

Copper Deposition in a Trench Using the Phase Field Method

This model example is based on the Copper Deposition in a Trench Using the Level Set Method model, available in the Electrodeposition Module Application Library, which demonstrates that there is a nonuniform deposition along the trench surface, leading to formation of a cavity or void (Ref. 1). The model example presented here uses the Phase Field interface instead of the Level Set interface and computations are performed for up to 20 s, well beyond the time for cavity formation, similar to the Level Set model.

Model Definition

The deposition process is inherently time-dependent because the cathode boundary moves as the deposition process takes place. The model is defined by the material balances for the involved ions (copper, Cu^{2+} , and sulfate, SO_4^{2-}) and by the electroneutrality condition.

The model geometry is shown in Figure 1. The upper horizontal boundary represents the anode, while the cathode is placed at the bottom. The vertical walls are assumed to be insulating.

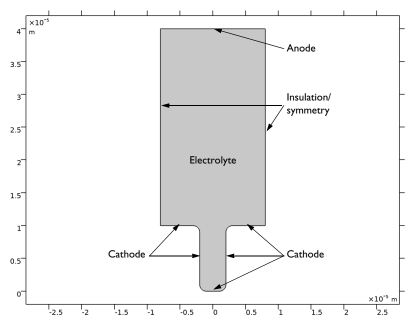


Figure 1: Model domain with boundaries corresponding to the anode, cathode, and vertical symmetry walls.

When using the Phase Field method, both the electrode and the electrolyte are described on the same domain. The Phase Field interface is used to keep track of the deformation at the cathode surface during deposition. For simplicity, the anode position is kept fixed in this model.

In the Phase Field interface the two-phase flow dynamics is governed by a Cahn-Hilliard equation. The equation tracks a diffuse interface separating the immiscible phases. The diffuse interface is defined as the region where the dimensionless phase field variable ϕ goes from -1, in the electrolyte domain, to 1, in the deposited region. to 1. When solved in COMSOL Multiphysics, the Cahn-Hilliard equation is split up into two equations

$$\frac{\partial \Phi}{\partial t} + \mathbf{u} \cdot \nabla \Phi = \nabla \cdot \frac{\gamma \lambda}{\varepsilon^2} \nabla \Psi$$

$$\psi = -\nabla \cdot \varepsilon^2 \nabla \phi + (\phi^2 - 1)\phi$$

where ${\bf u}$ is the fluid velocity (m/s), γ is the mobility (m³·s/kg), λ is the mixing energy density (N) and ϵ (m) is the interface thickness parameter. The ψ variable is referred to as the phase field help variable. The following equation relates the mixing energy density and the interface thickness to the surface tension coefficient:

$$\sigma = \frac{2\sqrt{2}}{3}\frac{\lambda}{\varepsilon}$$

In the present model, the interface thickness parameter is set to $\varepsilon = h_{\rm max}/16$, where $h_{\rm max}$ is the maximum mesh element size in the domain. The mobility parameter γ determines the time scale of the Cahn–Hilliard diffusion and must be chosen judiciously. It must be large enough to retain a constant interfacial thickness but small enough so that the convective terms are not overly damped. A suitable value for γ is the maximum velocity magnitude occurring in the model.

In the Phase Field interface, the volume fractions of the individual fluids are

$$V_{\rm f1} = \frac{1 - \phi}{2}, \qquad V_{\rm f2} = \frac{1 + \phi}{2}$$

The volume fraction, V_{f1} , varies from 1 in the electrolyte domain to 0 in the deposited region.

The phase field delta function is approximated by:

$$\delta = \frac{3}{4}(1 - \phi^2)|\nabla\phi|$$

The velocity field used in the transport equation for phase field variable is evaluated from the copper deposition reaction current density:

$$\mathbf{u} = \mathbf{n} \cdot \left(-\frac{i_{\text{loc}} M_{\text{Cu}}}{2F} \rho_{\text{Cu}} \right)$$

where i_{loc} is the local current density, M_{Cu} is the molar mass and ρ_{Cu} is the density of copper.

The interface normal \mathbf{n} is calculated as

$$\mathbf{n} = \frac{\nabla \phi}{|\nabla \phi|}$$

The phase field variable of value 1 enters the domain from the bottommost boundaries of the domain, which are prescribed using the Inlet boundary condition. The rest of the boundaries are prescribed using the Outlet boundary condition.

The usage of the Phase Field method means that the balance equations have to be modified in comparison with those used in the Copper Deposition in a Trench model. First, the flux for each of the ions in the electrolyte is given by the Nernst–Planck equation with effective diffusion coefficients and mobilities,

$$\mathbf{N}_i = -D_{i,\,\mathrm{eff}} \nabla c_i - z_i u_{i,\,\mathrm{eff}} F c_i \nabla \phi_l,$$

where \mathbf{N}_i denotes the transport vector (mol/(m²·s)), c_i the concentration in the electrolyte (mol/m³), z_i the charge for the ionic species, $u_{i,\text{eff}}$ the mobility of the charged species (m²/(s·J·mole)), F Faraday's constant (As/mole), and ϕ_l the potential in the electrolyte (V). The effective diffusion coefficients are defined using the electrolyte volume fraction:

$$D_{i, eff} = \varepsilon_l D_i$$

where ε_l is the electrolyte volume fraction; ε_l is defined in terms of the phase field volume fraction, V_{f1} , and varies from 1 in the electrolyte domain to 0 in the deposited region.

Furthermore, the material balances are expressed through

$$\frac{\partial \varepsilon_l c_i}{\partial t} + \nabla \cdot \mathbf{N}_i = R_i$$

one for each species, that is i = 1, 2.

The rate of electrochemical reaction is

$$R_i = -\frac{v_i i_v}{nF}$$

The electrolyte volume fraction compensation ensures that the electrode will not act as a reservoir for the copper and sulfate ions.

The electroneutrality condition is given by the following expression:

$$\sum_{i} z_i c_i = 0$$

The deposition process is assumed to take place through the following simplified mechanism:

$$Cu^{2+} + e^{-} = Cu^{+}$$

$$Cu^+ + e^- = Cu$$

where the first step is rate determining step, RDS, and the second step is assumed to be at equilibrium (Ref. 1). This gives the following the Butler–Volmer equation for the local current density as a function of potential and copper concentration:

$$i_{\rm loc} = i_0 \left(\exp \left(\frac{1.5F\eta}{RT} \right) - \frac{c_{\rm Cu^{2+}}}{c_{\rm Cu^{2+}} \operatorname{ref}} \exp \left(-\frac{0.5F\eta}{RT} \right) \right)$$

where η denotes the overpotential defined as

$$\eta = \phi_{\rm s} - \phi_{\rm l} - E_{\rm eq}$$

where ϕ_8 denotes the electronic potential of the respective electrode.

The copper deposition reaction current density at the cathode surface is added as a source term in the domain, using the phase field delta function to prescribe the deposition reaction along the deforming boundary:

$$i_v = i_{loc} \delta$$

The boundary condition at the anode surface is:

$$\mathbf{N}_{\mathrm{Cu}^{2+}} \cdot \mathbf{n} = -\frac{i_{\mathrm{loc}}}{2F}$$

where **n** denotes the normal vector to the boundary. For simplicity, the anode surface is not deformed in this model.

All other boundaries are insulating:

$$\mathbf{N}_{\mathbf{C}\mathbf{n}^{2+}} \cdot \mathbf{n} = 0$$

For the sulfate ions, insulating conditions apply everywhere:

$$\mathbf{N}_{\mathrm{SO}_4^{2-}} \cdot \mathbf{n} = 0$$

The initial conditions set the composition of the electrolyte according to

$$c_{\mathrm{Cu}^{2+}} = c_0$$
$$c_{\mathrm{SO}^{2-}_{+}} = c_0$$

The electrochemical model described above is set up using the Tertiary Current Distribution, Nernst–Planck Equations interface.

Results and Discussion

Figure 2 shows surface plots of the electrolyte potential and the phase field variable after 20 s of deposition operation. The surface plot of the phase field variable, plotted using a gray scale, represents the copper deposition region where its value is greater than 0. It can be seen that there is a nonuniform deposition along the trench (cathode) surface. The deposition rate is higher near mouth of the trench than at the bottom of the trench, leading to formation of a cavity or void. The isolated cavity can be detrimental to the quality of the deposition because a trapped electrolyte can later cause corrosion of components in the circuit board.

The electrolyte potential in the cavity relaxes so that the electrode potential $(\phi_s - \phi_l)$ corresponds to the resulting equilibrium potential $(E_{
m eq})$ for the concentration in the cavity, leading the local current density value to 0 inside the cavity.

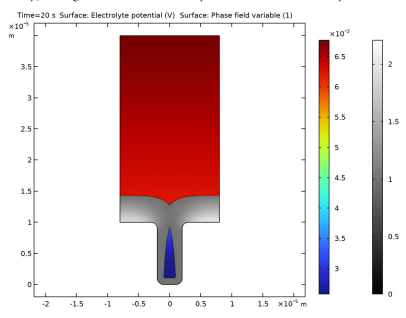


Figure 2: Surface plot of electrolyte potential along with the phase field variable contour of value 0.5 after 20 seconds of deposition operation.

Figure 3 shows the surface plots of the corresponding concentration of copper ions and the phase field variable after 20 s of deposition operation. The simulation shows substantial variations in copper ion concentration in the cell. Such variations eventually cause free convection in the cell. The nonuniform deposition rate at the cathode surface is attributed

to nonuniform electrolyte current distribution which is accentuated by the depletion of copper ions at the bottom of the trench.

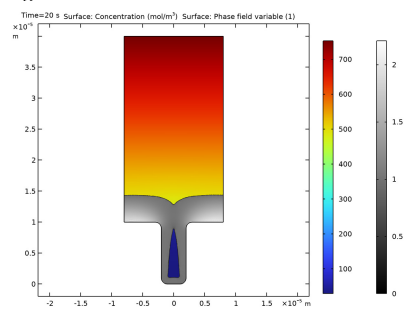


Figure 3: Surface plot of concentration of copper ions along with the phase field variable contour of value 0.5 after 20 seconds of deposition operation.

In Figure 4 and Figure 5 we now compare the results obtained from the Phase Field interface in this model with those obtained from the Deformed Geometry node in the

Copper Deposition in a Trench model, after 14 s of deposition operation (which corresponds to the time close to the cavity formation)

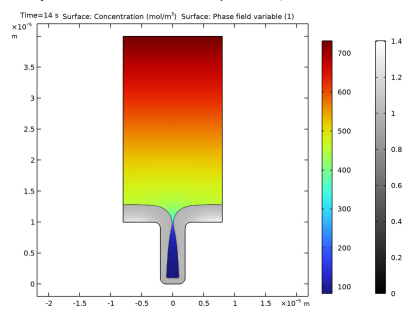


Figure 4: Surface plot of concentration of copper ions along with the phase field variable contour of value 0.5 after 14 seconds of deposition operation using the Phase Field interface.

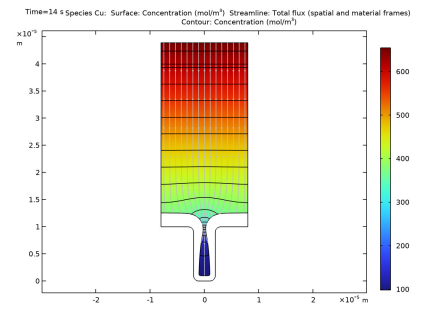


Figure 5: Surface plot of the concentration distribution of copper ions, the isopotential lines, the current density lines, and the displacement of the cathode and anode surfaces after 14 seconds of deposition operation using the Deformed Geometry node.

It can be seen from Figure 4 and Figure 5 that the deposition profile and the concentration distribution of copper ions obtained from the Phase Field interface match well with the same obtained from the Deformed Geometry node. Some minor differences can be seen in the plots with regards to concentrations and position of the moving cathode. These can mainly be attributed to the anode position, which is allowed to move in Figure 5, but is fixed in Figure 4.

In conclusion, electrodeposition models using the phase field method can be useful in identifying cavity or void regions formed during a deposition process in complex geometries.

Reference

1. E. Mattsson and J.O'M. Bockris, "Galvanostatic Studies of the Kinetics of Deposition and Dissolution in the Copper + Copper Sulphate System", *Trans. Far. Soc.*, vol. 55, p. 1586, 1959.

Application Library path: Electrodeposition_Module/Tutorials/ cu_trench_deposition_pf

Modeling Instructions

From the File menu, choose New.

NEW

In the New window, click Model Wizard.

MODEL WIZARD

- I In the Model Wizard window, click **2** 2D.
- 2 In the Select Physics tree, select Electrochemistry>Tertiary Current Distribution, Nernst-Planck>Tertiary, Electroneutrality (tcd).
- 3 Click Add.
- 4 In the Concentrations (mol/m³) table, enter the following settings:

cCu cS04

- 5 In the Select Physics tree, select Mathematics>Moving Interface>Phase Field in Fluids (pf).
- 6 Click Add.
- 7 Click 🕣 Study.
- 8 In the Select Study tree, select Preset Studies for Some Physics Interfaces>
 Time Dependent with Initialization.
- 9 Click M Done.

GEOMETRY I

Draw the geometry by making a union of two rectangles. Round off the corners of the trench using fillets.

Rectangle I (rI)

- I In the Geometry toolbar, click Rectangle.
- 2 In the Settings window for Rectangle, locate the Size and Shape section.
- 3 In the Width text field, type 1.6e-5.

- 4 In the Height text field, type 3e-5.
- 5 Locate the Position section. In the x text field, type -0.8e-5.
- 6 In the y text field, type 1e-5.

Rectangle 2 (r2)

- I In the Geometry toolbar, click Rectangle.
- 2 In the Settings window for Rectangle, locate the Size and Shape section.
- 3 In the Width text field, type 0.4e-5.
- 4 In the Height text field, type 1e-5.
- 5 Locate the Position section. In the x text field, type -0.2e-5.
- 6 Click **Build Selected**.
- 7 Click the **Zoom Extents** button in the **Graphics** toolbar.

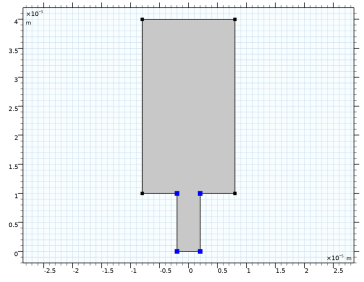
Union I (uni I)

- I In the Geometry toolbar, click Booleans and Partitions and choose Union.
- 2 In the Settings window for Union, locate the Union section.
- 3 Clear the Keep interior boundaries check box.
- 4 Click in the **Graphics** window and then press Ctrl+A to select both objects.

Fillet I (fill)

I In the Geometry toolbar, click / Fillet.

2 On the object unil, select Points 3–6 only.



- 3 In the Settings window for Fillet, locate the Radius section.
- 4 In the Radius text field, type 1e-6.

Form Union (fin)

- I In the Model Builder window, click Form Union (fin).
- 2 In the Settings window for Form Union/Assembly, click 📳 Build Selected.

GLOBAL DEFINITIONS

Load the model parameters from a text file.

Parameters 1

- I 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 cu_trench_deposition_ls_parameters.txt.

DEFINITIONS

Load the model variables from a text file.

Variables 1

- I In the Home toolbar, click \supseteq 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 cu_trench_deposition_pf_variables.txt.

TERTIARY CURRENT DISTRIBUTION, NERNST-PLANCK (TCD)

Now set up the electrochemical model, consisting of a Highly Conductive Porous Electrode domain and an electrode boundary.

Species Charges 1

- I In the Model Builder window, under Component I (compl)>Tertiary Current Distribution,
 Nernst-Planck (tcd) click Species Charges I.
- 2 In the Settings window for Species Charges, locate the Charge section.
- 3 In the z_{cCu} text field, type z_Cu.
- **4** In the z_{cSO4} text field, type z_S04.

Highly Conductive Porous Electrode I

- I In the Physics toolbar, click Domains and choose Highly Conductive Porous Electrode.
- 2 In the Settings window for Highly Conductive Porous Electrode, locate the Domain Selection section.
- 3 From the Selection list, choose All domains.
- **4** Locate the **Diffusion** section. In the $D_{\rm eCu}$ text field, type D_Cu.
- 5 In the D_{cSO4} text field, type D_SO4.
 Set the electrolyte volume fraction to eps1. This implies that all the volume of the modeled domain belongs to the electrolyte phase.
- 6 Locate the Porous Matrix Properties section. In the ε₁ text field, type eps1.
 Set the effective transport parameter correction for diffusion to user defined and set the correction factor to eps1 so that the correction is applied only in the electrolyte phase excluding the deposited region.
- 7 Locate the Effective Transport Parameter Correction section. From the Diffusion list, choose User defined. In the f_{Dl} text field, type eps1.
- 8 Locate the Electrode Phase Potential Condition section. In the $\phi_{s,ext}$ text field, type phis cathode.

Porous Electrode Reaction I

Set up the copper deposition electrode reaction kinetics parameters.

- I In the Model Builder window, click Porous Electrode Reaction I.
- 2 In the Settings window for Porous Electrode Reaction, locate the Stoichiometric Coefficients section.
- **3** In the *n* text field, type 2.
- 4 Locate the Equilibrium Potential section. From the E_{eq} list, choose User defined. In the associated text field, type Eeq_rel.
- 5 Locate the Electrode Kinetics section. From the Kinetics expression type list, choose Concentration dependent kinetics.
- **6** In the i_0 text field, type i0.
- 7 In the α_a text field, type alpha_a.
- **8** In the α_c text field, type alpha_c.
- **9** In the $C_{\rm O}$ text field, type cCu/Cinit.
- 10 Locate the Active Specific Surface Area section. In the $a_{
 m v}$ text field, type pf.delta.

Electrode Surface 1

Define the electrode kinetics at the anode surface.

- I In the Physics toolbar, click Boundaries and choose Electrode Surface.
- 2 Select Boundary 3 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 phis_anode.
- **5** Click to expand the **Dissolving-Depositing Species** section. Click + **Add**.
- **6** In the table, enter the following settings:

Species	Density (kg/m^3)	Molar mass (kg/mol)
cdep_anode	8960	0.06355

Electrode Reaction 1

- I In the Model Builder window, click Electrode Reaction I.
- 2 In the Settings window for Electrode Reaction, locate the Stoichiometric Coefficients section.
- 3 In the n text field, type 2.

- **4** Locate the **Electrode Kinetics** section. In the $i_{0,\mathrm{ref}}(T)$ text field, type i0.
- **5** In the α_a text field, type alpha_a.

Initial Values 1

Set the initial concentration of copper and sulfate ions.

- I In the Model Builder window, under Component I (compl)>Tertiary Current Distribution, Nernst-Planck (tcd) click Initial Values 1.
- 2 In the Settings window for Initial Values, locate the Initial Values section.
- **3** In the cSO4 text field, type Cinit.

PHASE FIELD IN FLUIDS (PF)

Now, set up the Phase Field model.

- I In the Model Builder window, under Component I (compl) click Phase Field in Fluids (pf).
- 2 In the Settings window for Phase Field in Fluids, click to expand the Discretization section.
- 3 From the Element order list, choose Quadratic.

Phase Field Model I

Next, specify the parameter controlling interface thickness, mobility tuning parameter, and velocity field.

- I In the Model Builder window, under Component I (compl)>Phase Field in Fluids (pf) click
 Phase Field Model I.
- 2 In the Settings window for Phase Field Model, locate the Phase Field Parameters section.
- 3 In the ε_{pf} text field, type pf.hmax/16.
- 4 From the Mobility tuning parameter list, choose Calculate from velocity.
- 5 In the U text field, type max(Vn, eps).
- **6** Locate the **Convection** section. Specify the **u** vector as

-Vn*pf.intnormx	x
-Vn*pf.intnormy	у

Inlet I

Define inlet and outlet boundaries for the phase field variable.

- I In the Physics toolbar, click Boundaries and choose Inlet.
- 2 Select Boundaries 2, 4–7, and 9–12 only.
- 3 In the Settings window for Inlet, locate the Phase Field Condition section.

4 From the list, choose Fluid 2 ($\varphi = 1$).

Outlet 1

- I In the Physics toolbar, click Boundaries and choose Outlet.
- 2 Select Boundaries 1, 3, and 8 only.

GLOBAL DEFINITIONS

Default Model Inputs

Set up the temperature value used in the entire model.

- I In the Model Builder window, under Global Definitions click Default Model Inputs.
- 2 In the Settings window for Default Model Inputs, locate the Browse Model Inputs section.
- 3 In the tree, select General>Temperature (K) minput.T.
- **4** Find the **Expression for remaining selection** subsection. In the **Temperature** text field, type T0.

MESH I

Now make the settings for the mesh.

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

Size

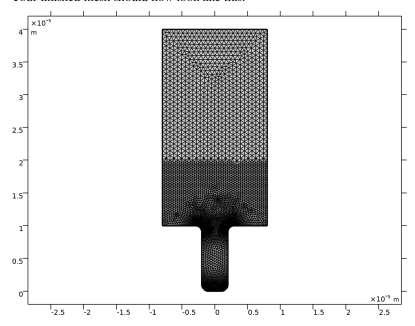
- I In the Model Builder window, under Component I (compl)>Mesh I click Size.
- 2 In the Settings window for Size, locate the Element Size section.
- 3 From the Predefined list, choose Normal.

Refine I

- I In the Mesh toolbar, click Modify and choose Refine.
- 2 In the Settings window for Refine, click to expand the Refine Elements in Box section.
- 3 Select the Specify bounding box check box.
- 4 In row x, set Lower bound to -1E-5.
- 5 In row x, set Upper bound to 1E-5.
- 6 In row y, set Lower bound to -8E-7.
- 7 In row y, set Upper bound to 2E-5.

8 Click Build All.

Your finished mesh should now look like this:



STUDY I

Modify the solver settings to simulate the deposition process during 20 s, storing the solution every 0.5 s. Use the initial valued based scaling and clear generate default plots option and then start the computation.

Step 2: Time Dependent

- I In the Model Builder window, under Study I 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 (0,0.5,20).

Solution I (soll)

- I In the Study toolbar, click Show Default Solver.
- 2 In the Model Builder window, expand the Solution I (soll) node, then click Dependent Variables 2.
- 3 In the Settings window for Dependent Variables, locate the Scaling section.
- 4 From the Method list, choose Initial value based.
- 5 In the Model Builder window, click Study 1.

- 6 In the Settings window for Study, locate the Study Settings section.
- 7 Clear the Generate default plots check box.
- 8 In the Study toolbar, click **Compute**.

RESULTS

The following steps reproduce the plots from the Results and Discussion section.

Electrolyte Potential (tcd)

- I In the Home toolbar, click **Add Plot Group** and choose **2D Plot Group**.
- 2 In the Settings window for 2D Plot Group, type E in the Label text field.
- 3 In the Label text field, type Electrolyte Potential (tcd).

Surface I

In the Electrolyte Potential (tcd) toolbar, click Surface.

Filter I

- I In the Electrolyte Potential (tcd) toolbar, click **Filter**.
- 2 In the Settings window for Filter, locate the Element Selection section.
- 3 In the Logical expression for inclusion text field, type phipf<0.

Surface I

In the Model Builder window, right-click Surface I and choose Duplicate.

Surface 2

- I In the Model Builder window, click Surface 2.
- 2 In the Settings window for Surface, locate the Expression section.
- **3** In the **Expression** text field, type phipf.
- 4 Locate the Coloring and Style section. Click Change Color Table.
- 5 In the Color Table dialog box, select Linear>GrayScale in the tree.
- 6 Click OK.

Filter I

- I In the Model Builder window, expand the Surface 2 node, then click Filter I.
- 2 In the Settings window for Filter, locate the Element Selection section.
- 3 In the Logical expression for inclusion text field, type phipf>0.

Electrolyte Potential (tcd)

Now, plot the concentration of copper ions by making use of the Duplicate functionality.

I In the Model Builder window, under Results right-click Electrolyte Potential (tcd) and choose Duplicate.

Concentration (tcd)

- I In the Model Builder window, expand the Results>Electrolyte Potential (tcd) I node, then click Electrolyte Potential (tcd) I.
- 2 In the Settings window for 2D Plot Group, type Concentration (tcd) in the Label text field.

Surface 1

- I In the Model Builder window, click Surface I.
- 2 In the Settings window for Surface, click Replace Expression in the upper-right corner of the Expression section. From the menu, choose Component I (compl)> Tertiary Current Distribution, Nernst-Planck>Species cCu>cCu Concentration mol/m³.
- 3 In the Concentration (tcd) toolbar, click Plot.

Concentration (tcd)

Now, plot the concentration of copper ions at 14 s to compare the results with the deformed geometry formulation.

- I In the Model Builder window, click Concentration (tcd).
- 2 In the Settings window for 2D Plot Group, locate the Data section.
- 3 From the Time (s) list, choose 14.
- 4 In the Concentration (tcd) toolbar, click Plot.