

UNCARIA GAMBIR AS NATURAL CORROSION INHIBITOR FOR MILD STEEL IN ACIDIC SOLUTION

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UNCARIA GAMBIR AS NATURAL CORROSION INHIBITOR FOR MILD STEEL IN ACIDIC SOLUTION

by

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LIST OF ABBREVIATIONS

FTIR	Fourier Transfrom Infrared Spectroscopy
HPLC	High Performance Liquid Chromatography
EIS	Electrochemical Impedance Spectroscopy
SEM	Scanning Electron Microscopy
EDX	Energy Dispersive X-Ray Spectroscopy
XRF	X-Ray Fluorescence Spectroscopy
PZC	Potential Zero Charge
EAG	ethyl acetate gambir extract
СН	(+)-catechin hydrate
HCl	hydrochloric acid
GAE	gallic acid equivalent
CE	catechin equivalent
IE	inhibition efficiency
CR	corrosion rate
CPE	constant phase element
Ea	activation energy
ΔH	enthalpy
ΔS	entropy
ΔG_{ads}	Gibbs free energy of adsorption
ppm	parts per million
g	gram xii

mg	milligram
mL	millilitre
cm	centimetre
mm	millimetre
min	minutes
wt	weight
w/v	weight per volume
v/v	volume per volume
mV	millivolt
mV s ⁻¹	millivolt per second
Hz	Hertz
kHz	kiloHertz
kHz $\Omega \text{ cm}^2$	kiloHertz Ohm's centimetre square
$\Omega \ { m cm}^2$	Ohm's centimetre square
$\Omega \text{ cm}^2$ mA cm ⁻²	Ohm's centimetre square milliampere per centimetre square
Ω cm2 mA cm ⁻² mpy	Ohm's centimetre square milliampere per centimetre square mils of penetration per year
Ω cm2 mA cm ⁻² mpy mm y ⁻¹	Ohm's centimetre square milliampere per centimetre square mils of penetration per year millimeters per year
$\Omega \text{ cm}^2$ mA cm ⁻² mpy mm y ⁻¹ E _{corr}	Ohm's centimetre square milliampere per centimetre square mils of penetration per year millimeters per year corrosion potential
$\Omega \text{ cm}^2$ mA cm ⁻² mpy mm y ⁻¹ E _{corr} E _r	Ohm's centimetre square milliampere per centimetre square mils of penetration per year millimeters per year corrosion potential Antropov 'rational' corrosion potential
Ω cm ² mA cm ⁻² mpy mm y ⁻¹ E _{corr} E _r I _{corr}	Ohm's centimetre square milliampere per centimetre square mils of penetration per year millimeters per year corrosion potential Antropov 'rational' corrosion potential corrosion current density

UNCARIA GAMBIR AS NATURAL CORROSION INHIBITOR FOR MILD STEEL IN ACIDIC SOLUTION

ABSTRACT

Uncaria gambir, a native Southeast Asia herbal plant has been characterized and studied as a mild steel corrosion inhibitor in acidic media. It was revealed that ethyl acetate gambir extract gave the highest condensed tannin, phenol and flavonoid content compared with other solvent extracts. Quantification studies by means of HPLC have shown that more than 80 % (wt) of gambir extract consists mainly of catechin. The effect of both ethyl acetate gambir extract and (+)-catechin hydrate as corrosion inhibitors for mild steel in 1 M HCl solution was done using various techniques such as weight loss, potentiodynamic polarization and electrochemical impedance spectroscopy (EIS) measurements. It was revealed that inhibition was concentration dependent as inhibition efficiency (IE) increases with the increase in the inhibitor concentration. Potentiodynamic polarization measurement indicated that the inhibitors act as "mixed-type" inhibitors with predominant anodic inhibition. EIS measurement indicated that the corrosion of mild steel in the absence and presence of inhibitor was mainly controlled by charge transfer process. The Randles-CPE model has been used as the equivalent circuit throughout this study. The optimum concentration obtained from all tests was at 1000 ppm (0.1 % wt). Temperature studies revealed that the corrosion rate of mild steel in the absence and presence of both inhibitors exponentially with temperature. Adsorption increases isotherm determination revealed that the inhibitor follows the Langmuir adsorption model. Surface analysis via SEM portrayed that there was a significant morphological improvement of mild steel surface in the presence of the inhibitor. The calculated value of free energy of adsorption, ΔG_{ads} and potential zero charge (PZC) for the inhibitors indicate that the inhibition mechanism process was spontaneous and the inhibitors were physically adsorbed (physiosorption) onto the mild steel surface.

UNCARIA GAMBIR SEBAGAI PERENCAT KAKISAN SEMULA JADI BAGI KELULI LEMBUT DI DALAM LARUTAN BERASID

ABSTRAK

Uncaria gambir, tumbuhan herba asli dari Asia Tenggara telah dicirikan dan dikaji sebagai perencat kakisan keluli lembut dalam medium berasid. Kajian menunjukkan bahawa ekstrak etil asetat memberikan kandungan tanin terkondensasi, fenol dan flavonoid tertinggi berbanding ekstrak dari bahan pelarut yang lain. Kajian secara kuantitatif menggunakan HPLC telah menunjukkan bahawa lebih 80 % (berat) dari ekstrak gambir mengandungi katekin. Kesan ekstrak gambir daripada pelarut etil asetat dan (+)-katekin hidrat sebagai perencat kakisan bagi keluli lembut di dalam larutan 1 M HCl telah dijalankan menggunakan pelbagai teknik seperti penentuan kehilangan berat, pengukuran kekutuban potensiodinamik dan pengukuran spektroskopi elektrokimia impedans (EIS). Kajian menunjukkan bahawa perencatan tersebut adalah bergantung kepada kepekatan dimana keberkesanan perencatan (IE) meningkat dengan peningkatan kepekatan perencat. Pengukuran kekutuban potensiodinamik menunjukkan bahawa perencat tersebut bertindak sebagai perencat jenis campuran dengan kesan perencatan anodik yang paling dominan. Pengukuran EIS telah menunjukkan bahawa pengaratan keluli lembut dalam ketidak hadiran atau kehadiran perencat adalah secara amnya dikawal oleh proses pertukaran cas. Model Randles-CPE telah digunakan sebagai litar yang bersesuaian sepanjang kajian ini. Kepekatan optimum yang diperoleh daripada kesemua ujian adalah pada 1000 ppm (0.1 % berat). Kajian suhu menunjukkan bahawa kadar kakisan keluli lembut dalam ketidak hadiran dan kehadiran kedua-dua perencat meningkat secara eksponen dengan suhu. Penentuan isoterma penjerapan menunjukkan bahawa perencat tersebut mengikuti model penjerapan Langmuir. Analisis permukaan melalui SEM telah menggambarkan bahawa terdapat pemulihan morfologi yang signifikan pada permukaan keluli lembut dalam kehadiran perencat. Nilai tenaga bebas penjerapan, ΔG_{ads} dan keupayaan cas sifar (PZC) yang dikira bagi perencat tersebut menunjukkan bahawa proses mekanisma perencatan yang berlaku adalah secara spontan dan terjerap secara fizikal (fizik-jerap) pada permukaan keluli lembut.

CHAPTER 1

INTRODUCTION

1.1 Introduction to Corrosion

Since the human existence, people have witness natural disaster around the world. One of the natural occurring disasters that still happen nowadays is corrosion. Corrosion or mostly known as rust were derived from Latin words '*rodere*' which means gnawing and '*corrodere*' which means gnawing into pieces (Sastri *et al.*, 2007a; Davis, 2000). Even though there are many terms that explain this phenomenon, but the fact is corrosion will cause physical damage, destroys the lustre and beauty of an object and thus shortens their life. An essay wrote by a Roman philosopher Pliny (AD 23-79) entitled "*Ferrum Corrumpitar*" was one of the earliest manuscript recorded that tells about the physical damage of an iron in a period of time.

Previously, there were only few scientist who show some interest to study the corrosion phenomenon until Robert Boyle describe about the corrosion mechanism in his essay "*Mechanical Origin of Corrosiveness*" (Zaki Ahmad, 2006a; Sastri *et al.*, 2007a). Early in 19th century, the corrosion concept was explained in more detail from the electrochemistry perspective. In the year 1833, Michael Faraday established a quantitative relationship between chemical action and electric current. This theory has been used to explain the mechanism that happened in corrosion process.

Faraday's first and second laws are the basis for calculation of corrosion rates in metal (Walsh, 1991).

1.1.2 Corrosion in definitions

Corrosion process occurs in nature like any other natural disasters. The term corrosion can leads us into many definitions. Generally, corrosion will form a solid wastage on the metal surface when metal is being exposed in reactive environment. According to Fontana (1986), corrosion is the deterioration of materials as a result of reaction with its environment. Corrosion also is the destructive attack of a metal by chemical or electrochemical reaction with environment (Uhlig, 1985). Despite of all its different definitions, corrosion is basically the result of interaction between materials and their environment. Some of the typical corrosive environments are (Zaki Ahmad, 2006a):

- > Air and humidity
- Fresh, distilled, salt and marine water
- Acids and alkalies
- Soil, steam, gases

Nowadays, corrosion is not confined to metals and alloys alone, but it also now encompasses with all types of natural and man-made material for example biomaterials and nanomaterials. The scope of corrosion is consistent with the revolutionary changes in materials development witnessed in recent years.

1.1.3 Corrosion of iron

Irons play an important role in the industrial sectors nowadays. Problems that mostly encountered by the industrial sectors are corrosion of metals. The driving force that causes iron to corrode is natural consequence of their temporary existence in ironic form (thermodynamically unstable). In order to produce irons starting from naturally occurring minerals and ores, it is necessary to provide a certain amount of energy (Fig. 1.1). It is therefore only natural that when these irons are exposed to their environments, they would revert back to original state in which they were found (Roberge, 2008a; Zaki Ahmad, 2006a).

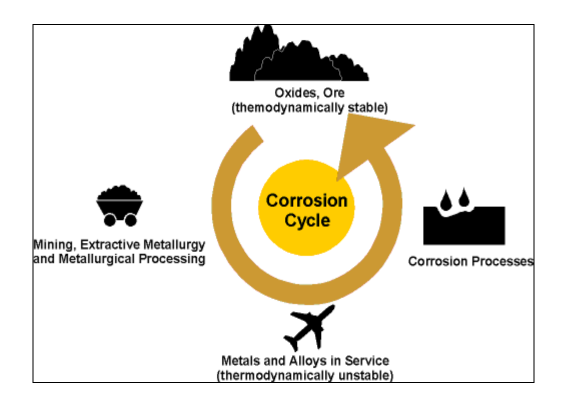


Figure 1.1: The corrosion cycle of steel.

In the presence of water and oxygen, metallic iron is thermodynamically unstable and corrosion proceeds according to the electrochemical mechanisms (Figure 1.2). As many electrochemical processes, the redox reaction of corrosion occurs both in anodic and cathodic terminal side of the metal surface. More than one oxidation and reduction process may occur during the process.

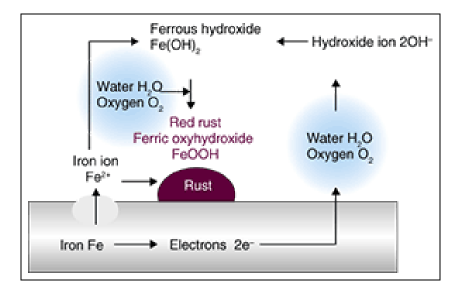


Figure 1.2: Electrochemical mechanism of corrosion of iron.

In the initial stage, iron is oxidized at the anode terminal to dissolve Fe^{2+} :

$$\mathbf{Fe} \to \mathbf{Fe}^{2+} + 2\mathbf{e}^{-} \tag{1.1}$$

Oxygen is reduced at the cathode terminal to form OH ions:

$$H_2 0 + \frac{1}{2} 0_2 + 2e^- \rightarrow 20H^-$$
 (1.2)

The cathodic reaction represented by Equation 1.2 exemplifies corrosion in natural environment where corrosion occurs at nearly neutral pH values (Kruger, 2001).

In de-aerated or acidic solution, the cathodic reaction is (Sheir et al., 1994):

$$2\mathrm{H}^{+} + 2\mathrm{e}^{-} \to \mathrm{H}_{2} \tag{1.3}$$

If there is no oxidation or reduction occurs, the overall corrosion reaction is of course the sum of the cathodic and anodic partial reactions. This will form insoluble iron (II) hydroxide or green rust:

$$Fe^{2+} + 2 OH^- \rightarrow Fe(OH)_2$$
 (1.4)

The colour of Fe(OH)₂, although white when the substance is pure, is normally green to greenish black because of incipient oxidation by air (Revie and Uhlig, 2008a). The unstable iron (II) ions can also further oxidize to produce stable iron (III) ions:

$$4Fe^{2+} + 4H^{+} + 0_2 \rightarrow 4Fe^{3+} + 4H_20$$
(1.5)

The iron (II) ions react with hydroxide ions to produce hydrated iron (III) oxides known as iron (III) hydroxides or ferric hydroxides or rust (reddish brown ferric hydroxide):

$$Fe^{3+} + 3OH^- \rightarrow Fe(OH)_3$$
 (1.6)

.

If iron comes into contact with hydrochloric acid, only the properties of the salt ferrous chloride and hydrogen gas can be detected (Groysman, 2010a):

$$Fe + 2HCl \rightarrow FeCl_2 + H_2$$
 (1.7)

The $FeCl_2$ will be oxidized by air transported through the anode channels to ferric hydroxide (Tamura, 2008):

$$\operatorname{FeCl}_2 + \frac{1}{4} \operatorname{O}_2 + \frac{5}{2} \operatorname{H}_2 \operatorname{O} \to \operatorname{Fe}(\operatorname{OH})_3 + 2\operatorname{HCl}$$
 (1.8)

The aging of $Fe(OH)_3$ leads to dehydration even in the presence of water and forms oxyhydroxides, FeOOH where no acid or base is consumed or generated and there would be no change in pH during the aging (Morgan and Stumm, 1981):

 $Fe(OH)_3 \rightarrow FeOOH + H_2O$ (1.9)

Or, the Fe(OH)₃ can slowly transform into a crystallized form written as Fe₂O₃.H₂O (red-brown in colour). There are many forms of corrosion products can be observed in the corrosion reaction of iron. Further oxidation and hydrolysis process in corrosion reaction of Fe(OH)₃ will produce α -FeOOH (goethite), β -FeOOH (akaganite), γ -FeOOH (lepidocrocite), Fe₃O₄ (magnetite) and so forth.

1.1.3 Consequences of corrosion

Corrosion of metal costs the Malaysian economy more than billions ringgit per year at current prices. For most industrialized nations, the average corrosion cost is 3.5-4.5 % of the gross domestic product (GDP) (Zaki Ahmad, 2006a; Davis, 2000). In a study of corrosion cost conducted jointly by C.C. Technologies Inc. USA, Federal Highway Agencies (FHWA) USA and National Association of Corrosion Engineers (NACE), the direct corrosion cost was estimated to be around 276 billion US dollars, approximately 3.1 % of the national gross domestic product (Zaki Ahmad, 2006a). Of far more serious consequences is how corrosion affects our daily lives. Some consequences are economic and social, and cause the following (Davis, 2000):

- Replacement of corroded equipment.
- > Preventive maintenance, for example, painting.
- Shutdown of equipment due to corrosion failure.
- Contamination of a product.
- Safety, for example, sudden failure can cause fire, explosion, release of toxic product and construction collapse.
- Health, for example, pollution due to escaping product from corroded equipment.

These costs can be reduced by broader application of corrosion-resistant materials and the application of best corrosion-related technical practices.

1.2 Corrosion protection

It is impossible for the corrosion scientist nowadays to completely eliminate corrosion. However, the types of electrochemical corrosion just described can be prevented or controlled by utilizing the current understanding of the principles underlying corrosion process. This understanding has been the basis for the development of a number of corrosion prevention measures. Three corrosion control measures are based on the electrochemical driving force as shown in the Pourbaix diagram in Figure 1.3 (Kruger, 2001):

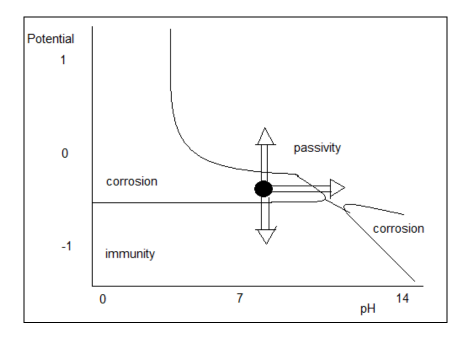


Figure 1.3: Pourbaix diagram for iron showing how corrosion protection can be achieved by using measures that bring a corroding system into either the immunity or the passivity region (Kruger, 2001).

The method chosen for corrosion prevention must consider the economical cost and the effectiveness since corrosion depends on both the specific water quality and material in a system. So, a particular method may be successful in one system, not in another (Singley *et al.*, 1985a). There are five primary methods of corrosion control that is material selection, coatings, inhibitors, cathodic protection and design (Singley *et al.*, 1985a; Kruger, 2001; Davis, 2000).

1.3 Corrosion inhibitors

Corrosion can be controlled by adding chemicals to the corrodent that form a protective film on the surface of a metal and provide a barrier between the water and the metal. This form of corrosion protection is called inhibition and the compound added to the system is called corrosion inhibitors. The inhibitors will reduce the corrosion rate but do not totally prevent it (Singley *et al.*, 1985a; Schweitzer, 2007b; Revie and Uhlig, 2008b; Kruger, 2001). The corrosion inhibitors are usually present in very low concentration (Groysman, 2010b; Sastri *et al.*, 2007b). According to Groysman (2010c), the concentration of corrosion inhibitors can be change from 1 to 15,000 ppm (0.0001 to 1.5 wt %). Besides, the inhibitors can be used at various ranges of pH, from acid, near neutral to alkaline. The most common and widely known use of inhibitors is their application in automobile cooling systems and boiler feedwaters.

It has been postulated that the inhibitors are adsorbed into the metal surface either by physical (electrostatic) adsorption or chemical adsorption. Physical adsorption is the result of electrostatic attractive forces between the organic ions and the electrically charged metal surface. Chemical adsorption is the transfer or sharing the inhibitor molecule's charge to the metal surface, forming a coordinate-type bond (Schweitzer, 2007b; Satri *et al.*, 2007b). Further explanations regarding the adsorption mechanism of inhibitors will be discussed on the next subchapter. The adsorbed inhibitor will reduces the corrosion rate of the metal surface either by retarding the anodic dissolution reaction of the metal by the cathodic evolution of hydrogen gas, or both.

According to Sastri *et al.* (2007b), a corrosion inhibitor can function in two ways. In some situations the added inhibitors can alter the corrosive environment into a noncorrosive or less corrosive species. In other cases, the corrosion inhibitor interacts with the metal surface and as a consequence, inhibits the corrosion of the metal. In the case of environment modifiers, the action and mechanism of inhibition is a simple interaction with the aggressive species in the environment, and thus reduce the attack of the metal by the aggressive species. Whilst, in the case of inhibitors which adsorb on the metal surface and inhibit the corrosion, there are two step namely: (i) transport of inhibitor to the metal surface and (ii) metal-inhibitor interaction.

In order to compare various corrosion inhibitors and select the most effective one for corrosion control, we have to calculate their efficiency (Groysman, 2010c; Schweitzer, 2007b; Roberge, 2008c; Sastri *et al.*, 2007b):

IE % =
$$\frac{CR_0 - CR_i}{CR_0} \times 100$$
 (1.10)

where CR_0 is the corrosion rate of metal in media without inhibitor, while CR_i is the corrosion rate of metal in media with inhibitor. The corrosion rate can be measured by any available method (weight loss or electrochemical method). According to Roberge (2008c), the efficiency of inhibitor increases with an increase of inhibitor concentrations until some point. Speaking about the efficiency, it is important to understand that there are some factors that can influence the efficiency of corrosion inhibitors. Groysman (2010c), stated that the common factors that influence the

efficiency of corrosion inhibitors are: the chemical position of aqueous solution, pH, flow rate, temperature and metal surface conditions (roughness, presence of corrosion product and other compounds). Inhibitors are most effective in definite pH range. For example, nitrites loses their effectiveness below a pH of 5.5 to 6.0, polyphosphate should be used between pH of 6.5 to 7.5, Zn-phosphate inhibitors are usually used at pH of 7.8 to 8.2.

Nowadays, there are various kinds of electrochemical instruments that commonly used to study the corrosion inhibition of a metal. Among those, the most popular once are gravimetric measurement or so called weight loss measurement, potentiodynamic polarization measurement and electrochemical impedance measurement. Numbers of research paper that have been published discuss about the metal corrosion inhibition behaviour using electrochemical measurements as stated above (Sherif and Park, 2006a; Obot *et al.*, 2009; Wahyuningrum *et al.*, 2008; Cheng *et al.*, 2007; Acharya and Upadhyay, 2004; Bouklah *et al.*, 2006; Morad and Kamal El-Dean, 2006; Talati *et al.*, 2005; Behpour *et al.*, 2007; Kustu *et al.*, 2007; Hosseini *et al.*, 2007; Hasanov *et al.*, 2007; Asan *et al.*, 2006).

1.3.1 The inhibitors

Generally, there are several classes of inhibitors, conveniently designated as follows (Schweitzer, 2007b; Revie and Uhlig, 2008b):

- Passivators, mainly chromates and nitrite.
- > Organic inhibitors, including slushing compound and pickling inhibitors.
- ➢ Vapour-phase inhibitors.

Each inhibitor that being used for corrosion inhibition study will give different results depending on the environment or method employed. Even though passivator inhibitors are less expensive compare with other inhibitor, but most of corrosion scientists or engineers tend to use organic inhibitor as their main ingredient especially in paint or radiator coolant formulations. The advantages of organic inhibitor are because these materials build up a protective film of adsorbed molecules on the metal surface that provide a barrier to the dissolution of the metal in the electrolyte, not like other inhibitor which only effective for certain conditions.

Organic inhibitor can be divided into two categories, synthetic organic inhibitor or natural organic inhibitor. There is a lot of paper published in using synthetic compound as an inhibitor, such an example azole derivatives (Sherif and Park, 2006a; Obot *et al.*, 2009; Wahyuningrum *et al.*, 2008), carboxymenthylchitosan (Cheng *et al.*, 2007), fluoroquinolones derivatives (Acharya and Upadhyay, 2004), pyridazine (Bouklah *et al.*, 2006), pyridine (Morad and Kamal El-Dean, 2006), Schiff bases derivatives (Talati *et al.*, 2005; Behpour *et al.*, 2007; Kustu *et al.*, 2007; Hosseini *et al.*, 2007; Hasanov *et al.*, 2007; Asan *et al.*, 2006) typtamine (Lowmunkhong *et al.*, 2010), quinine (Awad, 2006) and so forth. Unfortunately, even though these synthetic compounds showed very good anti-corrosive activity, but most of them are highly toxic to human and environment. Thus, synthetic inhibitors can cause reversible (temporary) or irreversible (permanent) damage to internal organ as its presence in drinking water tank or in a pipeline. The toxicity of synthetic inhibitors maybe arises during the synthesis of the compound or while undergoing its application (Raja and Sethuraman, 2008). Hence, there is now been a trend to use natural product as green or so called "eco-friendly" inhibitor.

Some of the natural product inhibitors give different range of efficiencies, depending on the type of plant, metal and corrosive media used. Valek and Martinez (2007) used the leaves extract of *Azadirachta indica* as corrosion inhibitor for copper in sulphuric acid solution. Corrosion inhibition has also been studied for the extracts of olive leaves or *Olea europaea* for carbon steel (El-Etre, 2007), natural honey and black radish juice for tin (Radojc[°]ic[′] *et al.*, 2008), *Ammi visnaga* or khillah for SX 316 steel (El-Etre, 2006), *Nypa fruticans* Wurmb for mild steel (Orubite and Oforka, 2004), *Sanseviera trifasciata* for aluminium (Oguzie, 2007), mangrove tannins (Rahim *et al.*, 2008) and so forth.

1.3.2 Corrosion inhibition of mild steel in acidic solution

Acid solution are widely used in industry, the most important fields of application being acid pickling, industrial acid cleaning, acid descaling and oil well acidizing. As a result of aggressivity of acidic solution, the inhibitors are commonly used to reduce the corrosive attack on metallic materials. The development of inhibitors of steels in acidic solution has been the subject of great interest especially from the point of view of their efficiency and applications. Acids are classified into two parts which is mineral acids (e.g. sulphuric acid, hydrochloric acid and phosphoric acid) and organic acids (e.g. acetic acid, malic acid and tartaric acid).

Most of corrosion inhibition studies were conducted in various concentration of acidic media since the use of acid solution in industrial sector is higher than others (alkaline or neutral) solution. Sulphuric acid and hydrochloric acid are aggressive towards the corrosion of iron and its alloys; hence most of corrosion chemists like to study the inhibitive properties in this environment. Eddy and Odoemelam (2009) reported about the corrosion inhibition behaviour of mild steel using ethanol extract of *Aloe vera* in sulphuric acid solution. Other research papers also use natural inhibitor as mild steel corrosion inhibitor, for example *Azadirachta indica* extract in hydrochloric acid and sulphuric acid solution (Oguzie, 2006), caffeic acid in sulphuric acid solution (de Souza and Spinelli, 2009), *Calendula officinalis* flower in hydrochloric acid solution (Raja and Sethuraman, 2009), *Carica papaya* in sulphuric acid solution (Okafor and Ebenso, 2007) and many more.

1.3.3 Methods for corrosion inhibition study

In every study, there must be significant methods to prove any theory arises. In order to study more detail about the corrosion inhibition phenomenon, various methods has been developed from past until now. Previously, there are only few tests or methods employed to study the corrosion inhibition, but as the world of research keep wheeling, more and more methods were being manipulate to understand the concept. Such methods are weight loss measurement, gasometric evaluation, potentiodynamic polarization measurement and electrochemical impedance spectroscopy (EIS). Surface and elemental analysis for example scanning electron microscope coupled with energy dispersive X-ray (SEM-EDX), X-ray diffraction spectroscopy (XRD), atomic force microscope (AFM) and X-ray florescence spectroscopy (XRF) are sometimes used to explain the surface morphology of metal and the formation of protective layer on metal surface. Below are few methods that commonly used in corrosion inhibition study.

1.3.3.1 Weight loss measurement

Weight loss measurements are considered as the simplest way to study corrosion phenomenon. Some refer it as gravimetric measurement (Okafor and Ebenso, 2006; Bendahou *et al.*, 2006). It gives useful information about the average corrosion rate over certain time period. In weight loss measurement, metal are usually being hanged or stand in a beaker that is loaded with corrosive medium and often covered with aluminium foil to avoid contamination and evaporation (Figure 1.4).

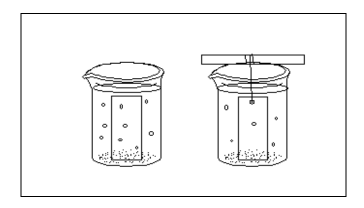


Figure 1.4: Typical weight loss measurement setup with metal being stand (left-hand side) and hanged (right-hand side).

There are certain things that we have to consider before undergo weight loss measurement. For example, the dimension or the size of metals, volume of electrolyte, pre-treatment before and after the test and time immersion. There is no restriction for the dimension of mild steel. El-Etre (2006) used a dimension of 2 cm x 5.6 cm x 0.2 mm (total surface area was 25 cm²) for his SX 316 steel while Valek and Martinez (2007) used a dimension of 4 cm x 5 cm x 3 mm (total surface area was 60 cm²) for their studies. Volume of electrolyte or solution that is used for the weight loss study must be sufficient in order to obtain a complete immersion of metal.

The pre-treatment of metal before and after weight loss test is important in order to get a precise value of inhibition. Metal surface is usually being treated before it is used in weight loss test. In this case, metal surface is been polished with emery paper or silicon carbide (SiC) paper from the most rough to less rough grade using a grinder polisher in order to obtain like a "mirror" surface. Wet polishing is commonly associated with this treatment where water will act as a lubricant.

Moreover, cleaning process is also one of the important procedures in weight loss measurement. This can divide into two categories, mechanical and chemical cleaning. Mechanical cleaning leads to the removal of substance that is attached on metal surface. Some methods used are scrubbing, scraping, sonicate, and brushing. While chemical cleaning is used to remove any material that forms or already exist on metal surface using suitable chemical solution. Examples of chemical solution used in chemical cleaning are acetone (El-Etre, 2006; Subha and Saratha, 2006; Raja and Sethuraman, 2009), alcohol (Oguzie, 2006; Valek and Martinez, 2007) and ASTM G1-90 or known as Clarke's solution (Singh and Kumar, 2003).

Time of immersion is another thing that we have consider while studying weight loss test. Time that is fixed in weight loss measurement is the time which gives the maximum corrosion rate.

1.3.3.2 Potentiodynamic polarization measurement

Polarization studies have been primarily laboratory electrochemical technique to study corrosion phenomena, especially pitting and passivity, by disturbing the natural corrosion potential of a system, frequently by substantial amounts of volts and measuring the external current flowing (Schweitzer, 2007c; Jain, 2010b). The characterization of a metal specimen is done via current-potential relationship (Jain, 2010b). According to Perez (2004a), polarization is a process when the electrode reaction are assumed to induced deviations from equilibrium due to the passage of an electrical current through an electrochemical cell, causing a change in the working electrode (WE). The specimen is polarized to within ± 25 mV of the corrosion potential is linear (Zaki Ahmad, 2006b). The term E_{corr} can be defined as the potential at which the rate of oxidation is exactly equal to the rate of reduction (Jain, 2010b). In general, the electrochemical and chemical rates of reaction due to anodic or cathodic overpotentials can be predicted using both Faraday (Eq. 1.12) and Arrhenius (Eq. 1.13) equation respectively (Perez, 2004a):

$$R_{\rm F} = \frac{iA_{\rm wj}}{zF} \tag{1.12}$$

$$R_{A} = \gamma_{a} \exp\left(-\frac{\Delta G^{*}}{RT}\right)$$
(1.13)

where i is applied current density (A cm⁻²), $A_{w,j}$ is the atomic weight of species j (g mol⁻¹), z is the oxidation state or valence number, γ_a is the chemical reaction constant and ΔG^* is the activation energy or free energy change (J mol⁻¹).

At equilibrium, Faraday's and Arrhenius rate equations become equal $(R_F = R_A)$ and consequently the current density becomes:

$$i = \gamma_a \exp\left(-\frac{\Delta G^*}{RT}\right) \tag{1.14}$$

On the other hand, if an electrode is polarized by an overpotential under steady-state conditions, then the rate of reaction are not equal ($R_F \neq R_A$) and consequently, the forward (i_f , cathodic) and reverse (i_r , anodic) current density components must be defined in terms of the free energy change ΔG deduce from Figure 1.5 where k'_f and k'_r are the forward and reverse rate. Hence;

$$i_{f} = k'_{f} \exp\left(-\frac{\Delta G_{f}^{*}}{RT}\right)$$
 (cathodic) (1.15)

$$i_r = k'_r \exp\left(-\frac{\Delta G_r^*}{RT}\right)$$
 (anodic) (1.16)

and,
$$\Delta G^*_{f} = \Delta G_{f} - \alpha z F \eta_{c}$$
 (1.17)

$$\Delta G^*_r = \Delta G_r + (1 - \alpha) z F \eta_a \tag{1.18}$$

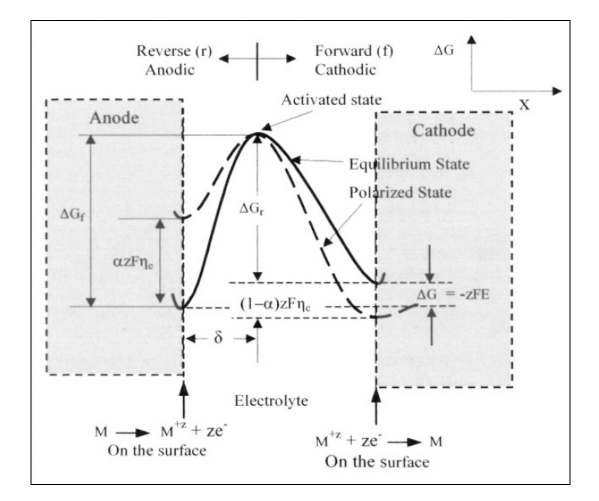


Figure 1.5: Schematic activation free energy distribution (Perez, 2004a).

 α is the symmetry coefficient, F is the Faraday's constant and η is the potential of the reaction or polarization (for both cathodic and anodic). For a cathodic case, the net current and the overpotential are $i = i_f - i_r$ and η_c respectively. Substituting equation (1.15) and (1.16) into this expression yields the net current density in a general form:

$$i = k'_{f} \exp\left(-\frac{\Delta G_{f}^{*}}{RT}\right) \exp\left(\frac{\alpha z F \eta}{RT}\right) - k'_{r} \exp\left(-\frac{\Delta G_{r}^{*}}{RT}\right) \exp\left(-\frac{(1-\alpha) z F \eta}{RT}\right)$$
(1.19)

From which the exchange current density is deduced as:

$$\mathbf{i_o} = \mathbf{k'_f} \exp\left(-\frac{\Delta \mathbf{G_f}^*}{\mathbf{RT}}\right) = \mathbf{k'_r} \exp\left(-\frac{\Delta \mathbf{G_r}^*}{\mathbf{RT}}\right)$$
(1.20)

Substituting equation (1.20) into (1.19) for one-step reaction yields the well known **Butler-Volmer equation** for polarizing an electrode from the open circuit potential E_0 under steady-state conditions:

$$i = i_{o} \left\{ \exp\left(\frac{\alpha z F \eta}{RT}\right)_{f} - \exp\left(-\frac{(1-\alpha) z F \eta}{RT}\right)_{r} \right\}$$
(1.21)

where $\eta = E - E_o$ and E is the applied potential.

If the overpotential is large and positive ($\eta > +0.052$ V), the second term in equation (1.21) can be neglected (Jain, 2010c):

$$\mathbf{i} = \mathbf{i}_{a} = \mathbf{i}_{o} \left\{ \exp\left(\frac{\alpha z F \eta}{RT}\right)_{f} \right\}$$
(1.22)

If the overpotential is large and negative ($\eta < -0.052$ V) the first term in equation (1.21) can be neglected:

$$\mathbf{i} = \mathbf{i}_{c} = \mathbf{i}_{o} \left\{ \exp\left(-\frac{(1-\alpha)\mathbf{z}\mathbf{F}\boldsymbol{\eta}}{\mathbf{R}\mathbf{T}}\right)_{\mathbf{r}} \right\}$$
(1.23)

Taking the logarithms in equation (1.22):

$$\begin{aligned} &\ln i_{a} = \ln i_{o} + \frac{\alpha z F \eta}{RT} \\ &\ln \left(\frac{i_{a}}{i_{o}}\right) = \frac{\alpha z F \eta}{RT} \\ &\eta = \frac{RT}{\alpha z F} \ln \left(\frac{i_{a}}{i_{o}}\right) \end{aligned}$$

or,

$$\eta_{a} = \frac{2.303 \text{RT}}{\alpha z F} \ \text{In} \ \left(\frac{i_{a}}{i_{o}} \right)$$

Then, by assuming that η_a = β_a log $(i_a\!/i_o)$ and η_b = β_c log $(i_c\!/i_o)$:

$$\beta_{a} = \frac{2.303 \text{RT}}{\alpha z F}$$
(1.24)

$$\beta_{\rm c} = \frac{2.303 \rm RT}{(1-\alpha) \rm zF} \tag{1.25}$$

 β_a and β_c are known as **Tafel slope**. These equations are used to calculate the anodic and cathodic current slope (Figure 1.6).

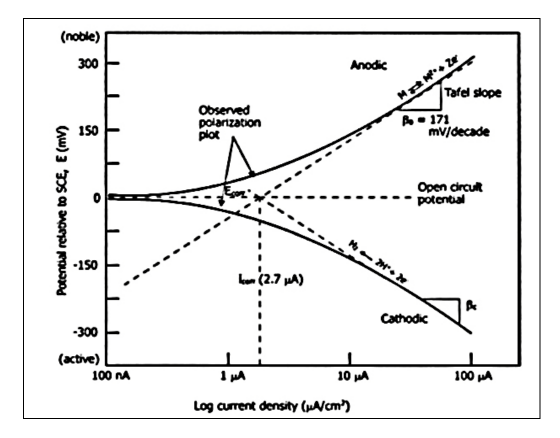


Figure 1.6: A hypothetical Tafel plot (Zaki Ahmad, 2006c).

Polarization resistance or so called linear polarization is usually being used in potentiodynamic polarization measurement. This technique is quick and reliable. According to Zaki Ahmad (2006c), polarization resistance (R_p) of a corroding metal is defined Ohm's Law as the slope of a potential (E) versus current density (log i) plot at the corrosion potential (E_{corr}). Here, $R_p = (\Delta E/\Delta I)$ at $\Delta E = 0$. By measuring this slope, the rate of corrosion can be measured. The correlation between i_{corr} and slope (dE/dI) is given by:

$$Rp = \frac{\Delta E}{\Delta I} = \frac{\beta_a \beta_c}{2.303 i_{corr} (\beta_a + \beta_c)}$$
(1.26)

where β_a and β_c are Tafel slope.

There are three types of inhibitor that we can observe in potentiodynamic polarization test, which is cathodic inhibitor, anodic inhibitor and mixed inhibitor. Cathodic inhibitors inhibit the hydrogen evolution in acidic solution or the reduction of oxygen in neutral or alkaline solutions. It is also observed that the cathodic polarization curve is affected when a cathodic inhibitor is added to the system. Substances with high overpotential for hydrogen in acidic solutions and those that form insoluble products in alkaline solution are generally effective cathodic inhibitors. Anodic inhibitors are generally effective in the pH range of 6.5 to 10.5 (near neutral to basic). Basically, oxyanions are very effective anodic inhibitors. Those oxyanions are thought to play a role of repairing the defects in the passive metal oxide film on the metal surface. Mixed type of inhibitors is generally represented by organic compounds. Irrespective of type of inhibitor, the inhibition processes involve transport of inhibitor to the metal site followed by interaction of the inhibitor with the surface of the metal, resulting a protection (Sastri *et al.*, 2007b).

1.3.3.3 Electrochemical impedance spectroscopy (EIS)

Electrochemical impedance spectroscopy (or AC impedance) is a now well established laboratory technique used to determine the electrical impedance of metalelectrolyte interface at various AC excitation frequencies (Schweitzer, 2007c). Impedance measurements combine the effect of DC resistance with capacitance and inductance. Furthermore, the AC impedance is capable of characterizing the corrosion interface more comprehensively in lower conductivity solution or highresistivity coatings (Schweitzer, 2007c; Revie and Uhlig, 2008c). In an alternating circuit, impedance determines the amplitude of current for a given voltage and the proportionality factor between voltage and current (Revie and Uhlig, 2008c; Groysman, 2010d). The response of an electrode to alternating potential signals of varying frequency is interpreted on the basis of circuit models of the electrode or electrolyte interphase (Revie and Uhlig, 2008c; Perez, 2004b).

In DC theory, resistance R is defined by Ohm's Law:

$$\mathbf{E} = \mathbf{I}\mathbf{R} \tag{1.27}$$

where I is the current (A) and E is the potential (V). Using the Ohm's law, one can apply a DC potential to a circuit and measure the resulting current, from which the resistance can be calculated (Zaki Ahmad, 2006b; Perez, 2004b; Groysman, 2010d). In AC theory:

$$\mathbf{E} = \mathbf{I}\mathbf{Z} \tag{1.28}$$

Here, Z is the magnitude of the impedance containing elements of an equivalent circuit, such as capacitors and inductors. Capacitors oppose or impede the current flow (Perez, 2004b). AC currents and voltages are vector quantities. Impedance can be expressed as a complete number where the resistance is the real component and combined capacitance and inductance is the imaging component (Zaki Ahmad, 2006b). The resulting vector for impedance is:

$$Z_{\text{total}} = Z' + Z''_{j}$$
 (1.29)

where Z' is the real impedance, Z" is the imaginary impedance and j is $\sqrt{(-1)}$. The absolute magnitude of impedance is:

$$|\mathbf{Z}| = \sqrt{(\mathbf{Z}')^2 + (\mathbf{Z}'')^2} \tag{1.30}$$

and,

$$\tan \theta = \frac{Z'}{Z''} \tag{1.31}$$

When the opposition to the current is capacitive resistance, the current leads the applied voltage in phase angle. But when the opposition to current flow is inductive reactance, the current lags behind the voltage in phase angle (Zaki Ahmad, 2006b). The phase angle (θ) is the difference between points on x-axis where current and voltage curve amplitudes are zero (Figure 1.7). Table 1.1 shows the transfer function for resistors, capacitors and inductors.

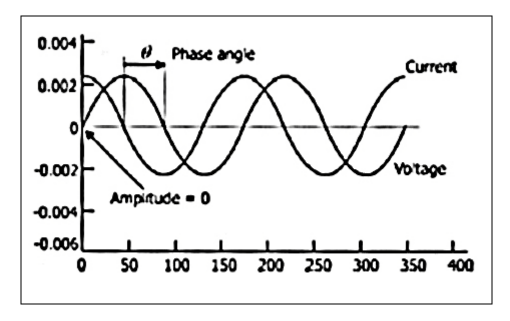


Figure 1.7: The voltage-current phase angle (Zaki Ahmad, 2006b).