

Microwave Cavity Plasma Reactor

This tutorial shows how to prepare a model of a hydrogen plasma sustained in a microwave cavity at moderate pressures. The model is based on the work in Ref. 1 and solves the plasma transport equations fully coupled with Maxwell's equations, fluid flow, and heat transfer. A microwave cylindrical chamber contains a bell jar were a hydrogen plasma is created. The reactor is carefully designed so that the electric field has its maximum intensity above a substrate and much lower intensity at the bell jar boundary. This is important in order for the plasma to keep its maximum density in the substrate region even at high power.

Note: The model requires the Plasma Module and the RF Module.

Model Definition

The electron density and mean electron energy are computed by solving a pair of driftdiffusion equations for the electron density and mean electron energy. For detailed information on electron transport, see Theory for the Drift Diffusion Interface in the Plasma Module User's Guide.

$$\frac{\partial}{\partial t}(n_e) + \nabla \cdot \left[-n_e(\boldsymbol{\mu}_e \bullet \mathbf{E}) - \mathbf{D}_e \bullet \nabla n_e \right] = R_e$$

$$\frac{\partial}{\partial t}(n_{\varepsilon}) + \nabla \cdot [-n_{\varepsilon}(\boldsymbol{\mu}_{\varepsilon} \bullet \mathbf{E}) - \mathbf{D}_{\varepsilon} \bullet \nabla n_{\varepsilon}] + \mathbf{E} \cdot \boldsymbol{\Gamma}_{e} \, = \, \boldsymbol{R}_{\varepsilon}$$

The electron source R_e and the energy loss due to inelastic collisions R_{ϵ} are defined later. The electron diffusivity, energy mobility, and energy diffusivity are computed from the electron mobility using:

$$\mathbf{D}_{e} = \mu_{e} T_{e}, \, \mu_{\varepsilon} = \left(\frac{5}{3}\right) \mu_{e}, \, \mathbf{D}_{\varepsilon} = \mu_{\varepsilon} T_{e}$$

The source coefficients in the above equations are determined by the plasma chemistry using rate coefficients. In the case of rate coefficients, the electron source term is given by:

$$R_e = \sum_{j=1}^{M} x_j k_j N_n n_e$$

where x_j is the mole fraction of the target species for reaction j, k_j is the rate coefficient for reaction j (SI unit: m^3/s), and N_n is the total neutral number density (SI unit: $1/m^3$). The electron energy loss is obtained by summing the collisional energy loss over all reactions:

$$R_{\varepsilon} = \sum_{j=1}^{P} x_{j} k_{j} N_{n} n_{e} \Delta \varepsilon_{j}$$

where $\Delta \varepsilon_j$ is the energy loss from reaction j (SI unit: V). The rate coefficients can be computed from cross section data by the following integral:

$$k_k = \gamma \int_0^\infty \varepsilon \sigma_k(\varepsilon) f(\varepsilon) d\varepsilon$$

where $\gamma = (2q/m_e)^{1/2}$ (SI unit: $C^{1/2}/kg^{1/2}$), m_e is the electron mass (SI unit: kg), ε is energy (SI unit: V), σ_k is the collision cross section (SI unit: m²), and f is the electron energy distribution function.

For nonelectron species, the following equation is solved for the mass fraction of each species. For detailed information on the transport of the nonelectron species, see *Theory for the Heavy Species Transport Interface* in the *Plasma Module User's Guide*.

$$\rho \frac{\partial}{\partial t}(w_k) + \rho(\mathbf{u} \cdot \nabla)w_k = \nabla \cdot \mathbf{j}_k + R_k$$

The electrostatic field is computed using the following equation:

$$-\nabla \cdot \varepsilon_0 \varepsilon_r \nabla V = \rho$$

The space charge density ρ is automatically computed based on the plasma chemistry specified in the model using the formula:

$$\rho = q \left(\sum_{k=1}^{N} Z_k n_k - n_e \right)$$

For detailed information about electrostatics see *Theory for the Electrostatics Interface* in the *Plasma Module User's Guide*.

In a microwave reactor, the high-frequency electric field is computed in the frequency domain through the equation

$$\nabla \times (\mu_{\mathbf{r}}^{-1} \nabla \times \mathbf{E}) - k_0^2 \left(\varepsilon_{\mathbf{r}} - \frac{j\sigma}{\omega \varepsilon_0} \right) \mathbf{E} = 0$$

The relationship between the plasma current density and the electric field becomes more complicated in the presence of a DC magnetic field. The following equation defines this relationship:

$$\sigma^{-1} \bullet J = \mathbf{E}$$

Here, σ is the plasma conductivity.

The fluid flow and heat transfer in the fluid are also solved for. The gas temperature plays an important role in the reactor operation since it can significantly influence the reduced electric field E/N.

PLASMA CHEMISTRY

The plasma chemistry is based on the simplified chemistry presented in table 4 of Ref. 1 and is presented in Table 1. The electron impact reactions are from Ref. 2 and retrieved from Ref. 3, and the rates are from Ref. 4 and Ref. 5. The model includes eight species: electrons, H₂, H, H₂⁺, H⁺, H₃⁺, and two excited states of hydrogen corresponding to the levels n = 2 and n = 3 that are represented by Hn2 and Hn3.

TABLE I: MODELED COLLISIONS AND REACTIONS.

Reaction	Formula	Туре	$\Delta\epsilon(eV)$	
1	e+H2=>e+H2	Elastic	-	
2	e+H2=>e+H2	Vibrational excitation	0.516-1.5	
3	e+H2=>e+2H	Dissociation	7.93-11.72 and17.22- 17.53	
4	e+H2=>e+H2	Excitation	12.4-14.6	
5	e+H2=>e+H+Hn2	Dissociative excitation	14.68	
6	e+H2=>e+H+Hn3	Dissociative excitation	16.57	
7	e+H2=>2e+H2+	Ionization	15.4	
8	e+H2=>2e+H+H+	Ionization	19	
9	e+H=>e+H	Elastic	-	
10	e+H=>e+Hn2	Excitation	10.2043	
11	e+H=>e+Hn3	Charge transfer	12.1	
12	e+H=>e+H	Excitation	12.755 and 13.0615	

TABLE I: MODELED COLLISIONS AND REACTIONS.

Reaction	Formula	Туре	$\Delta\epsilon(eV)$
13	e+H=>2e+H+	Ionization	13.6057
14	e+H3+=>3H	Recombination	0
15	e+H2+=>H+Hn2	Recombination	0.01
16	e+H2+=>H+Hn3	Recombination	0.01
17	e+H+=>Hn2	Recombination	0
18	e+H+=>Hn3	Recombination	0
19	Hn2+H2=>H3++e	Ionization	-
20	Hn3+H2=>H3++e	Ionization	-
21	H2+H2+=>H3++H	Ionization	-
22	H2+H2=>2H+H2	Dissociation	-
23	2H+H2=>H2+H2	Association	-
24	H2+H=>3H	Dissociation	-
26	3H=>H2+H	Association	-

In addition to the volume reactions, the surface reaction listed in Table 2.

TABLE 2: SURFACE REACTIONS.

Reaction	Formula	Sticking coefficient
1	H=>0.5H2	0.02
2	Hn2=>H	I
3	Hn3=>H	I
4	H+=>H	I
5	H2+=>H2	I
6	H3+=>H2+H	I

Results and Discussion

The figures in this section present model results for a hydrogen plasma sustained at 25 kPa with 5 kW. In general, the results agree well with results presented in figure 21 and figure 22 of Ref. 1. A detailed comparison is not attempted because many aspects of the models are different. The plasma chemistry is not exactly the same. Even if the reactions are mostly the same, the rates and cross sections were obtained independently. The reactor dimensions and configurations are only approximate. Additionally, there are differences in the model equations.

Using a global model fully coupled with a Boltzmann equation in the two-term approximation, the study performed in the model Hydrogen Global Model Coupled with the Two-Term Boltzmann Equation — also in the Plasma Module Application Library showed that for main quantities like electron density, H-atom density, and gas temperature there is not a significant difference when using a computed or a Maxwellian EEDF. This justified doing the space-dependent simulations using a Maxwellian EEDF. However, in this first attempt the results were wrong in many ways. An important aspect not captured was the plasma localization at the subtract. With a Maxwellian EEDF it is very easy to ionize in regions of low electron energy and the maximum of the plasma easily moves somewhere else. Further investigations with the global model showed significant differences in results for low mean electron energies. The EEDF was than adjusted to use a generalized distribution with a power law of 1.2.

The reactor dimensions were adjusted to have high electric field intensity at the substract and to have a minimum in reflected power without plasma. This part of the investigation is done with the Electromagnetic Waves, Frequency Domain interface only and it is not shown here. With the plasma present, the electromagnetic characteristics of the reactor change considerably. This is possible to observe in Figure 5 and Figure 6, which show the S-parameter and Smith plot when increasing the power from 500 to 5000 W. Having a high-dense plasma in a considerable part of the reactor strongly increases the reflected power.

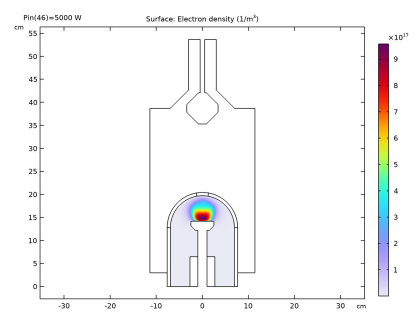


Figure 1: Electron density for a hydrogen plasma operating at $25\,\mathrm{kPa}$ with $5\,\mathrm{kW}$ of input power.

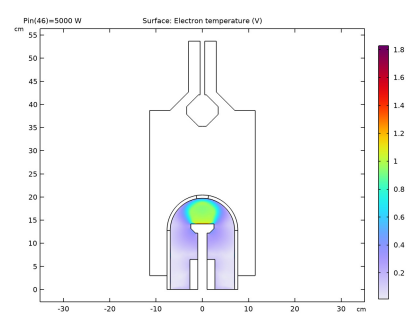


Figure 2: Electron temperature for a hydrogen plasma operating at 25 kPa with 5 kW of inputpower.

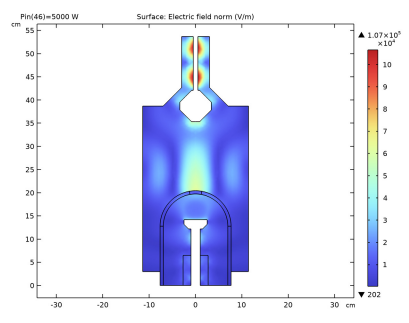


Figure 3: RF electric field norm for a hydrogen plasma operating at 25 kPa with 5 kW of input power.

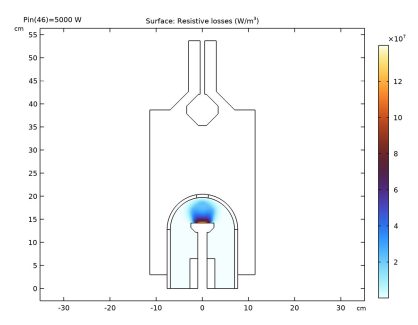


Figure 4: RF power absorbed by electrons for a hydrogen plasma operating at 25 kPa with 5 kW of input power.

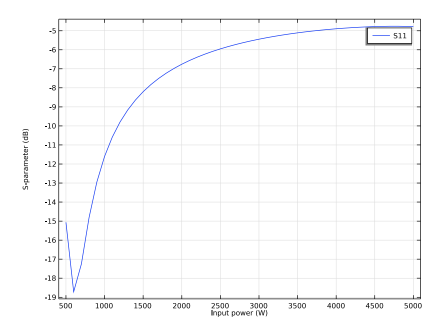


Figure 5: S-parameter as a function of the input power.

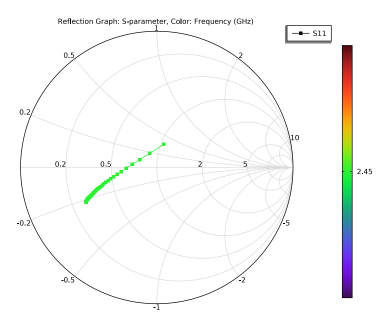


Figure 6: Smith plot as a function of the input power. For higher powers the curve moves away from the center.

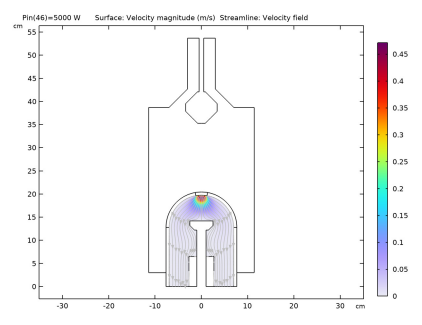


Figure 7: Fluid velocity for a hydrogen plasma operating at 25 kPa with 5 kW of input power.

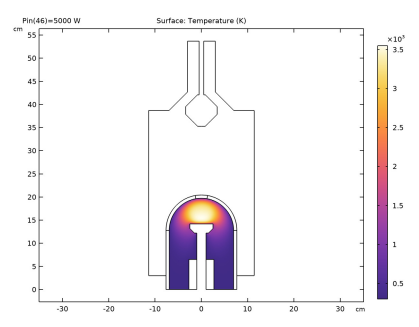


Figure 8: Gas temperature for a hydrogen plasma operating at 25 kPa with 5 kW of input power.

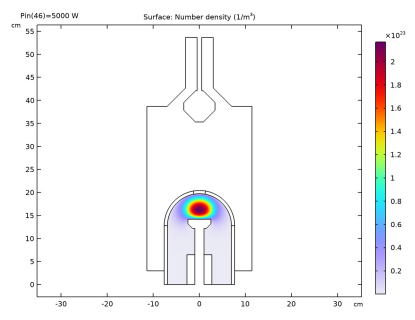


Figure 9: Number density of atomic hydrogen for a hydrogen plasma operating at 25~kPa with 5~kW of input power.

References

- 1. K. Hassouni, F. Silva, and A. Gicquel, "Modelling of diamond deposition microwave cavity generated plasmas," *J. Phys. D: Appl. Phys.*, vol. 43, p. 153001, 2010.
- 2. L. Marques, J. Jolly, and L.L. Alves, "Capacitively Coupled Radio-Frequency Hydrogen Discharges: The Role of Kinetics," *J. Appl. Phys.*, vol. 102, p. 063305, 2007.
- 3. IST-Lisbon database, www.lxcat.net, retrieved 2023.
- 4. M. Capitelli, C.M. Ferreira, B.F. Gordiets and A.I. Osipov, *Plasma Kinetics in Atmospheric Gases*, Springer, 2000.
- 5. R.K. Janev; W.D. Langer; K. Evans, Jr.; and D.E. Post, Jr., *Elementary Processes in Hydrogen-Helium Plasmas*, Springer-Verlag, 1987.

Application Library path: Plasma Module/Wave-Heated Discharges/ microwave_cavity_plasma

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 2D Axisymmetric.
- 2 In the Select Physics tree, select Plasma>Microwave Plasma.
- 3 Click Add.
- 4 Click 🔵 Study.
- 5 In the Select Study tree, select Preset Studies for Selected Multiphysics>Frequency-Transient.
- 6 Click M Done.

Import a file with parameters to be used in the model. Most of them are used to create the reactor geometry.

GLOBAL DEFINITIONS

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 microwave cavity plasma parameters.txt.

Create a profile to be used as initial conditions for the gas temperature.

DEFINITIONS

Variables 1

- I In the Model Builder window, under Component I (compl) right-click Definitions and choose Variables.
- 2 In the Settings window for Variables, locate the Variables section.
- **3** In the table, enter the following settings:

Name	Expression	Unit	Description
r0	0[cm]	m	Maximum r-coordinate
z0	14[cm]	m	Maximum z-coordinate
prf	exp(-((r0-r)/4[cm])^2 - ((z0-z)/2[cm])^2)		Initial temperature profile
Tinit	2500[K]*prf+1200[K]	K	Initial temperature

Create the geometry of the reactor. Some geometry elements are to be used later when creating the mesh.

GEOMETRY I

- I In the Model Builder window, under Component I (compl) click Geometry I.
- 2 In the Settings window for Geometry, locate the Units section.
- 3 From the Length unit list, choose cm.

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 R0.
- 4 In the Height text field, type H0.

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 Slw.
- 4 In the Height text field, type S1h.

Rectangle 3 (r3)

- 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 Shw.
- 4 In the Height text field, type Shh.
- 5 Locate the **Position** section. In the **z** text field, type S1h.

Rectangle 4 (r4)

- 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 Shw.
- 4 In the Height text field, type 2[mm].
- **5** Locate the **Position** section. In the **z** text field, type S1h+Shh.

Rectangle 5 (r5)

- 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 Bhw-Bth/2.
- 4 In the **Height** text field, type Bhh.
- 5 Locate the **Position** section. In the r text field, type RO-Bhw+Bth/2.

Circular Arc I (cal)

- I In the Geometry toolbar, click * More Primitives and choose Circular Arc.
- 2 In the Settings window for Circular Arc, locate the Center section.
- 3 In the z text field, type Bz.
- 4 Locate the Radius section. In the Radius text field, type Br.

Line Segment I (Is I)

- I In the Geometry toolbar, click * More Primitives and choose Line Segment.
- 2 In the Settings window for Line Segment, locate the Starting Point section.
- 3 From the Specify list, choose Coordinates.
- 4 In the r text field, type Br.
- 5 In the z text field, type Bz.
- **6** Locate the **Endpoint** section. From the **Specify** list, choose **Coordinates**.
- 7 In the r text field, type RO-Bhw.

Circular Arc 2 (ca2)

- I In the Geometry toolbar, click * More Primitives and choose Circular Arc.
- 2 In the Settings window for Circular Arc, locate the Center section.

- 3 In the z text field, type Bz.
- 4 Locate the Radius section. In the Radius text field, type Br.
- 5 Locate the Angles section. In the Start angle text field, type 80.
- **6** Click the **Zoom Extents** button in the **Graphics** toolbar.
- 7 Click **Build All Objects**.

Thicken I (thi I)

- I In the Geometry toolbar, click Conversions and choose Thicken.
- 2 Select the objects cal, ca2, and Isl only.
- 3 In the Settings window for Thicken, locate the Options section.
- 4 In the Total thickness text field, type Bth.

Rectangle 6 (r6)

- 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 Cxw.
- 4 In the Height text field, type Cxh.
- 5 Locate the Position section. In the z text field, type H0.

Polygon I (poll)

- I In the Geometry toolbar, click / Polygon.
- 2 In the Settings window for Polygon, locate the Coordinates section.
- **3** In the table, enter the following settings:

r (cm)	z (cm)		
Cxw	H0+Th		
Cxw+Tw	НО		
Cxw	НО		

Union I (uni I)

- I In the Geometry toolbar, click Booleans and Partitions and choose Union.
- 2 Select the objects poll, rl, and r6 only.
- 3 In the Settings window for Union, locate the Union section.
- 4 Clear the Keep interior boundaries check box.
- 5 Click Build All Objects.

Rectangle 7 (r7)

- 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 Dw.
- 4 In the Height text field, type Dw.
- 5 Locate the Position section. From the Base list, choose Center.
- 6 In the z text field, type H0.
- 7 Locate the Rotation Angle section. In the Rotation text field, type 45.

Chamfer I (chal)

- I In the Geometry toolbar, click Chamfer.
- 2 Click the **Zoom Extents** button in the **Graphics** toolbar.
- 3 On the object r7, select Points 1-4 only.
- 4 In the Settings window for Chamfer, locate the Distance section.
- 5 In the Distance from vertex text field, type Dcut.

Rectangle 8 (r8)

- 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 Cxiw.
- 4 In the Height text field, type Cxh.
- **5** Locate the **Position** section. In the **z** text field, type H0.

Difference | (dif1)

- I In the Geometry toolbar, click Booleans and Partitions and choose Difference.
- **2** Select the object **unil** only.
- 3 In the Settings window for Difference, locate the Difference section.
- 4 Click to select the Activate Selection toggle button for Objects to subtract.
- 5 Select the objects chal, r2, r3, r5, and r8 only.

Circular Arc 3 (ca3)

- I In the Geometry toolbar, click * More Primitives and choose Circular Arc.
- 2 In the Settings window for Circular Arc, locate the Center section.
- 3 In the z text field, type Shh+Slh.
- 4 Locate the Radius section. In the Radius text field, type 3[cm].

Line Segment 2 (Is2)

- I In the Geometry toolbar, click More Primitives and choose Line Segment.
- 2 In the Settings window for Line Segment, locate the Starting Point section.
- 3 Click to select the Activate Selection toggle button for Start vertex.
- 4 On the object ca3, select Point 1 only.
- 5 Locate the **Endpoint** section. Click to select the **Activate Selection** toggle button for **End vertex**.
- 6 On the object r4, select Point 2 only.

Line Segment 3 (Is3)

- I In the Geometry toolbar, click * More Primitives and choose Line Segment.
- 2 On the object Is2, select Point 1 only.
- 3 In the Settings window for Line Segment, locate the Endpoint section.
- 4 Click to select the Activate Selection toggle button for End vertex.
- **5** On the object **thil(3)**, select Point 2 only.

Chamfer 2 (cha2)

- I In the Geometry toolbar, click Chamfer.
- 2 On the object difl, select Point 9 only.
- 3 In the Settings window for Chamfer, locate the Distance section.
- 4 In the Distance from vertex text field, type 1 [cm].

Rectangle 9 (r9)

- 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 dielw.
- 4 In the Height text field, type dielh.
- **5** Locate the **Position** section. In the **r** text field, type S1w.

Mesh Control Edges I (mcel)

- I In the Geometry toolbar, click \times Virtual Operations and choose Mesh Control Edges.
- 2 On the object fin, select Boundaries 4, 22, 23, 26, and 38 only.
- 3 In the Geometry toolbar, click **Build All**.
- 4 Click the **Zoom Extents** button in the **Graphics** toolbar.

THE PLASMA CHEMISTRY ADD-IN

The next steps have instructions to first import the Plasma Chemistry add-in and then to use this add-in to import a file that automatically creates the hydrogen plasma chemistry.

The following is set or created automatically:

- a Species properties using Preset species data
- **b** Electron impact reactions for argon and oxygen
- c Heavy species reactions
- **d** Surface reactions

The documentation accompanying the Plasma Chemistry add-in contains more information about the file structure and what can be set automatically.

In the **Home** toolbar, click **Windows** and choose **Add-in Libraries**.

ADD-IN LIBRARIES

- I In the Add-in Libraries window, select Plasma Module>plasma_chemistry in the tree.
- 2 In the tree, select the check box for the node Plasma Module>plasma_chemistry (if it is not already selected).
- **3** Click **Done** to load the add-in and close the **Add-in Libraries** window.
- 4 In the Developer toolbar, click Add-ins and choose Plasma Chemistry> Plasma Chemistry.

GLOBAL DEFINITIONS

Plasma Chemistry I

- I In the Model Builder window, under Global Definitions click Plasma Chemistry I.
- 2 In the Settings window for Plasma Chemistry, locate the Plasma Chemistry Import section.
- 3 Click Browse.
- **4** Browse to the model's Application Libraries folder and double-click the file H2 plasma_chemistry.txt.
- 5 Click Import.

DEFINITIONS

In the Model Builder window, collapse the Component I (compl)>Definitions node.

GEOMETRY I

In the Model Builder window, collapse the Component I (compl)>Geometry I node.

Define material properties for air and quartz. The plasma conductivity is computed in the **Plasma Conductivity Coupling** multiphysics feature and overrides its definition set in the material properties.

ADD MATERIAL

- I In the Home toolbar, click **Add Material** to open the Add Material window.
- 2 Go to the Add Material window.
- 3 In the tree, select Built-in>Air.
- 4 Click Add to Component in the window toolbar.
- 5 In the tree, select Built-in>Glass (quartz).
- **6** Click **Add to Component** in the window toolbar.
- 7 In the Home toolbar, click **Add Material** to close the Add Material window.

MATERIALS

Glass (quartz) (mat2) Select Domains 2 and 4–6 only.

MATERIALS

In the Model Builder window, collapse the Component I (compl)>Materials node.

Set some properties of the plasma model and group the plasma chemistry elements.

PLASMA (PLAS)

- I In the Model Builder window, under Component I (compl) click Plasma (plas).
- 2 In the Settings window for Plasma, locate the Transport Settings section.
- 3 Find the Include subsection. Select the Calculate thermodynamic properties check box.
- 4 Select the Full expression for diffusivity check box.
- **5** Select the **Mixture diffusion correction** check box.
- **6** Select the **Convection** check box.
- 7 Locate the Plasma Properties section. Select the Use reduced electron transport properties check box.
- 8 Locate the Electron Energy Distribution Function Settings section. From the Electron energy distribution function list, choose Generalized.
- **9** In the g text field, type 1.2.
- 10 Locate the Domain Selection section. Click Clear Selection.

II Select Domain 1 only.

10: e+H2=>e+H2, 11: e+H2=>e+H2, 12: e+H2=>e+H2, 13: e+H2=>e+H2, 14: e+ H2=>e+H2, 15: e+H2=>e+H2, 16: e+H2=>e+H+Hn2, 17: e+H2=>e+H+Hn2, 18: e+H2=>e+H+Hn3, 19: e+H2=>e+H+H, 1: e+H2=>e+H2, 20: e+H2=>e+H+H, 21: e+H2=>2e+H2+, 22: e+H2=>2e+H+H+, 23: e+H=>e+H, 24: e+H=>e+Hn2, 25: e+ H=>e+Hn2, 26: e+H=>e+Hn3, 27: e+H=>e+H, 28: e+H=>e+H, 29: e+H=>2e+H+ . 2: e+H2=>e+H2, 30: e+H3+=>3H, 31: e+H3+=>H2+Hn2, 32: e+H2+=>H+Hn2, 33: e+H2+=>H+Hn3. 34: e+H+=>Hn2. 35: e+H+=>Hn3. 3: e+H2=>e+H2. 4: e+ H2=>e+H2, 5: e+H2=>e+H+H, 6: e+H2=>e+H+H, 7: e+H2=>e+H+H, 8: e+H2=> e+H+H. 9: e+H2=>e+H2

- I In the Model Builder window, under Component I (compl)>Plasma (plas), Ctrl-click to select 1: e+H2=>e+H2, 2: e+H2=>e+H2, 3: e+H2=>e+H2, 4: e+H2=>e+H2, 5: e+H2=>e+ H+H, 6: e+H2=>e+H+H, 7: e+H2=>e+H+H, 8: e+H2=>e+H+H, 9: e+H2=>e+H2, 10: e+ H2=>e+H2, I1: e+H2=>e+H2, I2: e+H2=>e+H2, I3: e+H2=>e+H2, I4: e+H2=>e+H2, 15: e+H2=>e+H2, 16: e+H2=>e+H+Hn2, 17: e+H2=>e+H+Hn2, 18: e+H2=>e+H+Hn3, 19: e+H2=>e+H+H, 20: e+H2=>e+H+H, 21: e+H2=>2e+H2+, 22: e+H2=>2e+H+H+, 23: e+H=>e+H, 24: e+H=>e+Hn2, 25: e+H=>e+Hn2, 26: e+H=>e+Hn3, 27: e+H=>e+H, 28: e+H=>e+H, 29: e+H=>2e+H+, 30: e+H3+=>3H, 31: e+H3+=>H2+Hn2, 32: e+H2+=> H+Hn2, 33: e+H2+=>H+Hn3, 34: e+H+=>Hn2, and 35: e+H+=>Hn3.
- **2** Right-click and choose **Group**.

Electron impact reactions

- I In the Settings window for Group, type Electron impact reactions in the Label text field.
- **2** In the **Model Builder** window, collapse the **Electron impact reactions** node.

Species: H, Species: H+, Species: H2+, Species: H3+, Species: Hn2, Species: Hn3, Species: e

- I In the Model Builder window, under Component I (compl)>Plasma (plas), Ctrl-click to select Species: e, Species: H2, Species: H, Species: Hn2, Species: Hn3, Species: H2+, Species: H+, and Species: H3+.
- 2 Right-click and choose **Group**.

Species

Set initial conditions for the heavy species.

I In the Settings window for Group, type Species in the Label text field.

Species: H2

- I In the Model Builder window, click Species: H2.
- 2 In the Settings window for Species, locate the Species Formula section.
- 3 Select the From mass constraint check box.

Species: H

- I In the Model Builder window, click Species: H.
- 2 In the Settings window for Species, locate the General Parameters section.
- **3** In the x_0 text field, type 1E-5.

Species: Hn3

- I In the Model Builder window, click Species: Hn3.
- 2 In the Settings window for Species, locate the General Parameters section.
- 3 In the x_0 text field, type 1E-10.

Species: H2+

- I In the Model Builder window, click Species: H2+.
- 2 In the Settings window for Species, locate the General Parameters section.
- **3** In the n_0 text field, type 1E7[1/m³].

Species: H+

- I In the Model Builder window, click Species: H+.
- 2 In the Settings window for Species, locate the General Parameters section.
- **3** In the n_0 text field, type 1E7[1/m³].

Species: H3+

- I In the Model Builder window, click Species: H3+.
- 2 In the Settings window for Species, locate the Species Formula section.
- 3 Select the Initial value from electroneutrality constraint check box.

PLASMA (PLAS)

Species

In the Model Builder window, collapse the Component I (compl)>Plasma (plas)>Species node.

- I: H=>0.5H2, 2: Hn2=>H, 3: Hn3=>H, 4: H+=>H, 5: H2+=>H2, 6: H3+=>H2+H
- I In the Model Builder window, under Component I (compl)>Plasma (plas), Ctrl-click to select I: H=>0.5H2, 2: Hn2=>H, 3: Hn3=>H, 4: H+=>H, 5: H2+=>H2, and 6: H3+=>H2+H.

2 Right-click and choose **Group**.

Surface reactions

In the Settings window for Group, type Surface reactions in the Label text field.

Set boundaries for surface reactions.

DEFINITIONS

Walls

- I In the **Definitions** toolbar, click **\(\bigcap_{\bigcap} \) Explicit**.
- 2 Select Domain 1 only.
- 3 In the Settings window for Explicit, locate the Output Entities section.
- 4 From the Output entities list, choose Adjacent boundaries.
- 5 In the Label text field, type Walls.

Walls neutrals

- I In the **Definitions** toolbar, click **\(\bigcap_{\text{a}} \) Explicit**.
- 2 In the Settings window for Explicit, type Walls neutrals in the Label text field.
- 3 Locate the Input Entities section. From the Geometric entity level list, choose Boundary.
- **4** Select Boundaries 2, 13–15, 17–19, 24, 32, and 34 only.

PLASMA (PLAS)

- 1: H=>0.5H2
- I In the Model Builder window, under Component I (compl)>Plasma (plas)> Surface reactions click 1: H=>0.5H2.
- 2 In the Settings window for Surface Reaction, locate the Boundary Selection section.
- 3 From the Selection list, choose Walls neutrals.
- 2: Hn2=>H
- I In the Model Builder window, click 2: Hn2=>H.
- 2 In the Settings window for Surface Reaction, locate the Boundary Selection section.
- 3 From the Selection list, choose Walls neutrals.
- 3: Hn3=>H
- I In the Model Builder window, click 3: Hn3=>H.
- 2 In the Settings window for Surface Reaction, locate the Boundary Selection section.
- 3 From the Selection list, choose Walls neutrals.

4: H+=>H

- I In the Model Builder window, click 4: H+=>H.
- 2 In the Settings window for Surface Reaction, locate the Boundary Selection section.
- 3 From the Selection list, choose Walls.

5: H2+=>H2

- I In the Model Builder window, click 5: H2+=>H2.
- 2 In the Settings window for Surface Reaction, locate the Boundary Selection section.
- 3 From the Selection list, choose Walls.

6: H3+=>H2+H

- I In the Model Builder window, click 6: H3+=>H2+H.
- 2 In the Settings window for Surface Reaction, locate the Boundary Selection section.
- 3 From the Selection list, choose Walls.

PLASMA (PLAS)

Surface reactions

In the Model Builder window, collapse the Component I (compl)>Plasma (plas)> Surface reactions node.

Species

Set an Outflow boundary condition for neutrals only. Ions are assumed to neutralized at the outlet.

Species: H

In the Model Builder window, expand the Component I (compl)>Plasma (plas)>Species node, then click Species: H.

Outflow I

- I In the **Physics** toolbar, click **Attributes** and choose **Outflow**.
- 2 In the Settings window for Outflow, locate the Boundary Selection section.
- 3 Click Clear Selection.
- 4 Select Boundary 20 only.

Species: Hn2

In the Model Builder window, under Component I (compl)>Plasma (plas)>Species click Species: Hn2.

Outflow I

- I In the Physics toolbar, click Attributes and choose Outflow.
- 2 In the Settings window for Outflow, locate the Boundary Selection section.
- 3 Click Clear Selection.
- 4 Select Boundary 20 only.

Species: Hn3

In the Model Builder window, under Component I (compl)>Plasma (plas)>Species click Species: Hn3.

Outflow I

- I In the Physics toolbar, click Attributes and choose Outflow.
- 2 In the Settings window for Outflow, locate the Boundary Selection section.
- 3 Click Clear Selection.
- 4 Select Boundary 20 only.

PLASMA (PLAS)

Species

In the Model Builder window, collapse the Component I (compl)>Plasma (plas)>Species node.

Wall I

- I In the Physics toolbar, click Boundaries and choose Wall.
- 2 In the Settings window for Wall, locate the Boundary Selection section.
- 3 From the Selection list, choose Walls.

Ground I

- I In the Physics toolbar, click Boundaries and choose Ground.
- **2** Select Boundaries 2, 13–15, 17–20, 24, 32, and 34 only.

Add physics interfaces to compute for the fluid flow and gas heating. The transport and thermodynamic properties are computed in the Plasma interface and will be linked to the interfaces below.

ADD PHYSICS

- I In the Physics toolbar, click and Physics to open the Add Physics window.
- 2 Go to the Add Physics window.
- 3 In the tree, select Heat Transfer>Heat Transfer in Fluids (ht).

- 4 Click Add to Component I in the window toolbar.
- 5 In the tree, select Fluid Flow>Single-Phase Flow>Laminar Flow (spf).
- 6 Click Add to Component I in the window toolbar.
- 7 In the Physics toolbar, click Add Physics to close the Add Physics window.

LAMINAR FLOW (SPF)

- I In the Settings window for Laminar Flow, locate the Physical Model section.
- 2 From the Compressibility list, choose Compressible flow (Ma<0.3).
- 3 Locate the Domain Selection section. Click | Clear Selection.
- 4 Select Domain 1 only.
- **5** Locate the **Physical Model** section. In the p_{ref} text field, type p0.

Fluid Properties 1

- I In the Model Builder window, under Component I (compl)>Laminar Flow (spf) click Fluid Properties I.
- 2 In the Settings window for Fluid Properties, locate the Model Input section.
- **3** From the T list, choose **Temperature** (ht).
- 4 Locate the Fluid Properties section. From the ρ list, choose Density (plas/pes1).
- **5** From the μ list, choose **Dynamic viscosity (plas/pes I)**.

Inlet I

- I In the **Physics** toolbar, click **Boundaries** and choose **Inlet**.
- 2 Select Boundary 32 only.
- 3 In the Settings window for Inlet, locate the Boundary Condition section.
- 4 From the list, choose Mass flow.
- 5 Locate the Mass Flow section. From the Mass flow type list, choose Standard flow rate (SCCM).
- **6** In the Q_{scem} text field, type Qs.
- 7 From the M_n list, choose Mean molar mass (plas/pes1).

Outlet I

- I In the Physics toolbar, click Boundaries and choose Outlet.
- 2 Select Boundary 20 only.

LAMINAR FLOW (SPF)

In the Model Builder window, collapse the Component I (compl)>Laminar Flow (spf) node.

HEAT TRANSFER IN FLUIDS (HT)

- I In the Model Builder window, under Component I (compl) click Heat Transfer in Fluids (ht).
- 2 In the Settings window for Heat Transfer in Fluids, locate the Domain Selection section.
- 3 Click Clear Selection.
- 4 Select Domain 1 only.

Fluid 1

- I In the Model Builder window, under Component I (compl)>Heat Transfer in Fluids (ht) click Fluid 1.
- 2 In the Settings window for Fluid, locate the Model Input section.
- **3** From the p_A list, choose **Absolute pressure (spf)**.
- 4 Locate the Heat Convection section. From the u list, choose Velocity field (spf).
- **5** Locate the **Heat Conduction, Fluid** section. From the k list, choose Thermal conductivity (plas/pes I).
- 6 Locate the Thermodynamics, Fluid section. From the Fluid type list, choose Gas/Liquid.
- 7 From the ρ list, choose Density (plas/pes I).
- 8 From the C_p list, choose Heat capacity at constant pressure (plas/pes1).
- **9** From the γ list, choose **User defined**.

Initial Values 1

- I In the Model Builder window, click Initial Values I.
- 2 In the Settings window for Initial Values, locate the Initial Values section.
- **3** In the *T* text field, type Tinit.

Temperature I

- I In the Physics toolbar, click Boundaries and choose Temperature.
- 2 Select Boundaries 2, 15, 17, and 18 only.
- 3 In the Settings window for Temperature, locate the Temperature section.
- **4** In the T_0 text field, type 1200[K].

Temperature 2

- I In the Physics toolbar, click Boundaries and choose Temperature.
- **2** Select Boundaries 13, 14, and 19 only.
- 3 In the Settings window for Temperature, locate the Temperature section.
- **4** In the T_0 text field, type 300[K].

Heat Flux 1

- I In the Physics toolbar, click Boundaries and choose Heat Flux.
- 2 In the Settings window for Heat Flux, locate the Heat Flux section.
- 3 From the Flux type list, choose Convective heat flux.
- **4** In the h text field, type 200.
- **5** In the T_{ext} text field, type 300[K].
- 6 Select Boundaries 24, 32, and 34 only.

The following heat source term includes all energy related with plasma volume reactions and is responsible to heat the gas.

Heat Source 1

- I In the Physics toolbar, click **Domains** and choose **Heat Source**.
- 2 In the Settings window for Heat Source, locate the Heat Source section.
- **3** From the Q_0 list, choose Heat source for gas (plas/pes1).
- **4** Select Domain 1 only.

Outflow I

- I In the Physics toolbar, click Boundaries and choose Outflow.
- 2 Select Boundary 20 only.

HEAT TRANSFER IN FLUIDS (HT)

In the Model Builder window, collapse the Component I (compl)>Heat Transfer in Fluids (ht) node.

ELECTROMAGNETIC WAVES, FREQUENCY DOMAIN (EMW)

In the Model Builder window, under Component I (compl) click Electromagnetic Waves, Frequency Domain (emw).

Port I

- I In the Physics toolbar, click Boundaries and choose Port.
- 2 Select Boundary 8 only.
- 3 In the Settings window for Port, locate the Port Properties section.
- 4 From the Type of port list, choose Coaxial.
- **5** In the $P_{\rm in}$ text field, type Pin.

ELECTROMAGNETIC WAVES, FREQUENCY DOMAIN (EMW)

In the Model Builder window, collapse the Component I (compl)>Electromagnetic Waves, Frequency Domain (emw) node.

The fluid velocity, pressure, and temperature are also linked to the plasma model making it fully self-consistent.

PLASMA (PLAS)

Plasma Model I

- I In the Model Builder window, under Component I (compl)>Plasma (plas) click Plasma Model I.
- 2 In the Settings window for Plasma Model, locate the Model Inputs section.
- 3 From the **u** list, choose **Velocity field (spf)**.
- **4** From the *T* list, choose **Temperature** (ht).
- 5 From the p_A list, choose Absolute pressure (spf).
- 6 Locate the Electron Density and Energy section. In the $\mu_e N_n$ text field, type 1.4E24[1/ (V*m*s)].

PLASMA (PLAS)

In the Model Builder window, collapse the Component I (compl)>Plasma (plas) node.

Create a mesh that is fine enough in the plasma region, specially at the subtract surface, and coarse elsewhere.

MESH I

Edge I

- I In the Mesh toolbar, click A Edge.
- 2 Select Boundary 32 only.

Size 1

- I Right-click **Edge** I and choose **Size**.
- 2 In the Settings window for Size, locate the Element Size section.
- 3 From the Calibrate for list, choose Fluid dynamics.
- 4 From the Predefined list, choose Extra fine.

Free Triangular 1

I In the Mesh toolbar, click Free Triangular.

- 2 In the Settings window for Free Triangular, click to expand the Control Entities section.
- 3 Clear the Smooth across removed control entities check box.

Size 1

- I Right-click Free Triangular I and choose Size.
- 2 In the Settings window for Size, locate the Element Size section.
- 3 From the Predefined list, choose Extra fine.
- 4 Locate the Geometric Entity Selection section. From the Geometric entity level list, choose Domain.
- **5** Select Domains 2, 3, 5, and 6 only.

Size 2

- I In the Model Builder window, right-click Free Triangular I and choose Size.
- 2 In the Settings window for Size, locate the Geometric Entity Selection section.
- 3 From the Geometric entity level list, choose Domain.
- **4** Select Domain 7 only.
- 5 Locate the Element Size section. From the Calibrate for list, choose Plasma.
- 6 From the Predefined list, choose Extremely fine.
- 7 Click the **Custom** button.
- 8 Locate the Element Size Parameters section.
- 9 Select the Maximum element size check box. In the associated text field, type 0.15.

Size 3

- I Right-click Free Triangular I and choose Size.
- 2 In the Settings window for Size, locate the Geometric Entity Selection section.
- 3 From the Geometric entity level list, choose Domain.
- 4 Select Domain 8 only.
- 5 Locate the Element Size section. From the Calibrate for list, choose Plasma.
- 6 From the Predefined list, choose Extremely fine.
- 7 Click the **Custom** button.
- 8 Locate the Element Size Parameters section.
- **9** Select the **Maximum element size** check box. In the associated text field, type **0.25**.

Size 4

I Right-click Free Triangular I and choose Size.

- 2 In the Settings window for Size, locate the Geometric Entity Selection section.
- 3 From the Geometric entity level list, choose Domain.
- **4** Select Domain 9 only.
- 5 Locate the Element Size section. From the Calibrate for list, choose Plasma.
- 6 From the Predefined list, choose Extremely fine.
- 7 Click the **Custom** button.
- 8 Locate the Element Size Parameters section.
- **9** Select the **Maximum element size** check box. In the associated text field, type 1.5.

Size 5

- I Right-click Free Triangular I and choose 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.
- **5** Select the **Maximum element size** check box. In the associated text field, type **0.025**.
- 6 Locate the Geometric Entity Selection section. From the Geometric entity level list, choose Domain.
- 7 Select Domain 1 only.

Boundary Layers 1

- I In the Mesh toolbar, click Boundary Layers.
- 2 In the Settings window for Boundary Layers, click to expand the Transition section.
- **3** Clear the **Smooth transition to interior mesh** check box.
- 4 Click to expand the Corner Settings section. Locate the Domain Selection section. From the Geometric entity level list, choose Domain.
- **5** Select Domains 1 and 7–9 only.

Boundary Layer Properties

- I In the Model Builder window, click Boundary Layer Properties.
- 2 In the Settings window for Boundary Layer Properties, locate the Layers section.
- 3 In the Number of layers text field, type 3.
- **4** Select Boundaries 13–15, 17–19, 24, and 34 only.
- 5 In the Model Builder window, right-click Mesh I and choose Build All.

First, solve a **Frequency-Transient** study for an input power of 500 W and without the fluid flow.

After, use the solutions from the previous study as initial conditions of a **Frequency-Stationary** study to ramp the power up to $5000 \ \mathrm{W}$.

STUDY I

Step 1: Frequency-Transient

- I In the Model Builder window, under Study I click Step I: Frequency-Transient.
- 2 In the Settings window for Frequency-Transient, locate the Physics and Variables Selection section.
- 3 In the table, clear the Solve for check box for Laminar Flow (spf).
- 4 Locate the Study Settings section. Click Range.
- 5 In the Range dialog box, choose Logarithmic from the Entry method list.
- 6 In the Start text field, type 1e-9.
- 7 In the **Stop** text field, type 100.
- 8 In the Steps per decade text field, type 3.
- 9 Click Replace.
- 10 In the Settings window for Frequency-Transient, locate the Study Settings section.
- II In the Output times text field, type 0 10^{range(log10(1.0e-9),1/3,
 log10(100))}.
- 12 In the Frequency text field, type f0.
- 13 In the Study toolbar, click $t_{=0}^{U}$ Get Initial Value.

RESULTS

Electric Field (emw), Electric Potential (plas), Electron Density (plas), Electron Temperature (plas), Temperature (ht)

- I In the Model Builder window, under Results, Ctrl-click to select Electron Density (plas), Electron Temperature (plas), Electric Potential (plas), Electric Field (emw), and Temperature (ht).
- 2 Right-click and choose Group.

Pin=500 W, Without Fluid Flow

In the **Settings** window for **Group**, type Pin=500 W, Without Fluid Flow in the **Label** text field.

PIN=500 W, WITHOUT FLUID FLOW

- I In the Model Builder window, click Study I.
- 2 In the Settings window for Study, type Pin=500 W, Without Fluid Flow in the Label text field.

Step 1: Frequency-Transient

- I In the Model Builder window, under Pin=500 W, Without Fluid Flow click Step 1: Frequency-Transient.
- 2 In the Settings window for Frequency-Transient, click to expand the Results While Solving section.
- **3** Select the **Plot** check box.
- 4 From the Update at list, choose Time steps taken by solver.
- 5 In the Study toolbar, click **Compute**.

ADD STUDY

- I In the Study 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 Preset Studies for Selected Multiphysics>Frequency-Stationary.
- 4 Click Add Study in the window toolbar.
- 5 In the Study toolbar, click Add Study to close the Add Study window.

STUDY 2

Step 1: Frequency-Stationary

- I In the Settings window for Frequency-Stationary, locate the Study Settings section.
- 2 In the Frequency text field, type f0.
- 3 Click to expand the Values of Dependent Variables section. Find the Initial values of variables solved for subsection. From the Settings list, choose User controlled.
- 4 From the Method list, choose Solution.
- 5 From the Study list, choose Pin=500 W, Without Fluid Flow, Frequency-Transient.
- 6 Click to expand the Study Extensions section. Select the Auxiliary sweep check box.
- 7 Click + Add.
- 8 Click Range.
- 9 In the Range dialog box, type 500 in the Start text field.

- 10 In the Step text field, type 100.
- II In the **Stop** text field, type 5000.
- 12 Click Replace.
- 13 In the Settings window for Frequency-Stationary, locate the Study Extensions section.
- 14 From the Run continuation for list, choose No parameter.
- 15 From the Reuse solution from previous step list, choose Yes.
- 16 In the Study toolbar, click t=0 Get Initial Value.

RESULTS

Electric Field (emw) I, Electric Potential (plas) I, Electron Density (plas) I, Electron Temperature (plas) 1, Pressure (spf), S-parameter (emw), Smith Plot (emw), Temperature (ht) I, Velocity (spf), Velocity, 3D (spf)

- I In the Model Builder window, under Results, Ctrl-click to select Electron Density (plas) I, Electron Temperature (plas) I, Electric Potential (plas) I, Electric Field (emw) I, Sparameter (emw), Smith Plot (emw), Temperature (ht) I, Velocity (spf), Pressure (spf), and Velocity, 3D (spf).
- 2 Right-click and choose **Group**.

Pin=500 To 5000 W

In the Settings window for Group, type Pin=500 To 5000 W in the Label text field.

PIN=500 TO 5000 W

- I In the Model Builder window, click Study 2.
- 2 In the Settings window for Study, type Pin=500 To 5000 W in the Label text field.

Step 1: Frequency-Stationary

- I In the Model Builder window, under Pin=500 To 5000 W click Step I: Frequency-Stationary.
- 2 In the Settings window for Frequency-Stationary, click to expand the Results While Solving section.
- **3** Select the **Plot** check box.
- 4 From the Plot group list, choose Electron Density (plas) 1.
- 5 In the Study toolbar, click **Compute**.

Add a mirror data set to better visualize the results.

RESULTS

Mirror 2D I

- I In the Model Builder window, expand the Results>Datasets node.
- 2 Right-click Results>Datasets and choose More 2D Datasets>Mirror 2D.
- 3 In the Settings window for Mirror 2D, locate the Data section.
- 4 From the Dataset list, choose Pin=500 To 5000 W/Solution 2 (sol2).
- 5 Click to expand the Advanced section. Select the Remove elements on the symmetry axis check box.

Electron Density (plas) I

- I In the Model Builder window, under Results>Pin=500 To 5000 W click Electron Density (plas) 1.
- 2 In the Settings window for 2D Plot Group, locate the Data section.
- 3 From the Dataset list, choose Mirror 2D 1.
- 4 In the Electron Density (plas) I toolbar, click **Plot**.

Electron Temperature (plas) I

- I In the Model Builder window, click Electron Temperature (plas) I.
- 2 In the Settings window for 2D Plot Group, locate the Data section.
- 3 From the Dataset list, choose Mirror 2D 1.
- 4 In the Electron Temperature (plas) I toolbar, click **Plot**.

Electric Potential (plas) 1

- I In the Model Builder window, click Electric Potential (plas) I.
- 2 In the Settings window for 2D Plot Group, locate the Data section.
- 3 From the Dataset list, choose Mirror 2D 1.
- 4 In the Electric Potential (plas) I toolbar, click **Plot**.

Electric Field (emw) I

- I In the Model Builder window, click Electric Field (emw) 1.
- 2 In the Settings window for 2D Plot Group, locate the Data section.
- 3 From the Dataset list, choose Mirror 2D 1.
- 4 In the Electric Field (emw) I toolbar, click **Plot**.

Velocity (spf)

I In the Model Builder window, click Velocity (spf).

- 2 In the Settings window for 2D Plot Group, locate the Data section.
- 3 From the Dataset list, choose Mirror 2D 1.
- 4 In the Velocity (spf) toolbar, click Plot.

Surface

- I In the Model Builder window, expand the Velocity (spf) node, then click Surface.
- 2 In the Settings window for Surface, locate the Coloring and Style section.
- 3 Click Change Color Table.
- 4 In the Color Table dialog box, select Rainbow>Prism in the tree.
- 5 Click OK.

Streamline I

- I In the Model Builder window, right-click Velocity (spf) and choose Streamline.
- 2 In the Settings window for Streamline, locate the Expression section.
- **3** In the **x-component** text field, type u.
- **4** In the **y-component** text field, type w.
- 5 Locate the Streamline Positioning section. From the Positioning list, choose Uniform density.
- 6 Locate the Coloring and Style section. Find the Point style subsection. From the Type list, choose Arrow.
- 7 From the Color list, choose Gray.
- 8 Locate the Streamline Positioning section. In the Separating distance text field, type 0.02.
- **9** In the **Velocity (spf)** toolbar, click **Plot**.

Temperature, 2D

- I In the Home toolbar, click Add Plot Group and choose 2D Plot Group.
- 2 In the Settings window for 2D Plot Group, type Temperature, 2D in the Label text field.
- 3 Locate the Data section. From the Dataset list, choose Mirror 2D 1.
- 4 From the Parameter value (Pin (W)) list, choose 5000.

Surface I

- I Right-click Temperature, 2D and choose Surface.
- 2 In the Settings window for Surface, locate the Expression section.
- **3** In the **Expression** text field, type T.
- 4 Locate the Coloring and Style section. Click Change Color Table.

- 5 In the Color Table dialog box, select Thermal>HeatCameraLight in the tree.
- 6 Click OK.
- 7 In the Temperature, 2D toolbar, click Plot.

H Number Density

- I In the Home toolbar, click Add Plot Group and choose 2D Plot Group.
- 2 In the Settings window for 2D Plot Group, type H Number Density in the Label text field.
- 3 Locate the Data section. From the Dataset list, choose Mirror 2D 1.
- 4 From the Parameter value (Pin (W)) list, choose 5000.

Surface I

- I Right-click H Number Density and choose Surface.
- 2 In the Settings window for Surface, locate the Expression section.
- 3 In the Expression text field, type plas.n wH.
- 4 In the H Number Density toolbar, click Plot.

Power Absorbed

- I In the Home toolbar, click **Add Plot Group** and choose **2D Plot Group**.
- 2 In the Settings window for 2D Plot Group, type Power Absorbed in the Label text field.
- 3 Locate the Data section. From the Dataset list, choose Mirror 2D 1.
- 4 From the Parameter value (Pin (W)) list, choose 5000.

Surface I

- I Right-click Power Absorbed and choose Surface.
- 2 In the Settings window for Surface, locate the Expression section.
- 3 In the Expression text field, type emw.Qrh.
- 4 Locate the Coloring and Style section. Click Change Color Table.
- 5 In the Color Table dialog box, select Thermal>ThermalWave in the tree.
- 6 Click OK.

Selection 1

- I Right-click Surface I and choose Selection.
- **2** Select Domain 1 only.
- 3 In the Power Absorbed toolbar, click Plot.