue aspect, this research indicated there to be a high possibility that serum reduces the corrosion fatigue life of Vitallium. However, the means by which the reduction is produced is not known. It is, therefore, suggested that more extensive studies be conducted to investigate the effects of serum on the initiation and propagation phases, with the aim of determining the mechanism by which the serum shifts the corrosion potential and reduces the corrosion fatigue lives.

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141 The following solutions have been used as simulated physiological saline solutions: Table A.1

 Compositio	Table A. ons of Various Solution	Inorganic Test	
Substance	Hank's [,]	Ringer's''	
	(g/l)	(g/l)	
NaCl	8.0	6.3	
CaCl ₂ ,	0.14		
KCl	0.40	0.37	
KH ₂ PO ₄	0,06	en en <u>de ser</u> de services de la service. Na esta de la service de la	
Na/HCO₃	0.35	2.4	
Na 2HPO 4	0.05	0.07	
NaH ₂ PO ₄		0.06	2
MgCl ₂	0.05	0.10	
MgSO ₄	0.03	0.06	
Sodium Lactate		3.10 (Optional)	

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APPENDIX B

Alberta Research Council

Chemical Analysis Results

REPORT OF ANALYSIS

Laboratory Sample Number: 1662-83G to 1664-83G

DESCRIPTION

Three samples of metal submitted September 1, 1983 by the University of Alberta, Mineral Engineering, 89th Avenue and 114th Street, Edmonton, Alberta, Attention: Mr. Bob Sutherby with request for the following analyses.

RE: PO# 18553NC

ANALYSIS

Laboratory Sample Number: Reference	<u>1662-836</u> Implant	<u>1663-83G</u> Fatigue	1664-83G Stainless
Elements detected by Induction Coupled Plasma, wt% (SL-ARC)			
Cobalt (Co)	58.6	51.5	0.31
Chromium (Cr)	27.1	24.0	18.0
Copper (Cu)	0.05	0.05	0.21
lron (Fe)	0.18	0.17	67.2
Manganese (Mn)	2.3	2.2	4.7
Molybdenum (Mo)	6.05	5.35	2.55
Nickel (Ni)	2.10	0.16	14.0
Phosphorus (P)	0.70	0.61	0.40
Sulfur (S)	0.16	0.15	0.15
Silicon (Si)	0.82	0.99	1.46

DATE September 20, 1983 Gasoline & Oil Laboratory Alberta Research Council APPENDIX C

Statistical Analyses of Corrosion Fatigue Life Data

Γį.

The purposes of these analyses were as follows:

Β.

С.

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- A. To determine the acceptability of the high data point in the 5% serum results,
 - To determine if the two data sets are from different populations; that is, to determine if the serum had any statistical effect on the corrosion fatigue life, and To determine the upper and lower limits of the data populations, and to determine the statistical parameters of the arithmetic data values.

The statistical analyses wede made based on the following two assumptions:

- 1. That the corrosion fatigue life data is log-normally distributed, and
- 2. While the means of the data may be shifted, the variance of the 5% serum data is equal to that of the 0% serum data. This assumption would be justified if the basic cracking mechanisms were the same in both cases.

		Table C.1		•••
Fatigue L	ife Data	in Natural	Logarithm	Form

<u>0% Serum</u>	<u>5%</u> Serum	• • •
10.692	10.162	
10.296	9.765	
10.509	10.254	
10.332	12.084	
10.568		
10.189		•
11.032		

If $E(\ln x) = \lambda$, and $(VAR(\ln x))^{1/2} = \zeta$

then $\lambda_0 = 10.517$ and $\zeta_0 = 0.286$ and $\lambda_5 = 10.566$ and $\zeta_5 = \zeta_0$

A. High Data Value in 5% Serum Results

Based on the assumptions listed above, what is the upper limit for the 99.8% confidence interval based on a t-distribution with 3 degrees of freedom?

```
1 - a = 0.998

\therefore 1 - (a/2) = 0.999 \Rightarrow t = 10.2

\therefore \text{ Upper limit} = \lambda_5 + t[\zeta_5/(n^{1/2})]

= 10.566 + 10.2 (0.286/(4^{1/2}))

= 12.025
```

Since ln(177000) = 12.084 > 12.025, then the data point can be rejected at a confidence interval of 80%.

Based on the rejection of this data point, the statistical parameters of the remaining data become as follows:

 $\lambda_5 = 10.060; \zeta_5 = 0.260$

Although assumption 2, above, indicates that the standard deviation of the two sets shall be considered to be the same ($\lambda = 0.286$), it is interesting to note how closely-the actual standard deviation values compare. This similarity may indicate that the cracking mechanisms active in both test sets are, in fact, fundamentally the same.

B. Statistical Effect of the Serum

To determine if the serum had any statistical effect, a t-test is performed to test the following hypothesis:

If $\lambda_0 \neq \lambda_5$, then $T \geq t$.

With 2 degrees of freedom, and a confidence interval of 80%:

1 - a = 0.80 $\therefore 1^{0} - (a/2) = 0.90 \Rightarrow t = 1.886$ $T = (\lambda_{o} - \lambda_{5})/(\zeta/(n^{1/2}))$ $= (10.517 - 10.060)/(0.286/(3^{1/2}))$ = 2.768

Since T is, in fact, greater than t, the hypothesis is valid at the 80% confidence interval.

Based on the confidence intervals used in the above analyses, the total confidence interval is:

 $(0.80)(0.998) \cong 0.80 \text{ or } 80\%$

```
0% Serum
```

G

degrees of freedom = 6

```
\lambda_{o} = 10.517

\zeta_{o} = 0.286

t = 1.440
```

lower limit = 10.361 (31603 cycles)
upper limit = 10.673 (43160 cycles)

5% Serum

Based on the equations given by Hald', the statistical parameters of the original data (assuming the log-normal distribution) are:

```
0% Serum
```

median $x_o = anti-ln (\lambda_o c)$

= 36897 cycles mode x_0 = anti-ln (λ_0 - 2.030 ζ_0^2)

= 34009 cycles

mean x_o = anti-ln $(\lambda_o + 1.1513\zeta_o^2)$ = 38432 cycles <u>5% Serum</u> Similarly:

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median $x_5 = 23388$ cycles mode $x_5 = 21857$ cycles mean $x_5 = 24194$ cycles