



Alternating Current-Induced Corrosion

Introduction

Corrosion induced by alternating currents (AC) is evident in the oil and gas industry, particularly when a pipeline is in close proximity to high power transmission lines.

The model presented here first evaluates the effect of a direct current (DC) applied potential on corrosion using a stationary analysis, and then evaluates the effect of AC on corrosion using a transient analysis. The model is subsequently extended to investigate the effect of frequency on the AC corrosion rate, thereby demonstrating the role of the capacitive double-layer at higher frequencies.

The model is based on a paper by Ghanbari and others (Ref. 1).

Model Definition

The model geometry is defined in 1D, where the model geometry length is set to equal to the diffusion layer thickness. A steel electrode surface is assumed to be located at the left-hand side, with the bulk electrolyte boundary placed at right, as shown in Figure 1.

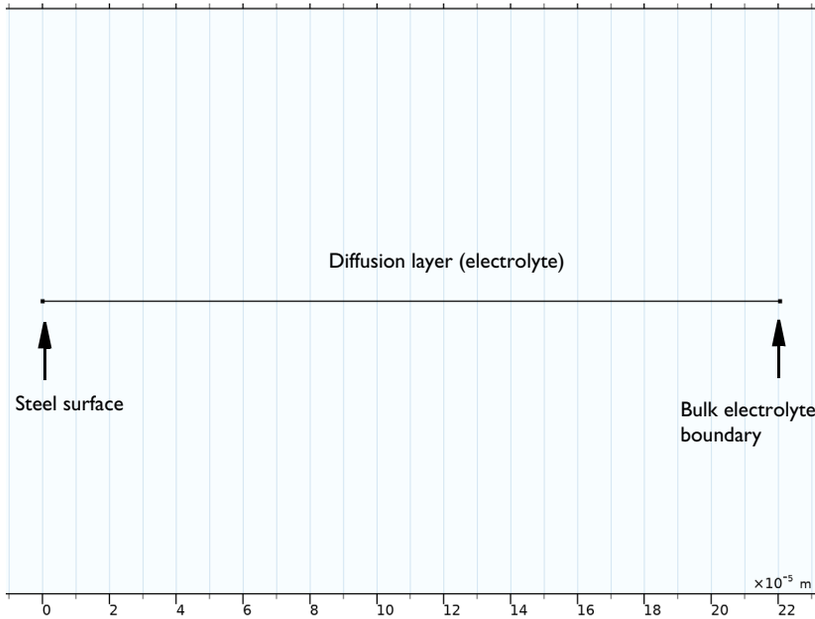


Figure 1: Model geometry

The concentration of dissolved oxygen is solved for across diffusion layer thickness, with the transport equation defined according to the Fick's law:

$$\frac{\partial c_i}{\partial t} + \nabla \cdot (-D_i \nabla c_i) = 0 \quad (1)$$

where c_i is the dissolved oxygen concentration in the electrolyte (mol/m^3) and D_i is diffusion coefficient of dissolved oxygen in the electrolyte (m^2/s).

The diffusion layer thickness is set based on the limiting current density for the oxygen reduction reaction, i_{lim} , across the diffusion layer.

$$L = \frac{4FDc_{\text{sat}}}{i_{\text{lim}}} \quad (2)$$

The oxygen concentration is set to equal the saturated concentration for dissolved oxygen when in equilibrium with air, c_{sat} , at the right hand side boundary. At the left hand electrode surface, the flux of oxygen is defined, based on the local current density for oxygen reduction in combination with Faraday's law.

The Electrode Surface boundary node is used to calculate the total current density, i_T , which includes contributions from the metal dissolution (anodic), oxygen reduction (cathodic) and hydrogen evolution (cathodic) electrochemical reactions, as well as double layer capacitance.

The electrode potential is assumed to be measured vs a reference electrode located outside the diffusion layer. The electrode potential vs the reference, E_T , is hence the sum of potential drop across the electrochemical interface, E , and potential drop across the solution resistance, R_s .

$$E_T = E + i_T R_s \quad (3)$$

The potential drop across the electrochemical interface, E , is solved using a Global Equations node.

The potential of the steel electrode is defined to be the sum of a DC potential, E_{DC} , and an AC potential perturbation, and is set according to

$$E_T = E_{\text{DC}} + E_{\text{RMS}} \sin(\omega t) \quad (4)$$

where E_{RMS} is the amplitude of AC potential and ω is the angular frequency which is equal to $2\pi f$. f is the frequency of AC potential and t is time.

Tafel kinetics are used to define all faradaic reactions. For the oxygen reduction reaction, the kinetics expression is concentration dependent according to

$$i_{\text{O}_2} = -\frac{c}{c_{\text{sat}}} i_{0, \text{O}_2} 10^{\eta/A_c} \quad (5)$$

where A_c is the Tafel slope, and $\eta = E - E_{\text{eq}}$ is the overpotential.

The non-faradaic double layer capacitance current density is defined as

$$i_{\text{dl}} = C_{\text{dl}} \frac{dE}{dt} \quad (6)$$

where C_{dl} is considered to be a combination of double layer and oxide capacitances. Different C_{dl} values are used in the model for different DC applied potentials, which are taken from [Ref. 1](#).

The model is solved in two steps. In the first step, a stationary solution for the pure DC problem is computed. In the second time-dependent step, the solution of the stationary step is used as initial values, and the simulation is performed for multiple consecutive periods. In order to compare the AC to the DC solution, the corrosion rate is calculated as an average for the last period of the simulation.

Results and Discussion

[Figure 2](#) shows the anodic (metal dissolution) current density for different applied DC potentials with and without AC. It can be seen that the anodic current density in both cases (with and without AC) increases with an increase in applied DC potential. Adding

the AC contribution generally increases the corrosion rate, with the effect of AC being most dominant near the open circuit potential of -0.67 V/SCE.

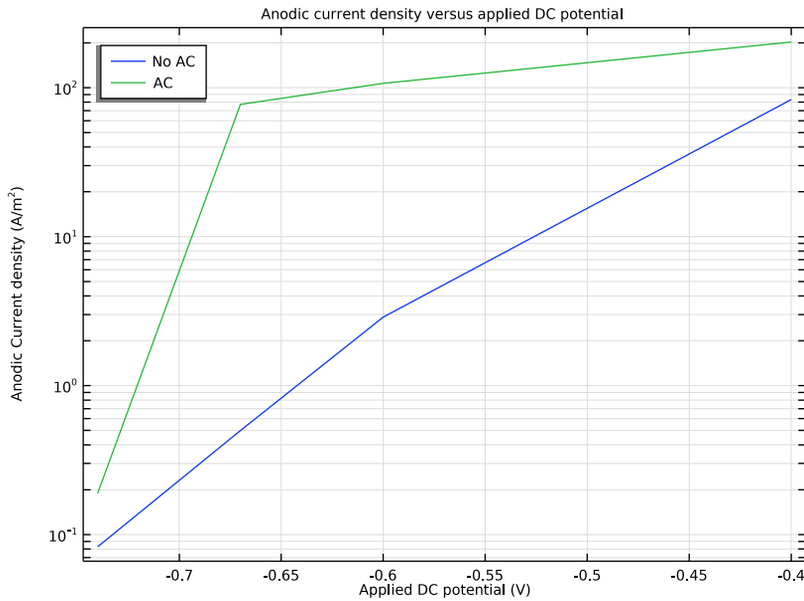


Figure 2: Anodic current density with and without AC for different applied DC potentials.

Figure 3 and Figure 4 show the change in total current density against time along with contributions from anodic, cathodic and double layer for frequencies 60 Hz and 0.01 Hz, respectively. It can be seen that double layer (non-faradaic) contribution to the total current density is quite significant at frequency of 60 Hz (Figure 3), whereas at lower

frequency of 0.01 Hz the total current density is mainly constituted of anodic current density (faradaic).

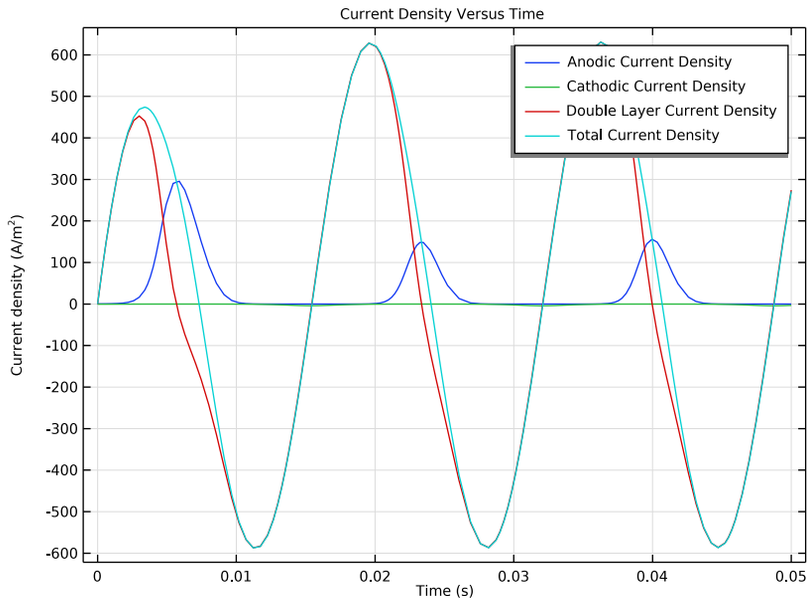


Figure 3: Transient distribution of the total current density along with anodic, cathodic and double layer contributions at frequency of 60 Hz.

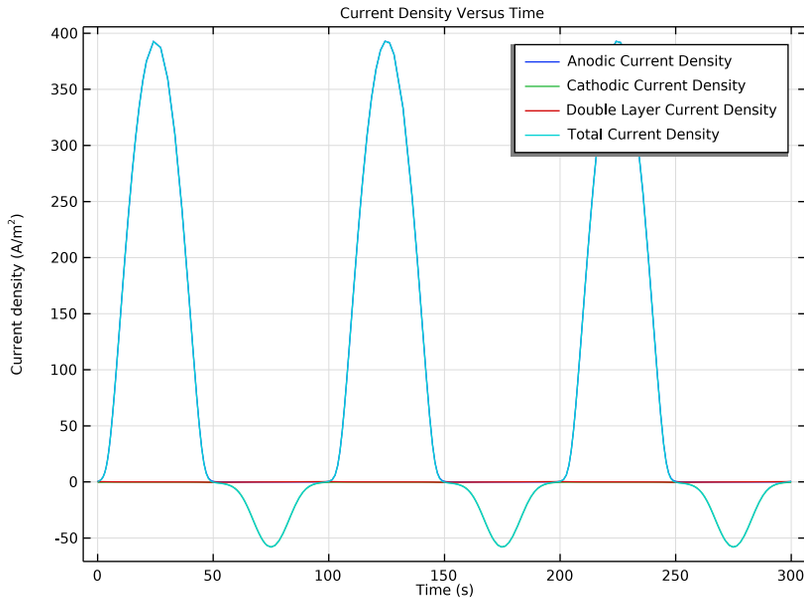


Figure 4: Transient distribution of the total current density along with anodic, cathodic and double layer contributions at frequency of 0.01 Hz.

References

1. E. Ghanbari, M. Iannuzzi, and R. S. Lillard, “The mechanism of alternating current corrosion of API grade X65 pipeline steel,” *Corrosion*, vol. 72, no. 9, pp. 1196–1210, 2016.

Application Library path: Corrosion_Module/General_Corrosion/ac_corrosion

Modeling Instructions

From the **File** menu, choose **New**.

NEW

In the **New** window, click  **Model Wizard**.

MODEL WIZARD

- 1 In the **Model Wizard** window, click  **ID**.
- 2 In the **Select Physics** tree, select **Electrochemistry>Electroanalysis (tcd)**.
- 3 Click **Add**.
- 4 In the **Number of species** text field, type 1.
- 5 In the **Concentrations** table, enter the following settings:

c02

- 6 In the **Select Physics** tree, select **Mathematics>ODE and DAE Interfaces>Global ODEs and DAEs (ge)**.
- 7 Click **Add**.
- 8 Click  **Study**.
- 9 In the **Select Study** tree, select **General Studies>Stationary**.
- 10 Click  **Done**.

GLOBAL DEFINITIONS

Load the model parameters from a text file.

Parameters 1

- 1 In the **Model Builder** window, under **Global Definitions** click **Parameters 1**.
- 2 In the **Settings** window for **Parameters**, locate the **Parameters** section.
- 3 Click  **Load from File**.
- 4 Browse to the model's Application Libraries folder and double-click the file `ac_corrosion_parameters.txt`.

GEOMETRY 1

Draw a 1D geometry representing the diffusion layer thickness.

- 1 From the **Geometry** menu, choose **Interval**.

Interval 1 (i1)

- 1 In the **Model Builder** window, under **Component 1 (comp1)** right-click **Geometry 1** and choose **Interval**.
- 2 In the **Settings** window for **Interval**, locate the **Interval** section.

3 In the table, enter the following settings:

| Coordinates (m) |
|-----------------|
| 0 |
| L |

4 Click  **Build All Objects**.

DEFINITIONS

Variables 1

Load the model variables from a text file.

- 1 In the **Home** toolbar, click  **Variables** and choose **Local Variables**.
- 2 In the **Settings** window for **Variables**, locate the **Variables** section.
- 3 Click  **Load from File**.
- 4 Browse to the model's Application Libraries folder and double-click the file `ac_corrosion_variables.txt`.

Integration 1 (intop1)

Add an integration operator.

- 1 In the **Definitions** toolbar, click  **Nonlocal Couplings** and choose **Integration**.
- 2 In the **Settings** window for **Integration**, locate the **Source Selection** section.
- 3 From the **Geometric entity level** list, choose **Boundary**.
- 4 Select Boundary 1 only.

ELECTROANALYSIS (TCD)

Set up the physics of the model.

Electrolyte 1

- 1 In the **Model Builder** window, under **Component 1 (comp1)>Electroanalysis (tcd)** click **Electrolyte 1**.
- 2 In the **Settings** window for **Electrolyte**, locate the **Diffusion** section.
- 3 In the D_{cO_2} text field, type `D_02`.

Initial Values 1

- 1 In the **Model Builder** window, click **Initial Values 1**.
- 2 In the **Settings** window for **Initial Values**, locate the **Initial Values** section.
- 3 In the cO_2 text field, type `c_02_sat`.

Concentration 1

- 1 In the **Physics** toolbar, click  **Boundaries** and choose **Concentration**.
- 2 Select Boundary 2 only.
- 3 In the **Settings** window for **Concentration**, locate the **Concentration** section.
- 4 Select the **Species cO2** check box.
- 5 In the $c_{0,cO2}$ text field, type `c_O2_sat`.

Electrode Surface 1

Next, set up the Electrode Surface node to describe the anodic, cathodic, and double-layer contributions to the total current density.

- 1 In the **Physics** toolbar, click  **Boundaries** and choose **Electrode Surface**.
- 2 Select Boundary 1 only.
- 3 In the **Settings** window for **Electrode Surface**, locate the **Electrode Phase Potential Condition** section.
- 4 In the $\phi_{s,ext}$ text field, type `E`.

Electrode Reaction: Anodic Reaction

- 1 In the **Model Builder** window, click **Electrode Reaction 1**.
- 2 In the **Settings** window for **Electrode Reaction**, locate the **Equilibrium Potential** section.
- 3 From the E_{eq} list, choose **User defined**. In the associated text field, type `Ecorr`.
- 4 Locate the **Electrode Kinetics** section. From the **Kinetics expression type** list, choose **Anodic Tafel equation**.
- 5 In the i_0 text field, type `icorr`.
- 6 In the A_a text field, type `ba`.
- 7 In the **Label** text field, type `Electrode Reaction: Anodic Reaction`.

Electrode Surface 1

In the **Model Builder** window, click **Electrode Surface 1**.

Electrode Reaction: Oxygen reduction cathodic reaction

- 1 In the **Physics** toolbar, click  **Attributes** and choose **Electrode Reaction**.
- 2 In the **Settings** window for **Electrode Reaction**, type `Electrode Reaction: Oxygen reduction cathodic reaction` in the **Label** text field.
- 3 Locate the **Stoichiometric Coefficients** section. In the n text field, type `4`.
- 4 In the ν_{cO2} text field, type `-1`.

- 5 Locate the **Equilibrium Potential** section. From the E_{eq} list, choose **User defined**. In the associated text field, type Ecorr.
- 6 Locate the **Electrode Kinetics** section. From the **Kinetics expression type** list, choose **Cathodic Tafel equation**.
- 7 In the i_0 text field, type icorr*cO2/ c_O2_sat.
- 8 In the A_c text field, type bc.

Electrode Surface 1

In the **Model Builder** window, click **Electrode Surface 1**.

Electrode Reaction: Hydrogen evolution cathodic reaction

- 1 In the **Physics** toolbar, click  **Attributes** and choose **Electrode Reaction**.
- 2 In the **Settings** window for **Electrode Reaction**, type Electrode Reaction: Hydrogen evolution cathodic reaction in the **Label** text field.
- 3 Locate the **Equilibrium Potential** section. From the E_{eq} list, choose **User defined**. In the associated text field, type E_H2.
- 4 Locate the **Electrode Kinetics** section. From the **Kinetics expression type** list, choose **Cathodic Tafel equation**.
- 5 In the i_0 text field, type i0_H2.
- 6 In the A_c text field, type b_H2.

Electrode Surface 1

In the **Model Builder** window, click **Electrode Surface 1**.

Double Layer Capacitance 1

- 1 In the **Physics** toolbar, click  **Attributes** and choose **Double Layer Capacitance**.
- 2 In the **Settings** window for **Double Layer Capacitance**, locate the **Double Layer Capacitance** section.
- 3 In the C_{dl} text field, type C.

GLOBAL ODES AND DAES (GE)

Add a Global Equation to solve for electric potential, E .

Global Equations 1

- 1 In the **Model Builder** window, under **Component 1 (comp1)>Global ODEs and DAES (ge)** click **Global Equations 1**.
- 2 In the **Settings** window for **Global Equations**, locate the **Global Equations** section.

3 In the table, enter the following settings:

| Name | f(u,ut,utt,t) (I) | Initial value (u_0) (I) | Initial value (u_t0) (I/s) | Description |
|------|--|----------------------------|-------------------------------|--------------------|
| E | E_app- tcd.itota vg_es1* Rs-E | E0 | 0 | Electric potential |

4 Locate the **Units** section. Click  **Define Dependent Variable Unit**.

5 In the **Dependent variable quantity** table, enter the following settings:

| Dependent variable quantity | Unit |
|-----------------------------|------|
| Custom unit | V |

6 Click  **Define Source Term Unit**.

7 In the **Source term quantity** table, enter the following settings:

| Source term quantity | Unit |
|----------------------|------|
| Custom unit | V |

STUDY : AC EFFECT

1 In the **Model Builder** window, click **Study 1**.

2 In the **Settings** window for **Study**, locate the **Study Settings** section.

3 Clear the **Generate default plots** check box.

4 In the **Label** text field, type Study : AC Effect.

Parametric Sweep

Use the parametric sweep to investigate the effect of applied DC potential for the respective double layer capacitance values.

1 In the **Study** toolbar, click  **Parametric Sweep**.

2 In the **Settings** window for **Parametric Sweep**, locate the **Study Settings** section.

3 Click  **Add**.

4 In the table, enter the following settings:

| Parameter name | Parameter value list | Parameter unit |
|-----------------------------|---------------------------------------|----------------|
| E_DC (Applied DC potential) | -0.74[V] -0.67[V] - 0.6[V] -0.4[V] | V |

5 Click  **Add**.

6 In the table, enter the following settings:

| Parameter name | Parameter value list | Parameter unit |
|------------------------------|--|------------------|
| C (Double layer capacitance) | 35[F/m ²] 2.61[F/m ²] 2.57[F/m ²] 3.09[F/m ²] | F/m ² |

Now solve the stationary study step to get results without AC effect first.

7 In the **Study** toolbar, click  **Compute**.

Solver Configurations

Store a copy of the solution for comparison purposes.

Solution 1 (sol1)

- 1 In the **Model Builder** window, expand the **Study : AC Effect>Solver Configurations** node.
- 2 Right-click **Solution 1 (sol1)** and choose **Solution>Copy**.

Solution 1 - No AC

- 1 In the **Model Builder** window, under **Study : AC Effect>Solver Configurations** click **Solution 1 - Copy 1 (sol2)**.
- 2 In the **Settings** window for **Solution**, type Solution 1 - No AC in the **Label** text field.

Time Dependent

Now add a Time Dependent study node to investigate the effect of AC.

- 1 In the **Study** toolbar, click  **Study Steps** and choose **Time Dependent>Time Dependent**.
- 2 In the **Settings** window for **Time Dependent**, locate the **Study Settings** section.
- 3 In the **Output times** text field, type range (0,tf/10,tf).
- 4 In the **Study** toolbar, click  **Compute**.

RESULTS

Plot the anodic current density to compare the effect of AC.

Effect of AC

- 1 In the **Home** toolbar, click  **Add Plot Group** and choose **ID Plot Group**.
- 2 In the **Settings** window for **ID Plot Group**, type Effect of AC in the **Label** text field.
- 3 Locate the **Data** section. From the **Dataset** list, choose **None**.
- 4 Click to expand the **Title** section. From the **Title type** list, choose **Manual**.

- 5 In the **Title** text area, type Anodic current density versus applied DC potential.
- 6 Locate the **Plot Settings** section.
- 7 Select the **y-axis label** check box. In the associated text field, type Anodic Current density (A/m^2).
- 8 Locate the **Axis** section. Select the **y-axis log scale** check box.
- 9 Locate the **Legend** section. From the **Position** list, choose **Upper left**.

Global 1

- 1 In the **Effect of AC** toolbar, click  **Global**.
- 2 In the **Settings** window for **Global**, locate the **Data** section.
- 3 From the **Dataset** list, choose **Study : AC Effect/Solution 1 - No AC (sol2)**.
- 4 Locate the **y-Axis Data** section. In the table, enter the following settings:

| Expression | Unit | Description |
|------------|------------------|------------------------|
| ia | A/m ² | Anodic current density |

- 5 Locate the **x-Axis Data** section. From the **Parameter** list, choose **Expression**.
- 6 In the **Expression** text field, type E_DC.
- 7 Click to expand the **Legends** section. From the **Legends** list, choose **Manual**.
- 8 In the table, enter the following settings:

| Legends |
|---------|
| No AC |

Effect of AC

In the **Model Builder** window, click **Effect of AC**.

Global 2

- 1 In the **Effect of AC** toolbar, click  **Global**.
- 2 In the **Settings** window for **Global**, locate the **Data** section.
- 3 From the **Dataset** list, choose **Study : AC Effect/Parametric Solutions 1 (sol4)**.
- 4 From the **Time selection** list, choose **Last**.

5 Locate the **y-Axis Data** section. In the table, enter the following settings:

| Expression | Unit | Description |
|---|------------------|--------------------------------------|
| $\text{timeint}(\text{tf}-1/\text{f},\text{tf},\text{ia})/(1/\text{f})$ | A/m ² | Time averaged anodic current density |

6 Locate the **x-Axis Data** section. From the **Axis source data** list, choose **Outer solutions**.

7 From the **Parameter** list, choose **Expression**.

8 In the **Expression** text field, type E_DC.

9 Locate the **Legends** section. From the **Legends** list, choose **Manual**.

10 In the table, enter the following settings:

| Legends |
|---------|
| AC |

11 In the **Effect of AC** toolbar, click  **Plot**.

The plot should look like [Figure 2](#).

ROOT

Next, add a new study to investigate the effect of the frequency on the corrosion using a Parametric Sweep.

ADD STUDY

1 In the **Home** toolbar, click  **Add Study** to open the **Add Study** window.

2 Go to the **Add Study** window.

3 Find the **Studies** subsection. In the **Select Study** tree, select **General Studies>Stationary**.

4 Click **Add Study** in the window toolbar.

5 In the **Home** toolbar, click  **Add Study** to close the **Add Study** window.

STUDY : FREQUENCY EFFECT

1 In the **Model Builder** window, click **Study 2**.

2 In the **Settings** window for **Study**, type Study : Frequency Effect in the **Label** text field.

3 Locate the **Study Settings** section. Clear the **Generate default plots** check box.

Time Dependent

In the **Study** toolbar, click  **Study Steps** and choose **Time Dependent>Time Dependent**.

Parametric Sweep

Use the parametric sweep to investigate the effect of two frequencies, 60[Hz] and 0.01[Hz].

- 1 In the **Study** toolbar, click  **Parametric Sweep**.
- 2 In the **Settings** window for **Parametric Sweep**, locate the **Study Settings** section.
- 3 Click  **Add**.
- 4 In the table, enter the following settings:

| Parameter name | Parameter value list | Parameter unit |
|-------------------------------|----------------------|----------------|
| f (Frequency of AC potential) | 60[Hz] 0.01 [Hz] | Hz |

Step 2: Time Dependent

- 1 In the **Model Builder** window, click **Step 2: Time Dependent**.
- 2 In the **Settings** window for **Time Dependent**, locate the **Study Settings** section.
- 3 In the **Output times** text field, type range (0,tf/10,tf).
- 4 In the **Study** toolbar, click  **Compute**.

RESULTS

Now, plot the total current density along with different contributions for different frequencies.

Effect of AC Frequency

- 1 In the **Home** toolbar, click  **Add Plot Group** and choose **ID Plot Group**.
- 2 In the **Settings** window for **ID Plot Group**, type Effect of AC Frequency in the **Label** text field.
- 3 Locate the **Data** section. From the **Dataset** list, choose **Study : Frequency Effect/ Parametric Solutions 2 (sol I)**.
- 4 From the **Parameter selection (f)** list, choose **From list**.
- 5 In the **Parameter values (f (Hz))** list, select **60**.
- 6 Locate the **Title** section. From the **Title type** list, choose **Manual**.
- 7 In the **Title** text area, type Current Density Versus Time.
- 8 Locate the **Plot Settings** section.
- 9 Select the **y-axis label** check box. In the associated text field, type Current density (A/m^2).

Point Graph: Anodic Current Density

- 1 In the **Effect of AC Frequency** toolbar, click  **Point Graph**.
- 2 In the **Settings** window for **Point Graph**, type Point Graph: Anodic Current Density in the **Label** text field.
- 3 Select Boundary 1 only.
- 4 Click **Replace Expression** in the upper-right corner of the **y-Axis Data** section. From the menu, choose **Component 1 (comp1)>Electroanalysis>Electrode kinetics>tcd.iloc_er1 - Local current density - A/m²**.
- 5 Click to expand the **Legends** section. Select the **Show legends** check box.
- 6 From the **Legends** list, choose **Manual**.
- 7 In the table, enter the following settings:

Legends

Anodic Current Density

Point Graph: Cathodic Current Density

- 1 Right-click **Point Graph: Anodic Current Density** and choose **Duplicate**.
- 2 In the **Settings** window for **Point Graph**, type Point Graph: Cathodic Current Density in the **Label** text field.
- 3 Locate the **y-Axis Data** section. In the **Expression** text field, type `tcd.iloc_er2+tcd.iloc_er3`.
- 4 Locate the **Legends** section. In the table, enter the following settings:

Legends

Cathodic Current Density

Point Graph: Double Layer Current Density

- 1 Right-click **Point Graph: Cathodic Current Density** and choose **Duplicate**.
- 2 In the **Settings** window for **Point Graph**, locate the **y-Axis Data** section.
- 3 In the **Expression** text field, type `tcd.id1`.
- 4 In the **Label** text field, type Point Graph: Double Layer Current Density.
- 5 Locate the **Legends** section. In the table, enter the following settings:

Legends

Double Layer Current Density

Point Graph: Total Current Density

- 1 Right-click **Point Graph: Double Layer Current Density** and choose **Duplicate**.
- 2 In the **Settings** window for **Point Graph**, click **Replace Expression** in the upper-right corner of the **y-Axis Data** section. From the menu, choose **Component 1 (comp1)> Electroanalysis>Electrode kinetics>tcd.itot - Total interface current density - A/m²**.
- 3 In the **Label** text field, type **Point Graph: Total Current Density**.
- 4 Locate the **Legends** section. In the table, enter the following settings:

Legends

Total Current Density

Effect of AC Frequency

- 1 In the **Model Builder** window, click **Effect of AC Frequency**.
- 2 In the **Effect of AC Frequency** toolbar, click  **Plot**.
The plot should look like [Figure 3](#).
Now plot current density contributions at 0.01 Hz.
- 3 In the **Settings** window for **ID Plot Group**, locate the **Data** section.
- 4 In the **Parameter values (f (Hz))** list, select **0.01**.
- 5 In the **Effect of AC Frequency** toolbar, click  **Plot**.
The plot should look like [Figure 4](#).