



Adsorption-Desorption Voltammetry

Introduction

For an electrochemical reaction to occur, the reacting species usually needs to adsorb to the electrode surface before undergoing reduction or oxidation, after which the resulting product species desorbs back into the electrolyte.

If the rate of adsorption or desorption is slow in comparison to the electrochemical charge transfer step, the adsorption-desorption phenomena may have to be accounted for in a model.

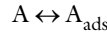
This example investigates the impact of various kinetic parameters for adsorption, desorption and electron transfer when performing cyclic voltammetry on a planar electrode.

The examples replicates the results of [Ref. 1](#).

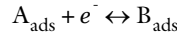
Model Definition

The model defines diffusion (by Fick's law) of an electrolyte species in a 1D geometry between $x = 0$ and $x = L$, and the local mass balances of two surface species at an electrode surface located at $x = 0$.

At the left-hand electrode boundary ($x = 0$) the electrolyte species A may adsorb according to



The adsorbed species A_{ads} may then undergo reduction to form B_{ads} in a charge transfer reaction according to



Langmuir isotherms are used for describing the kinetics, with the adsorption rate defined as

$$r_{\text{ads}} = k_{a1}c_A(1 - \Theta_{A_{\text{ads}}} - \Theta_{B_{\text{ads}}}) - k_{d1}\Theta_{A_{\text{ads}}}$$

where k_{a1} is the adsorption rate constant, c_A the electrolyte concentration of species A, $\Theta_{A_{\text{ads}}}$ the electrode surface coverage of species A_{ads} , $\Theta_{B_{\text{ads}}}$ the electrode surface coverage of species B_{ads} , and k_{d1} the desorption rate constant.

The charge transfer reaction is defined as

$$i_{\text{loc}} = k_0 \Gamma F \left(\Theta_{\text{B,ads}} e^{\frac{0.5 \eta F}{RT}} - \Theta_{\text{A,ads}} e^{-\frac{0.5 \eta F}{RT}} \right)$$

where k_0 is the charge transfer rate constant, Γ the density of surface sites at the electrode, F Faraday's constant, R the molar gas constant, and T the temperature.

The overpotential η is defined as

$$\eta = E - E_0$$

where E is the electrode potential and E_0 the formal potential.

The model is solved in a time-dependent simulation, ramping the potential from +0.5 V to -0.5 V and back, simulating a cyclic voltammogram.

When evaluating the voltammograms below, the total electrode current is defined as

$$I_{\text{tot}} = \pi r_d^2 i_{\text{loc}}$$

where r_d is the radius of the disk electrode.

The initial surface coverage of A_{ads} is set to $\Theta_{\text{A,ads}} = 0$, defining a situation where the cyclic voltammogram is recorded shortly after immersing the electrode in the electrolyte.

Two dimensionless parameters, K' and k_0' , are altered in a parametric sweep. They are defined as

$$K' = \frac{k_{a1}}{k_{d1} c_A}$$

and

$$k_0' = \frac{k_0 r_d^2}{D_A}$$

respectively. A high K' value hence represents a fast adsorption/slow desorption case, whereas a high k_0' value represents a case featuring fast charge transfer. The simulated cases are described in [Table 1](#).

TABLE 1: SIMULATION CASES IN THE PARAMETRIC SWEEP.

| | K' | k_0' | |
|--------|-----------|-----------|---------------------------------------|
| Case 1 | 10^5 | 10^2 | Fast adsorption, fast charge transfer |
| Case 2 | 10^{-5} | 10^2 | Slow adsorption, fast charge transfer |
| Case 3 | 10^5 | 10^{-2} | Fast adsorption, slow charge transfer |

Results and Discussion

Figure 1 shows the voltammograms for the three simulated cases. Case 1, with fast adsorption and fast kinetics results in a voltammogram fairly symmetric and centered around $E = 0$ V. The total integrated of the reduction current (that is, the negative peak) is significantly lower than the integrated oxidation current (the positive peak), this is a result of the adsorption-desorption reaction not being in equilibrium ($\Theta_{A_{ads}} = 0$) when the simulation is started.

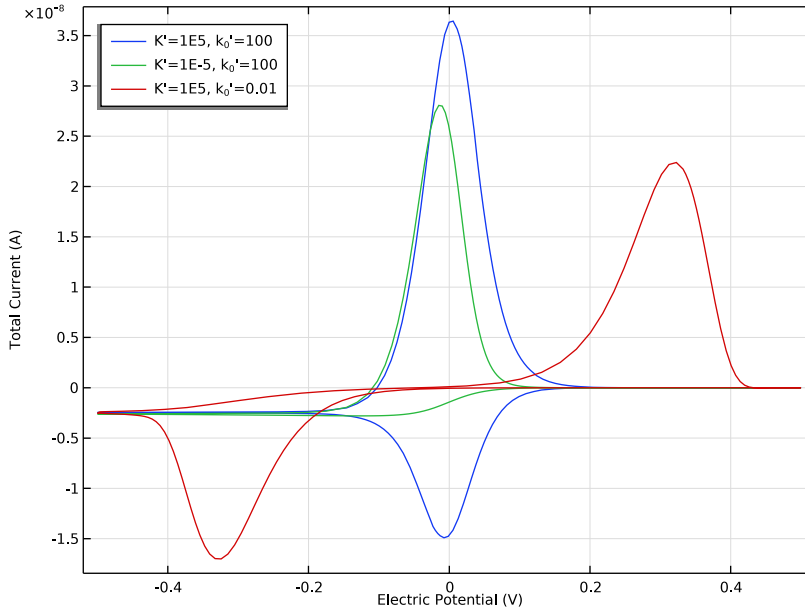


Figure 1: Cyclic voltammograms for the three investigated cases of 1) fast adsorption and charge transfer (blue), 2) slow adsorption and fast charge transfer (green), and 3) fast adsorption and slow charge transfer (red).

Case 2, with slower adsorption, features a limiting reduction current at $E < -0.1$ V, whereas the oxidation peak is fairly similar to that of Case 1.

Case 3, with slower charge transfer kinetics, features a more pronounced separation between the peaks.

More insights may be gained by inspecting the electrode surface coverages of A_{ads} and B_{ads} and the electrolyte concentration of species A at the electrode surface, as is shown in Figure 2 to Figure 4.

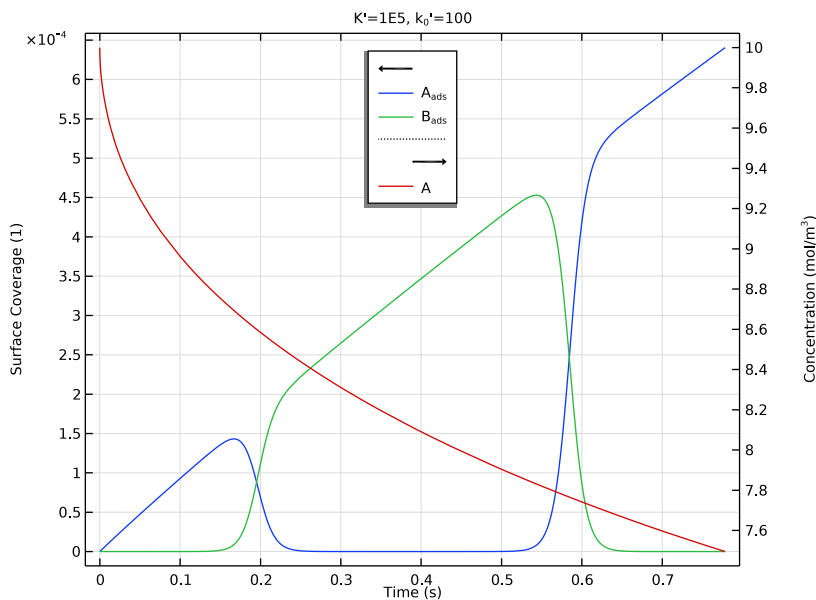


Figure 2: Surface coverages of species A_{ads} and B_{ads} and the concentration of A at the electrode surface for case 1) fast adsorption and fast charge transfer.

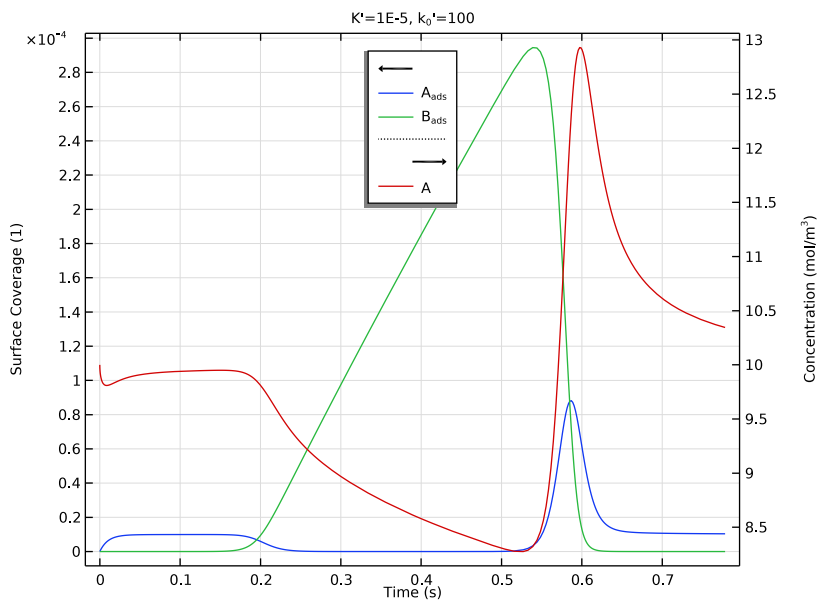


Figure 3: Surface coverages of species A_{ads} and B_{ads} and the concentration of A at the electrode surface for case 2) slow adsorption and fast charge transfer.

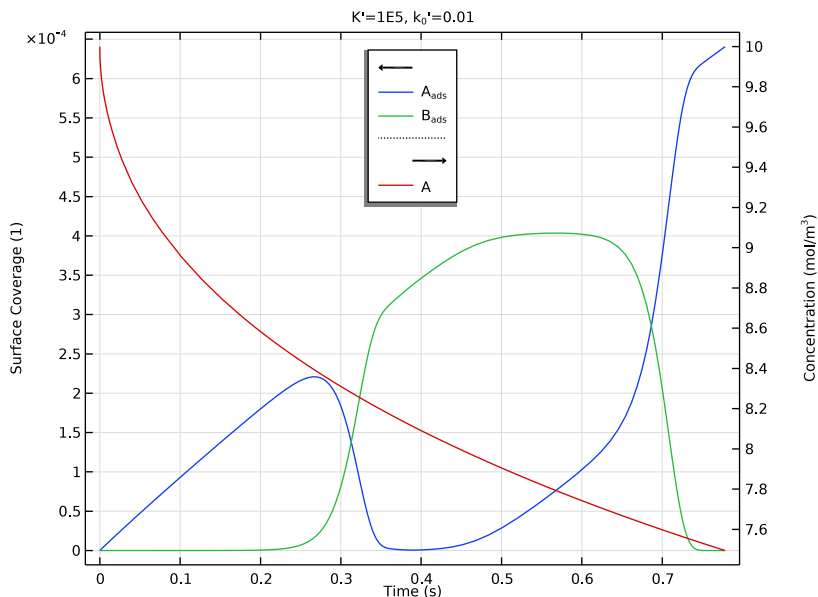


Figure 4: Surface coverages of species A_{ads} and B_{ads} and the concentration of A at the electrode surface for case 3) fast adsorption and slow charge transfer.

Reference


1. F. Chevallier, O. Klymenko, L. Jiang, T. Jones, and R. Compton, "Mathematical modelling and numerical simulation of adsorption processes at microdisk electrodes," *J. Electroanal. Chem.*, vol. 574, pp. 217–237, 2005.

Application Library path: Electrochemistry_Module/Electroanalysis/
adsorption_desorption_voltammetry


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**.

This model will model the transport of one species only (species A) in the electrolyte.


- 4 In the **Number of species** text field, type 1.
- 5 In the **Concentrations (mol/m³)** table, enter the following settings:

| c_A |
|------------|
| |

- 6 Click  **Study**.
- 7 In the **Select Study** tree, select **Preset Studies for Selected Physics Interfaces>Cyclic Voltammetry**.
- 8 Click  **Done**.

GLOBAL DEFINITIONS

Parameters I

- 1 In the **Model Builder** window, under **Global Definitions** click **Parameters I**.
- 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 `adsorption_desorption_voltammetry_parameters.txt`.

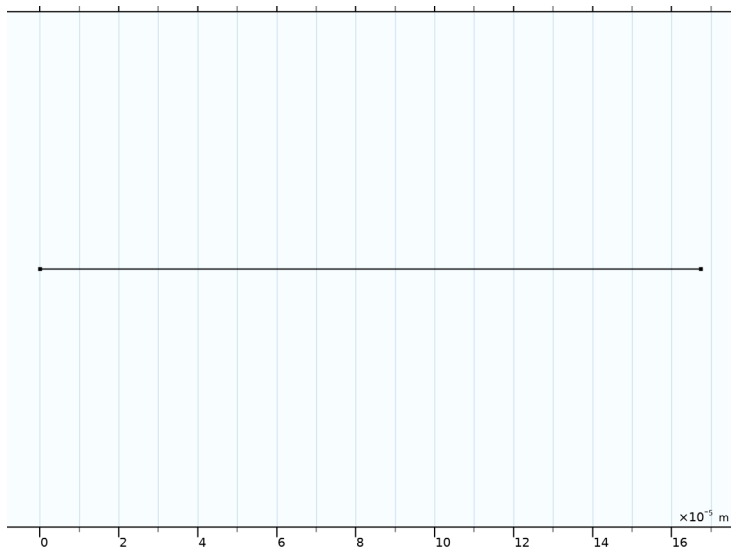
GEOMETRY I

Interval I (il)

- 1 In the **Model Builder** window, under **Component I (comp1)** right-click **Geometry I** 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**.



ELECTROANALYSIS (TCD)

- 1 In the **Model Builder** window, under **Component 1 (comp1)** click **Electroanalysis (tcd)**.
- 2 In the **Settings** window for **Electroanalysis**, locate the **Cross-Sectional Area** section.
- 3 In the A_c text field, type $A_{\text{electrode}}$.


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_{cA} text field, type D_A .

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 c_A text field, type c_{A_bulk} .

Electrode Surface 1

- 1 In the **Physics** toolbar, click  **Boundaries** and choose **Electrode Surface**.
- 2 Select Boundary 1 only.

3 In the **Settings** window for **Electrode Surface**, click to expand the **Adsorbing-Desorbing Species** section.

4 In the Γ_s text field, type Gamma.

5 Click **+ Add**.

At the electrode surface, we will model the surface coverages of two species (species A_{ads} and B_{ads}).

6 In the table, enter the following settings:

| Species | Site occupancy number (l) |
|------------------|---------------------------|
| A _{ads} | 1 |

7 Click **+ Add**.

8 In the table, enter the following settings:

| Species | Site occupancy number (l) |
|------------------|---------------------------|
| B _{ads} | 1 |

9 Locate the **Electrode Phase Potential Condition** section. From the **Electrode phase potential condition** list, choose **Cyclic voltammetry**.

10 In the **Linear sweep rate** text field, type nu.

11 In the **Vertex potential 1** text field, type E_{start}.

12 In the **Vertex potential 2** text field, type E_{vertex}.

Electrode Reaction 1

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

2 In the **Settings** window for **Electrode Reaction**, locate the **Model Input** section.

3 From the *T* list, choose **User defined**. In the associated text field, type T.

In the one-electron electrode charge transfer reaction, A_{ads} gets reduced to form B_{ads}. The electrolyte species A does not participate in the reaction.

4 Locate the **Stoichiometric Coefficients** section. In the **Stoichiometric coefficients for adsorbing-desorbing species** table, enter the following settings:

| Species | Stoichiometric coefficient (l) |
|------------------|--------------------------------|
| A _{ads} | - 1 |
| B _{ads} | 1 |

5 Locate the **Equilibrium Potential** section. From the *E_{eq}* list, choose **User defined**. In the associated text field, type E₀.

6 Locate the **Electrode Kinetics** section. From the **Kinetics expression type** list, choose **Concentration dependent kinetics**.

7 In the i_0 text field, type `k0*F_const*Gamma`.

8 In the C_R text field, type `tcd.theta_es1_B_ads`.

9 In the C_O text field, type `tcd.theta_es1_A_ads`.

`tcd.theta_es1_A_ads` and `tcd.theta_es1_B_ads` are the variable names for the surface coverage of A_ads and B_ads, respectively.

DEFINITIONS

Now define the adsorption-desorption rate and the reaction, involving species A and A_ads.

Variables I

1 In the **Model Builder** window, under **Component I (comp1)** right-click **Definitions** and choose **Variables**.

Define a variable expression for the adsorption rate.

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

3 In the table, enter the following settings:

| Name | Expression | Unit | Description |
|-------|---|-------------------------|-----------------|
| r_ads | <code>ka1* tcd.thetafree_es1* c_A-kd1* tcd.theta_es1_A_ads</code> | mol/(m ² ·s) | Adsorption rate |

In the expression above `tcd.thetafree_es1` is the surface fraction of free sites and `c_A` is the electrolyte concentration of species A. The values of the rate constants `ka1` and `kd1` are specified in the **Parameters** node by the text file you imported earlier.

ELECTROANALYSIS (TCD)

Electrode Surface I

In the **Model Builder** window, under **Component I (comp1)**>**Electroanalysis (tcd)** click **Electrode Surface I**.

Nonfaradaic Reactions I

1 In the **Physics** toolbar, click  **Attributes** and choose **Nonfaradaic Reactions**.

2 In the **Settings** window for **Nonfaradaic Reactions**, locate the **Reaction Rate** section.

3 Select the **Species c_A** check box.


- 4 In the $R_{0,cA}$ text field, type -r_ads.
- 5 In the **Reaction rate for adsorbing-desorbing species** table, enter the following settings:

| Species | Reaction rate (mol/(m ² *s)) |
|---------|---|
| A_ads | r_ads |

Electrode Surface 1

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

Initial Values for Adsorbing-Desorbing Species 1

- 1 In the **Physics** toolbar, click  **Attributes** and choose **Initial Values for Adsorbing-Desorbing Species**.
- 2 In the **Settings** window for **Initial Values for Adsorbing-Desorbing Species**, locate the **Initial Values for Adsorbing-Desorbing Species** section.
- 3 In the table, enter the following settings:

| Species | Surface coverage (1) |
|---------|----------------------|
| A_ads | theta_A_init |

The value of theta_A_init is 0, but may be changed later in **Parameters**.

MESH 1

Edit the mesh to get a very finely resolved mesh close to the electrode surface.

- 1 In the **Model Builder** window, under **Component 1 (comp1)** right-click **Mesh 1** and choose **Edit Physics-Induced Sequence**.

Size

- 1 In the **Model Builder** window, under **Component 1 (comp1)>Mesh 1** click **Size**.
- 2 In the **Settings** window for **Size**, locate the **Element Size** section.
- 3 Click the **Custom** button.
- 4 Locate the **Element Size Parameters** section. In the **Maximum element size** text field, type L/10.
- 5 In the **Maximum element growth rate** text field, type 1.1.

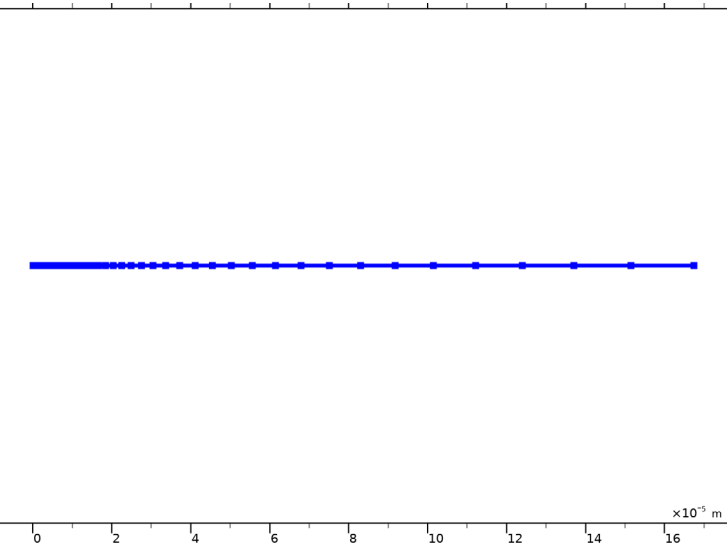
Size 1

- 1 In the **Model Builder** window, click **Size 1**.
- 2 In the **Settings** window for **Size**, locate the **Element Size** section.
- 3 Click the **Custom** button.

- 4 Locate the **Element Size Parameters** section.
- 5 Select the **Maximum element size** check box. In the associated text field, type $L/10000$.

Edge 1



In the **Model Builder** window, right-click **Edge 1** and choose **Build All**.



STUDY 1

The model is now ready for solving. Add a parametric sweep to solve for three different sets of parameters.

Parametric Sweep

- 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 |
|---|----------------------|----------------|
| K_prime (Dimensionless adsorption-desorption rate constant ratio) | 1e5 1e-5 1e5 | |

K_{prime} is a dimensionless parameter representing the ratio of the adsorption vs the desorption rate.

5 Click  **Add**.

6 In the table, enter the following settings:

| Parameter name | Parameter value list | Parameter unit |
|---|----------------------|----------------|
| k_0_prime (Dimensionless electron transfer rate constant) | 1e2 1e2 1e-2 | |

k_0_prime is a dimensionless parameter representing the charge transfer rate constant.

With the above settings we will now compute and compare: 1) a base case, 2) a reaction limited by adsorption and 3) a case limited by slow charge transfer.

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

RESULTS

Global I

1 In the **Model Builder** window, expand the **Cyclic Voltammograms (tcd)** node, then click **Global I**.

2 In the **Settings** window for **Global**, click to expand the **Legends** section.

Polish the legend as follows:

3 From the **Legends** list, choose **Evaluated**.

4 In the **Legend** text field, type $K' = \text{eval}(K_prime)$, $k_{0,0} = \text{eval}(k_0_prime)$.

Cyclic Voltammograms (tcd)

1 In the **Model Builder** window, click **Cyclic Voltammograms (tcd)**.

2 In the **Settings** window for **ID Plot Group**, locate the **Legend** section.

3 From the **Position** list, choose **Upper left**.

4 In the **Cyclic Voltammograms (tcd)** toolbar, click  **Plot**.

Compare the plot with [Figure 1](#).

Surface Coverages and Concentration

Plot the surface coverages of A_ads and B_ads, as well as the electrolyte concentration of A, at the electrode surface for the three cases as follows:

1 In the **Home** toolbar, click  **Add Plot Group** and choose **ID Plot Group**.


2 In the **Settings** window for **ID Plot Group**, type Surface Coverages and Concentration in the **Label** text field.

- 3 Locate the **Data** section. From the **Dataset** list, choose **Study 1/ Parametric Solutions 1 (sol2)**.
- 4 From the **Parameter selection (K_prime, k_0_prime)** list, choose **From list**.
- 5 In the **Parameter values (K_prime,k_0_prime)** list, select **1: K_prime=1E5, k_0_prime=100**.

Point Graph 1

- 1 Right-click **Surface Coverages and Concentration** and choose **Point Graph**.
- 2 Select Boundary 1 only.
- 3 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>Adsorbing-desorbing species>Surface coverage of adsorbing-desorbing species>tcd.theta_esI_A_ads - Surface coverage of adsorbing-desorbing species, 1-component**.
- 4 Click to expand the **Legends** section. Select the **Show legends** check box.
- 5 From the **Legends** list, choose **Manual**.
- 6 In the table, enter the following settings:


| Legends |
|-----------------|
| A_{ads} |

- 7 In the **Surface Coverages and Concentration** toolbar, click  **Plot**.
- 8 Right-click **Point Graph 1** and choose **Duplicate**.

Point Graph 2

- 1 In the **Model Builder** window, click **Point Graph 2**.
- 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>Adsorbing-desorbing species>Surface coverage of adsorbing-desorbing species>tcd.theta_esI_B_ads - Surface coverage of adsorbing-desorbing species, 2-component**.
- 3 Locate the **Legends** section. In the table, enter the following settings:

| Legends |
|-----------------|
| B_{ads} |

- 4 In the **Surface Coverages and Concentration** toolbar, click  **Plot**.
- 5 Right-click **Point Graph 2** and choose **Duplicate**.




Point Graph 3

- 1 In the **Model Builder** window, click **Point Graph 3**.
- 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>Species c_A>c_A - Concentration - mol/m³**.
- 3 Locate the **Legends** section. In the table, enter the following settings:

| Legends |
|---------|
| A |

Surface Coverages and Concentration

Polish the title, y-axis settings and legend position as follows:

- 1 In the **Model Builder** window, click **Surface Coverages and Concentration**.
- 2 In the **Settings** window for **ID Plot Group**, click to expand the **Title** section.
- 3 From the **Title type** list, choose **Manual**.
- 4 In the **Title** text area, type $K' = \text{eval}(K_prime)$, $k_{0'} = \text{eval}(k_0_prime)$.
- 5 Locate the **Plot Settings** section.
- 6 Select the **y-axis label** check box. In the associated text field, type Surface Coverage (1).
- 7 Select the **Two y-axes** check box.
- 8 In the table, select the **Plot on secondary y-axis** check box for **Point Graph 3**.
- 9 Locate the **Legend** section. From the **Position** list, choose **Upper middle**.
- 10 In the **Surface Coverages and Concentration** toolbar, click  **Plot**.
Compare the plot with [Figure 2](#).
- 11 Locate the **Data** section. In the **Parameter values (K_prime,k_0_prime)** list, select **2: K_prime=1E-5, k_0_prime=100**.
- 12 In the **Surface Coverages and Concentration** toolbar, click  **Plot**.
Compare the plot with [Figure 3](#).
- 13 In the **Parameter values (K_prime,k_0_prime)** list, select **3: K_prime=1E5, k_0_prime=0.01**.
- 14 In the **Surface Coverages and Concentration** toolbar, click  **Plot**.
Compare the plot with [Figure 4](#).