INTRODUCTION TO
Particle Tracing Module
Introduction to the Particle Tracing Module

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Introduction

The Particle Tracing Module is a general-purpose tool for computing the paths of particles as they move through a geometry and are subjected to various forces. The simulated particles could represent ions, electrons, cells, grains of sand, projectiles, planets, stars, and much more.

Most of the physics interfaces in COMSOL Multiphysics use the Finite Element Method (FEM) to compute fields such as temperature, fluid velocity, electric potential, concentration, or displacement. In contrast, particle tracing provides a Lagrangian description of a problem, in which the particles are treated as discrete entities that can interact with external fields, with boundaries in the surrounding geometry, and with each other. Their trajectories are computed in the time domain by solving a set of equations based on their laws of motion. The most common such system of equations are the second-order ordinary differential equations based on Newton’s Second Law of Motion, to which the forces based on external fields can easily be added.

As particles travel through the geometry of a model, they may also come into contact with boundaries in their surroundings. When this happens, each surface can affect the particle behavior in a different way. The built-in particle-boundary interactions include different types of absorption and combinations of diffuse and specular reflection. The Particle Tracing Module also offers a wide variety of release mechanisms to control how the particles are inserted into the geometry. After the simulation is complete, several dedicated tools for visualizing particle motion and reporting particle statistics are available.

A basic particle tracing model usually involves first solving for some fields, such as a fluid velocity field (left), then tracking particles in the field, using forces such as the built-in Drag Force (right).
Main Subject Areas

Particle Tracing Module applications can be loosely categorized into three subject areas:

- Charged particle tracing, the calculation of ion or electron paths under electric, magnetic, and collisional forces;
- Particle tracing in fluids, the tracking of micron-scale particles such as cells or airborne contaminants under the influence of drag, gravity, and other field forces; and
- Mathematical particle tracing, the setup of customized equations of motion, including the Hamiltonian and Lagrangian formulations from classical mechanics.

These three subject areas correspond to the three main physics interfaces included with the Particle Tracing Module: the Charged Particle Tracing interface ( ), the Particle Tracing for Fluid Flow interface ( ), and the Mathematical Particle Tracing interface ( ), respectively. In addition, several multiphysics interfaces are available, each with predefined couplings to other physics interfaces such as Electrostatics ( ) and Laminar Flow ( ).

Locations of Particle Tracing physics interfaces in the Model Wizard. The options shown are available with licenses for COMSOL Multiphysics and the Particle Tracing Module. Depending on other licensed products, the Model Wizard may include options other than those that are shown here.
# Physics Interface Guide by Space Dimension and Preset Study Type

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<th>ICON</th>
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<tr>
<td><strong>AC/DC</strong></td>
<td>![AC/DC Icon]</td>
<td></td>
<td>3D, 2D, 2D axisymmetric</td>
<td>bidirectionally coupled particle tracing; time dependent</td>
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<td>3D, 2D, 2D axisymmetric</td>
<td>bidirectionally coupled particle tracing; time dependent</td>
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<td><strong>Particle Tracing</strong></td>
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<td><strong>Particle Tracing for Fluid Flow</strong></td>
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<td><strong>Fluid-Particle Interaction</strong></td>
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<td><strong>Droplet Sprays in Fluid Flow</strong></td>
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<td><strong>Mathematical Particle Tracing</strong></td>
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<td>3D, 2D, 2D axisymmetric</td>
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Charged Particle Tracing

The Charged Particle Tracing interface is found under the AC/DC branch in the Model Wizard. It is primarily used to trace ions and electrons under the influence of electric and magnetic forces. These electric and magnetic forces can either be entered explicitly as user-defined expressions or computed using another physics interface. The magnetic force can also be taken from built-in data for the Earth’s magnetic field.

![Motion of trapped protons in the Earth’s magnetic field. The protons follow helical paths that oscillate between the northern and southern hemispheres.](image)

Included in the Electric Force and Magnetic Force features are built-in options to modulate AC fields. In this way, it is easy to include combinations of stationary and time-harmonic forces in a single model.
Quadrupole mass filter: Argon ions propagate through a quadrupole arrangement of electrodes (left). The stability of the design is determined by the magnitude of the DC and AC potentials used to guide the ions through the filter and reported as a function of the dimensionless Mathieu coefficient (right).

**Monte Carlo Collision Modeling**

In addition to electric and magnetic forces, energetic ions and electrons are often subjected to collisions with the molecules in a rarefied background gas. At sufficiently low pressure, these collisions can be modeled stochastically using a Monte Carlo approach. In each time step taken by a time-dependent solver, each model particle has a chance to collide with a background molecule and be scattered in a random direction.

The collision probability and the probability distribution function of scattered particle velocities are based on the particle kinetic energy, type of reaction, background gas density, and collision cross section data. Because the collision cross-section is usually a nontrivial function of energy, it is common to load cross-section data from a separate file and use it to define an interpolation function.

Some of the built-in collision types can release secondary particles of the same or a different species. For example, ionization reactions can yield secondary electrons, and charge exchange reactions can yield new combinations of ionized and neutralized species.

Because of its Monte Carlo collision modeling capability, the Charged Particle Tracing interface is sometimes extremely useful in modeling the motion of neutralized species, not just ions or electrons, in spite of the interface’s name.
Charge Exchange Cell: A fast neutral beam is produced by accelerating protons (light gray) and passing them through a vacuum chamber containing a rarefied argon gas. Collisions with the background gas produce energetic neutral molecules (dark gray). The remaining protons are deflected by an electrostatic field between two oppositely charged plates (red and blue).

**Multiphysics Interfaces for Particle Field Interaction**

If the number density of charged particles is sufficiently low, the Coulomb interaction between ions or electrons can sometimes be neglected, since it is likely to be dwarfed in magnitude by the external electric and magnetic fields. Such models use a unidirectional, or one-way, coupling from the electric and magnetic fields to the particle trajectories. In other words, first the fields are computed, and then these fields are used to exert forces on the charged particles.
Einzel Lens: The electrostatic force from the electrodes, illustrated by isosurfaces of electric potential, is much greater than the Coulomb interaction between the electrons, making this a unidirectionally coupled model.

**Coulomb Interaction**

If the density of charged particles is suitably high, then it may be necessary to include the Coulomb force that acts between the particles. When particle-particle interactions are included in a model, the computational requirements increase because each particle’s motion is affected by the position of every other particle in the model. When including the Coulomb force, it is often best to start with a small number of particles, solve the model, and then assess whether or not the effect is important.

**Bidirectional Particle-Field Interaction**

The Coulomb force is well-suited to models in which the interaction between particles may be significant but the total population of particles in the model remains constant over time. For beam simulations with space charge effects, however, ions or electrons are not just affected by particles that are released simultaneously, but also particles that are released earlier or later in time. This can lead to a very large number of model particles being required to accurately capture the Coulomb interaction.

An alternative to directly applying the Coulomb force is to allow particles to contribute to the space charge density in the domain for the purpose of computing the electric potential, then to use this modified electric potential to compute a
more accurate electric force on the particles. This is called a bidirectional, or two-way, coupling between the particles and fields. The bidirectional coupling is well-suited to simulations in which ions or electrons are released continuously over time such that the electric potential at any point in space remains approximately constant. The dedicated Particle Field Interaction, Nonrelativistic interface combines the Charged Particle Tracing interface with the Electrostatics interface to quickly set up the two-way interaction in which the particles perturb the electric potential in their surroundings.

A beam of relativistic ions or electrons can generate sufficient current to produce a significant magnetic force on the beam. This self-induced magnetic force can be implemented using the Particle Field Interaction, Relativistic interface, which automatically combines the Charged Particle Tracing, Electrostatics, and Magnetic Fields physics interfaces. This multiphysics coupling requires both the AC/DC Module and the Particle Tracing Module.

Releasing Nonlaminar Ion and Electron Beams
The Charged Particle Tracing interface includes a dedicated feature for releasing ion and electron beams. You can specify quantities such as the beam emittance and Twiss parameters to characterize the particle position and velocity distributions.
When one or more charged particle beams are released in a model, quantities such as beam emittance are automatically computed along the nominal beam trajectory and are available for postprocessing.

A beam of electrons is focused by a magnetic lens (left). The beam emittance can then be plotted as the thickness and color along the nominal beam path (right).
Particle Tracing for Fluid Flow

The Particle Tracing for Fluid Flow interface ( ), found under the Fluid Flow branch ( ) in the Model Wizard, computes the motion of particles in a background fluid. Particle motion can be driven by built-in forces including drag, gravitational, electric, magnetic, and acoustophoretic forces. User-defined forces can also be added. It is also possible to compute the particle mass and temperature as well as particle-particle interactions.

**Basic Requirements**

In order for the particle tracing approach to be valid, the system should be a *dilute* or *dispersed flow*. This means that the particles should occupy a very small fraction of the volume of the surrounding fluid, generally less than 1%. When the volume fraction of the particles is not small, the fluid system is categorized as a *dense flow* and a different modeling approach is required.

It is important to realize that with the particle tracing approach, particles do not displace the fluid they occupy. Furthermore, the finite radius of the particle is not usually taken into account when detecting and applying particle-wall interactions.

**Flow Regimes**

Motion of microscopic and macroscopic sized particles is typically dominated by the drag force acting on particles immersed in a fluid. There are two phases in the system: a *discrete phase* consisting of bubbles, particles, or droplets, and a *continuous phase* in which the particles are immersed. The velocity of the continuous phase is sometimes entered as an expression but is most often computed using a fluid flow physics interface such as Laminar Flow ( ), Turbulent Flow, k-ε ( ), or Creeping Flow ( ).

**Sparse Flow**

In a *sparse flow*, the continuous phase affects the motion of the particles via the drag force, but the particles don’t have enough inertia to significantly perturb the continuous phase. This is usually true when the particles in the discrete phase are very small or have relatively low number density. This is often referred to as unidirectional coupling or one–way coupling. When modeling such a system, it is usually most efficient to solve for the continuous phase first, then compute the trajectories of the discrete particles in a separate study. For example, in the following plot of a static mixer, the stationary velocity field and pressure were first solved using a stationary study, and then the particle trajectories were computed in a time–dependent study.
Comet tail plot of particle trajectories through a laminar static mixer (colored). In addition, Poincaré maps show the deviation of particle trajectories from their initial position (blue & red).

Dilute Flow
In a *dilute flow* the continuous phase affects the motion of the particles, and the particle motion in turn disrupts the continuous phase. This is often referred to as a bidirectional or two-way coupling. Compared to a sparse flow, the particles in a dilute flow typically have more inertia, due to the particles being bigger, denser, or more numerous.

The Fluid-Particle Interaction (\(\text{Fluid-Particle Interaction}\)) interface automatically sets up a two-way coupling between the Particle Tracing for Fluid Flow interface (\(\text{Particle Tracing for Fluid Flow}\)) and the Laminar Flow interface (\(\text{Laminar Flow}\)). The particles exert a volume force on the fluid at their location, equal in magnitude and opposite in direction to the total drag force exerted on the particles. The computational demand is significantly higher when modeling dilute flows than sparse flows. Before setting up a bidirectional coupling, it is often beneficial to first set up the model with a unidirectional coupling and then determine whether the effect of the particles on the surrounding fluid is significant.

Dispersed Flow
In a *dispersed flow*, the density of particles is greater than in the dilute flow described in the previous section, but their volume fraction is still lower than in a dense flow. The dispersed flow is the upper limit of the applicability of a
Lagrangian particle tracking approach. In addition to the bidirectional fluid-particle interaction mentioned in the previous section, particle-particle interactions may also need to be taken into account. This is sometimes referred to as four-way coupling. Particle-particle interactions can be included in models but the following limitations apply:

- Hard sphere collisions are not supported. Forces must vary continuously with respect to the distance between particles as in, for example, the Coulomb force between charged particles.
- The computation time scales as the number of particles squared. This is because every particle interacts with every other particle over all distances. It is possible to apply a cutoff distance for particle-particle interactions, but the interaction force must still be evaluated for every pair of particles.
- If the particle-particle interaction law is highly nonlinear, it may be necessary to use a very small time step. This is particularly true if the Lennard-Jones option is selected.

Particle Diffusion
Particle motion in a fluid is a combination of advective and diffusive transport. Advection is the bulk transport of particles by the mean fluid velocity via the drag force. Advection is typically deterministic; given a particle as a specified location in a laminar flow field, the motion of that particle at future times is completely predictable and reproducible (allowing, of course, some small differences due to the discrete time stepping used by the time dependent study). If 100 identical particles pass through the same point in a stationary flow field, each having the same velocity as it crosses that point, then all 100 of those particles will follow the same path.

Diffusion, on the other hand, is implemented as a stochastic term that can give different results for each particle in the simulation, even if the initial conditions are the same. There are two main mechanisms for diffusive transport in the Particle Tracing for Fluid Flow interface: molecular diffusion and turbulent dispersion.

Molecular Diffusion: The Brownian Force
Molecular diffusion is most significant for extremely small particles, on the order of 0.1 μm or smaller. When such small particles are in a fluid of nonzero absolute temperature, the thermal motion of individual atoms or molecules in that fluid causes them to collide randomly with the particles in directions other than that of the mean flow velocity. These random collisions can cause particles to spread out over time, a phenomenon known as Brownian motion.
Brownian motion of particles that are released from a single point. They diffuse outwards. Top left $t = 0$ s, top right $t = 10$ s, lower left $t = 30$ s, and lower right $t = 100$ s.

**Turbulent Dispersion**

Turbulence is the dominant mechanism for diffusive transport of macroscopic particles in a high Reynolds number flow. Unlike Brownian motion, turbulent dispersion can remain significant even for particles at the micron scale or larger.

For laminar flow, the drag force only contributes to the advective transport of the particles,

$$F_D = \frac{1}{\tau} m_p (\mathbf{u} - \mathbf{v})$$

However, if the surrounding fluid is turbulent, then the fluid velocity $\mathbf{u}$ at the particle’s position comprises a mean flow term and an instantaneous fluid velocity perturbation,
This is because the turbulence models in COMSOL Multiphysics based on the Reynolds Averaged Navier Stokes (RANS) model. In a RANS model, eddies in a turbulent flow are not modeled explicitly; instead, information about the magnitude and lifetime of eddies is only retained in a statistical sense, often through the use of transport equations for additional dependent variables. Several different expressions for the perturbation term exist, the simplest being

\[ \mathbf{u}' = \zeta \frac{\partial k}{\partial t} \]

where \( \zeta \) is a random unit vector and \( k \) is the turbulent kinetic energy. Some turbulence models, such as the \( k-\varepsilon \) model, solve for \( k \) as an additional degree of freedom; it represents the amount of energy associated with eddies in the flow. Unlike molecular diffusion, turbulent diffusion has a finite resolution in time. If the time step taken by the solver is made extremely small, then eventually the turbulent perturbation term between successive iterations no longer consists of uncorrelated random vectors. Therefore, to accurately model turbulent diffusion, it is important to know the average lifetime of eddies in the flow and to specify the maximum time step taken by the solver accordingly.

**Other Built-in Forces**

In addition to the drag force and Brownian force, the Particle Tracing for Fluid Flow interface includes a wide variety of other built-in forces that can be used to create high-fidelity multiphysics models, including the following:

- Electric and magnetic forces can be coupled to charged particles.
- Dielectrophoretic and magnetophoretic forces can affect particles in nonuniform electric and magnetic fields even if the particles are neutral. These forces are used in some biomedical industries to separate different cells based on their size or electrical properties.
- The acoustophoretic or acoustic radiation force affects particles in pressure waves.
- The thermophoretic force causes particles to migrate in a temperature gradient. This force is most significant in rarefied gases.
- Lift forces are exerted on particles in a shear flow.

In addition, some forces, such as the drag and thermophoretic forces, include high Knudsen number corrections that improve accuracy when the particles are extremely small or the surrounding fluid is a highly rarefied gas. The drag force
and lift force include correction terms to account for the presence of nearby walls, improving their accuracy for flows in channels and pipes.

Dielectrophoretic separation: the built-in dielectrophoretic force is used to separate red blood cells (red) and platelets (blue) in a flow channel with an applied electric potential (grayscale).

**Nozzles and Sprays**

The Particle Tracing for Fluid Flow interface is not just limited to solid particles in liquids or gases, but can also be used to model sprays of liquid droplets.

A dedicated Nozzle feature is available to release a spray of droplets from points in the geometry. In addition, the Droplet Breakup feature can be used to split the model particles into smaller particles over time. Typically the breakup occurs due to instability on the droplet surface created by external drag forces, or due to sudden acceleration of the droplet.

The Droplet Sprays in Fluid Flow interface ( ), found under the Fluid Flow branch ( ) in the Model Wizard, is a special case of the Particle Tracing for Fluid Flow interface ( ) that uses default settings and features that are frequently used to model sprays of liquid droplets. This includes options to specify particle viscosity and surface tension as well as an instance of the Droplet Breakup feature.
Mathematical Particle Tracing

The Mathematical Particle Tracing interface ( ) gives access to the underlying mathematical formalism on which the Charged Particle Tracing ( ) and Particle Tracing for Fluid Flow ( ) interfaces are built. The Mathematical Particle Tracing interface also allows for specification of particle motion in terms of either a Lagrangian or Hamiltonian. Often it is easier to write down an expression for the Lagrangian or Hamiltonian for particles rather than deriving the equations of motion. The Hamiltonian formulation solves for both the particle position and the particle momentum, so the number of degrees of freedom is doubled when the Hamiltonian formulation is activated.

Motion of stars in a galaxy based on a user-defined particle-particle interaction force for gravitational attraction.
Boundary Conditions

The following options are available for the boundary conditions for particles when they make contact with a wall: Bounce, Freeze, Stick, Disappear, Pass through, Diffuse scattering, Mixed diffuse and specular reflection, and General reflection. Some of the available options are illustrated in the figure below.

Demonstration of the Bounce, Freeze, Diffuse scattering, and Disappear Wall conditions.

- The Freeze, Stick, and Disappear options all stop a particle when it hits a boundary. The Stick option allows the particle to move if the boundary moves, as in the Moving Mesh interface ( ). The Disappear option completely removes particles from the simulation when they hit a surface.
- The Bounce option specularly reflects from the wall such that the particle kinetic energy is conserved.
- The Diffuse scattering option bounces particles off a wall according to the cosine law, sometimes called Knudsen’s cosine law or Lambertian reflection depending on the application area. The probability of a particle bouncing off the surface in a given direction within a differential solid angle $d\omega$ is given by $\cos(\theta)d\omega$, where $\theta$ is the angle between the direction of the reflected particle and the wall normal. The kinetic energy of the particle is conserved.
- The Mixed diffuse and specular reflection condition is a combination of the Bounce and Diffuse scattering conditions, each with a given probability.
- The Pass through option allows particles to cross boundaries unimpeded. By combining this with other Wall conditions, it is possible to define
boundaries that filter particles based on their properties or with a given probability.

- The General reflection option allows an arbitrary velocity to be specified after a particle makes contact with the wall. This can either be done in Cartesian coordinates or in the tangent-normal coordinate system. The velocity components can be functions of the incident particle velocity, energy or any other quantity.

**Conditional Particle-Wall Interactions**

The above boundary conditions for the particles striking the wall can all be conditional, either assigned a probability or contingent on a logical expression being satisfied.

A conditional Pass through Wall condition can be used to filter particles based on properties such as particle mass or diameter. Here the interior boundary acts as a screen that blocks large particles (marked mostly in shades of red and orange) but allows smaller particles to pass through unimpeded.

**Releasing Secondary Particles**

Secondary particles can be introduced into the modeling domain when a primary particle strikes the wall. Any number of secondary particles can be emitted when a particle strikes a wall. The released particles can be assigned initial velocity based on an expression or sampled from a Lambertian or isotropic distribution.
Particle Release Mechanisms

There are several ways to release particles, either from a grid, by selecting specific domains, or on specific boundaries. In each case the number of particles and the release frequency can be specified. There are also specific options available for the specific release features, described below.

**RELEASE FROM A GRID**

Particles can be released from a grid that can either be regular or graded. In addition to the initial coordinates of the particles, the initial velocity and times to release the particles can be specified. There are also predefined initial velocity distributions available to release particles with a Maxwellian, Isotropic spherical, hemispherical, conical, or Lambertian distribution. Particles can either be placed at all combinations of the initial coordinates or only at specified combinations of them.

The initial particle coordinates can be a list of values or expressions, like the sine curve shown above. At each release point, a single particle can be released or particle velocities can be sampled from a distribution. Here a cone of particles is released at each point.
**Release From a Domain**

There are several different mechanisms for choosing the release points from within a selected domain.

- When the particle release is Mesh based, there is an option to specify the Refinement factor. The higher this factor, the more particles are released from within each mesh element.
- When the particle release is Density based, the release of particles is weighted according to a user-defined expression. This is the most flexible option for releasing particles because the weighting function can be an analytic expression, a random function, or even a field variable computed by a previous study.
- The Random option allows the initial position of the particles to be chosen at random. New random positions are chosen for each release time.

![The mesh-based release uses the mesh element centers to initialize particle positions (right). A density-based release can be used to release particles with a user-defined position distribution, such as a bi-Gaussian distribution (center). The random release gives particles random initial positions on the selected domains (right).](image)

**Inlet**

The Inlet feature releases particles from a set of boundaries. The options for the domain-based Release are also available here, as well as an additional option to release particles uniformly from a selected boundary.

**Release From Edges and Points**

The Release from Edge feature can be used to release particles along edges in a 3D geometry. Similarly, the Release from Point feature can be used to release particles from selected points in the geometry.
When releasing particles along edges, particle positions can be mesh-based, weighted by a user-defined density, or uniformly distributed along the edges.

Using the Release from Edge feature, particles can be released from positions along arbitrary curves, such as the helix shown above.

RELEASE FROM DATA FILE

The Release from Data File feature allows the initial positions of particles to be specified by selecting columns from an imported text file. The initial velocity can also be read from the text file. If any Auxiliary Dependent Variable nodes are present, the initial values of auxiliary dependent variables can also be imported from the file.
Advanced Modeling Tools

The Particle Tracing Module provides a wide range of specialized modeling tools to assist in extracting specific quantities of interest.

**Particle Statistics**

The particle tracing interfaces define a number of special variables, some of which can only be used during results processing. These variables can be found in the Particle statistics list during results processing, as shown below:

An example of variables available from the particle statistics menu and available with the Mathematical Particle Tracing interface.
The variables defined include the following:

- **Particle index.** Each particle is assigned a unique index starting from 1 up to the total number of particles. This expression can be passed into a function, which can create, for example, random forces that are unique for each particle.

- **Particle release feature.** If there are multiple release features in a model, it is useful to be able to visualize how the particles mix together based on their initial release position. The Particle release feature variable takes a numeric value, starting at 1, which is unique to each release feature.

- **The release time of a given particle.** This works for both primary and secondary particles and thus allows for extraction of the time at which particles were released into the modeling domain.

- **The time at which a particle stopped at a boundary.** Using the difference between the release time and stop time, it is easy to compute the residence time of each particle.

There are also variables that are only available during results processing and can only be evaluated using the Global Evaluation node under Derived Values.

- **Total number of particles.** The total number of particles released may not be known in advance, especially in models that include secondary particle emission.

- **Total number of particles in selection.** If a selection has been applied to the Particle data set, the number of particles in that selection can be evaluated.

- **Transmission probability.** Often the transmission probability is the main quantity of interest in a particle tracing model. The Transmission probability variable is available on domains, boundaries, or combinations of both.

### Auxiliary Dependent Variables and Residence Time

Auxiliary dependent variables can be used to help keep track of things like residence time, particle trajectory length, integrated shear rate and so forth. When an Auxiliary Dependent Variable feature is added to the physics interface a first-order additional ordinary differential equation (ODE) is solved for each particle. This equation can be integrated either over time or along the particle trajectories.

The residence time can also be computed in a different way, by setting the Store particle status data property to On. This creates variables for the particle release time and the particle stop time (the time when a particle left the modeling domain). The residence time is then simply the difference between the two.
**Particle Counters**

Particle counters can be added to domains or boundaries in order to provide information about particles arriving on the set of selected domains or surfaces. It is possible to analyze the transmission of all particles or only those particles that are released by a specific feature. The Particle Counter computes quantities such as the number of particles transmitted, transmission probability, transmitted current, mass flow rate etc. The feature provides convenient expressions that can be used in the Filter node of the Particle Trajectories Plot, which allows only the particles which reach the Particle Counter selection to be visualized.

**Velocity Reinitialization in Domains**

It is possible to stop a particle or change its velocity when a specified logical expression is satisfied, even if the particle is not in contact with a boundary. For example, particles can be made to disappear after a certain amount of time, or to change direction if their \( z \)-coordinate exceeds a specified value. These interactions can also cause the emission of secondary particles; for example, a particle can be made to split into two or more smaller particles if its mass exceeds a threshold.
Results Processing Tools

Particle Trajectories Plots
A Particle Trajectories plot is available in both 2D and 3D. It is created automatically when a Study solving for a particle tracing interface is computed. There are options to set the particle trajectories to be rendered as lines, tubes, or ribbons. It is also possible to render the particles as points, comet tails, arrows, or with a number of ellipses along each trajectory. For models where the number of output times is small, an interpolation option is available that will create smoother looking particle trajectory plots.

The trajectory of an ion in a uniform magnetic field is plotted with no interpolation (left) and with interpolation (right).

Filters
Visualizing the trajectory of systems with a very large number of particles can consume a lot of computer resources and create cluttered-looking plots in which particles obscure each other from view. It is possible to filter the type of particle and the number of particles which should be rendered. To do this, right click the Particle Trajectories plot type and choose Filter.

The particles can be filtered so that only primary particles, secondary particles, or those satisfying a specified logical expression are plotted. It is also possible to plot only a specified fraction or number of particles.
Particle trajectories in a micromixer (left). It is possible to render only a fraction of the particles (middle) and to plot only the particles that are released by a certain inlet using a logical expression (right).

**PARTICLE PLOTS**

A Particle plot can be used to observe the value of a particle property over time. Built-in data series operations can be used, for example, to compute the sum or maximum of an expression over all particles.

Alternatively, the Particle plot can be used to compare two particle properties against each other for all particles at a set of selected time steps.

**Poincaré Maps and Phase Portraits**

Poincaré maps are available in both 2D and 3D. This plot type is useful to visualize the particle trajectories in a plot that represents the position of the particles in a section that is usually transversal to the particle trajectories. The
Poincaré map represents the particle trajectories in a space dimension that is one dimension lower than the original particle space.

Electrons in a magnetic lens (left). The Poincaré maps (right) show the electron distribution in beam cross-sections at three different axial positions.

Phase portraits are available as a 2D plot type under More Plots. Use a Phase Portrait plot to visualize large data sets of particle trajectories. The traditional use of a phase portrait is to plot the particle position on the $x$-axis and the particle velocity on the $y$-axis. Each dot in the $xy$-plane represents a particle.

**Particle Evaluation**

Information about expressions and variables along particle trajectories can be written to the Results Table using the Particle Evaluation option under Derived Values. Once the data has been written to the results table it can be manipulated and plotted accordingly. There are options to only write data to the table for a fraction or a specific number of particles.

**Operations on Particle Data**

It is possible to set the source data set for the Integration $\int$, Average $\bar{V}$, Maximum $\text{MAX}$, and Minimum $\text{MIN}$ to be a Particle data set. This allows operations to be performed on the particle data set, to compute average particle velocity, maximum energy, and so on.

**Histograms**

Statistical information about the behavior of the particles is often best visualized with a histogram. The histogram sorts the value of a variable into a specified
number of bins. The most obvious application of the histogram is that it allows for visualization of the velocity and energy distribution function of a set of particles.

Information about the ion energy distribution function in a plasma can be visualized using a 2D Histogram.
Tracing Particles through a Laminar Static Mixer

This section takes you through the modeling stages of computing particle trajectories based on a computed fluid velocity field. In static mixers, also called motionless or in-line mixers, a fluid is pumped through a pipe containing stationary blades. This mixing technique is particularly well suited for laminar flow mixing because it generates only small pressure losses in this flow regime. This example studies the flow in a twisted-blade static mixer. It evaluates the mixing performance by calculating the trajectory of suspended particles and displaying their positions at several cross-sections of the mixing blades.

Model Wizard

Note: These instructions are for the user interface on Windows but apply, with minor differences, also to Linux and Mac.

To start the software, double-click the COMSOL icon on the desktop. When the software opens, you can choose to use the Model Wizard to create a new COMSOL model or Blank Model to create one manually. For this tutorial, click the Model Wizard button.

If COMSOL is already open, you can start the Model Wizard by selecting New from the File menu and then click Model Wizard.

The Model Wizard guides you through the first steps of setting up an application. The next window lets you select the dimension of the modeling space.
2 In the Select Space Dimension window click the 3D button 

3 In the Select Physics tree under Fluid Flow>Single-Phase Flow, click Laminar Flow.

   Even though this is a particle tracing model, the first step is to solve for the fluid velocity field, which will then be used to exert a drag force on the particles.

4 Click Add then click the Study button.

5 In the tree under Preset Studies, click Stationary.

6 Click the Done button.

Global Definitions and Definitions

1 On the Home toolbar click Parameters button. Alternatively, click the Parameters node in the Model Builder.

   Note: On Linux and Mac, the Home toolbar refers to the specific set of controls near the top of the Desktop.

2 In the Settings window locate the Parameters section.

3 In the table, enter the following settings:

<table>
<thead>
<tr>
<th>Name</th>
<th>Expression</th>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drho</td>
<td>[mm]</td>
<td>0.001 m</td>
<td>Tube radius</td>
</tr>
<tr>
<td>Q_in</td>
<td>[l/min]</td>
<td>0.001 m</td>
<td>Mean velocity</td>
</tr>
</tbody>
</table>

   For these parameter values, and for water with an approximate density of 1000 kg/m³^3 and viscosity of 0.001 Pa s, the Reynolds number of the flow is approximately 60, meaning that the use of the Laminar flow physics interface is appropriate.

Geometry

The mixer geometry is quite complicated so start by importing it from a file.

1 On the Home toolbar click Import button.

2 In the Settings window locate the Import section. Click Browse.
3 Browse to the application library folder and double-click the file laminar_mixer_particle.mphbin. The location of the files used in this exercise varies based on your installation. For example, if the installation is on your hard drive, the file path might be similar to C:\Program Files\COMSOL\COMSOL53a\Multiphysics\applications\.

4 Click Import. The geometry is imported to the Graphics window.

5 Click the Wireframe Rendering button on the Graphics toolbar. This makes the surfaces of the mixer geometry transparent and makes the edges of the mixing blades visible. The geometry appears in the Graphics window as shown below.

The mixer geometry consists of three twisted blades, each making a 90 degree angle with the adjacent blades. The middle blade is twisted in the opposite direction from the first and last blades to improve the mixing performance.

**Materials**

1 On the Materials toolbar click Blank Material.

2 In the Settings window for Material locate the Material Contents section. In the table, enter the following settings:

```plaintext
<table>
<thead>
<tr>
<th>Property</th>
<th>Name</th>
<th>Value</th>
<th>Unit</th>
<th>Property group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density</td>
<td>rho</td>
<td>1000</td>
<td>kg/m³</td>
<td>Basic</td>
</tr>
<tr>
<td>Dynamic viscosity</td>
<td>mu</td>
<td>1e-3</td>
<td>Pa·s</td>
<td>Basic</td>
</tr>
</tbody>
</table>
```
Physics Interfaces

Now add an expression for the inflow velocity which is parabolic.

1. On the Physics toolbar click Boundaries and choose Inlet.  
2. Select Boundary 23 only.  
   **Note:** To add geometric entities to a selection, click on them in the Graphics window. Alternatively, Click Paste Selection and enter a list of numbers corresponding to the entities to select. Here, Boundary 23 is at the end of the modeling domain where the y-coordinate reaches its maximum value.

3. In the Settings window for Inlet locate the Velocity section. In the $U_0$ text field, type $2*(1-(x^2+z^2)/Ra^2)*u_{av}$.
   **Note:** The boundary condition, which was just added, is rather complicated but necessary to get a fully developed flow profile. The CFD, Microfluidics, and Plasma Modules all have a special Laminar Inflow boundary condition that ensures laminar flow. It is not necessary to enter a complicated expression for the velocity profile, just the average velocity or mass flow rate.

4. On the Physics toolbar click Boundaries and choose Outlet.  
5. Select Boundary 20 only. This is the surface opposite Boundary 23 which was selected for the Inlet boundary condition.

Mesh

The mesh needs to be quite fine to ensure that the particle motion is accurate through the modeling domain. In this case, take care to ensure that the mesh is sufficiently fine close to the mixing blades.

1. In the Model Builder right-click Mesh 1 and choose More Operations>Free Triangular.  
2. Select Boundaries 5, 16–18, and 53–55 only. These boundaries include the leading edges of all three mixing blades.
3. Right-click Free Triangular 1 and choose Size.  
4. In the Settings window for Size locate the Element Size section.  
   - From the Calibrate for list, choose Fluid dynamics.  
   - From the Predefined list, choose Extremely fine.  
5. Click the Size node above the Free Triangular node.
6 In the Settings window for Size locate the Element Size section.
   - From the Predefined list, choose Extremely fine.
   - Click the Custom button. By first choosing the Extremely fine option and then switching to a Custom mesh, you can begin customizing the mesh with a reasonable set of default values.
   - Locate the Element Size Parameters section. In the Curvature factor text field, type 0.15. The curvature factor is used to refine the mesh on curved surfaces while allowing the mesh to be somewhat coarser on planar surfaces.
7 Right-click Mesh 1 and choose More Operations>Free Triangular.
8 Select Boundary 23 only.
9 Right-click Free Triangular 2 and choose Size.
   - In the Settings window for Size locate the Element Size section. From the Calibrate for list, choose Fluid dynamics.
   - From the Predefined list, choose Extra fine.
10 Right-click Mesh 1 and choose Free Tetrahedral. Alternatively, select Free Tetrahedral from the Mesh toolbar.
11 Click Build All. The mesh should look like the following image. Transparency has been enabled so that the mesh on the interior boundaries can be seen.
Study

1 On the Home toolbar click Compute \(\Rightarrow\). In this model, the default solver sequence for the Stationary study is sufficient to model the flow. The study creates default plots of the fluid velocity and pressure as shown below.

![Default plots of the fluid velocity (left) and pressure (right).](image)

Model Wizard

Now that the fluid velocity has been computed, add a physics interface and a new study to compute the particle trajectories.

1 On the Home toolbar click Add Physics \(\Rightarrow\).

2 Go to the Add Physics window. In the Add Physics tree under Fluid Flow expand Particle Tracing \(\Rightarrow\) and click Particle Tracing for Fluid Flow \(\Rightarrow\).

3 Find the Physics interfaces in study subsection. In the Solve column, deactivate Study 1 by clicking on the check box containing the green check \(\checkmark\). The check box will be cleared \(\square\) indicating that this physics interface is not solved in Study 1. This subsection then looks like the following:

<table>
<thead>
<tr>
<th>Physics interfaces in study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Studies Solve</td>
</tr>
<tr>
<td>Study 1 (\square)</td>
</tr>
</tbody>
</table>

4 Click the Add to Component \(\Rightarrow\) button.

5 On the Home toolbar click Add Study \(\Rightarrow\).

6 Go to the Add Study window. Find the Studies subsection. In the tree under Preset Studies, click Time Dependent \(\Rightarrow\).
Find the Physics interface in study subsection. In the table, click on the green check box next to the Laminar Flow interface so that this subsection looks like the following:

<table>
<thead>
<tr>
<th>Physics Interfaces in study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physics</td>
</tr>
<tr>
<td>Laminar Flow (spf)</td>
</tr>
<tr>
<td>Particle Tracing for Fluid Flow (fp1)</td>
</tr>
</tbody>
</table>

It is not necessary to solve for the Laminar Flow physics in the new Time Dependent study step because it is a stationary field that was already solved for in a previous step.

8 Click the Add Study button.
9 On the Home toolbar, again click the Add Physics and Add Study buttons to remove the corresponding windows from view.

**Physics Interfaces**

In this section all of the relevant particle tracing physics features are set up. The default boundary condition, Wall, stops particles that touch a boundary. In the following steps, the drag force and the particle release mechanism are specified.

1 On the Physics toolbar click Domains and choose Drag Force.
2 Select Domain 1 only.
3 In the Settings window for Drag Force locate the Drag Force section.
   - The default Drag law is Stokes. There are other drag laws available, such as Schiller-Naumann, but Stokes drag is appropriate in this case because the particles are small enough that their relative Reynolds number is much less than 1.
   - From the $u$ list, choose Velocity field (spf/fp1) to take the fluid velocity from the Laminar Flow physics interface solved for in the previous study.
   - From the $μ$ list, choose Dynamic viscosity (spf/fp1).
The goal is to release particles with greater number density where the magnitude of the velocity field is higher. To do this you use the Density option for the particles’ Initial Position.
4 On the Physics toolbar click Boundaries and choose Inlet.
5 Select Boundary 23 only. This is the same surface as the Inlet for the Laminar Flow physics interface.
6 In the Settings window for Inlet locate the Initial Position section.
   - From the Initial position list, choose Density.
   - In the $N$ text field, type 3000. This is the total number of particles that will be released into the mixer.
   - In the $p$ text field, type $spf \cdot u$. The number density of particles at the Inlet will be approximately equal to the fluid velocity magnitude.
   - Locate the Initial Velocity section. From the $u$ list, choose Velocity field ($spf$).

7 On the Physics toolbar click Boundaries and choose Particle Counter.

   **Note:** The Particle Counter is used to compute the transmission probability of particles through the mixer. Alternatively, it is possible to compute the transmission probability in postprocessing by assigning a Selection to the Particle data set.

8 Select Boundary 20 only. This was the Outlet boundary for the Laminar Flow physics interface.

9 In the Settings window for Particle Counter locate the Particle Counter section.

10 From the Release feature list, choose Inlet 1. Because this model only includes one inlet, it is not strictly necessary to choose a specific release feature, although doing so can be useful in more complicated models.

11 In the Model Builder under Particle Tracing for Fluid Flow, click Particle Properties 1.

12 To change the diameter of the particles, in the Settings window for Particle Properties locate the Particle Properties section. In the $d_p$ text field, type $5E-7[m]$.

---

**Study 2**

1 In the Model Builder expand the Study 2 node and click Step 1: Time Dependent.

2 Expand the Values of Dependent Variables section. From the Settings list, choose User controlled.

3 From the Method list, choose Solution.

4 From the Study list, choose Study 1, Stationary.

5 Locate the Study Settings section. Click the Range button.
6 Go to the Range dialog box.
   - In the Step text field, type 0.2.
   - In the Stop text field, type 5.
7 Click the Replace button. The Settings window should look like this:

<table>
<thead>
<tr>
<th>Settings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time Dependent</td>
</tr>
<tr>
<td>Compute</td>
</tr>
<tr>
<td>Label: Time Dependent</td>
</tr>
<tr>
<td>Study Settings</td>
</tr>
<tr>
<td>Time unit:</td>
</tr>
<tr>
<td>Times: range(0,5)</td>
</tr>
<tr>
<td>Relative tolerance:</td>
</tr>
<tr>
<td>Results While Solving</td>
</tr>
<tr>
<td>Physics and Variables Selection</td>
</tr>
<tr>
<td>Modify physics tree and variables for study step</td>
</tr>
<tr>
<td>Physics interface</td>
</tr>
<tr>
<td>Laminar Flow (gpt)</td>
</tr>
<tr>
<td>Particle Tracing for Fluid Flow (fpt)</td>
</tr>
<tr>
<td>Values of Dependent Variables</td>
</tr>
<tr>
<td>Initial values of variables solved for</td>
</tr>
<tr>
<td>Settings: Physics controlled</td>
</tr>
<tr>
<td>Values of variables not solved for</td>
</tr>
<tr>
<td>Settings: User controlled</td>
</tr>
<tr>
<td>Method</td>
</tr>
<tr>
<td>Study</td>
</tr>
<tr>
<td>Selection</td>
</tr>
<tr>
<td>Store fields in output</td>
</tr>
<tr>
<td>Settings: All</td>
</tr>
<tr>
<td>Mesh Selection</td>
</tr>
<tr>
<td>Study Extensions</td>
</tr>
</tbody>
</table>

8 On the Home toolbar click Compute = .

**Results**

By default you end up in the settings window for the Particle Trajectories (fpt) plot group when the study is complete.
1 In the Settings window locate the Color Legend section.
2 From the Position list, choose Bottom.
3 In the Model Builder expand the Particle Trajectories (fpt) node then click Particle Trajectories 1.
4 In the Settings window for the Particle Trajectories plot locate the Coloring and Style section. This section controls how the particles and their paths are displayed.
   - Find the Line style subsection. From the Type list, choose Line.
   - Find the Point style subsection. From the Type list, choose None.
5 In the Model Builder expand the Results>Particle Trajectories (fpt) node and click Color Expression 1.
6 In the Settings window for Color Expression click Replace Expression in the upper-right corner of the Expression section. From the menu, choose Model>Component 1>Laminar Flow>Velocity and pressure>spf.sr – Shear rate.
7 Click the Plot button.
8 Click Zoom Extents in the Graphics toolbar.

Now evaluate the transmission probability using the Particle Counter that was created earlier. For this purpose, transmission probability is defined as the fraction of the released particles to reach the outlet by the final time.

1 On the Results toolbar click Global Evaluation.

2 In the Settings window for Global Evaluation locate the Data section.
   - From the Data set list, choose Study 2/Solution 2 (sol2).
   - From the Time selection list, choose Last.

3 Click Replace Expression in the upper-right corner of the Expression section. From the menu, choose Model>Component 1>Particle Tracing for Fluid Flow>Particle Counter 1>ft.pcnt1.alpha – Transmission probability.

4 Click the Evaluate button. The transmission probability should be about 0.8.

**Poincaré Plot**

One useful way of visualizing how particles mix is to use a Poincaré plot. The Poincaré plot places a colored dot for each particle at the location at which the particle passes through a cut plane (known as a Poincaré section). The first step is to define the cut planes where the intersections with the particle trajectories will be taken.
1 On the Results toolbar click Cut Plane.
2 In the Settings window for Cut Plane locate the Data section. From the Data set list, choose Particle 1.
3 Locate the Plane Data section.
   - From the Plane list, choose xz-planes.
   - In the y-coordinate text field, type 0.006.
   - Select the Additional parallel planes check box.
   - In the Distances text field, type 0.006 0.016 0.026 0.036 0.042.
4 Click the Plot button.

The cut planes will be shown in the Graphics window as shown below.

Cross-sections at which Poincaré maps will be plotted are shown in red.

Poincaré Map

1 On the Home toolbar click Add Plot Group and choose 3D Plot Group.
2 In the Settings window for 3D Plot Group locate the Data section. From the Data set list, choose Particle 1.

The plots only display particle positions in each cut plane, but the plot group should still use Particle 1 or one of the Solution data sets as its data source, since this is used to render the data set edges in the Graphics window.
On the 3D Plot Group 4 toolbar, click More Plots  and choose Poincaré Map.

In the Settings window for Poincaré Map locate the Data section. From the Cut plane list, choose Cut Plane 1.

Locate the Coloring and Style section.
- Select the Radius scale factor check box.
- In the associated text field, type \(6 \times 10^{-5}\). This makes the intersection points look smaller so that adjacent points in each cross section don’t obscure each other.

Click the Plot button .

Right-click Poincaré Map 1  and choose Color Expression .

In the Settings window for Color Expression locate the Expression section. In the Expression text field, type \(\text{at}(0,qx<0)\).

The \(\text{at}\) operator evaluates an expression at the time specified in the first argument. Here, it is a logical expression based on the initial particle position.

Locate the Coloring and Style section. Clear the Color legend check box.

Next add a Surface plot to render the cut planes where the intersection points are rendered.

**Surface Plot**

On the 3D Plot Group 4 toolbar click Surface .

In the Settings window for Surface locate the Data section. From the Data set list, choose Cut Plane 1.

Locate the Expression section. In the Expression text field, type 1.

Locate the Coloring and Style section.
- From the Coloring list, choose Uniform.
- From the Color list, choose Gray.

Click the Plot button .

Click the Go to Default 3D View button on the Graphics toolbar.
In the above figure, the location of the particles at 6 Poincaré sections are shown. The color represents the location of the particle at its initial position. So, particles marked as red had an initial position of \( x < 0 \) and particles marked as blue had an initial position of \( x > 0 \). The \( \text{at()} \) operator is used to mark the particles with the color of their initial position. The first Poincaré section (the one furthest to the left in the above figure) clearly indicates which particles start with coordinates of \( x < 0 \). As the particles follow the flow field, they begin to mix together. By the end of the mixer, the particles have not mixed completely—there are still significant pockets of only red and only blue particles.