COMSOL MULTIPHYSICS®

REFERENCE GUIDE

VERSION 3.4



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Commands Grouped by Function

User Interface Functions

FUNCTION	PURPOSE
comsol	Start the COMSOL Multiphysics graphical user interface or a COMSOL Multiphysics server

Solver Functions

FUNCTION	PURPOSE
adaption	Solve PDE problem using adaptive mesh refinement
femeig	Solve eigenvalue PDE problem
femlin	Solve linear stationary PDE problem
femnlin	Solve nonlinear stationary PDE problem
femstatic	Solve stationary PDE problem
femtime	Solve time-dependent PDE problem
poisson	Fast solution of Poisson's equation on a rectangular grid

Geometry Functions

FUNCTION	PURPOSE
chamfer	Create flattened corners in 2D geometry object
elevate	Elevate degrees of 2D geometry object Bézier curves
embed	Embed 2D geometry object as 3D geometry object
extrude	Extrude 2D geometry object to 3D geometry object
fillet	Create circular rounded corners in 2D geometry object
flcontour2mesh	Create boundary mesh from contour data
flim2curve	Create 2D curve object from image data
flmesh2spline	Create spline curves from mesh
geomanalyze	Decompose and analyze geometry of FEM problem
geomarrayr	Create rectangular array of geometry object
geomcoerce	Coerce geometry objects
geomcomp	Analyze (compose) geometry objects

FUNCTION	PURPOSE
geomcsg	General function for analyzing geometry objects
geomdel	Delete interior boundaries
geomedit	Edit geometry object
geomexport	Export geometry object to file
geomfile	Geometry M-file
geomgetwrkpln	Retrieve work plane information
geomgroup	Group geometry objects into an assembly
geomimport	Import geometry object from file
geominfo	Retrieve geometry information.
geomobject	Create geometry object
geomplot	Plot a geometry object
geomposition	Position 3D geometry object
geomspline	Spline interpolation
geomsurf	Surface interpolation
get	Get geometry object properties
getparts	Extract parts from an assembly object
loft	Loft 2D geometry sections to 3D geometry
mirror	Reflect geometry
move	Move geometry object
revolve	Revolve 2D geometry object to 3D geometry object
rotate	Rotate geometry object
scale	Scale geometry object
split	Split geometry object
tangent	Create a tangent line

Geometry Objects

FUNCTION	PURPOSE
arc1, arc2	Elliptical or circular arc/solid sector
block2, block3	Rectangular block face/solid object
circ1, circ2	Circle curve/solid object
cone2, cone3	Cone face/solid object.
curve2, curve3	2D/3D rational Bézier curve object
cylinder2, cylinder3	Cylinder face/solid object
econe2, econe3	Eccentric cone face/solid object
ellip1, ellip2	Ellipse curve/solid object
ellipsoid2, ellipsoid3	Ellipsoid face/solid object
face3	3D rational Bézier surface object
gencyl2, gencyl3	Straight homogeneous generalized cylinder face/solid object
geomO, geom1, geom2, geom3	0D/ID/2D/3D geometry object
helix1, helix2, helix3	Helix curve/face/solid object
hexahedron2, hexahedron3	Hexahedron face/solid object
line1, line2	Open curve/solid polygon
point1, point2, point3	ID/2D/3D point object
poly1, poly2	Closed curve/solid polygon
pyramid2, pyramid3	Pyramid face/solid object
rect1, rect2	Rectangle curve/solid object
solidO, solid1, solid2, solid3	0D/ID/2D/3D solid object
sphere3, sphere2	Sphere solid/face object
square1, square2	Square curve/solid object

FUNCTION	PURPOSE
tetrahedron2, tetrahedron3	Tetrahedron face/solid object
torus2, torus3	Torus face/solid object

Mesh Functions

FUNCTION	PURPOSE
femmesh	Create a mesh object
flcontour2mesh	Create boundary mesh from contour data
get	Get mesh object properties
mesh2geom	Create geometry from (deformed) mesh
meshbndlayer	Create boundary layer mesh
meshcaseadd	Add new mesh cases
meshcasedel	Delete mesh cases
meshcopy	Copy mesh between boundaries
meshdel	Delete elements in a mesh
meshembed	Embed a 2D mesh into 3D
meshenrich	Make mesh object complete
meshexport	Export meshes to file
neshextend	Extend a mesh to the desired finite element types
meshextrude	Extrude a 2D mesh into a 3D mesh
neshhex2tet	Convert hexahedral elements to tetrahedral elements
neshimport	Import meshes from file
neshinit	Build an initial mesh
neshmap	Build a mapped mesh
meshplot	Plot mesh
neshpoi	Make regular mesh on a rectangular geometry
meshquad2tri	Convert quadrilateral elements to triangular elements
neshqual	Mesh quality measure
neshrefine	Refine a mesh
neshrevolve	Revolve a 2D mesh into a 3D mesh
neshsmooth	Jiggle internal points of a mesh
meshsweep	Build a swept mesh
xmeshinfo	Get extended mesh information

Utility Functions

FUNCTION	PURPOSE
assemble	Assemble the stiffness matrix, right-hand side, mass matrix, and constraints of a PDE problem
asseminit	Compute initial value
femdiff	Symbolically differentiate general form
femsim	Create Simulink structure
femstate	Create state-space model for PDE problem
femstruct	FEM structure information
femwave	Extend FEM structure to a wave equation problem
flcompact	Compact equ/bnd/edg/pnt fields
flform	Convert between PDE forms
flload	Load a COMSOL Multiphysics file
flngdof	Get number of global degrees of freedom
flnull	Compute null space of a matrix, its complement, and the range of the matrix
flreport	Globally turn off progress window or show it
flsave	Save a COMSOL Multiphysics file
multiphysics	Multiphysics function
pde2draw	Convert a PDE Toolbox geometry description
pde2fem	Convert a PDE Toolbox model description to an FEM structure
pde2geom	Convert a PDE Toolbox decomposed geometry
solsize	Get number of solutions in a solution object

Postprocessing Functions

FUNCTION	PURPOSE	
femplot	Description of properties common to all plot functions	
meshintegrate	Compute integrals in arbitrary cross sections	
postanim	Shorthand command for animation	
postarrow	Shorthand command for arrow plot in 2D and 3D	
postarrowbnd	Shorthand command for boundary arrow plot in 2D and 3D	
postcont	Shorthand command for contour plot in 2D	
postcoord	Get coordinates in a model	

FUNCTION	PURPOSE	
postcrossplot	Cross-section plot	
posteval	Evaluate expressions on subdomains, boundaries, edges, and vertices	
postflow	Shorthand command for streamline plot in 2D and 3D	
postglobaleval	Evaluate globally defined expressions, such as solutions to ODEs	
postglobalplot	Plotting globally defined expressions, such as solutions to ODEs	
postgp	Extract Gauss points and Gauss point weights	
postint	Integrate expression over subdomains, boundaries, edges, and vertices	
postinterp	Evaluate expressions in arbitrary points	
postiso	Shorthand command for isosurface plot in 3D	
postlin	Shorthand command for line plot	
postmax	Compute maximum value for expression	
postmin	Compute minimum value for expression	
postmovie	Postprocessing animation function	
postplot	Postprocessing plot function	
postprinc	Shorthand command for subdomain principal stress/strain plot in 2D and 3D	
postprincbnd	Shorthand command for boundary principal stress/strain plot in 2D and 3D	
postslice	Shorthand command for slice plot in 3D	
postsurf	Shorthand command for surface plot in 2D and 3D	
posttet	Shorthand command for subdomain plot in 3D	

Low-Level Functions

FUNCTION	PURPOSE
dst	Discrete sine transform
idst	Inverse discrete sine transform

Shape Function Classes

FUNCTION	PURPOSE	
sharg_2_5	ifth-order Argyris shape function object in 2D	
shbub	Bubble shape function object	
shcurl	Vector shape function object	
shdens	Density element shape function object	
shdisc	Discontinuous shape function object	
shdiv	Divergence shape function object	
shgp	Gauss-point shape function object	
shherm	Hermite shape function object	
shlag	Lagrange shape function object	
shuwhelm	Scalar plane wave basis function object	

Element Syntax Classes

FUNCTION	PURPOSE
elsconstr	Coefficient and general form constraint element
elconst	Global expression variable element
elcontact	Contact map operator element
elcplextr	Extrusion coupling variable element
elcplgenint	Destination-aware integration coupling variable element
elcplproj	Projection coupling variable element
elcplscalar	Integration coupling variable element
elcurlconstr	Vector constraint element
elempty	Empty element which defines basic syntax
elepspec	Evaluation and constraint point pattern declaration element
eleqc	Coefficient and general form equation element
eleqw	Weak form equation element

FUNCTION	PURPOSE
elgeom	Geometric variable element
elgpspec	Integration point pattern declaration element
elinline	Inline function declaration element
elinterp	Interpolation function declaration element
elinv	Inverse matrix component variable element
elirradiation	Irradiation coupling variable element
elmapextr	Extrusion map operator element
elmesh	Mesh variable element
elode	Global scalar variable and equation element
elpconstr	Point-wise constraint element
elpiecewise	Piecewise function declaration element
elplastic	Plastic strain variable element
elpric	Principal component and vector variable element
elshape	Shape function declaration element
elshell_arg2	Shell equation element
elvar	Expression variable element

Mathematical Functions

FUNCTION	PURPOSE
flc1hs	Smoothed Heaviside function with continuous first derivative
fldc1hs	Derivative of flc1hs
flc2hs	Smoothed Heaviside function with continuous second derivative
fldc2hs	Derivative of flc2hs
flsmhs	Smoothed Heaviside function
fldsmhs	Derivative of smoothed Heaviside function
flsmsign	Smoothed sign function
fldsmsign	Derivative of smoothed sign function

Obsolete Functions in 3.3

FUNCTION	PURPOSE	REPLACEMENT
shvec	First-order simplex vector shape element	shcurl

Obsolete Functions in 3.2

FUNCTION	PURPOSE	REPLACEMENT
dxfread	Import geometry from DXF file	geomimport
dxfwrite	Export geometry to DXF file	geomexport
igesread	Import 3D geometry from IGES file	geomimport
stlread	Import 3D geometry from STL file	geomimport
vrmlread	Import 3D geometry from VRML file	geomimport

Obsolete Functions in 3.1

FUNCTION	PURPOSE	REPLACEMENT
flgetrules	Import differentiation rules from FEMLAB 1.1	
flgeomsf2	Set 2D geometry object weights on standard form	
flsde	Indices of edges in a set of subdomains	
flsdp	Indices of points in a set of subdomains	

FUNCTION	PURPOSE	REPLACEMENT
flsdt	Indices of elements in a set of subdomains	
fltrg	Triangle geometry data	

Obsolete Functions in FEMLAB 3.0

FUNCTION	PURPOSE	REPLACEMENT
appl2fem	Expand application mode data to FEM structure	multiphysics
change	Change 2D geometry object	
elemdefault	Return available default element types for an application mode	
faceprim3	Primitive 3D face object	
femiter	Solve stationary PDE problem by iterative methods	fem{n}lin
fldae	Implicit DAE solver	femtime
fldaek	Iterative implicit DAE solver	femtime
fldaspk	Direct or iterative implicit DAE solver	femtime
fleeceng	Energy norm error estimator function	adaption
fleel2	L ² norm error estimator function	adaption
fleelfun	Linear functional error estimator	adaption
fleig	Solve generalized sparse eigenvalue problem	femeig
flgbit	Good Broyden iterative solver	fem{n}lin
flgmres	GMRES iterative solver	fem{n}lin
flisop2p1	Matrix M-file for Navier-Stokes Iso P2-P1 element	
fllrq	Iterative real symmetric definite generalized eigenvalue solver	femeig
flngbit	Good Broyden iterative solver for use with fldaek	femtime
flngmres	GMRES iterative solver for use with fldaek	femtime
flntfqmr	TFQMR iterative solver for use with fldaek	femtime
fltfqmr	TFQMR iterative solver	fem{n}lin
fltpft	Minimize the error for a given number of elements	adaption
fltpqty	Refine a given fraction of the elements	adaption
fltpworst	Refine elements with error greater than a fraction of the worst error	adaption
multigrid	Linear or nonlinear (adaptive) multigrid solver	fem{n}lin
solidprim3	Primitive 3D solid object	

In FEMLAB 3.0, all FEMLAB 2.3 Element Class Methods, Element Library Low-Level Functions, and Shape Function Class Methods are obsolete.

Purpose	Solve PDE problem using adaptive mesh refinement.						
Syntax	fem = adaption(fem,) [fem.sol,fem.mesh] = adaption(fem,)						
Description	<pre>fem = adaption(fem) solves a linear or nonlinear stationary PDE problem or eigenvalue PDE problem. In addition, adaption performs adaptive mesh refinement.</pre>						
	[fem.sol,fem.mesh] = adaption(fem) explicitly returns the solution structure and the adapted mesh object.						
	The function adaption accepts the following property/value pairs:						
	TABLE I-I: VALUE PAIRS						
	PROPERTY Eefun	VALUE fleel2	DEFAULT fleel2	DESCRIPTION Error estimation function			
	Eigselect	vector of positive scalars	1	Weights for eigenmodes			
	Geomnum	integer	1	Geometry number			
	Hauto	positive integer	7	Mesh generation parameter for refinement method meshinit			
	L2scale	vector of positive scalars	1	Scale factors for the L2 error norm			
	L2staborder	vector of positive integers	1	Orders in the stability estimate for the L2 error estimate			
	Maxt	positive scalar	Inf	Maximum number of mesh elements			
	Ngen	scalar integer	5 (1D) 2 (2D) 1 (3D)	Maximum number of refinements			
	Out	fem sol u lambda mesh solcompdof Kc Lc Dc Null ud Nnp uscale stop cell array of these strings	fem [sol,mesh]	Output variables			
	Resmethod	weak coefficient	weak	Residual computation method			

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Resorder	scalar vector	0	Order of decrease of equation residuals
Rmethod	regular longest meshinit	longest	Refinement method
Solver	stationary eigenvalue	stationary	Solver type
Tpfun	fltpft fltpworst fltpqty	fltpft	Element selection method
Tppar	Nonnegative real number		Parameter to the element selection method

TABLE I-I: VALID PROPERTY/VALUE	PAIRS
---------------------------------	-------

In addition, the common solver properties listed under femsolver are available. Also, when using the stationary solver type, the properties listed under femstatic apply, and for the eigenvalue solver type the properties in femeig apply. See therefore the entries femsolver, femstatic, and femeig for more information about the property/values.

Algorithm

The algorithm solves a sequence of PDE problems using a sequence of refined meshes. The first mesh is obtained from the mesh field in the FEM structure. The following generations of meshes are obtained by solving the PDE problem, computing a mesh element error indicator based on the error estimate function, selecting a set of elements based on the element pick function, and then finally refining these elements. The solution to the PDE problem is then recomputed. The loop continues until the maximum number of element generations has been reached, or until the maximum number of elements is obtained.

The PDE problem is stored in the FEM structure fem. See femstruct for details. The adaptive solver works in one geometry at a time. The geometry number is specified in the property geomnum. The solver does only support simplex meshes. The residual computation method weak support all solution forms. The residual computation method coefficient does not support the solution form weak, weak contributions or constraints on subdomains.

First, the solver chosen by the property Solver is called.

Error Estimation

Then, the residuals in the equations are computed for all mesh elements. The *error estimation function* given by the property Eefun is called with parameter Eepar.

There is one predefined function available: fleel2. The function fleel2 computes the error indicator using the L^2 norm, the function fleeceng computes the error indicator using the energy norm, and the function fleelfun computes the error indicator using a linear functional (fleeceng and fleelfun are not yet implemented).

The error estimator gives local error indicators $(f(i,j) h(j)^{\beta(i)})^{\alpha} \operatorname{Vol}(j)$, where *i* is the equation number, *j* is the mesh element number, *h* is the mesh element size, and *Vol* is the mesh element volume. f(i,j) is the scaled absolute value of the *i*th equation residual on the *j*th mesh element. The mesh element error indicator is the sum of these local error indicators over the equation index *i*. The global error indicator is the α root of the sum of the mesh element error indicators over the mesh element index *j*. See the Users Guide for more information on the Adaption Solver Algorithm.

If the eigenvalue solver is used, the weighting of the error indicators for the different eigenfunctions is specified in the property Eigselect. The *n*th component of the Eigselect vector is the weight for the error indicator of the *n*th eigenfunction.

Mesh Refinement

Then, a refinement of the mesh is generated based on the local error indicators. The aim is to refine the mesh most where the errors are largest. The mesh refinements ratios are determined by the function given in the property Tpfun, with parameter Tppar. There are three predefined functions available: fltpft, fltpworst, and fltpqty. fltpft tries to minimize the total error for a prescribed mesh size, namely Tppar times the current number of mesh elements (the default Tppar is 1.7). Each element can be refined several times. fltpworst and fltpqty refine each element at most once. fltpworst refines the elements with an error greater than a fraction of the worst error, whereas fltpqty refines a given fraction of the elements (the fraction is given in the property Tppar, the default is 0.5).

The property Resorder is a scalar or vector which gives the order of decrease of the equation residuals as the mesh size h tends to 0. If it is a vector, the residual of the nth equation is $O(h^{\text{Resorder}(n)})$.

Once the mesh refinement ratios have been determined, the mesh is refined using the method given in the property Rmethod, and the algorithm starts a new iteration. The refinement method is either longest, regular, or meshinit. Details on the first two refinement methods can be found in the entry on meshrefine. Meshinit means that a new mesh is generated through a call to meshinit using the Hmesh property to control the element sizes.

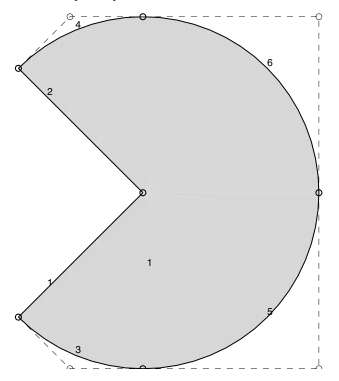
Convergence Control

No more than Ngen successive refinements are attempted. Refinement is also stopped when the number of elements in the mesh exceeds Maxt.

The property Stop makes it possible to return a partially converged solution when the nonlinear, iterative, or eigenvalue solver fails at some point. If a failure occurs, the result from the previous iteration is returned. The output value Stop is 0 if a complete solution was returned, 1 if a partial solution was returned, and 2 if no solution was returned.

When the Report property is on, a progress window is shown. Information about the progress of the adaptive process is printed after each adaptive step. You get a message on the number of mesh elements obtained by the adaptive step, and an error indicator. This error indicator is not an absolute error estimate. In favorable cases there is a constant C such that C times the error indicator is an upper bound of some norm of the error.

EXAMPLES



Adaptive Solver Versus Regular Refinement for the Laplace Equation Solve the Laplace equation over a circle sector

with Dirichlet boundary conditions

 $-\nabla \cdot \nabla u = 0$ u = 0in Ω

$$u = 0$$
 on Γ_1

$$u = \cos\left(\frac{2}{3}\operatorname{atan2}(x, y)\right)$$
 on Γ_2

and compare to the exact solution.

```
clear fem
fem.geom = circ2-poly2([-1 0 -1],[-1 0 1]);
fem.shape = 1;
fem.equ.c = 1; fem.bnd.h = 1;
fem.bnd.r = \{0 \ | \cos(2/3 \tan 2(y,x)) | \};
```

```
fem.bnd.ind = [1 1 2 2 2 2];
fem.solform = 'general';
```

We refine the elements using the L^2 -norm error estimator, with error minimization to achieve a factor of 1.3 in each step, and a maximum number of elements of 500.

We test how many refinements we have to use with an uniform element net:

```
tol = errmax;
fem.mesh = meshinit(fem, 'hmax', 0.3);
fem.xmesh = meshextend(fem);
errmax = 1;
i = 0;
while errmax>tol
 fem.sol = femstatic(fem);
 errmax = postmax(fem,errexpr);
  i = i+1;
 fprintf(1,'Refinement:%3d, error:%3.10f, using mesh:\n',...
          i,errmax);
  fem.mesh
 if errmax>tol
    fem.mesh = meshrefine(fem);
    fem.xmesh = meshextend(fem);
  end
end
```

Note that with uniform refinement, you need much more mesh elements to achieve the same absolute error as with the adaptive method. Also note that the error is reduced only by a small factor when the number of elements is quadrupled by the uniform refinement. For a problem with regular solution, you can expect a $O(h^2)$ error, but this solution is singular because $u \approx r^{1/3}$ at the origin.

Eigenmodes of a Nonconvex Geometry Solve the eigenvalue problem

$$\begin{cases} -\nabla \cdot \nabla u = \lambda u & \text{in } \Omega \\ u = 0 & \text{on } \partial \Omega \end{cases},$$

where the domain Ω is a polygon with some concave corners. Adapt the mesh for the first and second eigenpair and compare. Finally, adapt the mesh using a equally weighted sum of the error estimates of both these eigenpairs.

First set the adaptive solver to eigenvalue and use the L^2 -norm error estimator. Adapt for the first eigenvalue and solve the problem using a maximum number of 500 triangles:

Now solve the same problem but adapt for the second eigenvalue:.

```
fem.mesh = meshinit(fem);
fem.xmesh = meshextend(fem);
fem = adaption(fem,'solver','eigenvalue','eefun','fleel2',...
                                'eigselect',2,'maxt',500);
clf
subplot(211), postsurf(fem,'u','solnum',2), axis equal
subplot(212), meshplot(fem), axis equal
```

Finally, generate a mesh adapted for both these eigenvalues. This can be done by specifying a vector of weights for the errors in each eigenpair:

```
fem.mesh = meshinit(fem);
fem.xmesh = meshextend(fem);
fem = adaption(fem,'solver','eigenvalue','eefun','fleel2',...
                                'eigselect',[1 2],'maxt',500);
clf
subplot(211), postsurf(fem,'u'), axis equal;
subplot(212), meshplot(fem), axis equal;
```

Cautionary The change of solution form to weak in COMSOL Multiphysics 3.3 may make it necessary to add fem.solform = 'general'; to your script models before calling meshextend.

	The Coefficient residual computation method does not support weak equations and does not take other weak contributions into account.
Diagnostics	Upon termination, one of the following messages is displayed:
	• Maximum number of elements reached
	• Maximum number of refinements reached
Compatibility	COMSOL Multiphysics 3.2: the change to weak solution may make it necessary to add fem.solform = 'general'; to your script models before calling meshextend.
	FEMLAB 3.1: error estimators fleelfun, fleeceng, and property Stop are not supported.
	The property Variables has been renamed Const in FEMLAB 2.3.
	The properties epoint and tpoint are obsolete from FEMLAB 2.2. Use fem.***.gporder to specify integration order. See assemble for details.
	The properties toln and normn has been made obsolete from FEMLAB 1.2. Ntol replaces toln.
See Also	femstruct, meshinit, meshrefine, meshextend, femeig, femlin, femnlin

Purpose	Create elliptical or circular arc.
Syntax	<pre>c = arc1(cx,cy,a,b,theta,phi1,phi2) c = arc2(cx,cy,a,b,theta,phi1,phi2) c = arc1(cx,cy,r,phi1,phi2) c = arc2(cx,cy,r,phi1,phi2)</pre>
Description	c = arc1(cx,cy,a,b,theta,phi1,phi2) creates a 2D curve geometry object in the form of an elliptical arc, centered in the coordinate given by cx and cy. The lengths of the semi-axes are a and b, and they are rotated the angle theta. The start and end angles are phi1 and phi2, respectively, and are specified with respect to the semi-axes of the ellipse. The valid range of these angles is 0<=phi1, phi2<2*pi.
	c = arc2(cx,cy,a,b,theta,phi1,phi2) creates a 2D solid geometry object in the form of an elliptical sector.
	c = arc1(cx,cy,r,phi1,phi2) creates a 2D curve geometry object in the form of a circular arc, where r is the radius.
	c = arc2(cx,cy,r,theta,phi1,phi2) creates a 2D solid geometry object in the form of a circular sector.
Examples	The commands below create two circular arc objects, coerce them into one curve object and plot the result.
	<pre>c1 = arc1(0,-1,1,pi/2,3*pi/2); c2 = arc1(0,1,1,3*pi/2,pi/2); g = geomcsg({},{c1,c2}); c = curve2(g) geomplot(c) axis equal</pre>
Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported.
See Also	geom0, geom1, geom2, geom3,curve2, curve3

 Purpose
 Assemble the stiffness matrix, right-hand side, mass matrix, damping matrix, and constraints of a PDE problem.

Syntax

```
[K,L,M,N] = assemble(fem,...)
[K,L,M,N,D] = assemble(fem,...)
[K,L,M,N,D,E] = assemble(fem,...)
[D,M,...] = assemble(fem,'Out',{'D' 'M' ...}, ...)
```

Description assemble is a fundamental function in COMSOL Multiphysics. It assembles a PDE problem using a finite element discretization.

For time-dependent problems, the finite element discretization is the system of ODEs

$$0 = L(U, \dot{U}, \ddot{U}, t) - N_F(U, t)\Lambda$$
$$0 = M(U, t)$$

where L is the residual vector, M is the constraint residual vector, U is the solution vector, and Λ is the Lagrange multiplier vector. The linearization of this system uses the stiffness matrix K, the damping matrix D, the mass matrix E, and the constraint Jacobian matrix N given by

$$K = -\frac{\partial L}{\partial U}, \quad D = -\frac{\partial L}{\partial \dot{U}}, \quad E = -\frac{\partial L}{\partial \ddot{U}}, \quad N = -\frac{\partial M}{\partial U}$$

Here N_F is the constraint force Jacobian matrix. If only ideal constraints are used then

$$N_F = N^T$$
.

All these matrices can depend on the solution vector U. The matrices K, D, and E can also depend on the time derivatives \dot{U} and \ddot{U} .

For a stationary problem, the discretization is

$$0 = L(U) - N_F(U) \Lambda$$
$$0 = M(U)$$

and the linearized problem is

$$K(U - U_0) = L - N_F \Lambda$$
$$NU = M$$

where K, L, M, N and N_F are evaluated for some linearization "point" $U = U_0$. For an eigenvalue problem, the discretization reads

$$KU - (\lambda - \lambda_0)DU + (\lambda - \lambda_0)^2 EU = -N_F \Lambda$$
$$NU = M$$

where K, D, E, N and N_F are evaluated for an equilibrium "point" $U = U_0$. The eigenvalue is denoted by λ , and the linearization point for the eigenvalue by λ_0 .

Valid property/value pairs for the assemble function are given in the following table:

TABLE I-2: VALID PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Assemtol	scalar	1e-12	Assembly tolerance
Blocksize	Positive integer	5000	Assembly block size
Complexfun	on off	off	Use complex-valued functions with real input
Const	cell array		Definitions of constants
Eigname	string		Eigenvalue name
Eigref	string	0	Linearization point for the eigenvalue
Matherr	on off	on	Error for undefined operations
Mcase	non-negative integer	mesh case with largest number of DOFs	Mesh case
Out	K L M N NF D E Ksp A AL BE C DA EA F G GA H Q R cell array of these strings	[K,L,M,N] [K,L,M,N,D] [K,L,M,N,D,E]	Output matrices
Solcomp	cell array of strings		Degree of freedom names to solve for

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Т	scalar	0	Time for evaluation
U	solution object numeric vector scalar	0	Solution for evaluation

TABLE I-2: VALID PROPERTY/VALUE PAIRS

The property Assemtol affects the assembly process. If the local stiffness matrix elements result in a negligible global matrix entry, this element is replaced by a zero. These zeros are removed from the matrices after the assembly process, saving space and computational overhead (in a sparse matrix format the zero matrix entries does not have to be stored). The tolerance is used in a relative sense. Namely if the local matrix contribution (from one element) is $A_l = \{A_{l,ij}\}$ and the currently assembled global matrix is $A = \{A_{ij}\}$ then the entry A_{ij} is replaced by a zero if

 $|A_{ij} + [A_l]_{ij}| < \varepsilon_a \max_{kl} |A_{l,kl}|$

where $[]_{ij}$ denotes the contribution from a local matrix to the global matrix entry ij, and where ε_a is the assembly tolerance controlled by the property Assemtol. For certain types of shape functions the procedure described above is not always safe to perform. This is the case for shape function elements with degrees of freedoms that are of different types, for example when a field variable and its spatial derivative is combined (as in the shherm or sharg_2_5 elements), or when the displacement and displacement angles are combined (as in some Euler Beam elements in the Structural Mechanics Module). For this reason, the above process is never used for local matrix contributions from these types of shape function elements. If the Assemtol is zero then no elements are neglected, but the removal of zeros are still performed. If Assemtol is negative, no elements are neglected, and zeros are not removed from the assembled matrices.

The property Blocksize determines the number of mesh elements that are assembled together in a vectorized manner. A low value gives a lower memory consumption, while a high value might give a better performance.

The properties Complexfun and Matherr are described in femsolver.

The property Const gives a list of definitions of constants to be used in evaluations. This list is a row cell array with alternating constant names and numeric values. This list is appended to the list given in fem.const. If there is a conflict, the definition in Const is used.

The properties Eigname and Eigref are described in femeig. Note that assemble has an empty default value for the property Eigname. So if you want to assemble the matrices for an eigenvalue problem formulated using and eigenvalue name, then the Eigname property must be given. If the variable name lambda is used and if Eigname is not set (and if lambda is not defined in another way), then this variable is evaluated to zero.

The property Out determines which matrices to output. KSp is the sparsity pattern of K. The matrices A, AL, BE, C, and Q are the contributions to the K matrix that come from the coefficients a, α , β , c, and q, respectively. The vectors F, G, and GA are the contributions to the L vector that come from the terms f (or F), g (or G), and γ (or Γ), respectively. The matrix H is the contribution to the N matrix that comes from the h coefficient. The vector R is the contribution to the M vector that comes from the r (or R) coefficient. The matrix DA is the contribution to the D matrix that comes from the d_a coefficient. The matrix EA is the contribution to the E matrix that comes from the e_a coefficient.

The property Solcomp is a cell array of degree of freedom names to solve for. This means that the columns of the matrices D, K, and N correspond to these degrees of freedom. Similarly, the rows of the matrices D, K, and L correspond to these degrees of freedom.

The property T determines for which time the matrices are evaluated.

The property U determines the values of the degrees of freedom for which the matrices are computed (i.e., the linearization point), and also their first and second time derivatives if U is a time-dependent solution object. U can be a solution (femsol) object, a solution vector (this has to be a column vector with values for all the degrees of freedom in the discretized problem), or a scalar (which is expanded to a solution vector).

Sparsity Structure of Finite Element Discretization of Poisson's Equation Assemble the stiffness matrix, right-hand side, and constraint matrices of Poisson's equation

$$\begin{cases} -\Delta u = 1 & \text{in } \Omega \\ u = 0 & \text{on } \partial \Omega \end{cases}$$

where Ω is the unit disk.

clear fem fem.geom = circ2;

Examples

	<pre>fem.mesh = meshinit(fem); fem.shape = 2; fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = 1; fem.xmesh = meshextend(fem); [K,L,M,N] = assemble(fem); n = size(N,1); The sparsity structure of the FEM formulation of the PDE problem is: spy([K,N',L;N,sparse(n,n),M]); The selement of the right assemble to the right hand side. You can assign a descent of the selement of t</pre>
	The column to the right corresponds to the right-hand side. You can continue and solve the PDE problem by using the function femstatic:
	fem.sol = femstatic('In',{'K' K 'L' L 'M' M 'N' N}); postsurf(fem,'u')
Compatibility	FEMLAB 3.0: the properties Context and Sd are not supported and output matrices AS, ALS, BES, CS, DAS, FS, GAS not supported.
	In FEMLAB 2.3, the size of the matrices D, K, L was unaffected by Solcomp. In FEMLAB 3.0, the size of the matrices D, K, L, and N shrinks if Solcomp is a subset of all degree of freedom names.
	The property Variables has been renamed to Const in FEMLAB 2.3.
	The properties bdl, epoint, sdl, tpoint are obsolete from FEMLAB 2.2. Use fem.xxx.gporder to specify integration order.
	The outputs KM, LM, MM, NM, DM, MC, NC, NCL, MU, NU are no longer available in FEMLAB 2.2 and later versions.
	The default value for u and t is 0 in FEMLAB 1.1. In FEMLAB 1.0 it was an error to use u or t in a level 4 expression when the properties u or t were not passed to assemble.
See Also	femstruct, femsolver, femlin, femnlin, femtime, femeig

Purpose	Compute initial	value.		
Syntax		it(fem,) it(fem,'u',femsrc, it(fem,'init',fems		
Description		it(fem,) compute essions in the FEM stru		ject corresponding to the
		it(fem,'u',femsrc,. on femsrc.sol in the s		hese initial value expressions ucture femsrc.
				ers the solution femsrc.sol em, using interpolation.
	if Solnum has ler type as the source	ngth greater than 1, the	en the output so t or U). Otherw	f Init is a solution object or olution object is of the same rise, the output solution first time derivatives).
		seminit accepts the fo	c .	
	PROPERTY	VALUE	DEFAULT	DESCRIPTION
	Blocksize	integer	5000	Assembly block size
	Complexfun	on off	off	Use complex functions with real input
	Const	cell array		Definition of constants
	Framesrc	string cell array of strings	reference frame	Frame for source geometry
	Gmap	integer vector	0	Geometry map
	Init	solution object cell array solution vector string scalar FEM structure		Initial value specification
	Mcase	integer	lowest existing mesh case	Mesh case
	Mcasesrc	integer	Mcase	Mesh case for source solution
	Out	fem sol u	sol	Output

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Outcomp	cell array of strings		Solution components to output
Solnum	integer vector		Solution numbers to use in source solution
Т	real vector		Time for evaluation
U	solution object solution vector scalar FEM structure	0	Solution for evaluation
Xmesh	extended mesh object	fem.xmesh	Extended mesh for source solution

TABLE 1-3:	VALID	PROPERTY/VALUE PAIRS
------------	-------	----------------------

The properties Blocksize, Complexfun, Const, and Outcomp are described in femsolver.

The property Framesrc is needed when mapping a solution after remeshing in a moving mesh simulation. In such a case, the source geometry does not conform to the destination geometry. Rather, the deformed source mesh agrees with the destination geometry if the source mesh is viewed in a certain frame. The property Framesrc contains the name of this frame, or if there are several source geometries, a frame name for each source geometry.

The geometry map vector Gmap tells for each geometry g in the destination FEM structure fem the corresponding geometry number in the source FEM structure, namely Gmap(G). If Gmap(G)=0, there is no corresponding source geometry. The default is a trivial Gmap, that is, the geometry numbers are unchanged.

The initial values are given by the property Init. This can be:

- A solution object or a solution vector, corresponding to the extended mesh Xmesh. That solution will be mapped to the current xmesh (fem.xmesh).
- A cell array of alternating DOF names and expressions, or a single expression. The DOFs will be given the values of the expressions, evaluated for the solution U on the xmesh Xmesh. In this context, a DOF name can also be the name of the time derivative of a DOF. For example, ut is the time derivative of the DOF u.
- An FEM structure. The solution in that FEM structure will be mapped to the current xmesh.
- A scalar. The scalar will be expanded to a solution vector.

	If the property Init is not given, the initial value will be computed by evaluating the initial expressions in the FEM structure for the solution U on the xmesh Xmesh.
	If the source solution (Init or U) is a numeric vector, its corresponding mesh case number can be given in the Mcasesrc property.
	The property Solnum gives the solution numbers to use in the source solution. If the source solution is time-dependent, interpolation at the times in T can be used instead. By default, only the last solution (first solution for eigensolutions) is used.
Compatibility	The properties context, initmethod, and linsolver are obsolete from FEMLAB 3.0.
	The property Variables has been renamed to Const in FEMLAB 2.3.
See Also	assemble, femsolver, femlin, femnlin, femtime, adaption, meshextend

Purpose	Create a right-angled block geometry object.
Syntax	<pre>obj = block3 obj = block2 obj = block3(lx,ly,lz,) obj = block2(lx,ly,lz,)</pre>
Description	obj = block3 creates a right-angled solid block object with all side lengths equal to 1, one corner at the origin, and the local <i>z</i> -axis equal to the global <i>z</i> -axis. block3 is a subclass of solid3.

obj = block3(lx, ly, lz, ...) creates a right-angled solid block geometry object with positive side lengths 1x, 1y, and 1z. 1x, 1y, and 1z are positive real scalars, or strings that evaluate to positive real scalars, given the evaluation context provided by the property const.

The functions block3/block2 accept the following property/values:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object
base	corner center	corner	Positions the object either centered about pos or with one corner in pos
const	Cell array of strings	{}	Evaluation context for string inputs
pos	Vector of reals or cell array of strings	[0 0]	Position of the object
rot	real or string	0	Rotational angle about axis (radians)

TABLE I-4: VALID PROPERTY/VALUE PAIRS

axis sets the local z-axis, stated either as a directional vector of length 3, or as a 1-by-2 vector of spherical coordinates. axis is a vector of real scalars, or a cell array of strings that evaluate to real scalars, given the evaluation context provided by the property const. See gency13 for more information on axis.

pos sets the position of the object, either centered about the position or with one corner in the position. The corresponding values of base are center and corner. pos is a vector of real scalars, or a cell array of strings that evaluate to real scalars, given the evaluation context provided by the property const.

rot is an intrinsic rotational angle for the object, about its local z-axis provided by the property axis. rot is a real scalar, or a string that evaluate to a real scalar, given the evaluation context provided by the property const. The angle is assumed to be in radians if it is numeric, and in degrees if it is a string.

obj = block2(...) creates a right-angled surface block geometry object with properties as given for the block3 function. block2 is a subclass of face3.

Block objects have the following properties:

PROPERTY	DESCRIPTION		
lx, ly, lz	Side lengths		
base	Base point		
x, y, z, xyz	Position of the object. Components and vector forms		
ax2	Rotational angle of symmetry axis		
ax3	Axis of symmetry		
rot	Rotational angle		

TABLE I-5: BLOCK OBJECT PROPERTIES

In addition, all 3D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom3 for details.

Examples The following commands create a surface and solid block object, where the position is defined in the two alternative ways.

	b1 = block2(1,2.1,0.5,'base','center','pos',[1 0 1], 'axis',[0 0 1],'rot',0)
	get(b1,'xyz') b2 = block3(1,1,1,'base','corner','pos',[-1 -1 -1]) get(b2,'xyz')
Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported.
See Also	face3, hexahedron2, hexahedron3

Create flattened corners in 2D geometry object.

Purpose Syntax

g = chamfer(g1,...)

Description

g = chamfer(g1,...) creates flattened corners in 2D geometry object g1 according to given property values.

The function chamfer accepts the following property/values:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
angles	I-by- <i>m</i> vector		Angles (in radians) with respect to edges in first row of edges
dist1	I-by- <i>m</i> vector		Distances along edges in the first row of edges. A positive entry states the chamfered length and a negative entry states the remaining length of edge after chamfering
dist2	I-by- <i>m</i> vector		Distances along edges in second row of edges, using same format as dist1
edges	2-by- <i>m</i> matrix		Pairs of edge numbers
lengths	I-by- <i>m</i> vector		Lengths of line segments that make up the flattened corners
out	Cell array of strings	none	Determines the output
point	integers all none	all	out

TABLE I-6: VALID PROPERTY/VALUE PAIRS

The corners to chamfer is either specified with either the property point or edges. The default value is the all corners are chamfered.

The size of the chamfer is specified with any of the following combinations of properties: dist1 and dist2; dist1 and angles; dist1 and lengths; and length and angles. If only dist1 is supplied the chamfering distance is equal for both edges. All these properties can be given as a vector or as a single value.

Examples

Chamfer a rectangle in different ways.

```
r = rect2;
s1 = chamfer(r,'dist1',0.1);
s2 = chamfer(r,'edges',[1 2;2 3],'angles',pi/4,'lengths',0.5);
```

Diagnostics	If a chamfer cannot be created according to the specified properties this corner is ignored.
	When the chamfers generates intersections with other edges in the geometry, an error message is given.
Compatibility	FEMLAB 3.0: The property trim is no longer supported. Only pair of edges that have a common vertex can be chamfered. For edges that are not linear, the linear approximation of the edge in the corner is used to compute a chamfer.
See Also	curve2, curve3,fillet

Purpose

Create circle geometry object.

obj	=	circ2
obj	=	circ1
obj	=	circ2(r,)
obj	=	circ1(r,)

Description

obj = circ2 creates a solid circle geometry object with radius 1, centered at the origin. circ2 is a subclass of ellip2 and solid2.

obj = circ2(r,...) creates a circle object with radius r, centered at the origin. r is a positive real scalar, or a string that evaluates to a positive real scalar, given the evaluation context provided by the property const.

The functions circ2/circ1 accept the following property/values:

TABLE I-7: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
base	corner center	center	Positions the object either centered about pos or with the lower left corner of surrounding box in pos
const	Cell array of strings	{}	Evaluation context for string inputs
pos	Vector of reals or cell array of strings	[0 0]	Position of the object
rot	real or string	0	Rotational angle about pos (radians)

obj = circ1(...) creates a curve circle geometry object with properties as given for the circ2 function. circ1 is a subclass of ellip1 and curve2.

Circle objects have the following properties:

TABLE I-8: CIRCLE OBJECT PROPERTIES

PROPERTY	DESCRIPTION
r	Radius
base	Base point
х, у	Position of the object
rot	Rotational angle

In addition, all 2D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom2 for details.

Examples	The commands below create a unit solid circle geometry object and plot it.
	<pre>c1 = circ2(1,'base','center','pos',[0 0]); get(c1,'base') geomplot(c1)</pre>
Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported.
See Also	ellip1, ellip2,curve2, curve3

Purpose	Start COMSOL software products.
Syntax	comsol comsol server comsol reaction
Description	comsol starts the COMSOL Multiphysics graphical user interface from the COMSOL Script command prompt or from COMSOL Script or MATLAB.
	comsol server starts a COMSOL Multiphysics server within the COMSOL Script or MATLAB process. You can connect to the COMSOL Multiphysics server from a COMSOL Multiphysics client. The COMSOL Multiphysics client must be started outside COMSOL Script or MATLAB.
	comsol reaction starts the COMSOL Reaction Engineering Lab graphical user interface from within the COMSOL Script or a MATLAB process. You can connect
	There is also a comsol command available on the command prompt in Windows, UNIX/Linux, and Mac. Using the COMSOL command you can start COMSOL Multiphysics running stand alone. You can also start a COMSOL Multiphysics client for connecting to a COMSOL Multiphysics server. The options to this command are listed in the <i>Installation Guide</i> .

Purpose	Create a circular cone geometry object.				
Syntax	<pre>c3 = cone3 c2 = cone2 c3 = cone3(r,h) c2 = cone2(r,h) c3 = cone3(r,h,ang) c2 = cone2(r,h,ang) c3 = cone3(r,h,ang,) c2 = cone2(r,h,ang,)</pre>				
Description	c3 = cone3 creates a solid circular cone geometry object with bottom r height equal to 1, top radius equal to 0.5, and the center of the bottom at cone3 is a subclass of econe3.			• •	
	c3 = cone3(r,h) creates a solid circular cone geometry object, with bottom radius r, height h, and top radius $r/2$.				
	c3 = cone3(r,h,ang) creates a solid circular cone geometry object, with bottom radius r, height h, and the angle ang between the local z-axis and a generator of the conical surface. ang is given in radians in the interval [0,pi/2].				
	The functions cone3/cone2 accept the following property/values:				
	TABLE I-9: VALID PROPERTY/VALUE PAIRS				
	PROPERTY	VALUE	DEFAULT	DESCRIPTION	
	axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object	
	const	Cell array of strings	{}	Evaluation context for string inputs	
	pos	Vector of	[0 0]	Position of the object	

rot real or string 0 Rotational angle about axis (radians)

reals or cell array of strings

For more information on input arguments and properties see gency13.

c2 = cone2(...) creates a surface circular cone geometry object without bottom and top faces, according to the arguments as described for cone3. cone2 is a subclass of econe2. Cone objects have the following properties:

TABLE I-I	0: CONE	OBJECT F	PROPERTIES

	PROPERTY	DESCRIPTION
	r	Radius
	h	Height
	ang	Semi-angle
	x, y, z, xyz	Position of the object. Components and vector forms
	ax2	Rotational angle of symmetry axis
	ax3	Axis of symmetry
	rot	Rotational angle
	accessed using the sy	eometry object properties are available. All properties can be ntax get(object,property). See geom3 for details. om for more information on geometry objects.
Compatibility	•	ntax is obsolete but still supported. The numbering of faces, different from the numbering in objects created in 2.3.
Examples	Create a cone with a	n apex
	h = 2; r = 1; c3 = cone3(r,h, get(c3,'ang')	atan(r/h));
	Created truncated an	nd rotated cone
		,atan(0.7*r/h),'pos',[1 -2 4], is',[1 -1 0.3],'rot',pi/3);
See Also	cylinder2, cylind geom1, geom2, geo	der3,econe2, econe3,face3,gencyl2, gencyl3,geomO, om3,geomcsg

Purpose	Create a curve object.
Syntax	<pre>c3 = curve3(x,y,z) c3 = curve3(x,y,z,w) c3 = curve3(vtx,vtxpre,edg,edgpre,fac,mfdpre,mfd) [c3,] = curve3(g3,) c3 = curve3(g2) c2 = curve2(x,y) c2 = curve2(x,y,w) c2 = curve2(vtx,edg,mfd) [c2,] = curve2(g2,)</pre>
Description	c3 = curve3(x,y,z) creates a 3D curve object. The degree is determined from the number of control points given in the vectors, x, y, and z. Length 2 generates a straight line. Lengths 3 and 4 generates rational Bézier curves of degrees 2 and 3 respectively. Unit weights are used.
	c3 = curve3(x,y,z,w) works similarly to the above, but also applies the positive weights w to the control points of the curve.
	c3 = curve3(vtx,vtxpre,edg,edgpre,fac,mfdpre,mfd) creates 3D curve geometry object c3 from the arguments vtx, vtxpre, edg, edgpre, fac, mfdpre, and mfd. The arguments must define a valid 3D curve object. See geom3 for a description of the arguments.
	[c3,] = curve3(g3,) coerces the 3D geometry object g3 to a 3D curve object c3.
	c3 = curve3(g2) coerces the 2D geometry object g2 to a 3D curve object c3, by embedding it in the plane $z = 0$.
	c2 = curve2(x,y) creates a 2D curve object in the form of a Bézier curve, with the control points given by the vectors x and y of the same lengths. Length 2 generates a straight line. Lengths 3 and 4 generates rational Bézier curves of degrees 2 and 3, respectively. Unit weights are used.
	c2 = curve2(x, y, w) works similarly to the above, but also applies the positive weights w to the control points of the curve.
	c2 = curve2(vtx,edg,mfd) creates a 2D curve object from the properties vtx, edg, and mfd. The arguments must define a valid 2D curve object. See geom2 for a description of the arguments.
	[c2,] = curve2(g2,) coerces the 2D geometry object g2 to a 2D curve object.

The coercion functions [c2,...] = curve2(g2,...) and [c3,...] = curve3(g3,...) accept the following property/values:

TABLE I-II: VALID PROPERTY/V	ALUE PAIRS
------------------------------	------------

PROPERTY	VALUE	DEFAULT	DESCRIPTION
out	stx ftx ctx ptx	{}	Cell array of output names.

See geomcsg and geom for more information on geometry objects.

The nD geometry object properties are available. The properties can be accessed using the syntax get(object,property). See geom for details.

Examples The commands below compute the union of a unit circle and a unit square, coerce the solid object to a curve object, and plot the result.

```
s = circ2+square2;
c = curve2(s);
geomplot(c)
```

The following commands generate and plot an elliptic 3D arc:

```
c = curve3([0 1 2],[0 1 0],[0 1 2],[1 1/sqrt(2) 1]);
geomplot(c)
```

Compatibility The FEMLAB 2.3 syntax is obsolete but still supported.

See Also face3, geom0, geom1, geom2, geom3, geomcsg, point1, point2, point3

Purpose	Create a cylinder geometry object.
Syntax	<pre>c3 = cylinder3 c2 = cylinder2 c3 = cylinder3(r,h) c2 = cylinder2(r,h) c3 = cylinder3(r,h,) c2 = cylinder2(r,h,)</pre>
Description	c3 = cylinder3 generates a solid cylinder object, with radius and height equal to 1, axis along the z-axis and bottom surface centered at the origin. cylinder3 is a subclass of cone3.

c3 = cylinder3(r,h) generates a solid cylinder object with radius r and height h.

The functions cylinder3/cylinder2 accept the following property/values:

TABLE I-12: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object
const	Cell array of strings	{}	Evaluation context for string inputs.
pos	Vector of reals or cell array of strings	[0 0]	Position of the bottom surface
rot	real or string	0	Rotational angle about axis (radians)

For more information on input arguments and properties, see gency13.

c2 = cylinder2(...) creates a surface cylinder object, from arguments as described for cylinder3. cylinder2 is a subclass of cone2.

Cylinder objects have the following properties:

TABLE I-13: CYLINDER OBJECT PROPERTIES

PROPERTY	DESCRIPTION
r	Radius
h	Height
x, y, z, xyz	Position of the object. Components and vector forms
ax2	Rotational angle of symmetry axis

TABLE I-13: CYLINDER OBJECT PROPERTIES

	PROPERTY	DESCRIPTION		
	ax3	Axis of symmetry		
	rot	Rotational angle		
	In addition, all 3D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom3 for details.			
	See geomesg an	nd geom for more information on geometry objects.		
Examples	The following commands generates a surface cylinder object and a solid cylinder object.			
		der2(0.5,4,'pos',[1,1,0],'axis',[pi/2,0]); der3(20,40,'pos',[0,0,-100],'axis',[1,1,1]);		
Compatibility		2.3 syntax is obsolete but still supported. The numbering of faces, ces is different from the numbering in objects created in 2.3.		
See Also	gencyl2, gen geomcsg	cyl3,cone2, cone3,face3,geom0, geom1, geom2, geom3,		

Purpose	Discrete sine transform.
Syntax	y = dst(x) x = idst(y)
Description	$y = dst(x)$ computes the discrete sine transform (DST) of the columns of x. For best performance, the number of rows in x should be $2^{m}-1$, for some integer m.
	y = dst(x,n) pads or truncates the vector x to length n before transforming.
	If x is a matrix, the dst operation is applied to each column.
	$x = idst(y)$ calculates the inverse discrete sine transform of the columns of y. For best performance, the number of rows in y should be 2^m-1 , for some integer m.
	x = idst(y,n) pads or truncates the vector y to length n before transforming.
	If y is a matrix, the idst operation is applied to each column.
See Also	poisson

Purpose	Create eccentric cone geometry object.
Syntax	<pre>ec3 = econe3 ec2 = econe2 ec3 = econe3(a,b,h) ec2 = econe2(a,b,h) ec3 = econe3(a,b,h,rat) ec2 = econe2(a,b,h,rat) ec3 = econe3(a,b,h,rat,) ec2 = econe2(a,b,h,rat,)</pre>
Description	ec3 = econe3 creates a solid eccentric cone geometry object semi-axes of the elliptical bottom surface equal to one, axis al <i>z</i> -axis, and the center of the bottom surface at the origin. ecc

ct with height and along the coordinate cone3 is a subclass of gency13.

ec3 = econe3(a,b,h) creates a solid eccentric cone geometry object with semi-axes a and b, and height h.

ec3 = econe3(a,b,h,rat) creates a cone with the non-negative ratio rat between the top and bottom surface.

The functions econe3/econe2 accept the following property/values:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object
const	Cell array of strings	{}	Evaluation context for string inputs
displ	2-by- <i>nd</i> matrix	[0;0]	Displacement of extrusion top
pos	Vector of reals or cell array of strings	[0 0]	Position of the bottom surface
rot	real or string	0	Rotational angle about axis (radians)

TABLE I-14: VALID PROPERTY/VALUE PAIRS

For more information on input arguments and properties see gency13.

ec2 = econe2(...) creates a surface eccentric cone geometry object, without bottom and top faces, according to the arguments described for econe3. econe2 is a subclass of gency12.

Eccentric cone objects have the following properties:

TABLE I-15: ECCENTRIC CONE OBJECT PROPERTIES

PROPERTY	DESCRIPTION	
a, b	Semi-axes	
r	Radius	
h	Height	
rat	Ratio	
x, y, z, xyz	Position of the object. Components and vector forms	
ax2	Rotational angle of symmetry axis	
ax3	Axis of symmetry	
rot	Rotational angle	

In addition, all 3D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom3 for details.

Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported. The numbering of faces, edges and vertices is different from the numbering in objects created in 2.3.
Examples	Create a truncated eccentric cone with the basis surface in the <i>xy</i> -plane.
	e = econe2(10, 40, 20, 0.5)
	Create an eccentric cone with an apex, that is, a singular patch, on top.

e = econe3(1,2,4,0,'displ',[1,1],'pos',[100 100 100],... 'axis',[0 1 4],'rot',pi/4)

```
See Also cone2, cone3, gency12, gency13, face3
```

Purpose	Define coefficient or general form constraints.
Syntax	<pre>el.elem = 'elcconstr' el.g{ig} = geomnum el.form = 'coefficient' 'general' el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.dim{idim} = dimvarname el.geomdim{ig}{edim}.r{eldomgrp} = rvec el.geomdim{ig}{edim}.h{eldomgrp} = hmat el.geomdim{ig}{edim}.cpoints{eldomgrp}{ic} = cpind</pre>
Description	The elcconstr element adds a set of constraints specified in coefficient or general form, as specified by the el.form field, to the FEM problem. For the syntax of the ind field, see elempty. The coefficient rvec has the same syntax as the fem.bnd.r field, while hmat corresponds to an fem.bnd.h entry. See further the section "Specifying a Model" in the COMSOL Multiphysics Scripting Guide. The cpoints field differs from fem.bnd.cporder in that it contains pattern indices instead of orders, see elepspec.
	Dirichlet boundary conditions are implemented using elcconstr elements if the solution form is Coefficient or General. When assembling in the Weak solution form, an elpconstr elements replaces the elcconstr.
Examples	<pre>In a 2D model, add a Dirichlet boundary condition on u at boundary 1 and 2 using constraint point pattern 1: el.elem = 'elcconstr'; el.g = {'1'}; el.form = 'coefficient'; gd.ind = {{'1', '2'}}; gd.dim = {'u'}; gd.r = {{'0'}}; gd.h = {{'1'}}; gd.h = {{'1'}}; el.geomdim{1} = {{},gd,{}; fem.elem = [fem.elem {el}]; } }</pre>
See Also	elempty, elpconstr, elcurlconstr, elepspec, eleqc

Purpose	Define global expression variables.
Syntax	el.elem = 'elconst' el.var{2*ivar-1} = varname el.var{2*ivar} = varexpr
Description	The elconst element declares expression variables varname to be accessible across all geometries and dimensions. The defining expressions, varexpr, can contain any variables, including variables that are only present on some domains. Expressions are expanded in the context where evaluation is requested.
Examples	<pre>Add global expressions for the transformation between Cartesian and cylindrical polar coordinates. clear el; el.elem = 'elconst'; el.var = {'r','sqrt(x^2+y^2)','phi','atan2(y,x)'};</pre>
See Also	<pre>fem.elem = [fem.elem {el}]; elempty</pre>

Purpose	Define contact map operators.
Syntax	<pre>el.elem = 'elcontact' el.g{ig} = geomnum el.opname{iop} = opname el.mphname{iop} = mphname el.gapname{iop} = gapname el.contname{iop} = contname el.conttol{iop} = 'auto' abstol el.visname{iop} = visname el.method{iop} = 'direct' 'ball' el.checkdist{iop} = chkdist el.srcframe{iop} = frame el.srcn{iop}{idim} = srcnx_i el.dstx{iop}{idim} = dstx_i el.dstn{iop}{idim} = dstnx_i</pre>
	el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.src{iop} = eldomgrplist
Description	The elcontact element defines contact map operators and related gap distance, contact flag and visibility flag variables. When evaluated, a contact map operator searches for the closest source (or <i>master</i>) point found following a ray in the dstn direction from the point given by the dstx expressions evaluated at the destination (or <i>slave</i>) point.
	For each operator name, a set of master domains is specified by listing one or more domain group indices in the corresponding el.geomdim{ig}{edim}.src{iop} field. Normal direction expressions for the master domains are specified in the el.srcn{iop}{idim} fields. Coordinate expressions for the master are obtained indirectly from the frame specified in the el.srcframe{iop} field.
	The map operator allows evaluation of any expression at a corresponding master point, while the gap distance variable evaluates to the distance between master and slave point. The optional mphname field for each map operator gives the name of a corresponding multiphysics operator. This second operator evaluates to the same value as the main contact operator, but its Jacobian does not contain any contribution from the map (mesh position), only from the argument expression
	The contact flag variable evaluates to a nonzero value if th.e master point is well defined and the gap distance less than the conttol treshold value. The visibility flag variable is nonzero if a corresponding master point was found for the slave point where the flag is evaluated.

	There are two slightly different methods available to search for master points. The 'direct' method is a clean and stable direct search algorithm while the 'ball' method is faster by only treating master elements inside a given ball radius accurately. This ball radius can be set using the checkdist field, and should normally be larger than any mesh element taking part in the search.
Cautionary	The elcontact element is only implemented for boundaries. That is, no edge-edge, edge-boundary or similar contact can be detected and evaluated.
	The computation of complete Jacobians rely on the availability of spatial derivatives of the mapped expression with respect to local coordinate directions. These local derivatives cannot be calculated for all variables, a notable example being any global spatial derivatives. Therefore, for expressions like $map(uTx)$ some Jacobian contributions will be missing.
Examples	Evaluate and display the distance from a hard surface to the closest point of a cylinder lying on its side on the surface.
	clear fem; fem.geom = rect2(2,0.2,'pos',[-1,-0.2])+circ2(0.8,'pos',[0,0.8]); fem.mesh = meshinit(fem); fem.sshape = 2;
	<pre>clear el; el.elem = 'elcontact'; el.g = {'1'}; el.opname = {'map'}; el.gapname = {'gap'}; el.wisname = {'vis'}; el.method = 'ball'; el.checkdist = '1'; el.srcframe = {'xy'}; el.srcn = {{'nx', 'ny'}}; el.dstx = {{'x', 'y'}; el.dstn = {{'nx', 'ny'}};</pre>
	clear src11 src11.ind = {{'3','4'}}; src11.src = {{'1'}}; el.geomdim{1} = {{},src11};
	<pre>fem.elem = {el}; fem.xmesh = meshextend(fem); fem.sol = asseminit(fem);</pre>

```
postcrossplot(fem,1,[6 8],'lindata','if(vis,gap,0)',...
'linxdata','if(vis,map(x),sign(x))');
% compare to the theoretical value
hold on;
x=-1:0.05:1;
plot(x,sqrt(x.^2+0.8^2)-0.8,'ro')
See Also elmapextr
```

Purpose	Define extrusion coupling variables.
Syntax	<pre>el.elem = 'elcplextr' el.g{ig} = geomnum el.var{ivar} = varname el.map{imap} = linmap genmap unitmap el.usenan = 'true' 'false' el.extttol = tol el.src{ig}{edim}.ind{srcdomgrp} = domainlist el.src{ig}{edim}.map{ivar}{srcdomgrp} = imap el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.map{ivar}{eldomgrp} = imap linmap.type = 'linear' linmap.sg = srcig linmap.sv{ivtx} = srcvtx linmap.sframe = srcframe linmap.dy = dstig linmap.dv{ivtx} = dstvtx linmap.drame = dstframe genmap.type = 'local' genmap.expr{idim} = transexpr genmap.frame = frame unitmap.type = 'unit' unitmap.frame = frame</pre>
Description	The elcplextr element declares the extrusion coupling variable names listed in the var field to be accessible on domains where the corresponding destination map field geomdim{ig}{edim}.map{ivar}{eldomgrp} entry is nonempty. Both destination and source map fields contain indices into the map field which consists of a list of transformation specifications. The available transformation types are 'linear', 'local', and 'unit', each with its own syntax. The unit transformation takes one optional argument specifying which frame is to be used for evaluating the mesh position. If not given, the reference frame is assumed. Unit transformations without a frame field actually never has to be specified explicitly, since using index zero in the source and destination map fields is interpreted as an implicit unit transformation. The local transformation, which is called "general" in the COMSOL Multiphysics user interface, lets you specify an arbitrary expression for each source dimension.
	These expressions can contain spatial coordinate variables from any frame. The frame field decides which frame to use in the search operation when the local

transformation is used as source transformation. Therefore, choose a frame such that the transformation is as linear as possible relative to the given frame.

Linear transformations are described in the section on coupling variables in the "Analysis Guide" chapter of the *COMSOL Multiphysics User's Guide*. Note that the source geometry and vertex fields linmap.sg and linmap.sv refer to the vertices that are used as source for the *transformation*, which are usually related to the coupling variable destination domain. This is because linear maps are best used as destination maps, specifying a map from the destination domain into the source domain. The frame fields specify which coordinate set to use in evaluating the vertex positions.

The coupling variable source transformation and expression is set up using the src field. A separate domain grouping is specified for the source dimensions which does not contribute to the global domain group splitting. Source expressions in the expr{ivar}{srcdomgrp} field can be left as empty cell arrays to signify that the particular source domain group is not part of the source for a given variable.

CautionaryParameter or time dependency in the source transformation is not properly detected
by the solvers, which means that the source transformation will not be updated
between parameter or time steps in that case. Solution dependencies in the
transformation are properly detected, but do not give any Jacobian contributions
from the transformation.

Examples

Calculate the first ten eigenvalues of a 3-by-2 rectangle with periodic boundary conditions both left-right and top-bottom. Different map types are used.

```
fem.geom = rect2(3,2);
fem.mesh = meshinit(fem, 'hmax',0.05);
fem.equ.c = 1;
fem.equ.da = 1;
fem.bnd.ind = [0 1 2 0];
fem.bnd.constr = {'ucx-u', 'ucy-u'};
fem.elem = {};
el.elem = 'elcplextr';
el.g = {'1'};
el.var = {'ucx', 'ucy'};
clear map1;
map1.type = 'linear';
map1.sg = '1';
map1.sv = {'2','3'};
map1.dg = '1';
```

```
map1.dv = {'1','4'};
  clear map2;
  map2.type = 'local';
  map2.expr = \{ 'x' \};
  el.map = {map1 map2};
  clear src;
  src.ind = {{'1'},{'4'}};
  src.expr = {{{}, 'u'}, {'u', {}};
  src.map = {{{},'0'},{'2',{}}};
  el.src{1} = {{},src,{}};
  clear dst;
  dst.ind = {{'2'},{'3'}};
  dst.map = {{ '1', {}}, {{}, '2'}};
  el.geomdim{1} = {{},dst,{}};
  fem.elem = [fem.elem {el}];
  fem.xmesh = meshextend(fem);
  fem.sol = femeig(fem, 'neigs', 10, 'shift', 1);
  postplot(fem,'tridata','u','triz','u','refine',3,'solnum',8);
elempty, elcplproj
```

See Also

Purpose	Define destination-aware integration coupling variables.
Syntax	<pre>el.elem = 'elcplgenint' el.g{ig} = geomnum el.var{ivar} = varname el.global = varlist el.src{ig}{edim}.ind{srcdomgrp} = domainlist el.src{ig}{edim}.expr{ivar}{srcdomgrp} = srcexpr el.src{ig}{edim}.ipoints{ivar}{srcdomgrp} = ip el.src{ig}{edim}.iorders{ivar}{srcdomgrp} = io el.src{ig}{edim}.frame{ivar}{srcdomgrp} = frame el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.usage{ivar} = eldomgrplist</pre>
Description	The elcplgenint element accepts the same syntax as the elcplscalar element with the only notable exception that a destination operator dest(subexpr) can be used in the source expression. The destination operator's argument will be evaluated on the destination point instead of on the source domain. This can be used to evaluate convolution integrals.
Cautionary	The integral is evaluated for each destination point, whether the dest() operator is present in the source expression or not. Use an elcplscalar element if there is no destination dependence.
Examples	<pre>Plot part of the Fourier transform of g= x <1. clear fem; fem.geom = geom1([-10 10]); fem.mesh = meshinit(fem,'report','off','hmax',0.1); fem.equ.gporder = 4; fem.elem = {}; clear el el.elem = 'elcplgenint'; el.g = {'1'}; el.var = {'G'}; clear src; src.expr = {{'(abs(x)<1)*exp(-i*dest(x)*x)'}}; src.iorders = {'4'}; el.src = {{{},src}; clear dst; dst.usage = {{'1'}}; el.geomdim = {{},dst}; fem.elem = [fem.elem {el}]; fem.xmesh=meshextend(fem); postplot(fem,'lindata','G','liny','G');</pre>

elepspec, elcplscalar, elgpspec

Purpose	Define projection coupling variables.
Syntax	<pre>el.elem = 'elcplproj' el.g{ig} = geomnum el.var{ivar} = varname el.map{imap} = projmap linmap genmap unitmap el.src{ig}{srcdim}.ind{srcdomgrp} = domainlist el.src{ig}{srcdim}.expr{ivar}{srcdomgrp} = srcexpr el.src{ig}{srcdim}.iorder{ivar}{srcdomgrp} = intorder el.src{ig}{srcdim}.map{ivar}{srcdomgrp} = imap el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.map{ivar}{eldomgrp} = imap</pre>
	<pre>projmap.type = 'projection' projmap.sg = srcig projmap.sv{ivtx} = srcvtx projmap.sframe = srcframe projmap.dg = dstig projmap.dv{ivtx} = dstvtx projmap.dframe = dstframe</pre>
	<pre>linmap.type = 'linear' linmap.sg = srcig linmap.sv{ivtx} = srcvtx linmap.sframe = srcframe linmap.dg = dstig linmap.dv{ivtx} = dstvtx linmap.dframe = dstframe</pre>
	genmap.type = 'local' genmap.expr{idim} = transexpr genmap.frame = frame
	unitmap.type = 'unit unitmap.frame = frame'
Description	The elcplproj projection variable element is closely related to the elcplextr element. Both elements map the srcdim-dimensional source domain onto an intermediate srcdim-dimensional fictitious domain. While the destination transformation in the elcplextr element maps the destination domain into the intermediate domain, the elcplproj element destination transformation maps the destination only into the first srcdim-1 dimensions. The last dimension of the fictitious domain is collapsed by integration onto the first srcdim-1 dimensions. See the section on coupling variables in the "Analysis Guide" chapter of the <i>COMSOL Multiphysics User's Guide</i> for more information.

	All map types available for elcplextr can be used also in projection coupling variables. The common combination of a unit source map and a linear destination map is not very useful, though. Instead, there is a map type projection which specifies srcdim+1 vertices in the source geometry and srcdim vertices in the destination. The basis defined by vectors from the first source vertex to each of the remaining vertices is mapped onto a right-handed orthogonal system with unit axes. The basis described by the destination vertices is then mapped onto the first srcdim-1 dimensions of the same orthogonal basis. This means that the direction of integration is effectively from the first source vertex to the last. In addition to the fields present in the elcplextr element, the elcplproj requires an integration order for the line integrals evaluated for each destination point. The iorder field specifies the order of polynomials that should be exactly integrated.
Cautionary	Projection coupling is only implemented for simplex meshes. When finding integration limits, the elcplproj element works directly on the basic polyhedral mesh. Therefore, results can be inaccurate if the mesh does not properly resolve the geometry.
	Parameter or time dependency in the source transformation is not properly detected by the solvers, which means that the source transformation will not be updated between parameter or time steps in that case. Solution dependencies in the transformation are properly detected, but do not give any Jacobian contributions from the transformation.
	The automatic detection of nonlinear and time or parameter dependent problems does not work properly in that all problems containing projection coupling variables are considered to be nonlinear and time dependent.
Examples	<pre>Project the diagonal cross section distance on the left and bottom edges of a square. clear fem fem.geom = square2; fem.mesh = meshinit(fem); fem.elem = {}; el.elem = 'elcplproj'; el.g = {'1'}; el.var = {'d','d'}; clear map1; map1.type = 'projection'; map1.sg = '1'; rend out = {'11', '01', '01'};</pre>
	map1.sv = {'1','2','3'};

```
map1.dg = '1';
map1.dv = {'1', '2'};
clear map2;
map2.type = 'projection';
map2.sg = '1';
map2.sv = {'1', '4', '3'};
map2.dg = '1';
map2.dv = \{ '1', '4' \};
el.map = {map1 map2};
clear src;
src.ind = {{'1'}};
src.expr = {{'1'},{'1'}};
src.iorder = {{'1'},{'1'}};
src.map = {{'1'},{'2'}};
el.src{1} = {{},{},src};
clear dst;
dst.ind = {{'1'},{'4'}};
dst.map = {{'1',{}},{{},'2'}};
el.geomdim{1} = {{},dst,{}};
fem.elem = [fem.elem {el}];
fem.xmesh = meshextend(fem);
postint(fem, 'd/sqrt(2)', 'edim', 1, 'dl', [1 4])
```

elempty, elcplextr

Purpose	Define integration coupling variables.
Syntax	el.elem = 'elcplscalar' el.g{ig} = geomnum el.var{ivar} = varname el.global = varlist el.maxvars = maxvarlist
	<pre>el.src{ig}{edim}.ind{srcdomgrp} = domainlist el.src{ig}{edim}.expr{ivar}{srcdomgrp} = srcexpr el.src{ig}{edim}.ipoints{ivar}{srcdomgrp} = ip el.src{ig}{edim}.iorders{ivar}{srcdomgrp} = io el.src{ig}{edim}.frame{ivar}{srcdomgrp} = frame el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.usage{ivar} = eldomgrplist</pre>
Description	The elcplscalar element declares the integration coupling variable names listed in the var field to be accessible on domain groups specified as a, possibly empty, cell array for each variable in the usage field, or globally if the variable index is mentioned in the global field. The same variable cannot be defined as both global and local.
	If a variable index is included in the maxvars list, the elcplscalar element computes an approximate maximum value of the source expression over the source domains, instead of the integral. The expression is evaluated and compared only in the same quadrature points as would otherwise have been used for integration; see below.
	The source domain grouping specified in the src{ig}{edim}.ind field does not contribute to the global domain splitting. For each variable and source domain group, a possibly empty (no contribution) source expression is given in the expr field.
	The ipoint field specifies an integration point pattern using indices referring to an elgpspec element. If the ipoints field is not present, the iorders field will be read instead and assumed to contain Gauss-Legendre quadrature orders for the source expressions. When specifying an elcplscalar element in a script, the latter syntax is more convenient.
	The frame field selects the set of spatial variables with reference to which the integration is performed. For example, if the source expression is '1', using the reference frame makes the variable evaluate to the undeformed volume of the source domains, while choosing a moving frame gives you the deformed volume.

Examples

See Also

Make the average and maximum values of the solution available on the boundary of a circle.

```
fem.geom = circ2;
  fem.mesh = meshinit(fem);
  fem.shape = 2;
  fem.equ.c = 1; fem.equ.f = 1;
  fem.bnd.h = 1;
  fem.elem = {};
  clear el;
  el.elem = 'elcplscalar';
  el.g = {'1'};
  el.var = { 'area' 'mean' 'max' };
  el.global = { '1 '};
  el.maxvars = {'3'};
  clear src;
  src.ind = {{'1'}};
  src.expr = {{'1'}, {'u/area'}, {'u'}};
  src.iorders = {{'4'}, {'4'}, {'4'}};
  el.src{1} = {{},{},src};
  clear dst;
  dst.ind = {{'1','2','3','4'}};
  dst.usage = {{},{'1'},{'1'}};
  el.geomdim{1} = {{},dst,{}};
  fem.elem = [fem.elem {el}];
  fem.xmesh = meshextend(fem);
  fem.sol = femstatic(fem);
  postint(fem, 'u/pi')
  postint(fem, 'mean/(2*pi)', 'edim',1)
  postint(fem, 'max/(2*pi)', 'edim',1)
elepspec, elgpspec
```

Purpose	Define constraints compatible with first order vector elements.
Syntax	el.elem = 'elcurlconstr' el.g{ig} = geomnum el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.constr{eldomgrp}{ic} = vconstr
Description	The elcurlconstr element adds constraints on vector expressions, vconstr, which are given as cell arrays with one entry for each space dimension. The projection of the vector expression onto element edges is constrained to zero at element edge center points.
	For the syntax of the ind field, see elempty. The constr field has one entry for each domain group, each being a cell array of vector expressions to be constrained.
Cautionary	This element is currently tailored to fit the shcurl vector shape functions by constraining the actual degrees of freedom. Other uses may be possible, but performance will be unpredictable.
Examples	<pre>Add a PEC condition on boundaries 1 to 6 in a 3D electromagnetics model. el.elem = 'elcurlconstr'; el.g = {'1'}; gd.ind = {{'1', '2', '3', '4', '5', '6'}}; gd.constr = {{{'tEx', 'tEy', 'tEz'}}; el.geomdim{1} = {{},{},gd,{}; fem.elem = [fem.elem {el}]; When exporting the fem structure from a vector element electromagnetics model, similar elcurlconstr elements are added to the fem.elemph field.</pre>
See Also	elempty, elpconstr

Purpose	Define some basic functionality of the element syntax elements.
Syntax	el.elem = 'elempty' el.g{ig} = geomnum el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist
Description	The elempty element does not contribute anything directly to the problem description. It is described here because all element classes are derived from elempty and therefore share some basic syntax. This "element syntax" uses a very limited subset of the data structures available in COMSOL Script to describe the complete FEM problem. All models have to be converted into element syntax before solving, a conversion handled by the meshextend function. Additional elements can be added in the fem.elem field. Unless there are conflicts, the additional elements are added to the global problem description.
	The only building blocks allowed in the element syntax are strings, row cell arrays and structs. Note that pure numerical values are not allowed: both integers and decimal number have to be wrapped up as strings. Most elements accept empty cell arrays as placeholders to signify that the element does not wish to contribute anything for some variable or on some domain.
	At the top level, all elements are structs with at least the field elem containing the element class name as a string. Most elements define contributions that are in some way local to one or more geometries. Such elements have a g field, which contains a row cell array of geometry numbers (quoted as COMSOL Script strings). These geometries are in turn referred to internally in the element using the position in the g field as index, called ig.
	Most element classes specify their contributions per geometry, dimension and domain group. See further the section "Specifying a Model" in the <i>COMSOL Multiphysics Scripting Guide</i> for an explanation of the domain group concept. The geomdim field, where present, is a nested cell array where the outer level position corresponds to local geometry index, ig, and inner level position corresponds to space dimension plus one, called edim in the element context.
	The geomdim{ig}{edim} entries are used as structs with field names and syntax depending on the particular element class. There are, however, some common principles. Whenever there is an ind field present, it is a cell array where the position corresponds to element domain group number, called eldomgrp, each entry being a domain list. The domain lists are in turn cell arrays of quoted domain numbers.

	If the ind field is not present in an element structure for an element that accepts a domain grouping, it is defaulted as if all domains belong to group one. Other fields can usually be specified either per domain group or using one entry valid for all groups, whether explicitly specified or defaulted as one single group. Note that this behavior is not explicitly documented for each element type.
Cautionary	Adding element syntax contributions bypasses all high-level syntax checks, which can result in unintelligible error messages or even unexpected termination of a scripting session.
Examples	Create a simple model (Poisson's equation on unit circle) and extract the element syntax created by meshextend.
	<pre>clear fem; fem.geom = circ2; fem.mesh = meshinit(fem); fem.shape = 2; fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = 1; fem.xmesh = meshextend(fem); elstr = fem.xmesh.getElements; clear elem; for i=1:length(elstr) elem{i} = eval(elstr(i)); end</pre>
	<pre>elstr = fem.xmesh.getInitElements; clear eleminit; for i=1:length(elstr) eleminit{i} = eval(elstr(i)); end</pre>
	The element syntax can be studied, modified, and then fed back into the fem.elem
	and fem.eleminit fields. An additional call to meshextend with property
	'standard' set to 'off' updates the fem.xmesh field before solving.
	<pre>fem.elem = elem; fem.eleminit = eleminit; fem ymach = machartand(fem 'standard' 'sff');</pre>

```
fem.eleminit = eleminit;
fem.xmesh = meshextend(fem,'standard','off');
fem.sol = femstatic(fem);
postsurf(fem,'u');
```

An elempty element can be used to force a domain group split, which can be necessary in some cases where subdomain variables are accessed on boundaries. To split the boundary into two domain groups, add the following commands to the element syntax created above:

	<pre>clear el; el.elem = 'elempty'; gd.ind = {{'1', '2'}, {'3', '4'}}; el.geomdim{1} = {{}, {}, gd}; fem.elem = [fem.elem {el}];</pre>
See Also	elcconstr, elconst, elcplextr, elcplgenint, elcplproj, elcplscalar, elcurlconstr, elepspec, eleqc, eleqw, elgeom, elgpspec, elinline, elinterp, elinv, elirradiation, elmesh, elpconstr, elplastic, elpric, elshape, elshell_arg2, elvar

Purpose	Declare constraint point patterns.
Syntax	el.elem = 'elepspec' el.g{ig} = geomnum el.geom{ig}.ep = meshcases patterns
	<pre>meshcases.default = patterns meshcases.case{elmcase} = patterns meshcases.mind{elmcase} = caselist</pre>
	patterns{iptrn} = lagorder ptrnlist
	ptrnlist{2*itype-1} = bmtypename ptrnlist{2*itype} = lagorder ptrn
	<pre>ptrn{ipnt} = lcoords</pre>
Description	The elepspec element defines local evaluation point patterns typically used for pointwise constraints. In contrast to most elements, the elepspec does not have a geomdim field. The geom field has one entry per geometry listed in the g field and one subfield, ep, which lists a number of patterns that other elements refer to by position.
	If there are no alternate mesh cases specified, each entry in the ep field is either an integral number, which is interpreted as a Lagrange point order to be used on all basic mesh element types and dimensions, or a cell array of pairs of basic mesh element type and a Lagrange order or an explicit pattern. For a list of basic mesh element type names, see elshape. Explicit patterns are cell arrays of points in the local coordinate system on each element, each point being a cell array of the same dimension as the basic mesh element type.
	If there are multiple mesh cases present in the model, the ep field is a struct with fields default, case, and mind. The default field has the same syntax as described above for the ep field itself. Multiple alternate cases which need the same evaluation point patterns can be grouped together using the mind field. This field is a cell array containing groups of mesh case numbers, each group, caselist, given as a cell array. For each element mesh case group, elmcase, an alternate pattern specification is given in the case field.
Cautionary	Currently, there can only be one elepspec element for each geometry, which is generated by default when converting the standard syntax. Therefore, no additional elepspec can be added in the fem.elem field unless meshextend is called with property standard set to off.

Examples

Given a single-geometry fem structure with an xmesh field, extract elements and add an additional pattern that can be used by other elements and update the xmesh.

```
elstr = fem.xmesh.getElements;
  clear elem;
  for i=1:length(elstr)
    elem{i} = eval(elstr(i));
    if strcmp(elem{i}.elem,'elepspec')
      iepspec = i;
    end
  end
  elstr = fem.xmesh.getInitElements;
 clear eleminit;
  for i=1:length(elstr)
    eleminit{i} = eval(elstr(i));
  end
  newptrn = length(elem{iepspec}.geom{1}.ep)+1;
  elem{iepspec}.geom{1}.ep{newptrn} = {'s(1)' {{'0.5'}}};
  fem.elem = elem;
 fem.eleminit = eleminit;
Here, additional elements using the constraint pattern with index newptrn can be
added to the fem.elem field.
```

fem.xmesh = meshextend(fem,'standard','off');

See Also elempty, elgpspec, elshape, elpconstr, elcconstr

Purpose	Define coefficient form or general form equation contributions.
Syntax	<pre>el.elem = 'eleqc' el.g{ig} = geomnum el.form = 'coefficient' 'general' el.eqvars = 'on' 'off' el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.dim{idim} = dimvarname el.geomdim{ig}{edim}.ea{eldomgrp} = eacoeff el.geomdim{ig}{edim}.da{eldomgrp} = dacoeff el.geomdim{ig}{edim}.al{eldomgrp} = ccoeff el.geomdim{ig}{edim}.al{eldomgrp} = alcoeff el.geomdim{ig}{edim}.al{eldomgrp} = gacoeff el.geomdim{ig}{edim}.ga{eldomgrp} = becoeff el.geomdim{ig}{edim}.a{eldomgrp} = becoeff el.geomdim{ig}{edim}.a{eldomgrp} = acoeff el.geomdim{ig}{edim}.f{eldomgrp} = fcoeff el.geomdim{ig}{edim}.f{eldomgrp}{eldomgrp}{imm} = ipind</pre>
Description	The eleqc element adds equation contributions in coefficient form or general form as specified by the el.form field. It also defines variables that evaluate to various parts of the equations. These equation variables can be turned off using the eqvars field.
	For the syntax of the ind field, see elempty. The ea, da, c, al, ga, be, a, and f coefficients have the same syntax as the corresponding fem.equ fields. See further the section "Specifying a Model" in the <i>COMSOL Multiphysics Scripting Guide</i> . In contrast to the standard syntax, the eleqc coefficients have the same names on all dimensions. That is, the standard fem.bnd.g and fem.bnd.q fields correspond to a geomdim{ig}{sdim-1}.f and a geomdim{ig}{sdim-1}.a, respectively.
	The ipoints field differs from the gporder fields in the standard syntax (for example, fem.equ.gporder) in that it always contains pattern indices instead of orders (see elgpspec).
	The COMSOL Multiphysics user interface generates an eleqc element for geometries where the solution form is the coefficient or general form. When assembling using the weak solution form (the default), equations are converted to weak form and an eleqw element is generated instead.
Cautionary	Because of the naming convention for equation variables, at most one eleqc element per geometry can have equvars set to on. Because the default eleqc element has equation variables turned on, unless otherwise specified, it is proper procedure to turn them off for any additional eleqc elements added in the fem.elem field.

Make sure that the integration point pattern index you use really does exist and corresponds to a reasonable integration order. When adding an eleqc element to an existing model, it may be necessary to extract and modify also the default elgpspec element.

Examples

Because equation contributions are simply added, you can introduce additions to a single coefficient using the fem.elem field.

```
clear fem;
fem.geom = circ2;
fem.mesh = meshinit(fem);
fem.shape = 2;
fem.equ.c = 1; fem.equ.f = 0;
fem.bnd.h = 1;
fem.elem = {};
clear el;
el.elem = 'eleqc';
el.g = {'1'};
el.form = 'coefficient';
el.eqvars = 'off';
clear equ;
equ.dim = {'u'};
equ.ind = {{'1'}};
equ.f = {{'1'}};
equ.ipoints = {{'1'}};
el.geomdim{1} = {{},{},equ};
fem.elem = [fem.elem {el}];
fem.xmesh = meshextend(fem);
fem.sol = femstatic(fem);
postplot(fem, 'tridata', 'u', 'triz', 'u', 'refine',3);
```

See Also

elempty, eleqw, elgpspec

Purpose	Define weak form contributions.
Syntax	<pre>el.elem = 'eleqw' el.g{ig} = geomnum el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.coeff{eldomgrp}{iequ} = weak el.geomdim{ig}{edim}.tcoeff{eldomgrp}{iequ} = dweak el.geomdim{ig}{edim}.ipoints{eldomgrp}{iequ} = ipind el.geomdim{ig}{edim}.dvolname{eldomgrp}{iequ} = dvolname</pre>
Description	The eleqw element adds weak form contributions to the FEM problem. For the syntax of the ind field, see elempty. The weak and dweak coefficients have the same syntax as the corresponding fields in the standard fem struct syntax. See further the section "Specifying a Model" in the <i>COMSOL Multiphysics Scripting Guide</i> . The ipoints field differs from the gporder fields in the standard syntax (for example, fem.equ.gporder) in that it always contains pattern indices instead of orders (see elgpspec).
	The field dvolname specifies the name of the differential volume factor to be used in integrating the particular equation. If you are using multiple frames, this name effectively decides in which frame the equation is defined. There is normally a unique volume factor name tied to each frame, with dvol being the default for fixed meshes.
	The main difference between specifying equations using eleqw and using eleqc is that the former always gives a correct Jacobian if it is possible to automatically differentiate all functions called.
Cautionary	Make sure that the integration point pattern index you use really does exist and corresponds to a reasonable integration order. When adding an eleqw element to an existing model, it may be necessary to extract and modify also the default elgpspec element.
Examples	As equation contributions are simply added, it is easy to introduce additional weak form terms using the fem.elem field. clear fem;
	<pre>fem.geom = circ2; fem.mesh = meshinit(fem); fem.shape = 2; fem.equ.c = 1; fem.equ.f = 0; fem.bnd.h = 1; fem.elem = {};</pre>
	clear el;

```
el.elem = 'eleqw';
el.g = {'1'};
clear equ;
equ.ind = {{'1'}};
equ.coeff = {{'u_test'}};
equ.ipoints = {{'1'}};
el.geomdim{1} = {{},{},equ};
fem.elem = [fem.elem {el}];
fem.xmesh = meshextend(fem);
fem.sol = femstatic(fem);
postplot(fem,'tridata','u','triz','u','refine',3);
```

elempty, eleqc, elgpspec

Purpose	Elevate degrees of 2D geometry object Bézier curves.
Syntax	<pre>ge = elevate(g,en,d) [ge,t1] = elevate(g,en,d) ge = elevate(g,d1) [ge,t1] = elevate(g,d1)</pre>
Description	<pre>ge = elevate(g,en,d) elevates the degrees of edges en in the 2D geometry object g, using the degree steps d. en is a vector that specifies the edge numbers of the curves to be degree elevated, and d is the corresponding vector that specifies the degrees of elevation, so that curve number en(i) is elevated by d(i) degrees.</pre>
	<pre>ge = elevate(g,dl) degree elevates the Bézier curve defined by g.rb{i}(:,k) and g.wt{i}(:,k), by the number of degrees specified in dl{i}(:,k). dl is a cell array of the same size as rb and wt. See geom2 for details on these properties. The first and last entries in dl must be empty, since there are no curves of degree 0, and curves of maximum degree cannot be degree elevated.</pre>
	<pre>[ge,t1] = elevate(g,) additionally returns the cell array t1, of length 3, containing permutation vectors for vertices, edges and subdomains, respectively. Entry i of such a vector contains the entity number j of the geometry object g from which the entity i in ge originates.</pre>
Examples	<pre>Elevate the degree of edge 1 and 3 in a circle, by one degree. c1 = circ2; figure, geomplot(c1,'edgelabels','on','ctrlmode','on'); axis equal [c2,t1] = elevate(c1,[1 3],[1 1]); figure, geomplot(c2,'edgelabels','on','ctrlmode','on'); axis equal</pre>
	An alternative way of obtaining the same degree elevated circle, is to use the input argument dl, as is done below.
	c3 = elevate(c1,{[] [] [1 0 1 0] []}); figure, geomplot(c3,'edgelabels','on','ctrlmode','on'); axis equal
See Also	geom0, geom1, geom2, geom3

Purpose	Define geometrical variables.			
Syntax	<pre>el.elem = 'elgeom' el.g{ig} = geomnum el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.frame = frame el.sorder = sorder el.method = {'Lenoir'} 'f133'</pre>			
Description	The elgeom element evaluates geometrical variables for the (undeformed) geometry, notably coordinates, curve and face parameters, and tangential vectors. These variables are defined on the domains in domainlist. The variable names are derived from the space coordinate names of the frame frame. See further the section "Specifying a Model" in the COMSOL Multiphysics Scripting Guide.			
	The Lenoir method (the default method) provides a continuous piecewise polynomial interpolation of order sorder. The fl33 method, which was the default method until version 3.3a, provides a discontinuous nonpolynomial-based interpolation. The sorder field is only applicable when using the Lenoir method and has a default value of 1.			
Examples	<pre>The default generated elgeom element for a 2D model is: el.elem = 'elgeom'; el.g = {'1'}; el.frame = 'xy'; el.sorder = '1'; el.method = 'Lenoir';</pre>			
See Also	elempty, elmesh			

Purpose	Declare integration point patterns.		
Syntax	el.elem = 'elgpspec' el.g{ig} = geomnum el.geom{ig}.ep{iptrn} = order ptrnlist		
	<pre>meshcases.default = patterns meshcases.case{elmcase} = patterns meshcases.mind{elmcase} = caselist</pre>		
	patterns{iptrn} = lagorder ptrnlist		
	ptrnlist{2*itype-1} = bmtypename ptrnlist{2*itype} = order ptrn		
	<pre>ptrn{ipnt} = {lcoords,weight}</pre>		
Description	The elgpspec element defines local integration point patterns and weights for the numerical quadrature needed when assembling equations on each mesh element. In contrast to most elements, the elgpspec does not have a geomdim field. The geom field has one entry per geometry listed in the g field and one subfield, ep, which lists a number of patterns which other elements refer to by position.		
	If there are no alternate mesh cases specified, each entry in the ep field is either an integral number, which is interpreted as the order of polynomials that should be integrated exactly on all basic mesh element types and dimensions, or a cell array of pairs of basic mesh element type and a polynomial order or an explicit pattern. For a list of basic mesh element type names, see elshape. Explicit patterns are cell arrays of points and weights in the local coordinate system on each element, each point-weight pair being represented as a single cell array where the weight follows directly after the coordinates.		
	If there are multiple mesh cases present in the model, the ep field is a struct with fields default, case, and mind. The default field has the same syntax as described above for the ep field itself. Multiple alternate cases which need the same integration point patterns can be grouped together using the mind field. This field is a cell array containing groups of mesh case numbers, each group, caselist, given as a cell array. For each element mesh case group, elmcase, an alternate pattern specification is given in the case field.		
Cautionary	Currently, there can only be one elgpspec element for each geometry, which is generated by default when converting the standard syntax. Therefore, no additional		

elgpspec can be added in the fem.elem field unless meshextend is called with property 'standard' set to 'off'.

Note that the sum of weights in an explicit pattern specification are supposed to be equal to the element's volume in the element local coordinate system, that is, to 1/2 for triangles and 1/6 for tetrahedra.

Examples

Given an fem structure with an xmesh field, extract elements and add an additional pattern which uses fourth order integration on curved simplices and explicit zeroth order integration on other simplices. Finally, the xmesh is updated.

```
elstr = fem.xmesh.getElements;
clear elem;
for i=1:length(elstr)
 elem{i} = eval(elstr(i));
  if strcmp(elem{i}.elem, 'elgpspec')
    igpspec = i;
  end
end
elstr = fem.xmesh.getInitElements;
clear eleminit;
for i=1:length(elstr)
  eleminit{i} = eval(elstr(i));
end
newptrn = length(elem{igpspec}.geom{1}.ep)+1;
elem{igpspec}.geom{1}.ep{newptrn} = {'s(2)', '4', ...}
  'ls(2)', {{'0.333333333', '0.333333333', '0.5'}};
fem.elem = elem;
fem.eleminit = eleminit;
```

Here, additional elements using the integration point pattern with index newptrn can be added to the fem.elem field.

fem.xmesh = meshextend(fem, 'standard', 'off');

See Also

elempty, elepspec, elshape, eleqc, eleqw

Purpose	Declare functions and corresponding symbolic derivatives.			
Syntax	el.elem = 'elinline' el.name = fname el.args{iarg} = argname el.expr = evalexpr el.dexpr{iarg} = devalexpr el.complex = 'true' 'false' el.linear = 'true' 'false'			
Description	The elinline element declares a new function with differentiation rules in terms of other built-in or COMSOL Script or MATLAB functions. COMSOL Multiphysics calls the function using the MATLAB interpreter if you run COMSOL Multiphysics with MATLAB and it calls the COMSOL Script interpreter otherwise.			
	Declared inline functions can be used with the syntax fname(arg1,arg2,) anywhere except for in other inline function definitions. The args field contains a list of formal parameter names which can be used in the expr field defining the function itself and the dexpr field defining derivative expressions with respect to each of the formal arguments.			
	The evalexpr and devalexpr expressions can contain any valid expression in the formal arguments, including COMSOL Script function calls. Global constants and variables are not available with pi being the only noticeable exception. Note that differentiation with respect to some formal argument can be disabled by just specifying the corresponding devalexpr as '0'.			
	Functions which can generate complex values from real data must have the complex field set to 'true'. The linear property decides if the function is treated as linear when deciding whether to reassemble the Jacobian at each time step/iteration or not.			
Cautionary	Note that inline functions cannot depend on other inline functions, only on built-ins and functions defined on your COMSOL Script or MATLAB path. Inline functions can be used to override built-ins but can never override another inline function.			
Examples	<pre>Use an inline function to redefine the derivative of the sqrt function in such a way that the Jacobian of sqrt(u^2+v^2) will exist for u=v=0. el.elem = 'elinline'; el.name = 'sqrt'; el.args = {'a'}; el.expr = 'sqrt(a)';</pre>			

```
el.dexpr = {'1/(2*sqrt(a)+eps)'};
el.complex = 'true';
el.linear = 'false';
fem.elem = [fem.elem {el}];
```

elempty

Purpose	Declare interpolation functions.			
Syntax	el.elem = 'elinterp' el.name = fname el.x = xgrid el.y = ygrid el.z = zgrid el.data = fdata el.method = 'nearest' 'linear' 'cubic' 'spline'			
Description	The elinterp element declares an interpolation function based on a 1D, 2D or 3D data set provided by the user. Interpolation functions take one, two or three arguments, depending on the dimension of the data set.			
	The xgrid, ygrid, and zgrid parameters are cell arrays of points, sorted in increasing order, where the data is given. Depending on the dimensions, not all fields are used. For 1D interpolation, the fdata parameter is a cell array of values corresponding to the points in el.x. In the 2D case, the size of fdata is length(xgrid)*length(ygrid), with x increasing fastest, and similarly in 3D.			
	There are four interpolation methods to choose from. Nearest neighbor and linear interpolation are available in all dimensions, while 'cubic' and 'spline' can only be used for interpolation in 1D data sets. The difference between the latter two is, generally speaking, that 'cubic' preserves monotonicity and does not overshoot at the cost of discontinuous second derivatives at the tying points.			
Cautionary	Interpolation is provided only for real numbers. To interpolate complex numbers, real and imaginary parts have to be treated separately.			
Examples	Given the matrices x (1-by-m), y (1-by-n) and F (m-by-n), create a corresponding interpolation element declaring a function $f(x,y)$.			
	<pre>cellX = cell(1,m); for i=1:m cellX{i} = num2str(x(i)); end</pre>			
	<pre>cellY = cell(1,n); for i=1:n cellY{i} = num2str(y(i)); end</pre>			
	<pre>cellF = cell(1,m*n); for i=1:m*n cellF{i} = num2str(F(i)); end</pre>			

```
el.elem = 'elinterp';
el.name = 'f';
  el.x = cellX;
  el.y = cellY;
  el.data = cellF;
el.method = 'linear';
  fem.elem = [fem.elem {el}];
elempty
```



Purpose	Define matrix inverse component variables.			
Syntax	<pre>el.elem = 'elinv' el.g{ig} = geomnum el.matrixdim = mdim el.format = 'symmetric' 'hermitian' 'unsymmetric' el.basename = bname el.postname = pname el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.matrix{eldomgrp} = matexpr</pre>			
Description	The elinv element defines components of the inverse of an mdim-by-mdim matrix field. The matexpr is a cell array of expressions which specify the source matrix in column order. If the format is 'symmetric' or 'hermitian', only the upper triangle has to be given.			
	Matrix inverse component names, that is, the variable names defined by the element, are created by appending row and column indices to the bname parameter and, if the postname field is present, append '_pname'. If the format is specified as 'symmetric', variables are generated only for the upper triangular part of the inverse, otherwise for all components.			
Examples	<pre>Define variables to evaluate the Jacobian inverse components for an explicit variable transformation [X(x,y,z), Y(x,y,z), Z(x,y,z)]. el.elem = 'elconst'; el.var = {'X','2*x','Y','y+z','Z','z'}; fem.elem = [fem.elem {el}];</pre>			
	<pre>clear el; el.elem = 'elinv'; el.g = {'1'}; el.matrixdim = '3'; el.format = 'unsymmetric'; el.basename = 'd'; gd.ind = {{'1'}}; gd.matrix = {{'diff(X,x)', 'diff(X,y)', 'diff(X,z)', 'diff(Y,x)', 'diff(Y,y)', 'diff(Y,z)', 'diff(Z,x)', 'diff(Z,y)', 'diff(Z,z)'}; el.geomdim{1} = {{},{},gd}; fem.elem = [fem.elem {el}];</pre>			
See Also	elempty, elpric			

Purpose Define irradiation variables for radiative heat transfer. el.elem = 'elirradiation' Syntax el.g{ig} = geomnum el.method = 'area' | 'hemicube' el.iorder = order el.resolution = res el.sectors = nsectors el.cache = 'on' | 'off' el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.name{eldomgrp} = Gname el.geomdim{ig}{edim}.ambname{eldomgrp} = Fname el.geomdim{ig}{edim}.expr{eldomgrp} = Jexpr el.geomdim{ig}{edim}.opexpr{eldomgrp} = opacity el.geomdim{ig}{edim}.cavity{eldomgrp} = cavitylist Description The elirradiation element defines one variable Gname representing the local irradiation from other surfaces and one Fname, known as *ambient view factor*, representing the fraction of the field of view not covered by other surfaces. The variable names can differ between domain groups. The irradiation at each point depends on the radiosity at all other visible surface points. An expression for the radiosity is provided in the expr field. For interior boundaries, it has to be made clear in which direction the radiation goes. The opacity expression is evaluated on adjacent subdomains, and is expected to be nonzero on exactly one. To avoid unnecessary visibility checks, the surfaces can be manually assigned to one or more cavities, each of which exchanges radiation only with other surfaces in the same cavity. For each domain group, specify to which cavities the group belongs. There are currently two view factor evaluation methods which can be selected using the method field. The 'area' method implies direct area integration using a simple quadrature rule of order order and no visibility checks. Different convex cavities can be held apart using the cavity field. The 'hemicube' method, and its generalizations to lower dimensions, uses techniques borrowed from computer graphics to handle surfaces obstructing each other. Essentially, images of resolution res-by-res are generated from each evaluation point in 3D. Use the sectors field to specify that a 2D geometry shall be interpreted as axially symmetric and to set the azimuthal resolution when evaluating view factors. The value nsectors is the number of sectors to a full revolution in a virtual 3D

geometry created by revolving the 2D mesh about the axis.

	Due to the rather complex evaluation, it is usually beneficial to store the view factors between calls. This can, however, generate a lot of data, which can potentially be a limiting factor preventing a solution on a given system. Therefore, the cache field can be set to 'off', but this increases run times considerably.			
Cautionary	Radiation is currently only possible between boundaries, that is, between entities of dimension one lower than the space dimension. Also, radiation only works within one geometry.			
	The elirradiation element is available only if your license includes the Heat Transfer Module.			
Examples	Compare the irradiation calculated by the hemicube algorithm with an analytical solution in a known case.			
	<pre>clear fem; fem.geom = geomcsg({rect2(1,1,'pos',[0 -1]),</pre>			
	<pre>fem.xmesh = meshextend(fem); postint(fem,'abs(G-Gref)','edim',1,'dl',7)/ postint(fem,'abs(Gref)','edim',1,'dl',7)</pre>			
See Also	elempty			

Purpose	Create ellipse geometry object.		
Syntax	<pre>obj = ellip2 obj = ellip1 obj = ellip2(a,b,) obj = ellip1(a,b,)</pre>		
Description	 obj = ellip2 creates a solid ellipse geometry object with center at the origin and semi-axes equal to 1. ellip2 is a subclass of solid2. obj = ellip2(a,b,) creates an ellipse object with semi-axes equal to a and b, 		

obj = e111p2(a, b, ...) creates an ellipse object with semi-axes equal to a and b, respectively, centered at the origin. a and b are positive real scalars, or strings that evaluate to positive real scalars, given the evaluation context provided by the property const.

The functions ellip2/ellip1 accept the following property/values:

TABLE I-I6: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
base	corner center	center	Positions the object either centered about pos or with the lower left corner of surrounding box in pos
const	Cell array of strings	{}	Evaluation context for string inputs
pos	Vector of reals or cell array of strings	[0 0]	Position of the object
rot	Real or string	0	Rotational angle about pos (radians)

obj = ellip1(...) creates a curve circle geometry object with properties as given for the ellip2 function. ellip1 is a subclass of curve2.

Ellipse objects have the following properties:

TABLE I-17: ELLIPSE OBJECT PROPERTIES

PROPERTY	DESCRIPTION
a, b	Semi-axes
х, у	Position of the object
rot	Rotational angle

In addition, all 2D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom2 for details.

Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported.		
Examples	The commands below create an ellipse object and plot it.		
	e1 = ellip2(1,0.3,'base','center','pos',[0,0],'rot',pi/4 get(e1,'rot') geomplot(e1)		
See Also	circ1, circ2,curve2, curve3,geomcsg		

Purpose	Create an ellipsoid geometry object.
Syntax	<pre>obj = ellipsoid3 obj = ellipsoid2 obj = ellipsoid3(a,b,c) obj = ellipsoid2(a,b,c) obj = ellipsoid3(a,b,c,)</pre>
	obj = ellipsoid2(a,b,c,)
Description	obj = ellipsoid3 creates a solid ellipsoid geometry object v

obj = **ellipsoid3** creates a solid ellipsoid geometry object with center at the origin and semi-axes equal to 1. **ellipsoid3** is a subclass of **solid3**.

obj = ellipsoid3(a,b,c,...) creates a solid ellipsoid object with semi-axes a, b, and c. a, b, and c are positive real scalars, or strings that evaluate to positive real scalars, given the evaluation context provided by the property Const.

The functions ellipsoid3/ellipsoid2 accept the following property/values:

TABLE I-18: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object
Const	Cell array of strings	{}	Evaluation context for string inputs
Pos	Vector of reals or cell array of strings	[0 0]	Position of the object
Rot	real or string	0	Rotational angle about Axis (radians)

Axis sets the local z-axis, stated either as a directional vector of length 3, or as a 1-by-2 vector of spherical coordinates. Axis is a vector of real scalars, or a cell array of strings that evaluate to real scalars, given the evaluation context provided by the property Const. See gency13 for more information on Axis.

Pos sets the center of the object. Pos is a vector of real scalars, or a cell array of strings that evaluate to real scalars, given the evaluation context provided by the property Const.

Rot is an intrinsic rotational angle for the object about its local z-axis provided by the property Axis. Rot is a real scalar, or a string that evaluates to a real scalar given

the evaluation context provided by the property Const. The angle is assumed to be in radians if it is numeric, and in degrees if it is a string.

obj = ellipsoid2(...) creates a surface ellipsoid object with the properties as given for the ellipsoid3 function. ellipsoid2 is a subclass of face3.

Ellipsoid objects have the following properties:

TABLE I-19: ELLIPSOID OBJECT PROPERTIES

PROPERTY	DESCRIPTION
a, b, c	Semi-axes
x, y, z, xyz	Position of the object. Components and vector forms
ax2	Rotational angle of symmetry axis
ax3	Axis of symmetry
rot	Rotational angle

In addition, all 3D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom3 for details.

Examples	The following commands create a surface and solid ellipsoid object, where the position and semi-axis are defined in the two alternative ways.
	e2 = ellipsoid2(1,1,1,'pos',[0 1 0],'axis',[0 0 1],'rot',0) e3 = ellipsoid3(12,10,8)
Compatibility	The representation of the ellipsoid objects has been changed. The FEMLAB 2.3 syntax is obsolete but still supported. If you use the old syntax or open 2.3 models containing ellipsoids they are converted to general face or solid objects.
See Also	face3,geom0, geom1, geom2, geom3,sphere3, sphere2

Purpose	Fast evaluation of predefined convolution integrals
Syntax	<pre>el.elem = 'elkernel' el.g{ig} = geomnum el.name = opname opnamelist el.kernel = 'unit' 'helmholtz2D' 'helmholtz2Dinf' 'helmholtz3D' 'helmholtz3Dinf' 'helmholtz2Daxi' 'helmholtz2Daxiinf' 'maxwell3Dinf' 'maxwellTEinf' 'maxwellTMinf' 'maxwellTEaxiinf' 'maxwellTMaxiinf'</pre>
	el.frame = srcframe el.iorder = gporder el.k = wavenumber el.symflags{idim} = '-1' '0' '1'
	<pre>el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.srcn{idim}{eldomgrp} = nexpr el.geomdim{ig}{edim}.srcu{eldomgrp} = uexpr el.geomdim{ig}{edim}.srcnux{eldomgrp} = nuxexpr el.geomdim{ig}{edim}.srcnxe{eldomgrp} = nxeexpr el.geomdim{ig}{edim}.srcnxcurle{eldomgrp} = nxculreexpr</pre>
Description	The elkernel element is a wrapper which defines an operator that represents a predefined convolution integral. You can choose between plain integration of an arbitrary expression, Helmholtz-Kirchhoff integral solutions to Helmholtz' equation in various dimensions and a number of instances the Stratton-Chu formula for far-field evaluation of electromagnetic fields under various conditions. For Helmholtz equation, both the complete integral solution and the far-field limiting case are provided.
	The type of integral is specified in the kernel field where 'unit' gives you an operator which takes an expression as only argument, while the rest define operators which are functions of the evaluation point coordinates. The 3D Stratton-Chu formula requires a list of three operator names, one for each field component, while

The integrals are evaluated on the mesh with the integration order specified in iorder and using the source coordinate names corresponding to the given frame. Both Helmholtz-Kirchhoff and Stratton-Chu integral evaluation require a free-space wave number, which is expected to be a global constant. The former, in addition, needs expressions for the normal vector, solution value and its normal derivative on the source domains, specified in srcn, srcu, and srcnux, respectively. The Stratton-Chu formula requires the cross products of source normal with electric

TM waves require two operator names and the remaining only one.

	field and source normal with curl of electric field, specified in srcnxe and srcnxcurle, respectively.
	The source domains are expected to form a closed surface containing all sources and inhomogenities, and the specified normal vector must be facing into the domain enclosed by this surface. When exploiting symmetry to model only 1/2, 1/4, or 1/8 of the actual geometry, a closed surface can be recovered using the symmetry flags. In the symflags field, -1 in position idim is interpreted as antisymmetry with respect to the coordinate plane normal to the idim-axis, while +1 means that the plane is a symmetry plane. If the symflags field is not given, all entries are considered to be 0, which signifies that the model is neither symmetric nor antisymmetric.
Cautionary	The operators defined by elkernel are primarily intended for postprocessing and therefore do not define any Jacobian contributions.
Examples	The acoustic field from a baffled piston oscillating with specified velocity normally to an infinite rigid plane can be evaluated explicitly using the helmholtz2Daxi kernel with srcu set to zero since the terms proportional to u cancel out anyway.
	fem.geom = circ2*rect2; fem.mesh = meshinit(fem);
	fem.expr = {'SPL' '10*log10(0.5*abs(p(x,y))^2/2e-5^2)'};
	<pre>clear el; el.elem = 'elkernel'; el.g = {'1'}; el.kernel = 'helmholtz2Daxiinf'; el.name = 'p'; el.iorder = '20'; el.k = '100'; clear src11 src11.ind = {{'2'}}; src11.srcn = {'0', '-1'}; src11.srcn = {'0', '-1'}; src11.srcnu = '0'; src11.srcnux = {'1'}; el.geomdim{1} = {{},src11}; fem.elem = {el};</pre>
	<pre>fem.xmesh = meshextend(fem); fem.sol = asseminit(fem);</pre>
	postcrossplot(fem,1,3,'lindata','SPL','linxdata', '180/pi*atan2(y,x)','refine',10);

Purpose	Define extrusion map operators.
Syntax	<pre>el.elem = 'elmapextr' el.g{ig} = geomnum el.opname{iop} = opname el.flagname{iop} = flagname el.extttol = tol el.usenan = 'true' 'false' el.map{imap} = linmap genmap unitmap el.srcmap{iop} = imap el.dstmap{iop} = imap el.dstmap{iop} = imap el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.src{iop} = eldomgrplist</pre>
	<pre>linmap.type = 'linear' linmap.sg = srcig linmap.sv{ivtx} = srcvtx linmap.sframe = srcframe linmap.dg = dstig linmap.dv{ivtx} = dstvtx linmap.dframe = dstframe genmap.type = 'local' genmap.expr{idim} = transexpr genmap.frame = frame unitmap.type = 'unit' unitmap.frame = frame</pre>
Description	The elmapextr element defines extrusion map operators which can be used at any location where the source and destination transformations make sense. Each map operator takes its values from the source domain groups listed in the corresponding el.geomdim{ig}{edim}.src{iop} field. For each operator, transformations are specified in the srcmap and dstmap fields in the form of indices into the map field that consists of a list of transformation specifications. The available transformation types are 'linear', 'local', and 'unit'. They are described in detail under elcplextr on page 58. For each operator name a global variable flagname{iop} is also defined, which evaluates to 1 for all destination points where the map operation can find a corresponding source a point, otherwise it evaluates to 0. If the flagname field is not
	corresponding source point, otherwise it evaluates to 0. If the flagname field is not given, the flag variables will be given the same name as the corresponding operator. Therefore, statements like if (my_map,my_map(u),0) make perfect sense.

Cautionary	Parameter or time dependency in the source transformation is not properly detected by the solvers, which means that the source transformation will not be updated between parameter or time steps in that case. Solution dependencies in the transformation are properly detected but do not give any Jacobian contributions from the transformation.
Examples	Calculate the first ten eigenvalues of a 3-by-2 rectangle with periodic boundary conditions both left-right and top-bottom. Different map types are used. Note that this is the same example as used under elcplextr.
	<pre>fem.geom = rect2(3,2); fem.mesh = meshinit(fem, 'hmax',0.05); fem.equ.c = 1; fem.equ.da = 1; fem.bnd.ind = [0 1 2 0]; fem.bnd.constr = {'left2right(u)-u','lower2upper(u)-u'}; fem.elem = {};</pre>
	el.elem = 'elmapextr'; el.g = {'1'}; el.opname = {'left2right','lower2upper'};
	clear map1; map1.type = 'unit';
	<pre>clear map2; map2.type = 'linear'; map2.sg = '1'; map2.sv = {'2','3'}; map2.dg = '1'; map2.dv = {'1','4'};</pre>
	clear map3; map3.type = 'local'; map3.expr = {'x'};
	el.map = {map1 map2 map3};
	el.srcmap = {'1','3'}; el.dstmap = {'2','3'};
	<pre>clear src; src.ind = {{'1'},{'4'}}; src.src = {{'2'},{'1'}}; el.geomdim{1} = {{},src,{}};</pre>
	<pre>fem.elem = [fem.elem {el}]; fem.xmesh = meshextend(fem);</pre>

```
fem.sol = femeig(fem,'neigs',10,'shift',1);
postplot(fem,'tridata','u','triz','u','refine',3,'solnum',8);
```

elcplextr

Purpose	Define mesh variables and a frame.
Syntax	<pre>el.elem = 'elmesh' el.g{ig} = geomnum el.frame = frame el.xvars = 'on' 'off' el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.sizename{eldomgrp} = sname el.geomdim{ig}{edim}.qualname{eldomgrp} = qname el.geomdim{ig}{edim}.dvolname{eldomgrp} = dvolname el.geomdim{ig}{edim}.meshtypename{eldomgrp} = meshtypename el.geomdim{ig}{edim}.meshtypename{eldomgrp} = meshtypename el.geomdim{ig}{edim}.sshape{eldomgrp} = sshape sshape{2*i-1} = bmtypename sshape{2*i = params params.type = 'fixed' 'moving_abs' 'moving_rel' 'moving_expr' params.sdimdofs{idim} = dofname params.sdimexprs{idim} = expr params.refframe = refframename</pre>
Description	Concerning the syntax of the ind field, see elempty. For each domain group, the elmesh element defines the variable names sname, qname, dvolname, meshtypename, and meshelemname, which evaluate to local mesh element size, element quality, element volume, mesh type index, and mesh element number, respectively. Also, if xvars='on' the space coordinates, the space coordinate's reference time derivative, and the normal vector are defined. These variable names are derived from the space coordinates of the frame frame. The type of frame is determined by sshape.
Examples	<pre>By default, the mesh size variable h is available only on the top dimension. If evaluated on a boundary, h returns the size of the adjacent subdomain element. An additional elmesh element can be used to define h on the boundary to represent the size of the boundary element. el.elem = 'elmesh'; el.g = {'1'}; el.frame = 'xy' el.xvars = 'off' gd.sizename = 'h'; el.geomdim{1} = {{},gd,{}; fem.elem = [fem.elem {el}];</pre>
See Also	elempty

Define global scalar dependent variables and equations.
el.elem = 'elode' el.dim{idim} = depvarname el.f{idim} = rexpr el.weak{iweak} = wexpr
The elode element adds globally available scalar dependent variables (named degrees of freedom) and corresponding equations. The dim field lists unique variable names which are allocated on a fictitious 0D geometry and made available throughout the model. The optional f field has the same number of entries as the dim field, while the weak field, if present, can have any number of entries. These fields define scalar equations on the form rexpr=0 and wexpr=0, respectively. The f field requires the presence of a dim field. See further "fem.ode—Global Variables and equations" on page 50 in the <i>COMSOL Multiphysics Programming Guide</i> .
The fictitious geometry mentioned above is for most purposes equivalent to a real 0D geometry with one domain, a point. This geometry can be explicitly referenced using geometry index 0. Therefore, the expressions rexpr and wexpr can contain variables which are globally available or explicitly available on domain 1 of geometry 0.
For further examples of use of scalar dependent variables and equations, see the COMSOL Multiphysics User's Guide.
Though the elode element applies to the ever-present fictitious geometry 0, a real geometry also has to be defined for the solvers to work. Note also that elode can be used to define global weak contributions to existing equations. That is, the weak field may be used without the presence of a dim field.
<pre>Solve a simple scalar wave equation: clear fem fem.geom = geom0(zeros(0,1)); fem.mesh = meshinit(fem); clear el el.elem = 'elode'; el.dim = {'u'}; el.f = {'utt+u'}; fem.elem = {el}; clear elinit; elinit.elem = 'elconst'; elinit.var = {'u', '0', 'ut', '1'};</pre>

```
fem.eleminit = {elinit};
fem.xmesh = meshextend(fem);
fem.sol = femtime(fem,'tlist',linspace(0,4*pi,100),...
'maxorder',2,'rtol',1e-8,'atol',1e-8);
postcrossplot(fem,0,1,'pointdata','u')
See Also elempty, eleqc, eleqw, elshape
```

Purpose	Define general pointwise constraints.
Syntax	el.elem = 'elpconstr' el.g{ig} = geomnum el.nname = Nname el.nfname = NFname el.mname = Mname
	<pre>el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.constr{eldomgrp}{ic} = constrexpr el.geomdim{ig}{edim}.constrf{eldomgrp}{ic} = constrfexpr el.geomdim{ig}{edim}.cpoints{eldomgrp}{ic} = cpind</pre>
Description	The elpconstr element adds a set of pointwise constraints of type constrexpr=0. The constr field has the same syntax as the fem.bnd.constr field. See further the section "Specifying a Model" in the COMSOL Multiphysics Scripting Guide. The optional constrf field controls the way constraint forces are applied. You enter an expression such that its Jacobian with respect to the test functions decides on which degrees of freedom the reaction force is applied for each constraint. If constrf is omitted, constraints are <i>ideal</i> , which corresponds to setting constrf to test(constrexpr).
	The constraint and constraint force Jacobians are by default assembled to matrices called NT and NF, and the constraint residual is called M. This can be changed by assigning different names to the optional nname, nfname and mname fields. Only certain names are recognized by functions like assemble, though, see page 29.
	The constraints are enforced at the same local coordinates in all elements in one domain group. The constraint point pattern is specified as a pattern index in the cpoints field. Indices refer to patterns defined by an elepspec element.
	Compared to the elcconstr element, the elpconstr can implement a wider range of constraints, as a correct constraint Jacobian is always calculated on the fly. This is in contrast to the user-specified Jacobian matrix h, used in fem.bnd.h and the elcconstr element.
Examples	<pre>Solve a 1D biharmonic equation (related to Euler beams) with constraints on both value and normal derivative at the endpoints. clear fem; fem.geom = solid1([0,1]); fem.mesh = meshinit(fem); fem.shape = 'shherm(1,3,''u'')'; fem.form = 'weak'; fem.equ.weak = 'uxx_test*uxx';</pre>

```
fem.bnd.dim = {'u'};
  fem.bnd.cporder = 1;
  fem.elem = {};
  clear el;
  el.elem = 'elpconstr';
  el.g = {'1'};
  clear gd;
  gd.ind = {{'1'},{'2'}};
  gd.constr = {{'-u', '1-ux'},{'-u', '1+ux'};
  gd.cpoints = {{'1'},{'1'}};
  el.geomdim{1} = {gd,{}};
  fem.elem = [fem.elem {el}];
  fem.xmesh = meshextend(fem);
  fem.sol = femstatic(fem);
  postplot(fem,'liny','u');
elempty, elcconstr, elcurlconstr, elepspec
```

Purpose	Declare piecewise functions.
Syntax	<pre>el.elem = 'elpiecewise' el.subtype = 'poly' 'exppoly' 'general' el.name = fname el.args = argname el.intervals{ibnd} = interval_bound el.expr{iexpr} = poly_spec expr el.extmethod = 'extrap' 'const' double_value 'none' el.smoothzone = double_value el.smoothorder = '0' '1' '2' el.complex = 'true' 'false' el.linear = 'true' 'false' poly_spec{2*ipow} = exponent poly_spec{2*ipow+1} = coefficent</pre>
Description	The elpiecewise element declares a function fname which is of type subtype in each interval, the boundaries of which are given in the intervals field. The polynomials or general expressions are given for each subinterval in the expr field, which contains one pair of polynomial exponent and coefficient or one expression for each interval. Derivatives are calculated by automatic symbolic differentiation. Outside the intervals, the value of the function is either extrapolated, taken from the nearest interval boundary or given a fixed number, according to the extmethod field. 'none' indicates that extrapolation is deactivated, and will result in errors or NaN values for out-of-range values, depending on how and where the element is used.
	Since the given expressions may be discontinuous at the interval boundaries, elpiecewise includes an optional smoothing option. If given, the smoothzone field specifies a relative size of the smoothing zone, interpreted as the fraction of each interval length which should be smoothed at the intersections between intervals. The smoothorder field gives the number of continuos derivatives that must exist at the boundary between smoothing zone and interval.
	Functions which can generate complex values from real data must have the complex field set to 'true'. The linear property decides if the function is treated as linear when deciding whether to reassemble the Jacobian at each time step/iteration or not.
Examples	Setup a piecewise function element of the polynomials $0.2x^{-6} + 5.1x + 0.05x^{6}$ and 60x, defined from 1.7 to 4 and 4 to 5.2, respectively, with continuos first derivatives at the intersection:

```
el.elem = 'elpiecewise';
el.name = 'myfun';
el.subtype = 'poly';
el.expr = {{'-6' '0.2' '1' '5.1' '6' '0.05'} {'1' '60'}}
el.intervals = {'1.7' '4' '5.2'}
el.smoothzone = '0.1';
el.smoothorder = '1';
fem.elem = [fem.elem {el}];
```

elempty, elinterp, elinline

Purpose	Define plastic strain variables.
Syntax	<pre>el.elem = 'elplastic' el.g{ig} = geomnum el.vars{ivar} = varname el.varsToCache = cachevarlist el.varPairsToGpProcess{2*igpvar-1} = gpvarname el.varPairsToGpProcess{2*igpvar} = gpvarexpr el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.Yield{eldomgrp} = yieldexpr el.geomdim{ig}{edim}.EffStress{eldomgrp} = effstressexpr el.geomdim{ig}{edim}.G{eldomgrp}{ivar} = gexpr el.geomdim{ig}{edim}.gporder{eldomgrp} = iorder</pre>
Description	The elplastic element defines the plastic strain variable names specified in the vars field. For the syntax of the ind field, see elempty. The yield function is effstressexpr-yieldexpr. For each strain variable in vars, the right-hand side of the corresponding rate equation is gexpr times the plastic multiplier, lambda. See "Continuum Application Modes", section "Theory Background", in the <i>Structural Mechanics Module User's Guide</i> . The gporder field specifies a quadrature rule order, which should preferably be the same as the order used in the assembly of the main equation.
	The varsToCache field contains a list of variables that can be assumed not to depend explicitly on the plastic strains. By specifying variable names representing complicated material property expressions or interpolated data independent of the plastic strains, it is possible to avoid repeated evaluation in the inner, plastic, loop.
	In addition to the plastic strain variables, the varPairsToGpProcess field defines a number of postprocessing variable-expression pairs, which, when evaluated, are linearly extrapolated from the integration points. Use this feature, for example, to avoid problems with nonconvergent plastic strains at sharp geometry corners.
	For each variable and integration point, the elplastic element declares an additional degree of freedom, which appears in the solution vector. However, consider these degrees of freedom to be internal data of the elplastic element. They are updated only by a special procedure in the nonlinear solver.
Cautionary	Note that some of the field names are mixed case, and case matters. Also, the domain-dependent fields do not accept empty entries for any domain group.
	The elplastic element is available only if your license includes the Structural Mechanics Module or the MEMS Module.

Examples By faking a single plastic strain variable, the elplastic element can be used also as a pure postprocessing element to define variables extrapolated from the integration points. The following example works for a 2D plane strain model. el.elem = 'elplastic'; el.g = {'1'}; el.vars = {'foo'}; el.varPairsToGpProcess = {'ex', 'ex_smpn',... 'ey','ey_smpn',... 'exy','exy_smpn'}; gd.ind = {{'1'}}; gd.Yield = {'0'}; gd.EffStress = {'0'}; gd.G = {{'0'}}; gd.gporder = {'2'}; el.geomdim{1} = {{},{},gd}; fem.elem = [fem.elem {el}];

See Also

elempty

Purpose	Define variables which evaluate principal values and vector components.
Syntax	<pre>el.elem = 'elpric' el.g{ig} = geomnum el.basename = bname el.postname = pname el.sdim{idim} = dimname el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.tensor{eldomgrp} = matexpr</pre>
Description	The elpric element evaluates eigenvalues and eigenvectors of a 3-by-3 real symmetric matrix. The tensor field always has six components specifying the upper triangle of the source matrix as a cell array of expressions in column order. The basename field is compulsory and specifies a single name from which all output variable names are derived. The output variables are defined wherever the tensor field is nonempty.
	Eigenvalue variable names are created by appending numbers 1 to 3 to bname and, if postname is present, append '_pname'. Eigenvector component names are then created by inserting the space variable names given in sdim directly after the component number. Eigenvalues are sorted in decreasing order.
Cautionary	No Jacobian contribution is calculated even if the tensor expressions contain dependent variables. The reason is the condition that the eigenvalues are sorted, which makes the eigenvector components discontinuous functions of the input matrix components.
Examples	<pre>Define postprocessing variables for principal strains and directions, given strain components [ex,ey,ez,exy,exz,eyz]. el.elem = 'elpric'; el.g = {'1'}; el.basename = 'e'; el.sdim = {'x','y','z'}; gd.ind = {{'1'}}; gd.tensor = {{'ex','exy','ey','exz','eyz','ez'}}; el.geomdim{1} = {{},{},{},gd}; fem.elem = [fem.elem {el}];</pre>
See Also	elempty, elinv

Purpose	Define pointwise constraints controlled by shape functions.
Syntax	el.elem = 'elsconstr' el.g{ig} = geomnum el.nname = Nname el.nfname = NFname el.mname = Mname
	el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.shelem = meshcases shapelist
	<pre>meshcases.default = shapelist; meshcases.case{elmcase} = shapelist; meshcases.mind{elmcase} = caselist;</pre>
	shapelist{eldomgrp}{ishape}{3*i-2} = bmtypename shapelist{eldomgrp}{ishape}{3*i-1} = shapename shapelist{eldomgrp}{ishape}{3*i} = shapeparams
	el.geomdim{ig}{edim}.constr{eldomgrp}{ic}{j} = constrexpr el.geomdim{ig}{edim}.constrf{eldomgrp}{ic}{j} = constrfexpr el.geomdim{ig}{edim}.cshape{eldomgrp}{ic} = ishape
Description	The elsconstr element defines pointwise constraints where the type of constraint is determined by a shape function object. The idea is that the constraint points are selected to be appropriate for variables having the corresponding shape function. For the syntax of the ind field, see elempty. For the syntax of the shelem field, see elshape. A difference to the syntax in elshape is that the cell array shapelist has three levels instead of two.
	The expressions that are used to formulate the constraint are given in the constr field, and the index of the corresponding shape function object is given in the cshape field. More precisely, in element geometry ig, dimension edim, and element domain group eldomgrp, constraint number ic is defined by the shape function object with index ishape=el.geomdim{ig}{edim}.cshape{eldomgrp}{ic}. The expressions needed to formulate this constraint is given by the cell array el.geomdim{ig}{edim}.constr{eldomgrp}{ic}. The number of expressions ne in this cell array depends on the shape function object.
	The optional constrf field controls the way constraint forces are applied. You enter an expression such that its Jacobian with respect to the test functions decides on which degrees of freedom the reaction force is applied for each constraint. If constrf is omitted, constraints are <i>ideal</i> , which corresponds to setting the components of constrf to test(constrexpr).

The constraint and constraint force Jacobians are by default assembled to matrices called NT and NF, and the constraint residual is called M. This can be changed by assigning different names to the optional nname, nfname and mname fields. Only certain names are recognized by functions like assemble, though, see page 29.

The elsconstr constraint element is only implemented for the shape functions shlag, shcurl, and shdiv. For shlag, the number of expressions ne=1, and this expression is constrained to be zero in the node points of the shlag object. For shcurl, the number of expressions ne=sdim, and these expressions are considered as components of a vector. The tangential component of this vector is constrained to be zero in the node points for the shcurl shape function. For shdiv, the number of expressions ne=sdim, and these expressions are considered as components of a vector. The shcurl shape function. For shdiv, the number of expressions ne=sdim, and these expressions are considered as components of a vector. The normal component of this vector is constrained to be zero in the node points for the shdiv shape function.

Example

Impose a constraint on a vector field E represented using shcurl shape functions of order 2. The constraint is that the tangential component of E-(2,3) is zero.

```
clear fem;
  fem.aeom = circ2:
  fem.mesh = meshinit(fem);
  fem.shape = 'shcurl(2,''E'')';
  fem.dim = {'Ex' 'Ey'};
  fem.equ.weak = '-(Ex*Ex_test+Ey*Ey_test+dExy_test*dExy)';
  clear el gd;
  el.elem = 'elsconstr';
  el.g = {'1'};
  gd.ind = {{'1', '2', '3', '4'}};
  gd.shelem{1}{1} = ...
    {'s(1)','shcurl',struct('fieldname','E','order','2')};
  gd.constr{1}{1} = {'Ex-2', 'Ey-3'};
  gd.cshape{1}{1} = '1';
  el.geomdim{1} = {{},gd,{}};
  fem.elem = {el};
  fem.xmesh = meshextend(fem);
  fem.sol = femstatic(fem);
  postarrow(fem,{'Ex' 'Ey'});
elempty, elpconstr, elshape, shdiv, shlag, shcurl
```

See Also

Purpose	Define dependent variables and select shape functions.
Syntax	<pre>el.elem = 'elshape' el.g{ig} = geomnum el.tvars = 'on' 'off' el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.shelem = meshcases shapelist</pre>
	<pre>meshcases.default = shapelist; meshcases.case{elmcase} = shapelist; meshcases.mind{elmcase} = caselist;</pre>
	shapelist{eldomgrp}{3*ishape-2} = bmtypename shapelist{eldomgrp}{3*ishape-1} = shapename shapelist{eldomgrp}{3*ishape} = shapeparams
Description	The elshape element is responsible for allocating degrees of freedom and defining dependent variables. For the syntax of the ind field, see elempty. The tvars field turns the generation of time derivative variables on or off (default on).

The shelem field has a rather complicated syntax. If no alternate mesh cases are defined, it is a cell array which for each domain group contains a cell array of triplets bmtypename-shapename-shapeparams. The string bmtypename is a unique identifier for a basic mesh element shape with certain additional properties, see the table below.

NAME	DESCRIPTION
ls(0)	0D simplex (all elements are equivalent in 0D)
s(1)	ID simplex, higher-order shape generated on boundaries in 2D and 3D
ls(I)	ID simplex, linear shape generated in ID
s(2)	2D simplex (triangle), higher-order shape generated on boundaries in 3D and in a layer closest to boundaries in 2D
ls(2)	2D simplex (triangle), linear shape generated in the inner of 2D domains
b(2)	2D brick (quadrilateral, quad), higher-order shape generated on boundaries in 3D and in a layer closest to boundaries in 2D
lb(2)	2D brick (quadrilateral, quad), bilinear shape generated in the inner of 2D domains
s(3)	3D simplex (tetrahedron), higher-order shape generated in a layer closest to boundary surfaces and free edges

TABLE I-20: BASIC MESH ELEMENT TYPE IDENTIFIERS FOR ELEMENT TYPES GENERATED IN MESHES

NAME	DESCRIPTION
ls(3)	3D simplex (tetrahedron), linear shape generated away from boundaries and edges in 3D
b(3)	3D brick (hexahedron, hex), higher-order shape generated in a layer closest to boundary surfaces and free edges
lb(3)	3D brick (hexahedron, hex), trilinear shape generated away from boundaries and edges in 3D
prism	3D prism (pentahedron, wedge), higher-order shape generated in a layer closest to boundary surfaces and free edges
lprism	3D prism (pentahedron, wedge), bilinear shape generated away from boundaries and edges in 3D

TABLE I-20: BASIC MESH ELEMENT TYPE IDENTIFIERS FOR ELEMENT TYPES GENERATED IN MESHES

The shapename identifier selects a shape function for the base mesh type, and the format of the shapeparams parameter in turn depends on the particular shape function. Typically, the shape function expects a struct with fields specifying dependent variable name and order.

If there are multiple mesh cases present in the model, the shelem field is a struct with fields default, case, and mind. The default field has the same syntax as described above for the shelem field itself. Multiple alternate cases which use the same shape functions can be grouped together using the mind field. This field is a cell array containing groups of mesh case numbers, each group, caselist, given as a cell array. For each element mesh case group, elmcase, an alternate shape list is given in the case field.

Cautionary	Multiple shape functions can be specified by the same elshape element simply by
	repeating a basic mesh type name with different shape function and/or parameters
	in the shelem field. This means that there also has to be some conflict resolution.
	When multiple shape functions specify the same dependent variable name, the one
	with the highest interpolation order for the basic field prevails.

The field variables defined on a given domain are the union of the variables defined in the default case by shape functions on all basic mesh element types on that domain. This means that variables can at times be missing on certain mesh element types or for certain mesh cases.

Examples Add a dependent variable 1m on the boundary that can be used as a Lagrange multiplier in a weak constraint.

clear fem;

```
fem.geom = circ2;
  fem.mesh = meshinit(fem);
  fem.shape = 2;
  fem.equ.c = 1; fem.equ.f = 1;
  fem.bnd.weak = 'lm test*u+lm*u test';
  fem.elem = {};
  clear el;
  el.elem = 'elshape';
  el.g = {'1'};
  el.tvars = 'off';
  gd.ind = {{'1', '2', '3', '4'}};
  gd.shelem = ...
    {{'s(1)', 'shlag', struct('basename', 'lm', 'order', '2')}};
  el.geomdim{1} = {{},gd,{}};
  fem.elem = [fem.elem {el}];
  fem.xmesh = meshextend(fem);
  fem.sol = femstatic(fem);
  postplot(fem,'tridata','u','triz','u');
elempty, sharg_2_5, shbub, shdens, shdiv, shgp, shlag, shcurl
```

Purpose	Create a linear flat faceted shell element.
Syntax	<pre>el.elem = 'elshell_arg2' el.g{ig} = geomnum el.dim = depvarnames el.equation = equation type el.omega = frequency el.postname = postfix el.geomdim{ig}{3}.ind{eldomgrp} = domainlist el.geomdim{ig}{3}.E{eldomgrp} = E_expr el.geomdim{ig}{3}.ru{eldomgrp} = nu_expr el.geomdim{ig}{3}.ru{eldomgrp} = rho_expr el.geomdim{ig}{3}.rho{eldomgrp} = th_expr el.geomdim{ig}{3}.thickness{eldomgrp} = th_expr el.geomdim{ig}{3}.alphadM{eldomgrp} = alpha_expr el.geomdim{ig}{3}.alphadM{eldomgrp} = beta_expr el.geomdim{ig}{3}.xlocalx{eldomgrp} = xlx_expr el.geomdim{ig}{3}.xlocalx{eldomgrp} = xly_expr el.geomdim{ig}{3}.xlocalz{eldomgrp} = nx_expr el.geomdim{ig}{3}.nsidex{eldomgrp} = ny_expr el.geomdim{ig}{3}.nsidez{eldomgrp} = nz_expr</pre>
Description	The elshell_arg2 element describes a linear Mindlin theory shell made up of essentially constant-strain triangles with added drilling rotations. The element lives on a 2D surface embedded in a 3D geometry. Its material properties, constraints and loads are specified directly in the element syntax structure. An elshell_arg2 element implements tasks which are handled by an eleqc or eleqw element when using the standard syntax. That is, it directly assembles contributions to the stiffness and mass matrices and to the residual vector. In addition, it defines a number of postprocessing variables. The shell element structure contains global properties, common to the entire shell, as well as local material properties on the boundary level. Note that the shell exists only on the boundary level and below.
	Note: The elshell_arg2 element requires a triangular mesh and will not work with a quadrilateral mesh.

Global properties: The equation field specifies whether to treat the problem as stationary, time harmonic or time-dependent. The three displacement and three rotation field variable names must be specified in the dim field, for example,

e.dim = {'u','v','w','thx','thy','thz'}

The displacement fields are most easily defined using 6 separate shlag objects of order 1.

FIELD	MEANING	SYNTAX	DEFAULT VALUE
elem	Shell element name	'elshell_arg2'	-
g	Geometry index	scalar number	1
dim	Field variable names for displacements and rotations	I-by-6 cell vector of strings	-
equation	Affects how matrices are assembled	'static' 'freq' 'time' 'eigen'	
omega	Frequency for the 'freq' equation	string expression	'0'
postname	Name that is appended to postprocessing variables	string	empty strin

TABLE 1-21: GLOBAL PROPERTIES OF THE ELSHELL_ARG2 SHELL ELEMENT STRUCTURE

Material, loads and constraints: The shell described by the elshell_arg2 element can be considered a collection of discrete, homogeneous, flat triangles. The material properties, including damping factors as well as the element thickness, are taken to be constant within any triangle. The syntax of the material properties, loads and constraints is analogous to the syntax of the coefficient form *level 1 coefficients*. See further the section "Specifying a Model" in the *COMSOL Multiphysics Scripting Guide*. However note that all values as well as expressions must be strings.

TABLE 1-22: BND LEVEL PROPERTIES IN THE ELSHELL_ARG2 SHELL ELEMENT STRUCTURE

FIELD	MEANING	SYNTAX
E	Elasticity modulus / Young's modulus	level I coefficient
nu	Poisson's ratio	level I coefficient
rho	Density	level I coefficient
thickness	Shell thickness	level I coefficient
height	Postprocessing level	level I coefficient
alphadM	Mass damping coefficient (submodes 'time' and 'freq' only)	level I coefficient
betadK	Stiffness damping coefficient (submodes 'time' and 'freq' only)	level I coefficient

FIELD	MEANING	SYNTAX	
xlocalx, xlocaly, xlocalz	Vector, whose projection on the shell defines the local x direction	level I coefficient	
nsidex, nsidey, nsidez	Direction vector which defines the "up" side of the shell	level I coefficient	

TABLE I-22: BND LEVEL PROPERTIES IN THE ELSHELL_ARG2 SHELL ELEMENT STRUCTURE

Postprocessing variables: The postprocessing variables defined by the elshell_arg2 element have standard names that do not depend on the names of the space variables given in fem.sdim. The postname property of the shell element structure, if not the empty string, is appended to all postprocessing variables. For example, the direct x strain will be referenced as exs or exs_postname, depending on the value of the postname field.

TABLE 1-23: POSTPROCESSING VARIABLES DEFINED BY THE ELSHELL_ARG2 ELEMENT

VARIABLE	MEANING
exs, eys, ezs, exys, exzs, eyzs	Strain tensor components in global coordinates
exls, eyls, ezls, exyls, exzls, eyzls	Strain tensor components in local coordinates
Nxls, Nyls, Nxyls	In-plane forces in local coordinates
Qxls, Qyls	Out-of-plane forces in local coordinates
Mxls, Myls, Mxyls	In-plane moments in local coordinates
exlxs, exlys, exlzs	Local system x-axis expressed in global coordinates
eylxs, eylys, eylzs	Local system y-axis expressed in global coordinates

Theory: The elshell_arg2 shell element is a combination of an isotropic version of the TRIC element proposed by Argyris and others (Ref. 1) and the constant strain triangle with drilling rotations due to Allman (Ref. 2). As such, the element is essentially a constant strain triangle whose displacement field vary linearly in the direction tangential to each edge, and as a restricted third order polynomial in the normal direction.

The material properties are considered to be constant within any triangle, and therefore symbolic integration can be used to describe an element stiffness matrix and a consistent element mass matrix in terms of element geometry and material data.

Cautionary	The elshell_ar2 shell element is not multiphysics enabled. This means that there will be no contributions to the exact Jacobian from solution-dependent material data.
	The elshell_arg2 element is available only if your license includes the Structural Mechanics Module.
Compatibility	COMSOL Multiphysics 3.2: The tdim field and wave extension in the time-dependent case are no longer used.
See Also	elempty
References	1. J. Argyris, L. Tenek, and L. Olofsson, "TRIC: a simple but sophisticated 3-node triangular element based on 6 rigid-body and 12 straining modes for fast computational simulations of arbitrary isotropic and laminated composite shells," <i>Comput. Methods Appl. Mech. Engrg.</i> , vol. 145, pp. 11–85, 1997.
	2. D. J. Allman, "Evaluation of the constant strain triangle with drilling rotations," <i>Int. J. Numer. Meth. Engrg.</i> , vol. 26, pp. 2645–2655, 1988.

Syntax	<pre>el.elem = 'eluwhelm' el.g{ig} = geomnum el.basename = fieldname el.ndir = ndir el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.Rho{eldomgrp} = rho el.geomdim{ig}{edim}.K{eldomgrp} = k el.geomdim{ig}{edim}.Q{eldomgrp} = q</pre>
Description	The eleqw element uses the Ultraweak variational formulation (UWVF) to implement a Helmholtz equation for the acoustic pressure, p ,
	$\nabla \bullet \left(-\frac{\nabla p}{\rho_0} \right) - \frac{k^2}{\rho_0} p = 0$
	with boundary conditions of the form
	$n \cdot \frac{\nabla p}{\rho_0} + i \frac{k}{\rho_0} p = q \left(-n \cdot \frac{\nabla p}{\rho_0} + i \frac{k}{\rho_0} p \right) + g$
	The density ρ_0 , wave number k , and parameter q are supplied directly to the element, while the boundary term g and all volume, point, and edge sources must be implemented separately, outside the element.
	The acoustic field variable fieldname must be represented by an shuwhelm shape function with fixed number of directions, ndir, throughout the domains where the eluwhelm element is active.
Cautionary	The eluwhelm element does not currently account for curved boundaries—all element edges and faces are assumed to be planar. This may change, which can possibly affect future element syntax.
See Also	elempty, shuwhelm

Assemble acoustic Helmholtz equation on ultraweak variational form.

Purpose

Purpose	Define expression variables.
Syntax	el.elem = 'elvar' el.g{'ig'} = geomnum el.geomdim{ig}{edim}.ind{eldomgrp} = domainlist el.geomdim{ig}{edim}.var{2*ivar-1} = varname el.geomdim{ig}{edim}.var{2*ivar}{eldomgrp} = varexpr
Description	The elvar element declares expression variables varname to be accessible on domain groups for which the defining expression varexpr is nonempty. For the syntax of the ind field, see elempty.
Examples	<pre>Redefine the space derivatives of u on an interior boundary to be evaluated on the "up" side instead of being averaged. clear fem; fem.geom = geomcsg({rect2(1,1,'pos',[-1 0]),rect2}); fem.mesh = meshinit(fem); fem.equ.ind = [1 2]; fem.equ.ind = [1 2]; fem.bnd.ind = [1 0 0 0 0 0 2]; fem.bnd.h = 1; fem.bnd.h = 1; fem.bnd.r = {0 1}; fem.elem = {}; clear el; el.elem = 'elvar'; el.g = {'1'}; clear gd; gd.ind = {{'4'}}; gd.var = {'ux',{'up(ux)'},'uy',{'up(uy'}}; el.geomdim{1} = {{},gd,{}}; fem.elem = [fem.elem {el}]; fem.xmesh = meshextend(fem); fem.sol = femstatic(fem);</pre>
See Also	<pre>postplot(fem,'lindata','ux','linz','ux'); elempty</pre>

Purpose	Embed a 2D geometry object as a 3D geometry object.
Syntax	g3 = embed(g2) g3 = embed(g2,p_wrkpln)
Description	g3 = embed(g2) embeds the 2D geometry object as a 3D geometry object. A 2D solid object becomes a 3D face object, a 2D curve object becomes a 3D curve object, and a 2D point object becomes a 3D point object.
	g3 = embed(g2,p_wrkpln) additionally, p_wrkpln specifies the position in the 3D space. See geomgetwrkpln for more information on p_wrkpln.
See also	extrude,curve2, curve3,face3,geom0, geom1, geom2, geom3,point1, point2, point3

Purpose	Extrude a 2D	Extrude a 2D geometry object into a 3D geometry object.			
Syntax	g3 = extruc	g3 = extrude(g2,)			
Description	g3 = extrude(g,) extrudes the 2D geometry object g into a 3D geometry object g3 according to given parameters.				
The function extrude accepts the following property/values: TABLE 1-24: VALID PROPERTY/VALUE PAIRS			pperty/values:		
	PROPERTY VALUES DEFAULT DESCRIPTION			DESCRIPTION	
	Displ	2-by- <i>nd</i> matrix	[0;0]	Displacement of extrusion top	
	Distance	k-by- nd matrix	1	Extrusion distances	
	Face	string	'all'	Cross-sectional faces to delete	
	Polres	scalar	50	Polygon resolution	
	Scale	2-by- nd matrix	[1;1]	Scale of extrusion top	
	Twist	I-by- <i>nd</i> vector	0	Twist angle (in radians)	
	Wrkpln	3-by-3 matrix	[0 1 0; 0 0 1; 0 0 0]	Work plane for 2D geometry cross section	

The 3D object g3 is an extruded object, where Distance is the extrusion distance in the normal direction of the bottom plane, defined by the property Wrkpln.

The properties Displ, Scale, and Twist defines the translation displacements, scale factors and rotation of the top with respect to the bottom of the extruded object. They are defined in the local system of the work plane.

To define a piecewise linear extrusion, Distance is given as a row vector, of size 1-by-*nd*, of displacements with respect to the bottom work plane. Scale, Displ, and Twist need to have the same number of columns as Distance.

To define a cubic extrusion Distance is given as a 3-by-*nd* matrix where rows 2 and 3 contain weights of the extrusion segments. The weights are given in the interval [0 1] and specifies the influence of the tangential continuity at the junctions. The weights of rows 2 and 3 specifies the influence from the first- and second-junction, respectively, of each segment. If the weight is close to 0, the influence of the junction is weak, and if it is close to 1, the influence is strong.

Polres defines the resolution in the polygon representations of the edges.

Face specifies if cross-sectional faces are removed: 'all' removes them, 'none' keeps them.

Compatibility	The numbering of faces, edges and vertices is different from the numbering in objects created in 2.3.
Examples	Creation of a cylinder of height 1.3.
	g3 = extrude(circ2,'distance',1.3);
	Extrusion of rectangle from a <i>zx</i> -plane.
	<pre>p_wrkpln = geomgetwrkpln('quick',{'zx',10}); g3 = extrude(rect2(1,2),'distance',1.3,'displ',[0.4;0], 'scale',[2;2],'wrkpln',p_wrkpln); geomplot(g3);</pre>
	Cubic extrusion of a circle.
	g3 = extrude(circ2,'distance',[1 3 4;0.3 0.3 0.3;0.3 0.3 0.3], 'scale',[1 1.5 2;1 1 2],'twist',[0 pi/6 pi/6], 'displ',[0 0 0;0 1 1]);
See Also	geomO, geom1, geom2, geom3,geomcsg,geomgetwrkpln

Purpose	Create 3D surf	ace geometry o	bject.	
Syntax	<pre>f3 = face3(x,y,z) f3 = face3(x,y,z,w) f3 = face3(vtx,vtxpre,edg,edgpre,fac,mfdpre,mfd) [f3,] = face3(g3,) f3 = face3(g2)</pre>			
Description	surfaces is deter	rmined from the equal size. If s i	e size of the r Lze(x,2)>1	f3. The degree of the rational Bézier natrices x, y, and z. The arrays x, y, and then a rectangular patch is created. If eated.
	triangular plana	r patches, 2-by	-2, 3-by-2, 4	, y, and z are: 3-by-1 for creating -by-2, 3-by-3, 4-by-3, 4-by-4 for 3,1), (2,2), (3,2), and (3,3) respectively.
	f3 = face3(x positive weight		•	the above, but also applies arbitrary e.
	geometry objec	t f3 from the f	ields vtx, vt	ac,mfdpre,mfd) creates a 3D surface xpre, edg, edgpre, fac, mfdpre, and e object. See geom3 for a description of
	[f3,] = f object f3.	ace3(g3,)	coerces the a	3D geometry object g3 to a 3D face
	object f3 is the	n embedded in	the plane z=	v object g2 to a 3D face object f3. The 0 and is a trimmed planar patch since it ılar or triangular Bézier patch.
	The coercion fu property/value		.] = face3	(g3,) a accepts the following
	TABLE I-25: VALID	PROPERTY/VALUE PA	AIRS	
	PROPERTY	VALUE	DEFAULT	DESCRIPTION
	Out	stx ftx ctx ptx	{}	Cell array of output names.

See geomesg and geom for more information on geometry objects.

The 3D geometry object properties are available. The properties can be accessed using the syntax get(object,property). See geom3 for details.

Examples	Create an untrimmed triangular patch in the plane, $y = 1$.
	f1 = face3([0 1 0]',[1 1 1]',[0 0 1]');
	Create a circular face as a trimmed patch in the plane, $z = 0$.
	f2 = face3(circ2(0,20,10));
	A patch that can constitute the wall of a cylinder, is created by setting the control weights explicitly, as in the command below.
	f3 = face3([-1 -1;-1 -1;0 0],[0 0;-1 -1;-1 -1], [0 1; 0 1; 0 1],[1 1;1/sqrt(2) 1/sqrt(2);1 1]);
	To generate a third degree rectangular patch, the following commands can be given.
	<pre>[x,y] = meshgrid(-3:3:6,0:2:6); z = rand(size(x)); f4 = face3(x,y,z);</pre>
Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported.
See Also	curve2, curve3,geom0, geom1, geom2, geom3,geomcsg,point1, point2, point3

Purpose	Symbolically differentiate a PDE in general form.
Syntax	<pre>fem1 = femdiff(fem,) xfem1 = femdiff(xfem,)</pre>
Description	fem1 = femdiff(fem,) symbolically differentiates the Γ , F , G , and R coefficients in a PDE given in general form. xfem can also be an extended FEM structure. In this case, femdiff differentiates all FEM structures in general form. The coefficients are obtained from the ga, f, and g, r fields from the fem.equ and fem.bnd structures, respectively. It returns an FEM structure where the fields equ and bnd have been updated with the fields c, al, be, a, and q, h according to "The Linear or Linearized Model" on page 366 in the <i>COMSOL Multiphysics User's Guide</i> .
	The expressions in the coefficients Γ , F , G , and R can contain expressions containing the binary operators +, -, *, /, ^, ==, ~=, >, >=, <, <=, , and &; the unary operators +, -, and ~; and the functions abs, acos, acosh, acot, acoth, acsc, acsch, asec, asech, asin, asinh, atan, atanh, cos, cosh, cot, coth, csc, csch, erf, exp, lambw, log, log10, log2, sec, sech, sign, sin, sinh, sqrt, tan, and tanh.

The function femdiff accepts the following property/value pairs:

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Defaults	off on	off	Return default fields
Diff	off on cell array containing a selection of ga, g, f, r, var, and expr	on	List of fields which can be differentiated in order to evaluate the Out values. The default string 'on' is equivalent to the cell array {'ga' 'g' 'f' 'r' 'expr'}. Note: not 'var'
Shrink	off on	on	List of differentiation rules
Simplify	off on	on	Simplify differentiated expressions

TABLE I-26: VALID PROPERTY/VALUE PAIRS

The properties Diff, Rules, and Simplify can alternatively be given as fields in the FEM structure: fem.diff, fem.rules, and fem.simplify.

	Use the field fem.rules to specify additional differentiation rules. The derivative of the inverse hyperbolic tangent function atanh can, for example, be specified as
	{'atanh(x)','1/(1-x^2)'}.
	It can also be stored as a field in the FEM structure.
	Assume a user-defined function $foo(a,b)$ has been written, implementing the analytical expression $a^2+a*sin(b^3)$. The derivatives of this function are specified as:
	'rules', {'foo(a,b)', '2*a+sin(b^3),3*a*b^2*cos(b^3)'}
	femdiff does not support functions with string arguments.
Cautionary	The relational and Boolean operators have been included for convenience only, and must be used with extreme caution. Their symbolic derivative is considered to be identically 0.
	Coefficient M-files are not allowed in the input.
Compatibility	The function flgetrules for converting FEMLAB $1.0/1.1$ differentiation rules is no longer available.
	The properties bdl, out, rules, and sdl are obsolete in FEMLAB 3.0.
	The fields fem.equ.varu and fem.bnd.varu, etc. are no longer generated in FEMLAB 3.0.
	The precedence rules for the operators and & have been changed to comply with MATLAB 6.0 precedence.
	The differentiation algorithm is new in FEMLAB 1.2. The @fldiffobj class is obsolete.
See Also	femnlin

Purpose	Solve eigenvalue PDE problem.
Syntax	fem.sol = femeig(fem,) [u,lambda] = femeig(fem,) fem = femeig(fem,'Out',{'fem'},) fem.sol = femeig('In',{'D' D 'K' K 'N' N},)
Description	<pre>fem.sol = femeig(fem,) assembles and solves the eigenvalue PDE problem described by the (possibly extended) FEM structure fem.</pre>
	fem.sol = femeig('In', {'D' D 'K' K 'N' N},) solves the eigenvalue problem given by the matrices D, K, and N.
	For both linear and nonlinear PDE problems, the eigenvalue problem is that of the linearization about a solution U_0 . If the eigenvalue appears nonlinearly, COMSOL

linearization about a solution U_0 . If the eigenvalue appears nonlinearly, COMSOL Multiphysics reduces the problem to a quadratic approximation around a value λ_0 specified by the property eigref. The discretized form of the problem reads

$$KU - (\lambda - \lambda_0)DU + (\lambda - \lambda_0)^2 EU = -N_F \Lambda$$
$$NU = M$$

where K, D, E, N and N_F are evaluated for $U = U_0$ and $\lambda = \lambda_0$. A is the Lagrange multiplier vector, λ is the eigenvalue. The eigenvalue name can be given by the property eigname. The linearization point U_0 can be given with the property U. The shift, described below, is compensated according to the linearization point for the eigenvalue. Therefore, changing the linearization point has no effect at all for linear or quadratic eigenvalue problems.

The function femeig accepts the following property/value pairs:

TABLE I-27: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Eigname	string	lambda	Name of eigenvalue variable
Eigref	string	0	Linearization point for the eigenvalue
Etol	positive scalar	0	Eigenvalue tolerance
In	cell array of names and matrices K N D E	N is empty E=0, D=0	Input matrices
Krylovdim	positive integer		Dimension of Krylov space

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Neigs	positive integer	6	Number of eigenvalues sought
Out	fem sol u lambda stop solcompdof Kc Dc Ec Null Nullf Nnp uscale nullfun symmetric	sol	Output variables
Shift	scalar	0	Eigenvalue search location

TABLE I-27: VALID PROPERTY/VALUE PAIRS

In addition, the properties described in the entry femsolver are supported.

Specify where to look for the desired eigenvalues with the property shift. Enter a real or complex scalar; the default value is 0, meaning that the solver tries to find eigenvalues close to 0.

When using an iterative method as linear system solver, the tolerance used in the convergence criterion is Itol times a saftey factor of 10^{-4} . When a direct method is used as linear system solver, with error checking enabled, Itol is used in the convergence criterion without any extra safety factor.

Using the property In you can specify explicit values for the matrices in the eigenvalue problem. The value for this property is a cell array with alternating matrix names and matrix values. The matrix names can be D, E, K, or N. If the N matrix is not given, it is taken to be empty. If the D or E matrix is not given, it is taken to be 0.

The property Out defines the output variables and their order. The output fem means the FEM structure with the *solution object* fem.sol added. sol is a femsol object containing the fields lambda and u. lambda is a row vector containing the eigenvalues. U is a *solution matrix*. Each column in the solution matrix is the solution vector of the eigenfunction for the corresponding eigenvalue in lambda. The output value stop returns nonzero if the solution process was not completed. Stop is 1 if a partial solution was returned, and 2 if no solution was returned. For the other outputs, see femlin.

For more information about the eigenvalue solver, see "The Eigenvalue Solver" on page 376 in the *COMSOL Multiphysics User's Guide*.

 Example
 Eigenmodes and Eigenvalues of the L-shaped Membrane

 Compute eigenvalues corresponding to eigenmodes for the PDE problem

```
\begin{cases} -\Delta u = \lambda u & \text{ in } \Omega \\ u = 0 & \text{ on } \partial \Omega \end{cases}
```

where Ω is the L-shaped membrane. Start by setting up the problem:

```
clear fem
fem.geom = poly2([-1 0 0 1 1 -1],[0 0 1 1 -1 -1]);
fem.mesh = meshinit(fem, 'hmax',0.1);
fem.shape = 2;
fem.equ.c = 1; fem.equ.da = 1;
fem.bnd.h = 1;
fem.xmesh = meshextend(fem);
fem.sol = femeig(fem, 'neigs',16);
```

Display the first and sixteenth eigenmodes. The membrane function is available in MATLAB and not in COMSOL Script.

postsurf(fem,'u')	%	first eigenmode
membrane(1,20,9,9)	%	the MATLAB function
<pre>postsurf(fem,'u','solnum',16)</pre>	%	sixteenth eigenmode

Cautionary	Consider the case of a linear eigenvalue problem, $E = 0$.Write the generalized eigenvalue problem as $(A - \lambda B)u = 0$. In the standard case the coefficients c and d_a are positive in the entire region. All eigenvalues are positive, and 0 is a good choice for the shift (eigenvalue search location). The cases where either c or d_a is zero are discussed below.
	• If $d_a = 0$ in a subregion, the mass matrix B becomes singular. This does not cause any trouble, provided that $c > 0$ everywhere. The pencil (A,B) has a set of infinite eigenvalues.
	• If $c = 0$ in a subregion, the stiffness matrix A becomes singular, and the pencil (A,B) has many zero eigenvalues. Choose a positive shift below the smallest nonzero eigenvalue.
	• If there is a region where both $c = 0$ and $d_a = 0$, we get a singular pencil. The whole eigenvalue problem is undetermined, and any value is equally plausible as an eigenvalue.
Compatibility	The property Variables has been renamed to Const in FEMLAB 2.3. The properties Epoint and Tpoint are obsolete from FEMLAB 2.2. Use fem.***.gporder to specify integration order.

See Also femsolver, assemble, femlin

Purpose	Solve linear or linearized stationary PDE problem.
Syntax	<pre>fem.sol = femlin(fem,) fem = femlin(fem,'Out', {'fem'},) [Ke,Le,Null,ud] = femlin(fem,) [Kl,Ll,Nnp] = femlin(fem,) [Ks,Ls] = femlin(fem,) fem.sol = femlin('In',{'K' K 'L' L 'M' M 'N' N 'NF' NF},)</pre>
Description	<pre>fem.sol = femlin(fem) solves a linear or linearized stationary PDE problem described by the (possibly extended) FEM structure fem. See femstruct for details on the FEM structure.</pre>
	<pre>fem.sol = femlin(fem, 'pname', 'P', 'plist',list,) solves a linear or linearized stationary PDE problem for several values of the parameter P. The values of the parameter P are given in the vector list.</pre>
	<pre>fem = femlin(fem, 'out', {'fem'}) modifies the FEM structure to include the solution structure, fem.sol.</pre>
	[Ke,Le,Null,ud] = femlin(fem) partially solves the PDE problem by eliminating the constraints. The solution of PDE problem can be obtained by the scripting command u = u0+Null*(Ke\Le)+ud, where u0 is the linearization point.
	$[Kl,Ll,Nnp] = femlin(fem)$ partially solves the PDE problem by using the Lagrange method. The solution can then be obtained by $u = Kl\Ll$, and then $u = u0+u(1:Nnp)$.
	$[Ks,Ls] = femlin(fem)$ partially solves the PDE problem by approximating the constraints with stiff springs. The solution to the PDE problem is $u = u0+Ks\Ls$.
	fem.sol = femlin('in',{'K' K 'N' N 'NF' NF 'L' L 'M' M}) solves a pre-assembled PDE problem.
	<pre>u = femlin('in', {'K' K 'L' L}, 'out', 'u') is equivalent to solving the linear system using u = K\L, with the important difference that you have access to all linear system solvers (except Geometric multigrid) using the Linsolver property.</pre>
	Consider the finite element discretization of a stationary PDE problem:
	$0 = \begin{bmatrix} L - N_F \Lambda \\ M \end{bmatrix},$
	where L, N_F , and M depend on the solution vector U . femlin solves the linearized form of this problem:

$$\begin{bmatrix} K N_F \\ N & 0 \end{bmatrix} \begin{bmatrix} U - U_0 \\ \Lambda \end{bmatrix} = \begin{bmatrix} L \\ M \end{bmatrix},$$

where K, N_F, N, L , and M are evaluated for $U = U_0$. Thus, if the original problem is linear and K is the correct Jacobian, femlin computes the solution of the original problem. The linearization "point" U_0 can be specified with the property U.

femlin can also partially solve the eigenvalue problem:

$$KU - (\lambda - \lambda_0)DU + (\lambda - \lambda_0)^2 EU = -N_F \Lambda$$
$$NU = M$$

in that it transforms the problem using one of the constraint-handling methods. Here λ is the eigenvalue, the name can be controlled by the property eigname. λ_0 is the eigenvalue linearization point, the value can be controlled by the property eigref.

The function femlin accepts the following property/values:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Eigname	string	lambda	Eigenvalue name
Eigref	string	0	Linearization point for the eigenvalue
In	cell array of names and matrices K L M N NF D E	N and M are empty, D=E=0, NF=N ^T	Input matrices
Кеер	string containing K, N auto	auto	Parameter-independent quantities
Oldcomp	cell array of strings		Old parameter components
Out	<pre>fem sol u plist stop solcompdof Kc Lc Dc Ec Null Nullf Nnp ud uscale nullfun symmetric cell array of these strings</pre>	sol	Output variables

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Pinitstep	positive real		Initial stepsize for parameter
Plist	real vector		List of parameter values
Pmaxstep	positive real		Maximum stepsize for parameter
Pminstep	positive real		Minimum stepsize for parameter
Pname	string		Parameter name
Porder	0	0	Predictor order for parameter stepping
Stopcond	string with expression		Stop parameter stepping before expression become negative

TABLE I-28: VALID PROPERTY/VALUE PAIRS

In addition, the properties described in the entry femsolver are supported.

The parametric solver properties Oldcomp, Pinitstep, Plist, Pmaxstep, Pminstep, Pname, Porder, and Stopcond are described under femnlin.

The property In explicitly provides assembled matrices. Its value is a cell array with alternating matrix names and matrix values. The allowed matrix names are K, L, M, N, NF, D, and E.

The property Out explicitly sets output variables and their order. The output variable fem means the FEM structure with the solution object fem.sol added. The outputs sol, u, and plist are the solution object, the solution matrix (sol.u), and the parameter list (sol.plist), respectively. The output value stop is 0 if a complete solution was returned, 1 if a partial solution was returned, and 2 if no solution was returned. The output solcompdof is a vector containing the indices of the degrees of freedom solved for. The output matrix Kc and the vector Lc are the matrix and right-hand side of the linear system after constraint handling; see "Constraint Handling" on page 499. The matrices Dc and Ec are the corresponding damping matrix and mass matrix after constraint handling for an eigenvalue or time-dependent problem. The outputs Null, Nullf and ud are related to the eliminate constraint handling method. The outputs Nnp and uscale are the number of degrees of freedom solved for and the scale factors used in the rescaling of the degrees of freedom; see "Scaling of Variables and Equations" on page 497. The

outputs nullfun and symmetric can be useful in finding out the result of the automatic null function or the automatic symmetric mechanisms.

Example

The L-Shaped Membrane with Three Subdomains

Take a look at the geometry of the L-shaped membrane for examples of what you can do. First create the L-shaped membrane and examine the subdomain labels and edge segment labels by plotting:

```
clear fem
sq1 = square2(0,0,1);
sq2 = move(sq1,0,-1);
sq3 = move(sq1,-1,-1);
fem.geom = sq1+sq2+sq3;
fem.mesh = meshinit(fem);
geomplot(fem,'edgelabel','on','sublabel','on')
```

Say you want to use c = 1, 1/2, and 1/3 and f = x, y, and x^2+1 in subdomains 1, 2, and 3, respectively. Use Dirichlet boundary conditions on the outer boundaries:

```
fem.shape = 2;
fem.equ.c = {1 1/2 1/3};
fem.equ.f = {'x' 'y' 'x^2+1'};
fem.bnd.h = 1;
fem.xmesh = meshextend(fem);
fem.sol = femlin(fem);
postsurf(fem,'u')
```

Using anisotropic $c = \begin{bmatrix} 2 & x+y \\ x+y & 10 \end{bmatrix}$ and f = 1 in all subdomains can be done by

```
fem.equ.c = {{{2 'x+y' 10}}};
fem.equ.f = 1;
fem.xmesh = meshextend(fem);
fem.sol = femlin(fem);
postsurf(fem,'u')
```

Cautionary

When using the general form it is assumed that the coefficients c, α , β , a, q, h have been computed using fem=femdiff(fem). In the user interface, this is done automatically.

Compatibility The property Variables has been renamed to Const in FEMLAB 2.3.

	If scaling is used, the matrix outputs from femlin are derived from the rescaled system. This means that the scale factors uscale have to be taken into account if a solution is computed from the matrices. See "Scaling of Variables and Equations" on page 497.
	The properties Epoint and Tpoint are obsolete from FEMLAB 2.2. Use fem.***.gporder to specify integration order.
	The properties u and t have been made obsolete in FEMLAB 1.1.
See Also	femsolver, femstruct, assemble, asseminit, femnlin, femeig, flnull

Purpose	Create a mesh object.
Syntax	<pre>fem.mesh = femmesh(p, el)</pre>
Description	<pre>fem.mesh = femmesh(p, el) creates a mesh object from the mesh data stored in p and el.</pre>
	n is an sqim-by-nn matrix containing the coordinates of the mesh vertices. The r_{-}

p is an sdim-by-np matrix containing the coordinates of the mesh vertices. The x-, y-, and z-coordinates are stored in the first, second, and third row, respectively. np is the number of mesh vertices.

el is a cell array of structures with mesh element information. Each structure stores information on elements of a specific type.

STRUCTURE FIELD	VALUE	DESCRIPTION	
type	one of the strings vtx edg tri quad tet prism hex	Element type. The valid element types are: vertex element (vtx), edge element (edg), triangular element (tri), quadrilateral element (quad), tetrahedral element (tet), prism element (prism), and hexahedral element (hex)	
elem	matrix of size nNodes-by-nElem	Mesh vertex indices for the element points. nElem is the number of elements and nNodes is the number of element points	
dom	matrix of size I-by-nElem	Geometry domain numbers	
param	matrix of size nParam-by-nElem	Geometry parameter values	
ud	matrix of size 2-by-nElem	Up- and down-side subdomain numbers. The first row contains the up-side subdomain numbers and the second row the down-side subdomain numbers	

The field param is only valid for elements of dimension 1, that is, edge elements, in 2D and 3D, and for elements of dimension 2, that is, triangular or quadrilateral elements, in 3D. For each edge element, the first and second row contain the starting and ending parameter value, respectively, in 2D, and starting and ending arc length value, respectively, in 3D. For each triangular and quadrilateral element, the rows contain the first and second parameter values for each element corner.

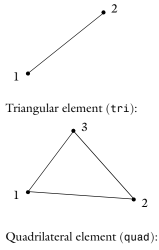
The field ud is only valid for elements of dimension sdim-1, also referred to as *boundary elements*. The direction of the normal vector of a boundary element

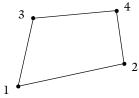
defines the up-side and down-side of the boundary element. For a 1D boundary element, the normal points to the left, considering the direction of the boundary element. For a 2D boundary element, the normal is defined as the cross product of the vector going from the first to the second element corner and the vector going from the third element corner.

The properties p and el can be accessed using the syntax get(object, property).

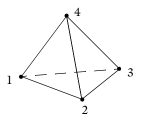
The (local) numbering of the corners of an element is defined according to the following.

Edge element (edg):

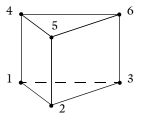




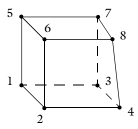
Tetrahedral element (tet):



Prism element (prism):



Hexahedral element (hex):

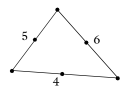


For second-order mesh element types, the strings edg2, tri2, quad2, tet2, prism2, and hex2 are used. The second-ordered nodes are numbered after the corner vertices according to the following.

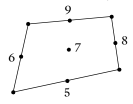
Edge element (edg2):

3

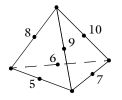
Triangular element (tri2):



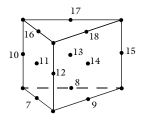
Quadrilateral element (quad2):



Tetrahedral element (tet2):



Prism element (prism2):

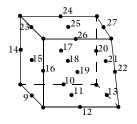


The mid node number for each quadrilateral face of the prism element can also be seen in the following table.

FACE (EDGE NODES)	FACE MID NODE
7,10,12,16	11

FACE MID NODE
13
14

Hexahedral element (hex2):



The mid-node number for each quadrilateral face of the hexahedral element can also be seen in the following table.

FACE (EDGE NODES)	FACE MID NODE
9,10,12,13	11
9,14,16,23	15
10,14,20,24	17
12,16,22,26	19
13,20,22,27	21
23,24,26,27	25

The mid-node number for the hexahedral element is 18.

When importing meshes with reduced second-order elements, also called *serendipity elements*, the mid node of quadrilateral elements (or quadrilateral faces) and the mid node of hexahedral elements must be added manually. The coordinates for the mid node for a second order quadrilateral element (or quadrilateral face) is calculated from the surrounding nodes according to

0.5*edgNodes-0.25*vertNodes, where edgNodes is the sum of the surrounding (4) edge mid nodes and vertNodes is the sum of the surrounding (4) vertex nodes. For a second order hexahedron, the coordinates of the mid node is calculated from the surrounding nodes according to 0.25*edgNodes-0.25*vertNodes, where edgNodes is the sum of the surrounding (12) edge mid nodes and vertNodes is the sum of the surrounding (8) vertex nodes.

	Degenerated elements (or collapsed elements), that is, elements where two or more nodes refer to the same mesh point, are not allowed.
Compatibility	The FEMLAB 2.3 (and earlier) mesh structure format is a valid input to femmesh as well.
See also	meshinit, meshrefine, meshplot

Purpose	Get mesh object properties.		
Syntax	get(m,prop)		
Description	get(m,prop) returns the value of a property prop for a mesh object m.		
	prop is a string that contains a valid property name. The following tables list the valid property names for mesh objects:		
	TABLE I-29: MESH O	BJECT PROPERTY NAMES	
	PROPERTY NAME	DESCRIPTION	
	Ρ	Mesh vertex coordinates	
	el Element information		
	p is a matrix where each column contains the coordinates for the corresponding mesh vertex. For example, p(:,34) returns the coordinates for Vertex 34.		
	el is a cell array of structures with mesh element information. See femmesh on page 134 for information about the field in these structures.		
	For information about the formats for the vtx property, see "1D Geometry Object Properties" on page 207.		
Example	<pre>Create a triangular mesh and determin the vertices that form mesh element 100: m = meshinit(rect2); el = get(m, 'el');</pre>		
	el{3}.elem	(:,100);	
See Also	femmesh		

Purpose Solve nonlinear stationary PDE problem.

Syntax	fem.sol = femnlin(fem,)
-	<pre>fem = femnlin(fem,'Out',{'fem'},)</pre>

Description

fem.sol = femnlin(fem) solves a stationary PDE problem.

fem.sol = femnlin(fem, 'pname', 'P', 'plist', list,...) solves a stationary PDE problem for several values of the parameter P. The values of the parameter P are given in the vector list.

The PDE problem is stored in the (possibly extended) FEM Structure fem. See femstruct for details.

The solver is an affine invariant form of the damped Newton method. The solver can optionally be combined with Uzawa iterations, often used to solve problems with the augmented Lagrangian technique.

The function femnlin accepts the following property/value pairs:

TABLE 1-30		PROPERTY/VALUE PAIRS
I/(DEL 1-30.	17 (210)	

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Augcomp	cell array of strings		Augmented Lagrange components
Augmaxiter	positive integer	25	Max number of augmentation iterations
Augsolver	umfpack spooles taucs_llt_mf taucs_ldlt luinc taucs_llt gmres fgmres cg amg gmg ssor ssoru sor soru jac lumped	umfpack	Linear system solver for augmented Lagrange components
Augtol	positive real	le-6	Tolerance for augmented Lagrange
Damping	on off	on	Use the damped Newton method
Hnlin	on off	off	Indicator of a highly nonlinear problem
Initstep	non-negative scalar	see below	Initial damping factor

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Кеер	string containing K, N auto	auto	Parameter and iteration-independent quantities
Maxiter	positive integer	25	Maximum number of Newton iterations
Minstep	positive scalar	see below	Minimum damping factor
Ntol	positive scalar	1e-6, 1e-3 (segregated solver groups)	Relative tolerance
Oldcomp	cell array of strings	{}	Old parameter components
Out	<pre>fem sol u plist stop solcompdof Kc Lc Null Nnp ud uscale nullfun symmetric cell array of these strings</pre>	sol	Output variables
Pinitstep	positive real		Initial stepsize for parameter
Plist	real vector		List of parameter values
Pmaxstep	positive real		Maximum stepsize for parameter
Pminstep	positive real		Minimum stepsize for parameter
Pname	string		Parameter name
Porder	0 1	1	Predictor order for parameter stepping
Rstep	real scalar > I	10	Restriction for step size update
Stopcond	string with expression		Stop parameter stepping before expression becomes negative

TABLE I-30: VALID PROPERTY/VALUE PAIRS

In addition, the properties described in the entry femsolver are supported.

The property Augcomp control the augmentation components. If this property is set to a subset of the solution components, then the solution procedure is split into two substeps, that are repeated until a convergence criterion is met or until the

maximum number of iterations is reached. The main components, which are the solution components excluding the augmentation components, are first solved for, while the augmentation components are held fix. After this, the augmentation components are solved for while the main components are held fix. In this second solution step, a linear solution approach is taken. Therefore, with the main components fixed, the augmentation components are assumed to fulfill a linear equation. The property Augtol control the tolerance for the augmentation components and the property Ntol the tolerance for the main components in the convergence criterion for the combined iteration (two substeps each). The convergence criterion is that the relative increment, from one iteration to the next, for the augmented components and the main components must not be larger than their tolerances. The maximum number of combined iterations is controlled by the property Augmaxiter. The linear system solver used for the solution of the augmentation components can be controlled by the property Augsolver. For a more detailed description of this solution procedure, see the "Nonlinear Solver Settings" on page 368 in the COMSOL Multiphysics User's Guide.

Setting Hnlin to on causes the solver to treat the problem as being highly nonlinear. This option can be tried if there is no convergence with Hnlin set to off. Depending on this parameter, certain standard values are selected for the Initstep and Minstep properties. Moreover, certain internal control structures are adapted. Especially, the error control is biased from a more absolute norm towards a relative norm. So this parameter is also useful if a solution with components of highly varying orders of magnitudes are present. In the context of parameter stepping, you can also try this option if the step sizes in the parameter seem to be too small.

Initstep is the initial damping factor for the step length. The default is 1 if Hnlin is off and 1e-4 if Hnlin is on.

Maxiter and Minstep are safeguards against infinite Newton iterations. They bound the number of iterations and the damping factor used in each iteration. Minstep defaults to 1e-4 if Hnlin is off and 1e-8 if Hnlin is on.

The tolerance Ntol gives the criterion for convergence, see "Nonlinear Solver Settings" on page 368 in the *COMSOL Multiphysics User's Guide*.

The property Out explicitly sets output variables and their order. The output variable fem means the FEM structure with the solution object fem.sol added. The solution object sol has a field sol.u, which is the solution vector for the FEM formulation of the PDE problem. The solution vector u is a column vector with one component for each degree of freedom of the discretized problem. If the parameter

variation feature is used, then sol.u is a matrix, and there are additional fields sol.pname and sol.plist. The field sol.pname is the name of the parameter, and sol.plist is a row vector with parameter values for which a solution was computed. The corresponding solution vectors are stored as columns in the matrix sol.u. The output variable Stop is 0 if a complete solution was returned, 1 if a partial solution was returned, and 2 if no solution was returned. For the other outputs, see femlin.

femlin and femnlin can solve a stationary problem for a number of values of a parameter. The name of the parameter is specified with the property Pname, and the values of the parameter is specified with the property Plist. The vector in Plist can be an increasing or decreasing sequence of parameter values. If more than two parameter values are given, then solutions are delivered for these parameter values (though the algorithm may internally compute the solution for intermediate values). If only two parameter values are given, the algorithm also delivers the solutions for the intermediate values determined by the algorithm. The algorithm tries to follow a continuous path of solutions when varying the parameter, and adjusts the step size in the parameter in order achieve this. If the algorithm detects that some sort of singularity or turning point is approached, then the stepsize is reduced, and the algorithm terminates. In this case, if the property Stop is set to on, the solutions for the visited parameter values are delivered.

When going from one parameter value to another, the initial guess at the new parameter value is by default obtained by following the tangent to the solution curve at the old parameter value. If the property Porder is set to 0, then the initial guess is instead taken as the solution for the old parameter value. In very simple cases, Porder = 0 may give better performance than the default Porder = 1.

The property Pinitstep specifies the initial parameter stepsize that will be tried. The algorithm terminates if the Newton method diverges and the parameter step is less than Pminstep. The property Pmaxstep provides an upper bound on the parameter step. If any of the properties Pinitstep, Pminstep, or Pmaxstep are 0 or not given, they are given default values.

For some applications the access to the solution at a previous parameter value is needed. Such an application is for example contact problems with friction in Structural Mechanics. The solution components controlled by the property Oldcomp are treated in a separate linear solution step, or updating step, performed after the solver for the parameter step has finished. These components are subtracted

	from the solution components and are not included in the main parametric solver step. The linear system solver used in the update solver step is UMFPACK.
	For more information on the parameter-stepping feature, see "The Parametric Solver" on page 379 in the COMSOL Multiphysics User's Guide.
	The property Rstep sets a restriction for the damping factor update in the Newton iteration. Each time the damping factor is updated, it is allowed to change at most by a factor Rstep.
	If the property Stop is set to on, the solver gives an output even if the algorithm fails at some point. If the parameter stepping feature is used with more than one parameter value, the output contains the solutions for the parameters that were successfully computed. Otherwise, the output is the nonconverged solution corresponding to the iteration where the failure occurred. If Stop is set to off, the solver terminates with an error if the algorithm fails.
	Use the property Stopcond to make sure the solver stops before a certain event. You provide a scalar expression that is evaluated after each parameter step. The parameter stepping is stopped if the real part of the expression is evaluated to something negative. The corresponding solution, for which the expression is negative is not returned.
	For more information about the nonlinear stationary solver, see "The Stationary Solver" on page 365 in the COMSOL Multiphysics User's Guide.
Diagnostics	If the Newton iteration does not converge, the error messages Maximal number of iterations reached or Damping factor too small are displayed. If during the solution process NaN or Inf elements are encountered in the solution even after reducing the damping factor to the minimum, the error message Inf or NaN repeatedly found in solution is printed. The message Underflow of parameter step length means that the Newton iterations did not converge, even after reducing the parameter step length to the limit given in Pminstep. This probably means that the curve of solutions has a turning point or bifurcation point close to the current parameter value and solution.
Compatibility	The property Variables has been renamed to Const in FEMLAB 2.3.
	The properties Epoint and Tpoint are obsolete from FEMLAB 2.2. Use fem.***.gporder to specify integration order.
	The property/value Jacobian/Lumped has been made obsolete from FEMLAB 1.1.

The properties Toln and Normn have been made obsolete from FEMLAB 1.2. Ntol replaces Toln.

See Also

assemble, asseminit, femlin, femsolver, femstatic, femstruct

 Purpose
 Description of properties common to all plot functions.

Description

Valid property/value pairs:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Axis	numeric vector		Axis limits
Axisequal	on off	on	Axis equal
Axislabel	cell array of strings		X-, Y- and Z-axis labels
Axisvisible	on off	on	Axis visible
Camlight	on off	off	Light at camera position
Campos	I-by-3 numeric vector		Position of camera
Camprojection	orthographic perspective	perspective	Projection
Camtarget	I-by-3 numeric vector		Camera aiming point
Camup	I-by-3 numeric vector		Rotation of the camera
Cam∨a	numeric between 0 and 180	90	Field of view in degrees
Grid	on off	off	Grid visible
Lightmodel	flat gouraud phong none	phong	Lighting algorithm
Lightreflection	dull shiny metal default I-by-3,4or5 numeric vector	default	Reflectance of surfaces
Parent	axes handle		Handle to axes object
Renderer	auto painters zbuffer opengl	auto	Rendering algorithm
Scenelight	on off	off	Create scene light
Scenelightpos	I-by-3 numeric vector		Location of scene light object
Title	string	empty	Plot title
Titlecolor	color	k	Title color

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Transparency	number between 0 and 1	1	Transparency (has only effect using OpenGL)
View	2, 3, or numeric pair	2 or 3	3D view point

Purpose	Create a Simulink structure.
Syntax	<pre>sct = femsim(fem,)</pre>
Description	<pre>sct = femsim(fem,) creates a Simulink structure sct for the FEM structure fem.</pre>
	To use the exported Simulink structure in Simulink, open the Blocksets &

To use the exported Simulink structure in Simulink, open the Blocksets & Toolboxes library in Simulink, double-click on the COMSOL Multiphysics icon, and drag the COMSOL Multiphysics Subsystem block to your Simulink model. Double click on your copy of the block, and enter the name of your Simulink Structure. This sets up the input and the output ports of the block.

The function femsim accepts the following property/value pairs:

TABLE I-31: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Input	cell array of strings	fem.const	Input variable names
Кеер	string containing K, L, M, N, D, E auto	auto	Time-independent quantities
Nonlin	off on auto	auto	Use nonlinear stationary solver
Outnames	cell array of strings	y1,y2,	Output data names
Output	cell array	{}	Output data (see below)
Redcomp	cell array of strings		Degrees of freedom in model reduction of wave equations
Redmodes	integer	10	Number of eigenmodes in reduction
Redstatic	off on	on	Use static modes in reduction
Reduction	off on	off	Model reduction
State	off on	off	Use linearized state-space model
Static	off on	off	Use static solver
Т	scalar	0	Time for evaluation of linearized model

In addition, the common solver properties described in the entry femsolver are supported, with modifications described below.

The names of the input variables are given in the property Input. The default is all names in fem.const.

The output values are determined by the entries in the Output cell array. Each entry in this cell array should be a cell array of the following form:

```
{'expr' xx pvlist}
```

where 'expr' is an expression, xx is a column vector containing global coordinates, and pvlist is a (possibly empty) list of property/value pairs (see postinterp). The function postinterp is used to evaluate the output value as

```
postinterp('expr', xx, pvlist)
```

The names of the output ports of the COMSOL Multiphysics Subsystem block are given in the property Outnames. The default names are y1, y2, etc.

The COMSOL Multiphysics Subsystem block can act in four different modes, chosen by the properties State and Static:

General Dynamic Export (State=off, Static=off)

Export a dynamic model, where the COMSOL Multiphysics degrees of freedom are part of the Simulink state vector. The COMSOL Multiphysics solver is called several times for each time step to compute the time derivative of the state vector, with inputs from Simulink. Only linear, time-independent constraints and the eliminate constraint handling method is supported. The Itol property is supported, see femstatic. The Const property is not supported.

General Static Export (State=off, Static=on)

Export a static model, where the COMSOL Multiphysics degrees of freedom are not part of the Simulink state vector. To compute the outputs of the COMSOL Multiphysics Subsystem block, the COMSOL Multiphysics linear or nonlinear stationary solver is called for each time step, with inputs from Simulink. If Nonlin=off, the linear solver is used and the property Itol is supported, see femstatic. If Nonlin=on, the nonlinear solver is used and the properties Hnlin, Initstep, Maxiter, Minstep, Ntol, and Rstep are supported, see femnlin.

Linearized Dynamic Export (State=on, Static=off)

Export a dynamic linearized model. The model is linearized about an equilibrium solution, and the matrices in the state-space form are computed. The COMSOL Multiphysics degrees of freedom are part of the Simulink state vector. At each time step, the matrices in the state-space form are used to compute the time derivative of the state vector, instead of calling the COMSOL Multiphysics solver. Only linear,

time-independent constraints and the eliminate constraint handling method is supported. The Const property is not supported.

The linearization point should be an equilibrium point (stationary solution) and is controlled by the property U, see femsolver. The inputs and the outputs are deviations from the equilibrium values.

Model reduction can be used to approximate the linearized model with a model that has fewer degrees of freedom, by using Reduction=on. A number of eigenmodes (given by the property Redmodes) and static modes (if Restatic=on) will then be computed, and the linearized model will be projected onto the corresponding subspace. The properties Etol, Itol, Krylovdim, and Shift of the eigenvalue solver are supported, see femeig.

When using model reduction on wave equation models that have been rewritten as a system of first-order equations (wave extension), the algorithm needs to know the names of the original (non time derivative) solution components. The names of the non-time derivative solution components should be specified using the property Redcomp. Since COMSOL Multiphysics 3.2, wave equations are usually formulated without wave extension; then the Redcomp property should not be used.

Linearized Static Export (State=on, Static=on)

Export a static linearized model. The model is linearized about an equilibrium solution, and a transfer matrix is computed. The COMSOL Multiphysics degrees of freedom are not part of the Simulink state vector. To compute the outputs of the COMSOL Multiphysics Subsystem block, the transfer matrix is used.

The linearization point should be an equilibrium point (stationary solution) and is controlled by the property U, see femsolver. The inputs and the outputs are deviations from the equilibrium values.

Example	Heat equation with heat source Q as input.
	<pre>fem.geom = solid1([0 1]); fem.mesh = meshinit(fem); fem.shape = 2; fem.equ.da = 1; fem.equ.c = 1; fem.equ.f = 'Q'; fem.xmesh = meshextend(fem); % Temperature u at x = 0.5 is output sct = femsim(fem, 'input',{'Q'}, 'outnames',{'Temp'}, 'output',{{'u' 0.5}});</pre>
Compatibility	For backward compatibility, the Input property can also be a vector of indices into fem. const

Most of the FEMLAB 2.3 data types in the Output cell array are still supported:

TABLE I-32: FEMLAB 2.3 OUTPUT DATA TYPES

ENTRY IN OUTPUT CELL ARRAY	INTERPRETATION		
Cell array {N iName}	The solution component fem.dim{iName} at mesh vertex number N in fem.mesh or fem.fem{g}.mesh, where g is the geometry given in the Geomnum property (default is 1)		
Integer N	Shortcut for $\{N \mid 1\}$. That is, solution component fem.dim $\{1\}$ at mesh vertex number N		
Struct lfun	Linear functional, is no longer supported. Use an integration coupling variable instead		
Cell array {N 'expr'}	Value of expression 'expr' at mesh vertex number N in the geometry given in the Geomnum property		
String func	Value of function func(fem,u,t,indata), is no longer supported		

The properties Mass and Timescale are no longer supported.

A Simulink structure with State=off can no longer be saved to file using the commands save or flsave.

See Also femsolver, femlin, femnlin, femeig, femtime, femstate

Purpose	Create a solution object.		
Syntax	<pre>fem.sol = femsol(u) fem.sol = femsol(u,'tlist',tlist) fem.sol = femsol({u ut},'tlist',tlist) fem.sol = femsol(u,'plist',plist,'pname',plist) fem.sol = femsol(u,'lambda',lambda)</pre>		
Description	<pre>fem.sol = femsol(u) creates a stationary solution object from a column vector u. The length of u must equal number of degrees of freedoms in the extended mesh object, fem.xmesh, (see flngdof).</pre>		
	<pre>fem.sol = femsol(u,) stores a matrix corresponding to a time-dependent, parametric, or eigenvalue solution in the solution object. The number of rows must equal the number of degrees of freedoms in the extended mesh object, fem.xmesh, (see flngdof) and the number of columns of u must equal the number of time steps, parameter values, or eigenvalues, respectively (see solsize).</pre>		
	<pre>fem.sol = femsol({u ut},) creates a time-dependent solution object containing also the first time derivative. The matrix u is the usual solution matrix, and ut is its time derivative.</pre>		
	<pre>fem.sol = femsol(u, 'mcase', mcase) sets the mesh case of the created solution object to mcase. The default mesh case is 0. Access Functions The following access functions lets you fetch properties from the solution object.</pre>		
	TABLE I-33: VALID PROPERTY/VALUE PAI		
	PROPERTY	VALUES	
	fem.sol.u Solution vector or matrix fem.sol.ut Matrix containing time derivative (time-dependent solutions)		
	fem.sol.tlist	List of time steps (time-dependent solutions)	
	fem.sol.plist List of parameter values (parametric solutions)		
	fem.sol.pname Parameter name (parametric solutions)		
	fem.sol.lambda	List of eigenvalues (eigenvalue solutions)	
	fem.sol.mcase Mesh case		

If the time-derivatives have not been stored in the femsol object, fem.sol.ut is computed as the slope of the linear interpolation between the time steps. To store the time derivatives in the solution, use the Outcomp property, see femsolver.

Example	Create a solution object:
	<pre>fem.geom = rect2; fem.mesh = meshinit(fem); fem.shape = 2; fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = 1; fem.xmesh = meshextend(fem); fem.sol = femtime(fem,'tlist',0:0.1:1)</pre>
	Fetch the solution vector and the list of time steps.
	u = fem.sol.u; tlist = fem.sol.tlist;
	Multiply the solution by 2 and recreate a solution object.
	<pre>fem.sol = femsol(2*u,'tlist',tlist);</pre>
	Postprocess the solution.
	postplot(fem,'tridata','u')
Compatibility	In FEMLAB 2.3 the solution was represented with a MATLAB structure. The solution object does not allow exactly the same type access as the structure. The solution object has been designed to be compatible with the MATLAB structure.
See also	asseminit, femeig, femlin, femnlin, femtime

Purpose Description of properties common to all solvers.

Description

In addition to the properties in the table below, the solvers accept properties controlling the linear system solvers, see the sections starting with "Linear System Solvers" on page 160.

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Assemtol	scalar	1e-12	Assembly tolerance
Blocksize	positive integer	5000	Assembly block size
Complexfun	off on	off	Use complex-valued functions with real input
Conjugate	off on	off	Use complex conjugate Hermitian transpose
Const	cell array of alternating strings and values, or a structure		Definition of constants
Constr	auto ideal nonideal	auto	Constraint force Jacobian
Init	solution object numeric vector scalar		Initial value
Кеер	string containing K, L, M, N, D, E auto	auto	Manual control of reassembly
Linsolver	umfpack spooles pardiso taucs_llt_mf taucs_ldlt luinc taucs_llt gmres fgmres cg amg gmg ssor ssoru sor soru jac vanka	umfpack	Linear system solver
Matherr	off on	on	Error for undefined operations
Mcase	non-negative integer (or vector for GMG)	mesh case with largest number of DOFs	Mesh case to solve for

TABLE I-34: COMMON SOLVER PROPERTY/VALUE PAIRS

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Method	eliminate elimlagr lagrange spring	eliminate	Constraint handling method
Nullfun	flnullorth flspnull auto	auto	Null space function
Outcomp	cell array of strings		Solution components to store in output
Report	on off	on	Show progress dialog box
Rowscale	on off	on	Equilibrate rows
Solcomp	cell array of strings		Solution components to solve for
Solfile	on off	off	Store solution on file
Solfileblock	positive scalar	16	Max size of solution block (MB)
Solfilename	string		Name of solution file
Stop	on off	on	Deliver partial solution when failing
Symmetric	on off auto	auto	Symmetric matrices
Symmtol	non-negative scalar	1e-10	Symmetry detection tolerance
U	solution object numeric vector scalar		Values of variables not solved for and linearization point
Uscale	auto init none cell array solution vector	auto	Scaling of variables

TABLE I-34: COMMON SOLVER PROPERTY/VALUE PAIRS

In addition to the constants in fem.const, you can define constants using the Const property, see assemble.

The parameter Constr controls how the constraint force Jacobian is computed. For auto and nonideal, the constraint force Jacobian matrix N_F is assembled independently of the constraint Jacobian matrix N. When auto is selected, a comparison between N_F and N^T is performed. If these matrices are found equal (up to a tolerance), then N_F is cleared and $N_F = N^T$. For ideal, only the constraint Jacobian matrix N is assembled and $N_F = N^T$.

If the solver fails to find a complete solution, it returns a partial solution if the property **Stop** is **on** (this is the default).

You can use the property Symmetric to tell the solver that the model is symmetric or you can use the automatic feature to find out. The symmetry detection tolerance Symmtol is used for this automatic feature (see "Which Problems are Symmetric?" on page 397 in the COMSOL Multiphysics User's Guide). If the model is Hermitian, you should set both the Symmetric and Conjugate properties to on.

PROPERTIES CORRESPONDING TO THE SOLVER MANAGER

The properties Init, U, Solcomp, and Outcomp correspond to settings in the Solver Manager (see "The Solver Manager and Solver Scripting" on page 401 in the *COMSOL Multiphysics User's Guide*).

The property Init determines the initial value for the solution components you solve for. For possible syntaxes, see asseminit. If you omit this property, the solver computes the initial value by evaluating the initial value expressions in fem.equ.init, fem.equ.dinit, fem.bnd.init, fem.bnd.dinit, etc. If any of these expressions depend on a solution component, the value 0 is used for that solution component.

The property U determines the value of solution component you do not solve for and the linearization point. A scalar value is equivalent to a solution vector containing that value in all its components.

The property Solcomp is a cell array containing the names of the degrees of freedom to solve for. The default is all degrees of freedom.

The property Outcomp is a cell array containing the names of the degrees of freedom to store in the output solution object. If the solution is time dependent, the property Outcomp can contain also the time derivatives of the DOF names. The default is all degrees of freedom (excluding the time derivatives).

THE PROGRESS WINDOW

By default, a progress window appears when a solver is called. This is similar to the progress window that appears in the COMSOL Multiphysics user interface (see "Solution Progress" on page 411 in the *COMSOL Multiphysics User's Guide*). The progress window gives you the possibility to cancel or stop the solver. Also, when running in MATLAB, it gives you the possibility to examine the convergence in a plot. Using the properties specified under Probe Plot Parameters below, in addition, the progress window gives you the possibility to plot values of certain quantities

during the solution process for the time dependent and parametric solver. If you do not want the progress window, use the Report property or the flreport command (see flreport).

PROBE PLOT PARAMETERS

Properties to the time-dependent solver and the parametric solver:

PROPERTY NAME	PROPERTY VALUE	DESCRIPTION
plotglobal	String or cell array of strings	Global plot expressions
plotglobalpar	Cell array or cell array of cell arrays	Property/values to postglobaleval
plotint	String or cell array of strings	Integration plot expressions
plotintpar	Cell array or cell array of cell arrays	Property/values to postint
plotinterp	String or cell array of strings	Probe plot expressions
plotinterppar	Cell array or cell array of cell arrays	Property/values to postinterp

The properties plotglobalpar, plotinpar, and plotinterppar must be a cell array containing property/values to the corresponding evaluating function postglobaleval, postint, and postinterp, respectively. In addition, the property title can be specified. Also, plotinterppar must contain the property probecoord with the value being an sdim-by-n coordinate matrix. If, for example, plotint is a cell array of expressions to plot, plotintpar can either be a cell array of property/values, or, if different property/values are to be specified for the different expressions, a cell array of cell arrays of property/values.

An example of a femstatic call for a parametric problem specifying one global expression, one interpolation expression, and two integration expressions.

```
fem.sol = femstatic(fem, ...
    'solcomp',{}, ...
    'outcomp',{}, ...
    'plotglobal',{'w1','w2'}, ...
    'plotglobalpar',{'phase',pi,'title','Global'}, ...
    'plotinterp','u', ...
    'plotinterppar',{'probecoord',[0.1; 0.2]}, ...
    'plotint',{'u','ux','uy'}, ...
    'plotintpar',{{'phase',0, 'edim',0,'d1',4}, ...
```

```
{'phase',pi,'edim',1,'intorder',5},...
{'phase',pi,'edim',2,'dl',[3,6,7]}});
```

ADVANCED SOLVER PARAMETERS

The section "Advanced Solver Settings" on page 492 describes the features corresponding to the properties Blocksize, Complexfun, Conjugate, Keep, Method, Nullfun, Rowscale, Solfile, and Uscale.

The property Assemtol affects the assembled matrices, see assemble for details.

By default, COMSOL Multiphysics gives an error message if the solver encounters an undefined mathematical operation when solving the model, for instance 0/0 or log(0). If you instead want the solver to proceed, put the property Matherr=off. Then 0/0=NaN (not a number) and log(0)=-Inf.

The Symmetric and Conjugate properties correspond to the Solver Parameters dialog box settings Matrix symmetry and Use Hermitian transpose in constraint matrix and in symmetry detection according to the following table:

MATRIX SYMMETRY	USE HERMITIAN TRANSPOSE	SYMMETRIC	CONJUGATE
Automatic	cleared	auto	off
Automatic	selected	auto	on
Nonsymmetric	cleared	off	off
Nonsymmetric	selected	off	on
Symmetric	n.a.	on	off
Hermitian	n.a.	on	on

The property Keep corresponds to the manual control of reassembly feature. Its value can be a string containing the letters D, E, K, L, M, N, or the string auto. These letters have the following meaning: E=constant mass, D=constant damping, K=constant Jacobian, L=constant load, M=constant constraint, N=constant constraint Jacobian (see "Manual Control of Reassembly" on page 496).

For the Nullfun property, flnullorth is the orthonormal null-space function, and flspnull is the sparse null-space function.

If Solfile=on, the solution is stored on a temporary file. The Solfilename property can be used to give the full path name of the file. A part of the solution is stored in memory in a few *blocks* (usually 1–5 blocks reside in memory). The maximum block size (in megabytes) can be controlled with the property Solfileblock.

The property Uscale determines a scaling of the degrees of freedom that is applied in order to get a more well-conditioned system; see "Scaling of Variables and Equations" on page 497. The possible values are:

TABLE I-35: VALUES FOR THE PROPERTY USCALE

VALUE	MEANING
auto	The scaling is automatically determined
init	The scaling is determined from the initial value. Use this if the sizes of the components of the initial value give a good estimate of the order of magnitude of the solution
none	No scaling is applied
cell array	A cell array with alternating degree of freedom names and positive numbers. The numbers specify the expected magnitude of the corresponding degree of freedom
solution vector	A numeric vector with positive components that specify the expected magnitude of the solution

The default is auto, except when using one of the syntaxes

```
[Ke,Le,Null,ud] = femstatic(fem,...)
[Kl,Ll,Nnp] = femstatic(fem,...)
[Ks,Ls] = femstatic(fem,...)
fem.sol = femstatic('In',{'K' K 'L' L 'M' M 'N' N},...)
```

which assume that the property Out is not given in the first three cases. In these cases the default is none. The resulting vector of scale factors is contained in the output variable uscale. The scaling of the degrees of freedom is applied symmetrically to the Jacobian matrix, that is, both the rows and columns are scaled.

LINEAR SYSTEM SOLVERS

The properties Linsolver, Prefun, Presmooth, Postsmooth, and Csolver select the linear system solver, preconditioner, presmoother, postsmoother, and coarse solver, according to the following table.

NAME	ALGORITHM
umfpack	UMFPACK direct solver
spooles	SPOOLES direct solver
pardiso	PARDISO direct solver
taucs_llt_mf	TAUCS direct Cholesky solver
taucs_ldlt	TAUCS direct LDLT solver (not recommended)

TABLE 1-36: LINEAR SYSTEM SOLVERS/PRECONDITIONERS/SMOOTHERS

NAME	ALGORITHM
luinc	Incomplete LU preconditioner/smoother
taucs_llt	TAUCS Incomplete Cholesky preconditioner
gmres	GMRES iterative solver
fgmres	FGMRES iterative solver
cg	Conjugate Gradients iterative solver
amg	Algebraic Multigrid iterative solver/preconditioner
gmg	Geometric Multigrid iterative solver/preconditioner
ssor	SSOR preconditioner/smoother
ssoru	SSORU preconditioner/smoother
sor	SOR preconditioner/smoother
soru	SORU preconditioner/smoother
jac	Jacobi (diagonal scaling) preconditioner/smoother
ssorvec	SSOR vector preconditioner/smoother
sorvec	SOR vector preconditioner/smoother
soruvec	SORU vector preconditioner/smoother
ssorgauge	SSOR gauge preconditioner/smoother
sorgauge	SOR gauge preconditioner/smoother
sorugauge	SORU gauge preconditioner/smoother
vanka	Vanka-type preconditioner/smoother

TABLE 1-36: LINEAR SYSTEM SOLVERS/PRECONDITIONERS/SMOOTHERS

For a description of these solvers, see the section "The Linear System Solvers" on page 392 in the *COMSOL Multiphysics User's Guide*.

DIRECT LINEAR SYSTEM SOLVER PROPERTIES

The umfpack, spooles, pardiso, taucs_llt_mf, luinc, and taucs_llt direct linear solvers/preconditioners/smoothers have the following properties.

TABLE I-37: DIRECT LINEAR SOLVERS PROPERTY/VALUE PAIRS

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Droptol	scalar between 0 and 1	0.01 when used as pre- conditioner or smoother, 0 when used as solver	Drop tolerance (luinc, taucs_llt, umfpack, spooles)
Errorchk	on off auto	on	Check error estimate (pardiso)
Errorchkd	on off	off	Check error estimate (umfpack, spooles)
Fillratio	non-negative scalar	2	Column fill-ratio (luinc)
Maxdepth	positive integer	10000	Maximum recursion depth (taucs_llt_mf)
Modified	on off	off	Modified incomplete Cholesky (taucs_1lt)
Pivotperturb	scalar between 0 and 1	1e-8	Pivot perturbation threshold (pardiso)
Pivotrefines	non-negative integer	0	Number of forced iterative refinements
Preorder	mmd nd ms both	mmd	Preordering algorithm (spooles)
Pardreorder	mmd nd	nd	Preordering algorithm (pardiso)
Pardrreorder	on off	on	Row preordering algorithm (pardiso)
Respectpattern	on off	on	Do not drop original nonzeros (luinc)
Rhob	scalar > I	400	Factor in linear error estimate (pardiso)

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Thresh	scalar between 0 and 1	0.1 (umfpack, spooles), 1.0 (luinc)	Pivot threshold (umfpack, spooles, luinc)
Umfalloc	non-negative scalar	0.7	Memory allocation factor (umfpack)

TABLE I-37: DIRECT LINEAR SOLVERS PROPERTY/VALUE PAIRS

ITERATIVE LINEAR SYSTEM SOLVER PROPERTIES

The iterative linear solvers/preconditioners/smoothers luinc, gmres, fgmres, cg, amg, gmg, ssor, ssoru, sor, soru, jac, ssorvec, sorvec, soruvec, ssorgauge, sorgauge, sorugauge, vanka have the following properties.

TABLE 1-38: ITERATIVE LINEAR SOLVERS PROPERTY/VALUE PAIRS

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Iluiter	non-negative integer	1	Fixed number of iterations (when used as preconditioner, smoother, or coarse solver) (luinc)
Iter	non-negative integer	2	Fixed number of iterations (when used as preconditioner, smoother, or coarse solver) (all except luinc)
Itol	positive real	1e-6, 0.1 (coarse solver)	Relative tolerance (note that when used as preconditioner or smoother a fixed number of iterations is default)
Itrestart	positive integer	50	Number of iterations before restart (gmres, fgmres)
Maxlinit	positive integer	10000, 500 (coarse solver)	Maximum number of linear iterations (when used with a tolerance)

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Prefun	<pre>luinc taucs_llt umfpack spooles gmres fgmres cg amg gmg ssor ssoru sor soru jac ssorvec sorvec soruvec ssorgauge sorgauge sorugauge vanka none</pre>	luinc	Preconditioner (gmres, fgmres, cg)
Prefuntype	left right	left	Left or right preconditioning (gmres, cg)
Prepar	cell array of property/ value pairs or structure		Preconditioner properties (gmres, fgmres, cg)
Relax	scalar between 0 and 2	1	Relaxation factor (Jacobi, SOR-based algorithms, incomplete LU, and Vanka)
Rhob	scalar >= 1	400, 1 (coarse solver)	Factor in linear error estimate (when used with a tolerance)
Seconditer	nonnegative integer	1	Number of secondary iterations (SOR vector and SOR gauge algorithms), number of SSOR updates (vanka)
Sorblocked	on off	off	Blocked SOR method
Sorvecdof	cell array of strings		Vector element variables (SOR vector and SOR gauge algorithms)
Vankablocked	on off	off	Blocked Vanka method
Vankarelax	scalar between 0 and 2	0.8	Relaxation factor for Vanka update
Vankarestart	positive integer	100	GMRES restart value (vanka)

TABLE I-38: ITERATIVE LINEAR SOLVERS PROPERTY/VALUE PAIRS

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Vankasolv	gmres direct	gmres	Local block solver (vanka)
Vankatol	positive scalar	0.02	GMRES tolerance (vanka)
Vankavars	cell array of strings	{}	Lagrange multiplie variables (vanka)

MULTIGRID SOLVER PROPERTIES

The multigrid solvers/preconditioners amg and gmg accept the following properties in addition to those in table 1-38.

TABLE I-39: MULTIGRID SOLVERS PROPERTY/VALUE PAIRS

PROPERTY	VALUES	DEFAULT	DESCRIPTION
amgauto	integer between I and IO	3	Quality of multigrid hierarchy (amg)
csolver	umfpack spooles pardiso taucs_llt_mf taucs_ldlt luinc taucs_llt gmres fgmres cg amg ssor ssoru sor soru jac ssorvec sorvec soruvec ssorgauge sorgauge sorugauge vanka	umfpack	Coarse solver
csolverpar	cell array with property/ value pairs	{}	Coarse solver properties
maxcoarsedof	positive integer	5000	Maximum number of DOFs at coarsest level (amg)
meshscale	vector of positive numbers	2	Mesh scale factor (gmg)
mgassem	on off numeric vector	on	Assembly on coarse levels (gmg)

PROPERTY	VALUES	DEFAULT	DESCRIPTION
mgauto	off explicit meshscale shape both meshrefine		Method for mesh case generation (gmg)
mgcycle	v w f	v	Cycle type
mggeom	vector of positive integers	all	Geometry numbers for multigrid hierarchy (gmg)
mgkeep	on off	off	Keep generated mesh cases (gmg)
mglevels	integer>1	6 (amg), 2 (gmg)	Maximum number of multigrid levels
postsmooth	ssor ssoru sor soru jac ssorvec sorvec soruvec ssorgauge sorgauge sorugauge luinc gmres fgmres cg amg vanka	soru	Postsmoother
postsmoothpar	cell array with property/ value pairs	{}	Postsmoother properties
presmooth	ssor ssoru sor soru jac ssorvec sorvec soruvec ssorgauge sorgauge sorugauge luinc gmres fgmres cg amg vanka	sor	Presmoother
presmoothpar	cell array with property/ value pairs	{}	Presmoother properties
rmethod	regular longest	regular	Mesh refinement method (gmg)
shapechg	vector of integers	- 1	Change in shape function orders (gmg)

TABLE I-39: MULTIGRID SOLVERS PROPERTY/VALUE PAIRS

For the Geometric multigrid solver/preconditioners, the construction of the multigrid hierarchy is controlled by the properties Mgauto, Mcase, Mglevels, Meshscale, Shapechg, and Mgassem:

- If Mgauto=both, shape, or meshscale, then the multigrid hierarchy is automatically constructed starting from the mesh case given in the property Mcase. This process is described in the section "Constructing a Multigrid Hierarchy" on page 521, where the methods are called **Coarse mesh and lower** order (both), Lower element order first (shape), and **Coarse mesh** (meshscale). The mesh coarsening factor is given in the scalar Meshscale, the shape function order change amount is given in the scalar Shapechg, and the number of multigrid levels (including the finest level) is given in the property Mglevels (default 2).
- If Mgauto=explicit, then the multigrid hierarchy is automatically constructed starting from the mesh case given in the property Mcase. The properties Meshscale and Shapechg should be vectors of the same length (however if one is scalar, it is expanded to the same length as the other). A number of coarse levels are constructed, where level i has a mesh that is coarsened with the factor Meshscale(i), and shape functions orders incremented with Shapechg(i) relative to mesh case Mcase. Shapechg(i) should be negative or zero.
- If Mgauto=meshrefine, then the multigrid hierarchy is automatically constructed by refining the mesh in mesh case Mcase repeatedly. The number of multigrid levels (including the original, coarsest level) is given in the property Mglevels (default 2). The refinement method can be specified using the property Rmethod, see meshrefine.
- If Mgauto=off, then only existing mesh cases are used in the hierarchy. If the property Mcase is a scalar, then all mesh cases that have fewer degrees of freedom than the mesh case Mcase are used as coarse levels. If Mcase is a vector with more than one component, the mesh cases in that vector are used. However, if the property Mglevels is given, no more than Mglevels levels are used. The solver sorts the list of mesh cases according to decreasing number of DOFs, and the solution is delivered for the mesh case with the largest number of DOFs. This corresponds to the Manual option in the COMSOL Multiphysics user interface.

The default for Mgauto is as follows: If the FEM structure has several mesh cases, then Mgauto=off, otherwise Mgauto is the same as the default in meshcaseadd.

The construction of coarse level matrices is controlled by the property Mgassem. If Mgassem is a vector, Mgassem(i) should be a 0 or 1. Mgassem(i)=1 means that matrices should be assembled in mesh case Mcase(i), rather than being projected from the next finer level. The length of Mgassem should be (at least) the number of mesh cases used, including the finest level. The value of Mgassem(i) for the finest

level i is ignored, because matrices are always assembled on the finest level. A scalar Mgassem applies to all coarse mesh cases.

When an iterative solver is used as preconditioner, smoother, or coarse solver you can choose whether to solve using a tolerance or to perform a fixed number of iterations. When used as a coarse solver the default is to solve using a tolerance. When used as a preconditioner or smoother the default is to perform a fixed number of iterations. If both properties Itol and Iter (or Iluiter for luinc) are given, the program will solve using a tolerance.

Four Examples How to Construct the Geometric Multigrid Hierarchy Assume that fem only contains the mesh case 0, and no xmesh.

Alternative 1:

fem.xmesh = meshextend(fem); fem.sol = femstatic(fem, 'linsolver', 'gmg');

This alternative uses a temporary hierarchy that is constructed by the solver. Since the solver also constructs a temporary xmesh, this alternative wastes some memory.

Alternative 2:

fem = femstatic(fem, 'linsolver', 'gmg', 'out', 'fem');

Here, the solver uses a temporary hierarchy, but there is only one xmesh. If another such solver call is made, fem.xmesh should first be deleted to save some memory.

Alternative 3:

```
fem =
femstatic(fem,'linsolver','gmg','mgkeep','on','out','fem');
```

Now the generated hierarchy is kept, which means that you can reuse it in a subsequent call:

fem.sol = femstatic(fem, 'linsolver', 'gmg');

Alternative 4:

```
fem = meshcaseadd(fem);
fem.xmesh = meshextend(fem);
fem.sol = femstatic(fem,'linsolver','gmg');
```

The meshcaseadd call adds mesh cases to the FEM structure. These form the multigrid hierarchy in the solver.

COMSOL Multiphysics 3.2: The default of the Conjugate property has been changed to off.

Compatibility

The following FEMLAB 2.3 general solver properties are obsolete in FEMLAB 3.0:

TABLE I-40: OBSOLETE PROPERTY/VALUE PAIRS

PROPERTY	VALUES	IMPLICATION
Initmethod	weak pointwise local dof	No longer supported
Itsolv	gbit gmres tfqmr	Use Linsolver property.
Jacobian	lumped numeric	No longer supported
Linsolver	matlab superlu	Uses default direct solver
Maxlinit	vector with 2 components	Ignores second component and warns
Nullfun	name of user-defined function	No longer supported
Sd		Use streamline diffusion, see the chapter "Stabilization Techniques" on page 433 in the Modeling Guide.

The default of the property Stop has been changed to On.

Purpose	Compute state-space form of a time-dependent PDE problem.
Syntax	<pre>[A,B,C,D] = femstate(fem,) [M,MA,MB,C,D] = femstate(fem,) [M,MA,MB,C,D,Null,ud,x0] = femstate(fem,) state = femstate(fem,)</pre>
Description	[A,B,C,D] = femstate(fem,) calculates the linearized state-space form of the dynamic PDE model fem on the format

$$\begin{cases} \frac{dx}{dt} = Ax + Bu\\ y = Cx + Du \end{cases}$$

where x are the state variables, u are the input variables, and y are the output variables.

[M,MA,MB,C,D] = femstate(fem,...) calculates the state-space form on the format

$$M\frac{dx}{dt} = MAx + MBu$$
$$y = Cx + Du$$

The matrices *M* and *MA* are usually much sparser than the matrix *A*.

[M,MA,MB,C,D,Null,ud,x0] = femstate(fem,...) also returns the null-space matrix Null, the constraint contribution ud, and the initial state x0. The full solution vector U can be obtained from the state variables by U = Null*x+u0, where u0 is the linearization point.

state = femstate(fem,...) returns the structure state containing the fields M, MA, MB, C, D, Null, and x0.

state = femstate(fem, 'out', 'statenom',...) returns the structure state
containing the fields A, B, C, D, Null, and x0.

s = femstate(fem, 'out', 'ss',...) returns the Control System Toolbox state-space object s = ss(A,B,C,D).

The output from femstate is intended for use from Simulink or the Control System Toolbox. The function femstate with the output state is equivalent to femsim with the property State=on. In addition to the properties of femsim, femstate accepts the following property/value pairs:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Out	A B C D M MA MB Null ud x0 state statenom ss cell array of these strings	[A,B,C,D] [M,MA,MB,C,D] [M,MA,MB,C,D, Null,ud,x0] state	Output variables
Sparse	off on	off	Sparse matrices

TABLE I-41: VALID PROPERTY/VALUE PAIRS

The property Sparse controls whether the matrices A, B, C, D, M, MA, MB, and Null are stored in the sparse format. See femsim for a description of the other properties.

The matrices M and MA are produced by the same algorithms that do the finite-element assembly and constraint elimination in COMSOL Multiphysics. M and MA are the same as the matrices D_c (eliminated mass matrix) and $-K_c$ (K_c is the eliminated stiffness matrix), respectively, from a call to femlin (see femlin on page 129). The matrices are produced from an exact residual vector Jacobian calculation (that is, differentiation of the residual vector with respect to the degrees of freedoms x) plus an algebraic elimination of the constraints. The matrix C is produced in a similar way; that is, the exact output vector Jacobian matrix plus constraint elimination.

The matrices MB and D are produced by a numerical differentiation of the residual and output vectors, respectively, with respect to the input parameters (the algorithm systematically perturbs the input parameters by multiplying them by a factor $(1+10^{-8})$).

When exporting the *A* and *B* matrices, *A* and *B* are computed by $A = M \setminus MA$ and $B = M \setminus MB$ (that is, from an LU factorization of *M* using the UMFPACK solver.

Compatibility See the femsim entry.

femsim

Purpose	Solve stationa	ry PDE problem with a	nonlinear or l	inear solver.	
Syntax	<pre>fem.sol = femstatic(fem,) fem = femstatic(fem,'Out',{'fem'},)</pre>				
Description	fem.sol = fe or nonlinear s		a stationary PD	DE problem using either a linea	
	stationary PD		alues of the pa	st',list,) solves a rameter P. The values of the	
	The PDE pro femstruct fo		ossibly extende	ed) FEM Structure fem. See	
		femstatic accepts the PROPERTY/VALUE PAIRS	following proj	perty/value pairs:	
	PROPERTY	VALUE	DEFAULT	DESCRIPTION	
	Llimitdof	cell array of strings		Lower limit dofs	
	Llimitval	vector	0	Lower limit vals	
	Maxsegiter	positiv integer	100	Maximum number of segregated iterations	
	Nonlin	off on auto	auto	Use the nonlinear solver	
	Out	<pre>fem sol u plist stop solcompdof Kc Lc Null Nnp ud uscale nullfun symmetric nonlin cell array of these strings</pre>	sol	Output variables	
	Segcomp	cell array of strings		Segregated group components	
	Segdamp	real vector	0.5	Segregated substep damping factors	
	Seggrps	cell array of cell array		Segregated group properties	
	Segorder	integer vector		Segregated substep group numbers	
	Subiter	integer vector	1	Segregated substep	

iterations

This solver uses the nonlinear solver described in femnlin if nonlin is on, and it uses the linear solver described in femlin if nonlin is off. If nonlin is set to auto an analysis is performed to automatically detect if the problem can be solved with the linear solver.

In addition to the properties listed above, also the properties for femlin or femnlin are supported, depending on which solver is used. For example, the linear solver does not support the properties Augcomp, Augsolver, Augtol, Augmaxiter, Porder, and Ntol. Similarly, the nonlinear solver does not support the properties In. Furthermore, the properties described in the entry femsolver are supported.

The automatic nonlinear/linear detection works in the following way. The linear solver is called if the residual Jacobian matrix (the stiffness matrix, K) and the constraint Jacobian matrix (the constraint matrix, N) are both found not solution dependent and if these matrices are detected as complete. In all other situations the nonlinear solver is used. The analysis is performed by a symbolic analysis of the expressions contributing to these matrices. Complete here means that in the residual and constraint vectors, only expressions where found for which COMSOL Multiphysics will compute the correct Jacobian contribution.

Therefore, if you want to solve a linearized (nonlinear) problem, you must select nonlin to off. Furthermore, there are variables for which COMSOL Multiphysics is conservative and will flag these, and their Jacobian contribution, as solution dependent even though they not always are. For these situations, the nonlinear solver will be used even though the linear solver could be used. This should only result in some extra computational effort, and should not influence the result. The opposite situation however, where the linear solver is used for a nonlinear problem is more dangerous. So, select nonlin to off with great care.

If the property Augcomp is given then the augmented lagrange solver is used, if the property Seggrps is given then the segregated solver is used and otherwise the standard nonlinear solver is used. These solvers are described in the COMSOL Multiphysics Users Guide; see The Stationary Solver on page 365, The Parametric Solver on page 379, The Stationary Segregated Solver on page 384 and The Parametric Segregated Solver on page 387.

The segregated solver group properties are given through the Seggrps property with one list for each group. The only mandatory property is the property Segcomp defining which solution components to be used for the group. Optional group properties are the standard solver properties; Ntol, Linsolver, Prefun, Uscale, etc. The segregated solution scheme is controlled by four properties not given through the Seggrps property. These are Maxsegiter, Segorder, Segdamp and Segiter. The segregated substeps are controlled by the property Segorder where the segregated group numbers should be given in the preferred solution order by an integer vector. The number of substeps is thereby determined by the length of the the given integer vector. If this property is not given the groups are solved for, from first to last. The damped Newton method used for the substeps, where the number of iterations and the damping factor is held fixed. The properties Segiter and Segdamp control the number of iterations and the damping factor respectively for the substeps. The maximum number of segregated iterations is controlled by the property Maxsegiter.

The linear solver and the segregated solver use the property Itol for termination of iterative linear system solvers and for error checking for direct solvers (if enabled). The nonliner solver uses an adaptive tolerance for termination of iterative linear system solvers. This adaptive tolerance is based on the maximum of Ntol and Itol. During the nonliner iterations, it can, however, be larger or smaller than this number. The parametric solver uses the same tolerance as the corresponding stationary solver.

See Also femlin, femnlin, femsolver, femstruct, assemble, asseminit

Purpose	FEM structure.
Syntax	help femstruct
Description	The <i>FEM structure</i> is a container for the full description of a PDE problem. See "FEM Structure Overview" on page 10 in the COMSOL Multiphysics Scripting Guide.
Compatibility	The fields fem.equiv and fem.mat are no longer supported. The fem.rules field is obsolete and replaced by fem.functions.
	The field fem.variables has been renamed to fem.const in FEMLAB 2.3.

Purpose	Solve time-depen	ndent PDE problem.			
Syntax	fem = femtime	time(fem,'Tlist',[t (fem,'Tlist',[t1 time('in',{'K' K 'N 'Tlist',[t1	. tn],'Out',{ ' N 'L' L 'M'	'fem'},)	
Description	fem.sol = fem	time(fem,) solves a	time-dependent	t PDE problem.	
	The PDE problem is stored in the (possibly extended) FEM structure fem. See femstruct for details. The time interval and possible intermediate time values are given in the property Tlist. The output times are controlled by the property Tout.				
	fem.sol = femtime('in',{'K' K 'N' N 'L' L 'M' M 'D' D 'E' E}) solves the pre-assembled linear problem				
	$E\ddot{U} + D\dot{U} + KU = L - N_F \Lambda$				
	NU = M				
	The function femtime accepts the following property/values:				
	TABLE I-43: VALID PROPERTY/VALUE PAIRS				
	PROPERTY	VALUES	DEFAULT	DESCRIPTION	
	Atol	See below	1E-3	Absolute tolerance	
	Complex	on off	off	Complex numbers	
	Consistent	off on bweuler	bweuler	Consistent initialization of DAE systems	
	Estrat	0 1	0	Error estimation strategy	

cell array of names and

matrices K | L | M | N |

string containing K, L,

integer between I and

positive scalar

M, N, D, E | auto

yes | maybe

positive scalar

DE

5

1 | 2

N and M are

empty,

D=E=0

auto

maybe

5

1

Input matrices

Initial time step

quantities

Time-independent

Singular mass matrix

Maximum BDF order

Maximum time step

Minimum BDF order

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In

Кеер

Initialstep

MassSingular

Maxorder

Maxstep

Minorder

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Out	<pre>fem sol u tlist solcompdof stop Kc Lc Dc Ec Null Nnp ud uscale nullfun symmetric cellarray of these strings</pre>	sol	Output variables
Rtol	numeric	0.01	Relative tolerance
Stopcond	string with expression		Stop before the expression becomes negative
Tlist	numeric vector		Time list
Tout	tout tsteps	See below	Output times
Tsteps	free intermediate strict		Time-stepping mode

TABLE I-43: VALID PROPERTY/VALUE PAIRS

In addition, the properties described in the entry femsolver are supported.

The maximum allowed relative error in each time step (the local error) is specified using Rtol. However, for small components of the solution vector U, the algorithm tries only to reduce the absolute local error in U below the tolerance given in Atol. The absolute tolerance Atol can be given for each degree of freedom separately. The value for the property Atol can be:

- A scalar.
- A solution vector.
- A solution object.
- A row cell array with alternating degree of freedom names and definitions. The definitions can be numeric scalars or string expressions. The string expressions may only depend on constants defined in fem.const or Const. Unspecified degree of freedom names are given the default value 0.

There is no guarantee that the error tolerances are met strictly, that is, for hard problems they can be exceeded.

For the tolerance parameter in the convergence criterion for linear systems, the maximum of the numbers Rtol and Itol is used.

Use Complex=on if complex numbers occur in the solution process.

The property Consistent controls the consistent initialization of a *differential algebraic equation* (DAE) system. The value Consistent=off means that the initial values are consistent (seldom the case since the initial value of the time derivative is 0). Otherwise, the solver tries to modify the initial values so that they become consistent. The value Consistent=on can be used for index-1 DAEs. Then the solver fixes the values of the differential DOFs, and solve for the initial values of the algebraic DOFs and the time derivative of the differential DOFs. The value Consistent=bweuler can be used for both index-1 and index-2 DAEs. Then the solver perturbs the initial values of all DOFs by taking a backward Euler step.

For a DAE system, if Estrat=1, then the algebraic DOFs are excluded from the error norm of the time discretization error.

You can suggest a size of the initial time step using the property Initialstep.

By default, the solver determines whether the system is differential-algebraic by looking after zero rows or columns in the mass matrix. If you have a DAE where the mass matrix has no zero rows or columns, put Masssingular=yes.

The property Maxorder gives the maximum degree of the interpolating polynomial in the BDF method.

The property Maxstep puts an upper limit on the time step size.

The property Out determines the output arguments and their order. The solution object fem.sol contains the output times and the corresponding solutions, see femsol. By default, the time derivatives are not stored in the solution. To store them, use the Outcomp property, see femsolver. This will also give a more accurate value in postprocessing of values interpolated in time. The output u is a matrix whose columns are the solution vectors for the output times. The output tlist is a row vector containing the output times. The output variable Stop is 0 if a complete solution was returned, 1 if a partial solution was returned, and 2 if no solution was returned. For the other outputs, see femlin.

The property Stop makes it possible to return a partial solution when the time stepping fails at some point. If a failure occurs, the computed time steps are returned in sol.

Use the property Stopcond to make sure the solver stops before a certain event. You provide a scalar expression that is evaluated after each time step. The time stepping is stopped if the real part of the expression is evaluated to something negative. The corresponding solution, for which the expression is negative is not returned.

You can use the property Keep to tell femtime that certain quantities are constant in time, which sometimes can speed up the computation, see "Manual Control of Reassembly" on page 496. The corresponding value is a string or a cell array of strings.

The property Tlist must be a strictly monotone vector of real numbers. Commonly, the vector consists of a start time and a stop time. If more than two numbers are given, the intermediate times can be used as output times, or to control the size of the time-steps (see below). If just a single number is given, it represents the stop time, and the start time is 0.

The property Tout determines the times that occur in the output. If Tout=tsteps, then the output contains the time steps actually taken by the solver. If Tout=tlist, then the output contains interpolated solutions for the times in the Tlist property. The default is Tout=tsteps, except when the property Tsteps=free and Tlist has length greater that 2, in which case Tout=tlist.

The property Tsteps controls the selection of time steps. If Tsteps=free, then the solver selects the time steps according to its own logic, disregarding the intermediate times in the Tlist vector. If Tsteps=strict, then time steps taken by the solver contain the times in Tlist. If Tsteps=intermediate, then there are at least one time step in each interval of the Tlist vector.

For more information about the time-dependent solver; see "The Time-Dependent Solver" on page 370 in COMSOL Multiphysics User's Guide.

Example

Solve the heat equation

$$\frac{\partial u}{\partial t} - \Delta u = 0$$

on a square geometry $-1 \le x, y \le 1$. Choose u(0) = 1 on the disk $x^2 + y^2 < 0.4^2$, and u(0) = 0 otherwise. Use Dirichlet boundary conditions u = 0. Compute the solution at times linspace(0,0.1,20).

```
clear fem
fem.geom = square2(2,'pos',[-1 -1])+circ2(0.4);
fem.mesh = meshinit(fem);
fem.shape = 2;
fem.equ.c = 1; fem.equ.da = 1;
fem.bnd.h = 1;
fem.equ.init = {0 1};
fem.xmesh = meshextend(fem);
fem.sol = femtime(fem,'report','on','tlist',linspace(0,0.1,20));
```

	postanim(fem,'u')
Cautionary	In structural mechanics models, the displacements are often quite small, and it is critical that the Atol property is chosen to be smaller than the actual displacements.
Compatibility	The property Variables has been renamed to Const in FEMLAB 2.3.
	The properties Epoint and Tpoint are obsolete from FEMLAB 2.2. Use fem.***.gporder to specify integration order.
See Also	femsolver, assemble, asseminit, femstruct, femlin, femnlin

Purpose	Extend FEM structure to a wave equation problem.
Syntax	fem1 = femwave(fem) xfem1 = femwave(xfem)
Description	<pre>fem1 = femwave(fem) extends the coefficients of the PDE problem to a wave equation problem. xfem can also be an extended FEM structure. In the latter case, use geomnum to specify the geometry number for the wave extension.</pre>

Note: Since COMSOL Multiphysics 3.2, wave equations can be more easily and efficiently formulated using the e_a coefficient or the second time derivative variable.

When fem is given in coefficient form, fem1 contains extended PDE and boundary coefficients to solve the wave equation problem

$$d_a \frac{\partial^2 u}{\partial t^2} - \nabla \cdot (c \nabla u + \alpha u - \gamma) + \beta \nabla u + a u = f$$

where the d_a coefficient was stored in the fem.equ.da field of the FEM structure fem. In the same way, when fem is given in general or weak form, fem1 contains extended PDE and boundary coefficients to solve the wave equation problem obtained by replacing the first time derivative term in the standard problem by a second time derivative.

The function introduces a set of new variables, v, such that

$$v = \frac{\partial u}{\partial t}$$

and then transforms the PDE problem by doubling the size of the system, and rewriting the coefficients according to the table below:

PDE COEFFICIENT	COEFFICIENT FORM	GENERAL FORM
С	$\begin{bmatrix} 0 & 0 \\ c & 0 \end{bmatrix}$	$\begin{bmatrix} 0 & 0 \\ c & 0 \end{bmatrix}$
α	$\begin{bmatrix} 0 & 0 \\ \alpha & 0 \end{bmatrix}$	$\begin{bmatrix} 0 & 0 \\ \alpha & 0 \end{bmatrix}$
γ	O Y	O Y
a	$\begin{bmatrix} 0 & -I \\ a & 0 \end{bmatrix}$	$\begin{bmatrix} 0 & -I \\ a & 0 \end{bmatrix}$
f	$\begin{bmatrix} 0\\f \end{bmatrix}$	$\begin{bmatrix} v \\ f \end{bmatrix}$
d_a	$\begin{bmatrix} I & 0 \\ 0 & d_a \end{bmatrix}$	$\begin{bmatrix} I & 0 \\ 0 & d_a \end{bmatrix}$
e_a	$\begin{bmatrix} e_a & 0 \\ 0 & 0 \end{bmatrix}$	$\begin{bmatrix} e_a & 0 \\ 0 & 0 \end{bmatrix}$
q	$\begin{bmatrix} 0 & 0 \\ q & 0 \end{bmatrix}$	$\begin{bmatrix} 0 & 0 \\ q & 0 \end{bmatrix}$

PDE COEFFICIENT	COEFFICIENT FORM	GENERAL FORM
g	$\begin{bmatrix} 0\\ g \end{bmatrix}$	$\begin{bmatrix} 0\\g \end{bmatrix}$
h	$\begin{bmatrix} \frac{\partial h}{\partial t} & h \\ h & 0 \end{bmatrix}$	$\begin{bmatrix} \frac{\partial h}{\partial t} & h \\ h & 0 \end{bmatrix}$
r	$\begin{bmatrix} \frac{\partial r}{\partial t} \\ r \end{bmatrix}$	$\begin{bmatrix} \frac{\partial r}{\partial t} - hv \\ r \end{bmatrix}$

TABLE I-44: TRANSFORMATION DONE BY FEMWAVE

When the property Tdiff is off, the following modifications to the table applies:

TABLE I-45: TRANSFORMATION DONE BY FEMWAVE

h	$\begin{bmatrix} 0 & 0 \\ h & 0 \end{bmatrix}$	$\begin{bmatrix} 0 & 0 \\ h & 0 \end{bmatrix}$
r	$\begin{bmatrix} 0\\ r \end{bmatrix}$	$\begin{bmatrix} 0\\ r\end{bmatrix}$

For a PDE problem in general form, femwave produces h using femdiff, when the Diff property is on and the field h does not exist.

If the coefficients weak, dweak, and constr are present, either in addition to the above coefficients, or on their own because the problem is in weak form, they also become doubled in size, but their treatment is very special because they may contain explicit references to the n dependent variables u, their time derivatives u_time and the test functions u_test , ux_test , etc.

Weak coefficients are cell arrays of length n. They become cell arrays of length 2n, by moving the existing entries down to the second half, then replacing all references therein to u_test, ux_test , etc. with references to v_test, vx_test , etc. The first half of the vector contains n entries of the form $v_i^*u_i_test$.

Dweak coefficients are cell arrays of length n. They become cell arrays of length 2n, by moving the existing entries down to the second half, then replacing all references therein to u_time , ux_time , u_test , ux_test , etc. with references to v_time , vx_time , v_test , vx_test , etc. The first half of the vector contains n entries of the form $u_i_time*u_i_test$.

Constr coefficients are cell arrays of length n. They become cell arrays of length 2n, by moving the existing entries down to the second half. If Tdiff is on, then the top half of the vector is filled by the following entries. If one of the coefficients is c_i , one of the new entries in the top half of the new vector is

$$\frac{\partial c_i}{\partial t} + v_j \frac{\partial c_i}{\partial u_j} + (v_j)_{x_k} \frac{\partial c_i}{\partial (u_j)_{x_k}}$$

where $(v_j)_{x_k}$ represents the partial derivative of v_j with respect to space coordinate x_k , and where there is implicit summation over repeated indices.

The function femwave accepts the following property/value pairs:

TABLE I-46: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Defaults	off on	off	Return default fields
Diff	cell array with strings that may contain the strings r, var, and/or expr or the strings off or on	on (for general form)	Differentiate constraints. Describes which fields that are differentiated if Tdiff is on. See also femdiff.
Geomnum	integer	1	Geometry number
Shrink	off on	off	Shrinks coefficients to most compact form.
Simplify	off on	on	Simplify differentiated expressions
Tdiff	on off	on	Differentiate constraints with respect to time

	Use fem.rules to specify additional differentiation rules. The derivative of the Inverse hyperbolic tangent function $atanh$ can, for example, be specified as {'atanh', '1./(1-x.^2)'}. It can also be stored as a field in the fem structure.
Cautionary	The properties bdl, out, rules, and sdl are obsolete in FEMLAB 3.0.
	The h and r coefficients at level 4 of the syntax must be given either as a scalar numeric value, or a string containing an expression.
	You should set the ODE Suite parameter maxorder to 2 for the solver ode15s for wave type problems. This is automatically done by the graphical user interface.
Compatibility	In FEMLAB 1.0, when using general form, you had to apply femdiff before femwave. This was because the h coefficient in fem affects the result of the r coefficient in the output fem1, and h had to be computed by symbolic differentiation by femdiff. In FEMLAB 1.1, h is automatically computed by femwave if not provided. Therefore femdiff can be applied after the femwave call in FEMLAB 1.1.
See Also	femtime

Purpose	Create circular rounded corners in geometry object.			
Syntax	<pre>g = fillet(g1,)</pre>			
Description	g = fillet(g	1,) creates ro	unded corn	ers in 2D geometry object.
	The function fillet accepts the following property/values:			property/values:
	PROPERTY	VALUE	DEFAULT	DESCRIPTION
	out	Cell array of strings	none	Determines the output
	point	integers all none	all	Specifies which vertices are filleted
	radii	I-by- <i>m</i> vector		Curvature radii of the fillet
Examples	for all corners. Fillet a rectang r = rect2; s1 = fille	le object:	1);	e corner then the single radius is used
Diagnostics	If fillet does radius, the corr		eating a rour	nded corner according to the specified
	When a fillet ir	ntersects another e	dge, the fun	action generates an error message.
Compatibility	have a commo	n vertex can be fill	eted. For ed	er supported. Only pair of edges that lges that are not linear, the linear sed to compute an approximate fillet.
See Also	chamfer, curv	ve2, curve3		

Purpose	Compact equ/bnd/edg/pnt fields.			
Syntax	fem = flcompact(fem)			
Description	<pre>fem = flcompact(fem) removes unused and duplicated coefficients in the fem.equ, fem.bnd, fem.edg, and fem.pnt fields. The resulting structures always have numeric ind fields. Coefficients are considered equal if they represent the same expression, that is, equivalent short-hand and expanded forms are compacted.</pre>			
		ICOMPACT accepts the fo PROPERTY/VALUE PAIRS	llowing prope	rty/value pairs:
	PROPERTY	VALUES	DEFAULT	DESCRIPTION
	defaults	off on	off	Return default fields
	shrink	off on	off	Shrinks coefficients to most compact form.
Compatibility	The syntaxes equ = flcompact(equ,'equ',nsd) bnd = flcompact(bnd,'bnd',nbnd) edg = flcompact(edg,'edg',nedg) pnt = flcompact(pnt,'pnt',npnt) field = flcompact(field,fldnames,nelem) are no longer supported in FEMLAB 3.1.			
See Also	multiphysics			

Purpose	Create boundary mesh from contour data.	
Syntax	<pre>m = flcontour2mesh(c)</pre>	
Description	m = flcontour2mesh(c) creates a boundary mesh m with fields m.p and m.e from the contour data c. The contour matrix c is a two-row matrix of contour lines. Ea contiguous drawing segment contains the value of the contour, the number of (x drawing pairs, and the pairs themselves. The segments are appended end-to-end of the contour segment contains the value of the contour, the number of (x) drawing pairs, and the pairs themselves. The segments are appended end-to-end of the contour segment contains the value of the contour, the number of (x)	
	c = [level1 x1 x2 x3 level2 x2 x2 x3; pairs1 y1 y2 y3 pairs2 y2 y2 y3]	
	The contour matrix format is used by the COMSOL Script or MATLAB function contourc.	
	By using the contour matrix format, you can convert geometry data defined by a point set, to a COMSOL Multiphysics geometry object. Firstly, define a contour matrix c corresponding to your point set and use flcontour2mesh to convert the contour matrix c to a 2D boundary mesh m. Then, use flmesh2spline to convert the mesh object m to a curve2 object.	
Examples	Create a mesh from contour data.	
	<pre>[x,y] = meshgrid(linspace(-3,3,50)); z = (x.^2+y.^2).*exp(-x.^2-y.^2)+cos(y)+sin(x); figure c = contour(z); m = flcontour2mesh(c); figure meshplot(m);</pre>	
See Also	contourc,flmesh2spline,flim2curve	

Purpose	Smoothed step functions.	
Syntax	<pre>y = flc1hs(x,scale) y = flc2hs(x,scale) y = fldc1hs(x,scale) y = fldc2hs(x,scale)</pre>	
Description	y = flc1hs(x,scale) and $y = flc2hs(x,scale)$ compute the values of a smoothed version of the Heaviside function $y = (x>0)$. The function is 0 for x<-scale, and 1 for x>scale.	
	In the interval -scale <x<scale, a="" by="" continuous="" defined="" derivative="" fifth-degree="" first="" flc1hs="" function="" heaviside="" is="" it="" overshoot.="" polynomial.<="" smoothed="" th="" with="" without=""></x<scale,>	
	In the interval -scale <x<scale, a="" by="" continuous="" defined="" derivative="" flc2hs="" function="" heaviside="" is="" it="" overshoot.="" polynomial.<="" second="" sixth-degree="" smoothed="" th="" with="" without=""></x<scale,>	
	The input x can be an array. The input scale must be positive scalar.	
	<pre>yp = fldc1hs(x,scale) and yp = fldc2hs(x,scale) compute the derivative of the functions flc1hs and flc2hs, respectively.</pre>	
See Also	flsmhs, flsmsign, fldsmhs, fldsmsign	

Purpose	Convert between PDE forms.
Syntax	fem1 = flform(fem,'outform',form,) [equ,bnd] = flform(fem,'outform',form,)
Description	<pre>fem1 = flform(fem, 'outform', form,) converts the FEM structure fem to an FEM structure fem1 on form. The fields in fem1.equ and fem1.bnd contain the corresponding fields from fem.equ and fem.bnd converted to the form form. fem1.form is set to form. All other fields in fem are copied to fem1.</pre>

[equ,bnd] = flform(fem,'outform',form,...)

is an alternative syntax, returning only the equ and bnd fields of the FEM structure.

Conversion from coefficient to general form is performed according to

$$\begin{split} \Gamma_{lj} &= -c_{lkji} \frac{\partial u_k}{\partial x_i} - \alpha_{lkj} u_k + \gamma_{lj} \\ F_l &= f_l - \beta_{lki} \frac{\partial u_k}{\partial x_i} - a_{lk} u_k \\ G_l &= g_l - q_{lk} u_k \\ R_m &= r_m - h_{ml} u_l \end{split}$$

using a notation where there is an implicit summation over the k (or l) and i indices in each product. Affected fields are therefore ga, c, al, f, be, and a from equ and g, q, r, and h from bnd, with c, al, be, a, g, and q removed and ga, f, r, and g remaining. Other fields within equ and bnd, such as shape, weak, init, var, etc., remain unchanged.

Conversion from general form to weak form is performed according to

$$\begin{split} W_l^{(n)} &= W_l^{(n)} + \Gamma_{lj} \frac{\partial v_l}{\partial x_j} + F_l v_l \\ W_l^{(nt)} &= W_l^{(nt)} + d_{alk} \frac{\partial u_k}{\partial t} v_l + e_{alk} \frac{d^2 u_k}{dt^2} v_l \\ W_l^{(n-1)} &= W_l^{(n-1)} + G_l v_l \\ R_m^{(n)} &= R_m \end{split}$$

where there is an implicit summation over the k and i indices in each product. n is the space dimension. Affected fields are therefore ga, f, weak, da, ea, and dweak from equ and g, weak, r, and constr from bnd, with weak, dweak, and constr the only fields remaining. Other fields within equ and bnd, such as shape, init, var, etc., remain unchanged.

In addition, when converting to weak form, flform tries to take fem.border into account. That is to say that if fem.border is not 1 or on, there may be interior boundaries on which boundary conditions should not be applied. This process is carried out because meshextend and the solvers pay no attention to fem.border when considering weak, dweak, and constr, unlike ga, c, f, q, r, h, etc.

The function flform accepts the following property/value pairs:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
defaults	off on	off	Return default fields
outform	coefficient general weak	coefficient	Output form
out	fem equ bnd edg pnt	fem	Output variables
shrink	off on	off	Shrinks coefficients to most compact form.
simplify	off on	on	Simplify expressions

TABLE I-49: VALID PROPERTY/VALUE PAIRS

Cautionary

Conversion from general form to coefficient form, or from weak form to general or coefficient form is not supported.

ExampleThe following code shows how the convergence can be improved for a stationary
solution of the model "Resistive Heating". The system is converted to general form,
the symbolic derivatives are computed using femdiff, and the system is solved with
femstatic.

```
% !!! First run the example under the multiphysics entry
fem = flform(fem, 'outform', 'general');
fem = femdiff(fem);
fem.xmesh = meshextend(fem);
fem.sol = femstatic(fem, 'report', 'on');
postsurf(fem, 'T');
```

Alternatively, change the outform to weak and remove the femdiff call.

See Also multiphysics, meshextend

Purpose	Create 2D curve object from image data.
Syntax	<pre>[c,r] = flim2curve(I,fmt,)</pre>
Description	[c,r] = flim2curve(I,fmt,) creates a curve2 object c and small curve2 objects r from the image I (gray-scale or RGB) operated upon by parameters contained in the cell-array fmt. c is a curve2 object that approximates the contours of I. r is a cell-array of curve2 objects containing small curves detected by the argument. This is very useful for images containing noise. I is either an <i>m</i> -by- <i>n</i> intensity image matrix or an <i>m</i> -by- <i>n</i> -by-3 RGB-image matrix as typically obtained from the function imread. fmt is a cell-array of length 2 used to create contours from I. If fmt{2} is empty then the scalar fmt{1} is used as a threshold value. If instead, fmt{1} is empty, then the vector fmt{2} is used to specify the contour levels of interest. See the function contourc for an explanation of the contour level syntax in fmt{2}. All flmesh2spline properties are supported.
	$c = flim2curve(i,fmt,)$ is an alternative syntax and is equivalent to $c = geomcsg({},{c,r{:}})$ where the arguments c and r are those obtained from the other call. This is less stable whenever i contain small structures.
Examples	<pre>Create contour-curves from function. [x,y] = meshgrid(linspace(-3,3,50)); z = (x.^2+y.^2).*exp(-x.^2-y.^2)+cos(y)+sin(x); figure imagesc(z) g = flim2curve(z,{[],[-1.5:0.5:2]}); figure geomplot(g,'Pointmode','off');</pre>
	<pre>Create curves from noisy picture. This example does not work in COMSOL Script because it uses the mri.mat file available in MATLAB. load mri pic = D(:,:,1,10); figure image(pic) v = axis; [c,r] = flim2curve(pic,{[],1:30:91},'KeepFrac',0.10); figure geomplot(c,'Pointmode','off'); Plot all small curves in a green color. for j = 1:length(r) hold on geomplot(r{j},'Pointmode','off','edgecolor','g')</pre>

end axis(v) axis ij

See Also

flcontour2mesh, flmesh2spline

Purpose	Load a COMSOL Multiphysics file.
Syntax	flload filename fem = flload('filename')
Description	flload(filename) retrieves FEM structures, geometry objects, or mesh objects from a COMSOL Multiphysics file. If filename has no extension it is assumed to be a Model MPH-file.
	flload supports Model MPH-files (.mph) for retrieving complete FEM structures and COMSOL Multiphysics text and binary files (.mphtxt, .mphbin) for retrieving geometry and mesh objects.
See Also	flsave

Purpose	Create spline curves from mesh.
Syntax	[g2,r2] = flmesh2spline(msh,)
Description	[g2,r2] = flmesh2spline(msh,) creates spline curves g2 and filtered small curves r2. The structure msh is a valid mesh, where only the fields msh.p and msh.e are needed. The object g2 is a curve2 object containing spline curves approximating the edge of msh. The variable r2 is a cell-array containing small curves filtered away by the algorithm. This is a useful feature when trying to

g2 = flmesh2spline(msh,...) is an alternative syntax and is equivalent to $g2 = geomcsg({},{g2,r2{:}})$ where the arguments g2 and r2 are those obtained from the other call. This is less stable whenever msh contain small (ill-conditioned) structures.

generate curves from meshes that originate from noisy contour data.

TABLE I-50: VALID PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
KeepFrac	real scalar >0, <=1	0.2	Fraction of points to keep
Smooth	on off	on	Curve smoothing on or off
SplineMethod	uniform chordlength centripetal foley	chordlength	Method for spline parametrization

The property KeepFrac provides a useful way to reduce the complexity of the resulting geometry object. If the algorithm fails to produce the desired result, try to lower the value of this property. The smoothing algorithm used is a simple anti-aliasing filter, and is controlled by the property Smooth. For an explanation of the property SplineMethod, see geomspline.

Note: You might need to refine the boundary mesh data using meshrefine to be able to get reasonable results using this function. Alternatively, you can create a finer mesh using meshinit by manipulating the mesh parameters.

Examples

Create spline curves from a full mesh:

msh = meshinit(circ2+rect2(1,1,'pos',[0.5 0.5]));

```
figure
meshplot(msh)
c = flmesh2spline(msh,'keepfrac',0.3)
figure
geomplot(c)
```

Create spline curves from contour data:

```
[x,y] = meshgrid(linspace(-3,3,50));
z = (x.^2+y.^2).*exp(-x.^2-y.^2)+cos(y)+sin(x);
figure
c = contour(z);
m = flcontour2mesh(c);
figure
meshplot(m)
g = flmesh2spline(m);
figure
geomplot(g)
```

See Also

flcontour2mesh, flim2curve, geomspline

Purpose	Get number of global degrees of freedom.
Syntax	n = flngdof(fem) n = flngdof(fem, mcase)
Description	The returned number n is the number of degrees of freedom in the FEM structure fem. This is the same as the length of the solution vector. If the mesh case mcase is not given, it is taken to be the mesh case with the greatest number of degrees of freedom in the extended mesh.
See Also	meshextend

Purpose	Compute null space of a matrix, its complement, and the range of the matrix.			
Syntax	<pre>Range = flnull(N,) [Null,Compl] = flnull(N,) [Null,Compl,Range] = flnull(N,) [] = flnull('in',{},'out',{},)</pre>			
Description	Range = flnull(N,) computes the range of N .			
	[Null,Compl complement.] = flnull(N,) c	computes the null	space of N and its
	[Null,Compl and the range		,) computes	null space, its complement,
		ull('in', $\{\ldots\}$,'ou and the range of N .	t',{},)	compute null space, its
	The function	flnull accepts the foll	lowing property/	value pairs:
	TABLE I-51: VALID	PROPERTY/VALUE PAIRS		
	PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
	In	N NT	Ν	Input matrices
	Nullfun	flnullorth flspnull auto	auto	Null-space function
	Out	Null Range Compl		Output variables
	The property Nullfun selects the null-space algorithm. The algorithm flnullorth computes a orthonormal basis for the null space by using singular value decomposition in a block-wise pattern. The method flspnull handles constraint matrices with non-local couplings by employing a sparse algorithm. The auto method automatically selects the most appropriate of flnullorth and flspnull. See the sections "Advanced Solver Settings" on page 492 and "Constraint Handling" on page 499 for further information on the use of these matrices.			
Example	The Poisson E	quation on the Unit D	isk	
		•		rates the way femstatic
	handles the co eliminate.	nstraints internally by	the default constr	aint handling method:
	clear fem			

clear tem
fem.geom = circ2;

```
fem.mesh = meshinit(fem);
fem.shape = 2;
fem.equ.c = 1; fem.equ.f = 1;
fem.bnd.h = 1;
fem.xmesh = meshextend(fem);
[K,L,M,N] = assemble(fem);
[Null,Compl,Range] = flnull(N);
ud = Compl*((Range'*N*Compl)\(Range'*M));
Ke = Null*((L-K*ud);
vn = Ke\Le;
u = Null*vn+ud;
fem.sol = femsol(u);
postplot(fem,'tridata','u')
```

See Also

femlin, assemble

Purpose	Globally turn off the report progress window or show it.
Syntax	flreport('off') flreport('on') flreport('show')
Description	flreport ('off') disables the use of the progress window that is normally shown during meshing and solution.
	flreport('on') enables the use of the progress window again. This means that COMSOL Multiphysics uses the value of the 'report' property to determine if the progress window should be shown.
	flreport('show') shows the progress window if it has been closed.

Purpose	Save a COMSOL Multiphysics file.
Syntax	flsave filename fem
Description	flsave filename arg1 arg2 saves FEM structures, geometry objects, and mesh objects to a COMSOL Multiphysics file. flsave supports Model MPH-file (.mph) and the COMSOL Multiphysics text and binary formats (.mphtxt, .mphbin).
	flsave filename saves all valid COMSOL data in the workspace.
	flsave('filename',arg1,) is an alternative syntax.
Compatibility	Since FEMLAB 3.0, flsave is obsolete for saving a MAT-file. Use save to save an FEM structure or any part of the FEM structure.
See Also	flload

Purpose	Smoothed step functions and their derivatives.
Syntax	<pre>y = flsmhs(x,scale) y = flsmsign(x,scale) yp = fldsmhs(x,scale) yp = fldsmsign(x,scale)</pre>
Description	y = flsmhs(x, scale) computes the values of a smoothed version of the Heaviside function $y = (x>0)$. The function is 0 for x<-scale, and 1 for x>scale.
	<pre>y = flsmsign(x,scale) computes the values of a smoothed version of the sign function y = sign(x). The function is -1 for x<-scale, and 1 for x>scale.</pre>
	In the interval -scale <x<scale, and="" are="" defined<br="" flsmhs="" flsmsign="" functions="" the="">by a seventh-degree polynomial, which is chosen so that the second derivative is continuous. Moreover, the moments of order 0, 1, and 2 agree with those for the Heaviside function and the sign function, respectively. This implies that the functions have small overshoots.</x<scale,>
	<pre>yp = fldsmhs(x,scale) and yp = fldsmsign(x,scale) compute the derivative of the functions flsmhs and flsmsign, respectively.</pre>
	The input x can be an array. The input scale must be positive scalar.
See Also	flc1hs, flc2hs, fldc1hs, fldc2hs

Purpose	Create straight homogeneous generalized cylinder geometry object.
Syntax	<pre>s3 = gency13 s2 = gency12 s3 = gency13(base) s2 = gency12(base) s3 = gency13(base,h) s2 = gency12(base,h) s3 = gency12(base,h,rat) s2 = gency12(base,h,rat) s3 = gency12(base,h,rat,) s2 = gency12(base,h,rat,)</pre>
Description	 s3 = gency13 creates a solid straight homogeneous generalized cylinder geometry object s3, with a solid circle base surface, cylinder axis of length 1 along the z-axis, and size of top surface equal to base surface. gency13 is a subclass of solid3. s3 = gency13(base) creates a solid straight homogeneous generalized cylinder geometry object with base surface base.
	 s3 = gency13(base,h) also sets the height of the generalized cylinder to h. s3 = gency13(base,h,rat) additionally specifies top surface with the scale factor

s3 = gency13(base,h,rat) additionally specifies top surface with the scale factor rat with respect to the origin, that is, all 2D points in the top plane are obtained by multiplying the points in the base plane with rat.

The functions gency13 and gency12 accept the following property/values:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object.
Const	Cell array of strings	{}	Evaluation context for string inputs.
Displ	2-by- <i>nd</i> matrix	[0;0]	Displacement of extrusion top
Pos	Vector of reals or cell array of strings	[0 0]	Position of the bottom surface.
Rot	real or string	0	Rotational angle about Axis (radians).

TABLE 1-52: VALID PROPERTY/VALUE PAIRS

s2 = gency12(...) creates a surface straight homogeneous generalized cylinder, from the same arguments as described for gency13. gency12 is a subclass of face3.

Generalized cylinder objects have the following properties:

TABLE I-53: GENERALIZED CYLINDER OBJECT PROPERTIES

PROPERTY	DESCRIPTION
base	Base 2D geometry object.
h	Height.
rat	Ratio.
dx, dy	Semi-axes.
x, y, z, xyz	Position of the object. Components and vector forms.
ax2	Rotational angle of symmetry axis.
ax3	Axis of symmetry.
rot	Rotational angle.

In addition, all 3D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom3 for details.

CompatibilityThe FEMLAB 2.3 syntax is obsolete but still supported. The numbering of faces,
edges and vertices is different from the numbering in objects created in 2.3.

Examples Creation of a 3D solid with two circular edges, and with a top face that is smaller than the bottom face.

See Also econe2, econe3, extrude, face3

Purpose	Low-level constructor functions for geometry objects.
Syntax	<pre>g = geom3(vertex,pvertex,edge,pedge,face,mfd,pcurve,) [g,] = geom3(g3,) g = geom2(vertex,edge,curve) [g,] = geom2(g2,) g = geom1(p,ud) [g,] = geom1(g1,) g = geom0(p)</pre>
Description	g3 = geom3(vertex,pvertex,edge,pedge,face,mfd,pcurve,) creates a geom3 object.
	$[g, \ldots] = geom3(g3, \ldots)$ coerces any 3D geometry object g3 to a geom3 object.
	c = geom2(vertex,edge,curve,) creates a geom2 object.
	[g,] = geom2(g2,) coerces any 2D geometry object to a geom2 object.
	c = geom1(vtx) creates a 1D geometry object from the property vtx.
	$[g, \ldots] = geom1(g1, \ldots)$ coerces any 1D geometry object to a geom1 object.
	g = geom0(p) creates a 0D geometry object, where p is a matrix of size 0-by-1.
	g = geom0(g1,) coerces any 0D geometry object to a geom0 object.
	The coercion functions accept the following property/values:
	TABLE I-54: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Out	stx ftx ctx ptx	{}	Cell array of output names.

3D Geometry Object Properties

vertex is a 5-by-nv matrix representing the vertices of the 3D geometry. Rows 1, 2, and 3 provide the 3D coordinates of the vertices. Row 4 provides the subdomain number. Row 5 contains a relative local tolerance for the entity. For nontolerant entities the tolerance is NaN.

pvertex is a 6-by-npv matrix containing embeddings of vertices in faces. Row 1 contains the vertex index (i.e. column in VERTEX), rows 2 and 3 contain (s, t) coordinates of the vertex on the face, row 4 contains a face index, and row 5 contains the manifold index into mfd. Row 6 contains a relative local tolerance for the entity.

edge is a 7-by-ne matrix representing the edges of the 3D geometry. Rows 1 and 2 contain the start and end vertex indices of the edge (0 if they do not exist). Rows 3 and 4 give the parameter values of the these vertices. Row 5 gives the index of a subdomain if the edge is not adjacent to a face. Row 6 gives a sign and an index to the underlying manifold. The sign indicates the direction of the edge relative the curve. Finally, row 7 contains a relative local tolerance for the entity.

pedge is a 10-by-npe matrix representing the embeddings of the edges in faces. The first row gives the index of the edge in edge. Rows 2 and 3 contain the start and end vertex indices in pvertex. Rows 4 and 5 give the parameter values of the these vertices. Row 6 and 7 give the indices of the faces to the left and right of the edge, respectively. Row 8 gives a sign and index to the parameter curve (if any), and row 9 gives the index to the surface. Row 10 contains a relative local tolerance for the entity.

face is a 4-by-nf matrix representing the faces of the 3D geometry. Rows 1 and 2 contain the up and down subdomain index of the face, and row 3 contains the manifold index of the face. Row 4 contains a relative local tolerance for the entity.

mfd is a cell array or Java array of 3D manifolds.

pcurve is a cell array or Java array of parameter curves.

TABLE I-55: JAVA 3D MANIFOLD CLASSES

		1
MANIFOLD	USAGE	DESCRIPTION
MfdBezierCurve	(xyzw)	Rational Bezier curve
MfdBezierTri	(xyzw)	Rational Bezier triangular surface
MfdBezierSurf	(xyzw)	Rational Bezier tensor-product surface
MfdBSplineCurve	(deg, knots, P, w)	B-spline curve
MfdBSplineSurf	(uDeg, vDeg, uKnots, vKnots, P, w)	B-spline surface
MfdMeshCurve	(coord, par)	Mesh curve
MfdMeshSurface	(coord, par, tri)	Mesh surface
MfdPolChain	(pol)	Polygon chain manifold

All properties can be accessed using the syntax get(object, property).

2D Geometry Object Properties

vertex is a 4-by-nv matrix representing the vertices of the 3D geometry. Rows 1 and 2 provide the 2D coordinates of the vertices. Row 3 provides the subdomain number. Row 4 contains a relative local tolerance for the entity. For non-tolerant entities the tolerance is NaN.

edge is a 8-by-ne matrix representing the edges of the 3D geometry. Rows 1 and 2 contain the start and end vertex indices of the edge, 0 if they do not exists. Rows 3 and 4 give the parameter values of the these vertices. 5 and 6 contain the left and right subdomain number of the edge. Row 7 gives a sign and an index to the array of underlying curves. The sign indicates the direction of the edge relative the curve. Row 8 contains a relative local tolerance for the entity.

curve is a cell array or Java array of 2D curves..

MANIFOLD	ARGUMENTS	DESCRIPTION
MfdBezierCurve	(xyzw)	Rational Bézier curve manifold
MfdBSplineCurve	(deg, knots, P, w)	B-spline curve
MfdFileCurve	(name,ind, s1,s2)	Geometry M-file manifold
MfdMeshCurve	(coord, par)	Mesh curve
MfdPolChain	(pol)	Polygon chain manifold

TABLE I-56: JAVA 2D MANIFOLD CLASSES

All properties can be accessed using the syntax get(object, property).

ID Geometry Object Properties

vtx is a 3-by-nvtx matrix representing the vertices of the 2D geometry. Row 1 provides the 1D coordinates of the vertices. Rows 2 and 3 provides the up and down subdomain.

All properties can be accessed using the syntax get(object, property).

OD Geometry Object Properties

A 0D geometry object g has the property p, a 0-by-ns double of empty coordinates. ns can be either 1 or 0, for a nonempty and empty object, respectively.

All properties can be accessed using the syntax get(object, property).

Compatibility The FEMLAB 3.0 syntax is obsolete but still supported.

See Also geom0/get, geom1/get, geom2/get, geom3/get, geomobject, geomedit, geominfo, point1, point2, point3, curve2, curve3, face3

Get geometry	object	properties.
--------------	--------	-------------

Syntax get(g,prop)

Description

Purpose

get (g, prop) returns the value of the property prop for a geometry object g, which can be a geometry object of type geom1 (1D geometry object), geom2 (2D geometry object), or geom3 (3D geometry object).

prop is a string that contains a valid property name. The following tables list the valid property names for geom1, geom2, and geom3 objects:

PROPERTY NAME	DESCRIPTION	
vtx	Vertices	
nv	Number of vertices	
Number of subdomains		
mp	Vertex coordinates	
sd	Vertex subdomain numbers	

TABLE I-57: GEOMI PROPERTY NAMES

For information about the formats for the vtx property, see "1D Geometry Object Properties" on page 207.

TABLE I-58: GEOM2 PROPERTY NAMES

PROPERTY NAME	DESCRIPTION
vertex	Vertices
edge	Edges
curve	2D manifolds
nv	Number of vertices
ne	Number of edges
ns	Number of subdomains
mp	Vertex coordinates
sd	Vertex subdomain numbers

For information about the formats for the properties vertex, edge, and curve, see "2D Geometry Object Properties" on page 207.

TABLE I-59: GEOM3 PROPERTY NAMES

PROPERTY NAME	DESCRIPTION
vertex	Vertices
pvertex	Parameter vertices

PROPERTY NAME	DESCRIPTION
edge	Edges
pedge	Parameter edges
face	Faces
mfd	3D manifolds
pcurve	Parameter curves
nv	Number of vertices
ne	Number of edges
nf	Number of faces
ns	Number of subvolumes
mp	Vertex coordinates
sd	Vertex subdomain numbers

TABLE I-59: GEOM3 PROPERTY NAMES

For information about the formats for the properties vertex, pvertex, edge, pedge, face, mfd, and pcurve, see "3D Geometry Object Properties" on page 205.

You can also use get to retrieve specific properties for primitives such as circ2, rect2, block3, cylinder3, ellipsoid3, and sphere3. For example, get(sph,'r') returns the radius of the sphere sph. In this case, type

help sphere3/get

to get a list of available property names.

For geom0 objects, the property ns returns either 1 or 0, for a nonempty and empty object, respectively (see "0D Geometry Object Properties" on page 207).

Example Create a cylinder object and return the number of subvolumes (1), the number of vertices (8), and the number of faces (6):

```
cyl = cylinder3;
ns = get(cyl,'ns')
nv = get(cyl,'nv')
nf = get(cyl,'nf')
geom0, geom1, geom2, geom3,geominfo
```

See Also

 Purpose
 Decompose and analyze geometry of FEM problem.

 Syntax
 fem = geomanalyze(fem, ...) fem = geomanalyze(fem, draw, ...) [fem, map] = geomanalyze(fem, ...)

 Description
 fem = geomanalyze(fem, ...) analyzes and updates the geometry data for the model defined by fem.

 fem = geomanalyze(fem, draw, ...) analyzes and updates the geometry data in the model defined by fem. draw is one or several input arguments given in the same ways as in geomcsg.

 [fem, map] = geomanalyze(fem, ...) additionally returns a cell array of vectors

[rem, map] = geomanalyze(rem, ...) additionally returns a cell array of vectors representing the mappings for the different domains. The vector map{k+1} describes the mapping of k-dimensional domains. Each element in this vector is the associated index of that domain in fem.geom before the call.

The function supports the following property/values:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Geomnum	integer	1	Geometry number
Imprint	on off	on	Make imprints when creating pairs
Ns	cell array of strings	{}	Name space
Paircand	all none cell array of strings	not specified	Specifies the geometries among which pairs are created
Repairtol	positive scalar	0.0	Tolerance for repairing gaps
Sf	string	union	Boolean expression
Solidify	on off	off	Coerce the result to a solid

TABLE I-60: VALID PROPERTY/VALUE PAIRS

If the property paircand is not specified, the geometry data in the model is the result of the boolean operation specified in the property sf. See geomesg for more details.

If the property paircand is specified, an assembly geometry is created. In addition identity pairs are created using the property imprint. See geomgroup for details.

The above properties are explained in geomcsg and geomgroup respectively.

Example The following is an example of a circle containing the source for the model, moving through two different subdomains. % Circle moving through two rectangles clear fem draw{1} = rect2(.5,1, 'pos',[0 0]); draw{2} = rect2(.6,1, 'pos', [0.5 0]); draw{3} = circ2(.1, 'pos', [0.2 0.5]); % Create analyzed geometry fem = []; fem = geomanalyze(fem,draw,'ns',{'R1','R2','C1'}); % Create mesh fem.mesh = meshinit(fem, 'report', 'off'); % Set source in circle fem.appl.mode = 'F1PDEC'; fem.appl.equ.f = $\{0 \ 0 \ 1\};$ fem.appl.equ.c = 1; fem.appl.bnd.h = 1; fem = multiphysics(fem); % Assemble and solve fem.xmesh = meshextend(fem); fem.sol = femstatic(fem, 'report', 'off'); % Plot solution postplot(fem, 'tridata', 'u') % Start loop and move geometry % number of steps in loop nSteps = 5;dist = .75/nSteps; % distance to move every step for i = 1:nSteps % Modify geometry in draw structure c1 = drawgetobj(fem, 'C1'); fem = drawsetobj(fem, 'C1', move(c1, dist, 0)); % Re/analyze geometry, and update boundary conditions fem = geomanalyze(fem); % Create mesh for new geometry fem.mesh = meshinit(fem, 'report', 'off'); % Update equation system fem = multiphysics(fem); % Assemble and solve new equation system fem.xmesh = meshextend(fem); fem.sol = femstatic(fem, 'report', 'off'); % Plot solution figure, postplot(fem, 'tridata', 'u') end

See Also

geomcsg, geomgroup, geomedit

Purpose	Create rectangular array of geometry objects.
Syntax	<pre>cg1 = geomarrayr(g1,dx) cg1 = geomarrayr(g1,dx,n) cg2 = geomarrayr(g2,dx,dy) cg2 = geomarrayr(g2,dx,dy,n) cg2 = geomarrayr(g2,dx,dy,nx,ny) cg3 = geomarrayr(g3,dx,dy,dz) cg3 = geomarrayr(g3,dx,dy,dz,n) cg3 = geomarrayr(g3,dx,dy,dz,nx,ny,nz)</pre>
Description	<pre>cg1 = geomarrayr(g1,dx) distributes copies of the 1D geometry object g1 with absolute displacements, dx with respect to geometry object g1. cg1 is a cell array with the distributed geometry objects. The cg1 cell array has the same size as dx. cg1 = geomarrayr(g1,dx,n) distributes copies of the 1D geometry object g1, n times with the relative scalar displacements dx.</pre>
	cg2 = geomarrayr(g2, dx, dy) distributes copies of the 2D geometry object g2 with absolute displacements, dx and dy with respect to geometry object g2. The displacements dx and dy are matrices of equal size. $cg2$ is a cell array with the distributed geometry objects. The $cg2$ cell array has the same size as dx and dy.
	cg2 = geomarrayr(g2, dx, dy, n) distributes copies of the 2D geometry object $g2$, n times with the relative scalar displacements dx and dy .
	cg2 = geomarrayr(g2,dx,dy,nx,ny) distributes copies of the 2D geometry object g2, nx, and ny times in corresponding directions, with the relative displacements dx and dy. The geometry g2 is included as the first item in the output cell array cg2.
	cg3 = geomarrayr(g3, dx, dy, dz) distributes copies of the geometry object g3 with absolute displacements, dx, dy, and dz with respect to geometry object g3. The displacements dx, dy, and dz are matrices of equal size. cg3 is a cell array with the distributed geometry objects. The cg3 cell array has the same size as dx, dy, and dz.
	cg3 = geomarrayr(g3, dx, dy, dz, n) distributes copies of the geometry object g3, n times with the relative scalar displacements dx, dy, and dz.
	cg3 = geomarrayr(g3,dx,dy,dz,nx,ny,nz) distributes copies of the geometry object g3, nx, ny, and nz times in corresponding directions, with the relative displacements dx, dy, and dz. The geometry g3 is included as the first item in the output cell array cg3.

	The input argument g1, g2, or g3 could also be a cell array of geometry objects. In that case the corresponding output argument cg1, cg2, or cg3 is a cell array of cell arrays.
Example	The following commands are used to create a block object with four equally sized holes.
	g=geomcsg(geomarrayr(cylinder3,4,4,0,2,2,1)); g2=block3(10,14,5,'corner',[-3 -5 -4])-g; geomplot(g2)
See Also	geomO, geom1, geom2, geom3,mo∨e

Decompose and coerce geometry objects.

Syntax

Description

Purpose

[g,...]=geomcoerce(class,ol,...)

g=geomcoerce(class,ol) decomposes the geometry objects in the cell array ol, coerces the composite object to the class class, and returns the coerced geometry object in g.

The function geomcoerce accepts the following property/values:

TABLE I-61: VALID PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Out	stx ftx ctx ptx		Output

Class is one of the strings: solid, face, curve, or point specifying the class of geometry object that is returned in g.

01 is a cell array of geometry objects.

For information on the geometry tables stx, ftx, ctx, and ptx, see geomcsg.

See Also geomcsg, geomanalyze

Purpose	Analyze geometry objects.
Syntax	[g,]=geomcomp(ol,)
Description	[g,]=geomcomp(ol,) analyzes the geometry objects in the cell array ol according to the specified properties and returns the analyzed geometry in g, where g is of the same geometry class as the objects in ol.

The function geomcomp accepts the following property/values:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Edge	integer vector all none	all	Specifies which edges that are deleted in the analyzed geometry
Face	integer vector all none	all	Specifies which faces that are deleted in the analyzed geometry
Ns	cell array of strings	none	Name space
Out	stx ftx ctx ptx		Output
Point	integer vector all none	none in 3D none in 2D all in ID	Specifies which vertices that are deleted in the analyzed geometry
Sf	text expression	union of objects	String with Boolean expression

TABLE I-62: VALID PROPERTY/VALUE PAIRS

For property descriptions of Ns and Sf, see geomcsg. For more information on the properties Edge, Face, and Point, see geomdel.

See Also

geomcsg, geomdel, geomanalyze

Purpose	Analyze geometry model.
Syntax	<pre>g = geomcsg(fem,) g = geomcsg(draw,) g = geomcsg(sl,) g = geomcsg(sl,fl,cl,) g = geomcsg(sl,fl,cl,pl,) g = geomcsg(sl,fl,cl,pl,) g = geomcsg(sl,cl,pl,) g = geomcsg(sl,cl,pl,) [g,st] = geomcsg(sl,fl,) [g,st,ft] = geomcsg(sl,fl,cl,) [g,st,ft,ct] = geomcsg(sl,fl,cl,l,l) [g,st,ft,ct,pt] = geomcsg(sl,fl,cl,pl,) [g,st,ct,pt] = geomcsg(sl,cl,pl,) [g,st,pt] = geomcsg(sl,pl,) [g,st,pt] = geomcsg(sl,pl,)</pre>
Description	 g = geomcsg(fem) analyzes the <i>geometry model</i> in fem.draw, by performing <i>Boolean operations</i> on the solid objects, and superimposing the objects of lower dimension on top of the result of the Boolean operations. The result is an <i>analyzed geometry</i> object g. g = geomcsg(draw) analyzes the <i>geometry model</i> draw and returns the geometry object g. g = geomcsg(sl) decomposes solid objects sl into the analyzed geometry object
	 g. g = geomcsg(sl,fl,) decomposes the 3D solid objects sl and the 3D face objects fl into the analyzed 3D geometry g. g = geomcsg(sl,fl,cl,) decomposes the 3D solid objects sl, the 3D face objects fl, and the 3D curve objects cl into the analyzed 3D geometry g. g = geomcsg(sl,fl,cl,pl,) decomposes the 3D solid objects sl, the 3D face objects fl, the 3D curve objects cl, and the 3D point objects pl into the analyzed 3D geometry g. g = geomcsg(sl,cl,) decomposes the 2D solid objects sl and the 2D curve objects cl into the analyzed 2D geometry g. g = geomcsg(sl,cl,pl,) decomposes the 2D solid objects sl, the 2D curve objects cl into the analyzed 2D geometry g.

g = geomcsg(sl,pl,...) decomposes the 1D solid objects sl and the 1D point objects pl into the analyzed 1D geometry g.

[g,st] = geomcsg(sl) additionally returns a solid table, st, that relates the original solid objects in sl to the subdomains in g.

[g,st,ft] = geomcsg(sl,fl,...) additionally returns a face table, ft, that relates the original face objects in fl to the face segments in g.

[g,st,ft,ct] = geomcsg(s1,f1,c1,...) additionally returns the curve table, ct, that relates curve objects in c1 to edge segments in g.

[g,st,ft,ct,pt] = geomcsg(sl,fl,cl,pl,...) additionally returns the point table, pt, that relates point objects in pl to vertices in g.

[g,ct] = geomcsg(sl,cl,...) additionally returns the curve table, ct, for 2D geometry objects.

[g,ct,pt] = geomcsg(sl,cl,pl,...) additionally returns the point table, pt, for 2D geometry objects.

[g,pt] = geomcsg(sl,pl,...) additionally returns the point table, pt, for 1D geometry objects.

s1, f1, c1, and p1 are cell arrays containing the geometry objects of different types.

st, ft, ct, and pt are matrices where each row corresponds to the number of the corresponding geometric entity in g, and each row corresponds to the object with the same index in s1, f1, c1, and p1, respectively.

The function geomesg accepts the following property/values:

TABLE I-63: VALID PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Sf	text expression	union of objects	String with Boolean expression
Ns	cell array of strings	none	Name space
Out	g st ft ct pt stx ftx ctx ptx	g	Output variables
Repairtol	scalar	0.0	Tolerance used for repairing gaps
Solidify	off on	off	Create all possible subdomains

Out specifies the output arguments from geomcsg. The outputs stx, ftx, ctx, and ptx correspond to st, ft, ct, and pt but contain more detailed information about the geometric entity relations.

stx is a cell array of matrices similar to st. Each matrix contains the relation between subdomains in each solid object in the inputs, and the final subdomains in the output g.

ftx is a cell array of matrices similar to ft. Each matrix contains the relation between face segments in each solid object and face object in the inputs, and the final face segments in the output g.

ctx is a cell array of matrices similar to ct. Each matrix contains the relation between edge segments in each solid object and curve object in the inputs, and the final edge segments in the output g.

ptx is a cell array of matrices similar to pt. Each matrix contains the relation between vertices in each solid object, face object, curve object, and point object in the inputs, and the final vertices in the output g.

Sf represents a *set formula* with variables names from ns. The operators +, *, and – correspond to the set operations union, intersection, and set difference, respectively. The precedence of the operators + and – are the same. * has higher precedence. You can control the precedence with parentheses.

Ns is a cell array of variable names that relates the elements in s1 to variable names in sf. Each element in ns contains a variable name. Each such variable assigns a name to the corresponding solid object in s1. This way you can refer to a solid object in s1 in the set formula sf.

Geometry Model

The geometry model fem.draw contains the following fields:

FIELD	ID	2D	3D	DESCRIPTION
s.objs	\checkmark	\checkmark	\checkmark	Cell array of solid objects
s.name	\checkmark	\checkmark	\checkmark	Cell array of names (default for the property ns)
s.sf	\checkmark	\checkmark	\checkmark	String with Boolean expression (default for the property sf)
f.objs			\checkmark	Cell array of face objects

TABLE I-64: GEOMETRY MODEL OR DRAW STRUCTURE

FIELD	ID	2D	3D	DESCRIPTION
f.name			\checkmark	Cell array of names (ignored by geomcsg)
c.objs		\checkmark	\checkmark	Cell array of curve objects.
c.name		\checkmark	\checkmark	Cell array of names (ignored by geomcsg)
p.objs	\checkmark	\checkmark	\checkmark	Cell array of point objects.
p.name	\checkmark	\checkmark	\checkmark	Cell array of names (ignored by geomcsg)

TABLE I-64: GEOMETRY MODEL OR DRAW STRUCTURE

Examples

3D Geometries

Perform a solid operation on two intersecting cylinders:

```
s1=cylinder3(2,2,[0.5 0.5 -1],[0 0 1]);
s2=cylinder3(1,1,[0.5 0.5 -0.5],[0 0 1]);
[g,st,stx]=geomcsg({s1,s2},'out',{'g','st','stx'},...
'ns',{'Cyl1','Cyl2'},'sf','Cyl1-Cyl2');
```

To easily create solid objects, use the overloaded operators +, *, and -, instead of calling geomcsg:

s=s1-s2; geomplot(s,'facemode','off')

s and g are equivalent, except that g is a geom3 object while s is a solid3 object.

2D Geometries

Create a unit circle solid object and a unit square solid object:

```
c1 = circ2;
sq1 = square2;
g = geomcsg({c1 sq1},{},'ns',{'a' 'b'},'sf','a-b');
```

Using object arithmetic for solid objects, the same result can be obtained by typing.

```
g = c1 - sq1;
```

You can plot the geometry object by

```
geomplot(g,'sublabel','on','edgelabel','on')
```

or just

```
geomplot(g)
```

You can obtain the number of subdomains and edge segment by just typing g or by explicitly getting the object properties.

g get(g,'nmr') get(g,'nbs')

There is one subdomain, with five edge segments, three circle edge segments, and two line edge segments.

ID Geometries

Create a simple 1D geometry by composing two 1D solids:

```
g1 = geomcsg({solid1([0 0.1 4]),solid1([3 4])});
geomplot(g1,'pointlabel','on','sublabel','on')
```

The resulting geometry consists of three subdomains and four vertices.

See Also geomanalyze, geomgroup, geomdel, geomcomp, geominfo, geomplot

Purpose	Delete points, edges, or faces in a geometry.
Syntax	<pre>[g,] = geomdel(g1,) g = geomdel(g1)</pre>
Description	[g,] = geomdel(g1,) deletes points, edges, or faces in the geometry object g1 according to the specified properties. The resulting object is of the same type (solid, face, curve, or point) as the original one.
	g = geomdel(g1) deletes all interior boundaries.
	The function geomdel accepts the following property/values:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Edge	integer vector all none	all	Specifies which edges that are deleted
Face	integer vector all none	all	Specifies which faces that are deleted
Out	stx ftx ctx ptx	none	Output variables (only in 3D). For more information, see the entry geomcsg
Point	integer vector all none	all in 3D none in 2D all in ID	Specifies which vertices that are deleted
Subdomain	integer vector all none	none	Specifies which subdomains that are deleted

TABLE I-65: VALID PROPERTY/VALUE PAIRS

In 1D, Point can either be an array of integers specifying which points that are deleted or one of the strings all or none. all means that all interior points are deleted and none that no points are deleted. The default value is all.

In 2D, Edge all means that all interior boundaries and edges outside any subdomain are deleted and none that no edges are deleted. The default value is all. Point all means that all vertices lying inside a subdomain are deleted and none that no vertices are deleted. The default value is none. Only isolated vertices can be deleted.

In 3D, Face all means that all faces inside or between subdomains are deleted and none that no faces are deleted. The default value is all. Edge all means that all edges lying inside faces are deleted and none that no edge segments are deleted. The

	default value is all. Only edge segments that are not face boundaries can be deleted. Point all means that all vertices lying in faces are deleted and none that no vertices are deleted. The default value is none. Only vertices that are not adjacent to an edge can be deleted.
Examples	The following command generates a block with a face inside, partitioning the subdomain into two parts:
	g = geomcsg({block3}, {face3([0.5 0.5;0.5 0.5],[0 1;0 1],[0 0;1 1])});
	Remove the inner face and all interior edge segments:
	g1=geomdel(g);
See Also	geomcsg

Purpose	Edit geometry	object.			
Syntax	[g,] = ge g1 = geomedi g2 = geomedi				
Descriptiong1 = geomedit(g,) splits the geometry object g into primitive can be edited. Each object in g1 is associated with the object g so to recreate the composite object again.					
	the cell array g		formation to g, t	y using the geometry objects in to create a new composite	
	geomedit only works for 2D geometries.				
	The function geomedit accepts the following property/values:				
	TABLE I-66: VALID	PROPERTY/VALUE PAIRS			
	PROPERTY	VALUE	DEFAULT	DESCRIPTION	
	Out	stx ctx ptx	none	Output variables. For more information, see the entry geomcsg	
Examples	The following	commands create a go	eometrv containi	ing eight curves, then splits the	
	geometry into primitive object g = curve2 gg = geome	<pre>primitive objects, and ct omitted: 2(rect2+circ2); edit(g);</pre>	d finally recreates	<pre>s the geometry with one , 'out', {'ctx'});</pre>	
See Also		nO, geom1, geom2,			

Purpose Export geometry objects to file.

Syntax geomexport(filename, geoms,...)

Description

geomexport(filename, geoms...) exports the geometry data in the cell array geoms of geometry objects to a file.

filename can be any of the following formats:

FILE FORMAT	NOTE	FILE EXTENSIONS
COMSOL Multiphysics Binary		.mphbin
COMSOL Multiphysics Text		.mphtxt
Parasolid Binary	1	.x_b
Parasolid Text	1	.x_t
DXF		.dxf

TABLE I-67: VALID FILE FORMATS

Note 1: This format requires a license for the COMSOL CAD Import Module. Only geometries imported using the CAD Import Module (cad3part objects) can be exported to Parasolid format.

The following properties are supported

TABLE I-68: VALID PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Report	on off	on	Display a progress window

Report determines if a progress window appears during the call.

Examples	r = rect2; geomexport('foo.dxf',{r}) b = block3; geomexport('bar.mphtxt',{b})
Diagnostics	geomexport replaces the functionality of the 3.1 function dxfwrite.
See Also	geomO, geom1, geom2, geom3,meshexport,geomimport

Purpose	Geometry M-file.			
Syntax	ne = geomfile d = geomfile(bs) [x,y] = geomfile(bs,s)			
Description	The Geometry M-file is a template format for a user-specified M-file that contains the complete geometry for a model. You can specify both 1D and 2D geometries by using the Geometry M-file format. The geomfile format is not supported in 3D.			
	<i>ID Geometry</i> For 1D geometries, the Geometry M-file essentially contains a set of points, and geometry information on the intervals between these point.			
	ne = geomfile returns the number of boundary points ne.			
	d = geomfile(bs) returns a matrix d with one column for each boundary point specified in bs, with the following contents:			
	• Row 1 contains the <i>x</i> -coordinate of the boundary point			
	• Row 2 contains the label of the "up" subdomain ("up" is the positive direction)			
	• Row 3 contains the label of the "down" subdomain ("down" is the negative direction)			
	The complement of the union of all subdomains is assigned the subdomain number 0.			
	<pre>2D Geometry 2D subdomains are represented by parameterized edge segments. Both the subdomains and edge segments are assigned unique positive numbers as labels. The edge segments cannot overlap. The full 2D problem description can contain several nonintersecting subdomains, and they can have common interior boundary segments. The boundary of a subdomain can consist of several edge segments. Each subdomain boundary need to consist of at least two edge segments. All edge segment junctions must coincide with edge segment endpoints. ne = geomfile returns the number of edge segments ne. d = geomfile(bs) returns a matrix d with one column for each edge segment specified in bs, with the following contents: • Row 1 contains the start parameter value • Row 2 contains the end parameter value</pre>			

- Row 3 contains the label of the left-hand subdomain (left with respect to direction induced by start and end from row 1 and 2)
- Row 4 contains the label of the right-hand subdomain

The complement of the union of all subdomains is assigned the subdomain number 0.

[x,y] = geomfile(bs,s) produces coordinates of edge segment points. bs
specifies the edge segments and s the corresponding parameter values. bs can be a
scalar.

Examples

The function cardg defines the geometry of a cardioid:

```
r = 2(1 + \cos(\phi))
```

```
function [x,y]=cardg(bs,s)
%CARDG Geometry File defining the geometry of a cardioid.
nbs=4;
if nargin==0
 x=nbs;
  return
end
     0
                    pi
                             3*pi/2
dl=[
             pi/2
      pi/2
             pi
                    3*pi/2
                             2*pi;
      1
             1
                    1
                             1
      0
             0
                    0
                             0];
if nargin==1
 x=dl(:,bs);
  return
end
x=zeros(size(s));
y=zeros(size(s));
[m,n]=size(bs);
if m==1 & n==1
 bs=bs*ones(size(s)); % expand bs
elseif m~=size(s,1) & n~=size(s,2),
  error('bs must be scalar or of same size as s');
end
r=2*(1+cos(s));
x(:)=r.*cos(s);
y(:)=r.*sin(s);
```

You can test the function by typing:

```
clear fem
fem.geom = 'cardg'
geomplot(fem), axis equal
fem.mesh = meshinit(fem);
meshplot(fem), axis equal
```

Then solve the PDE problem $-\Delta u = 1$ on the geometry defined by the cardioid. Use Dirichlet boundary conditions u = 0 on $\partial \Omega$. Finally plot the solution.

```
fem.equ.c = 1;
fem.equ.f = 1;
fem.bnd.h = 1;
fem.xmesh = meshextend(fem);
fem.sol = femstatic(fem);
postsurf(fem,'u')
The Geometry M-file format is not supported in 3D.
```

In 2D, each subdomain boundary must consist of at least two edge segments.

See Also geom0, geom1, geom2, geom3, geominfo, meshinit, meshrefine

Cautionary

Purpose	Retrieves coordinates for a work plane.
Syntax	<pre>[p_wrkpln,localsys] = geomgetwrkpln(type,args)</pre>
Description	<pre>[p_wrkpln,localsys] = geomgetwrkpln(type,args) returns the coordinate matrix p_wrkpln, spanning a plane, and the local coordinate system localsys of that plane, according to the string type and the cell array args.</pre>
	The columns of the 3-by-3 coordinate matrix p_wrkpln contain point coordinates for 3 non co-linear points spanning a work plane with a local z direction defined by lz = cross(p(:,2)-p(:,1),(p(:,3)-p(:,1)). The vector $lx =p(:,2)-p(:,1)$ is defined to be the local x-axis, and the local y-axis is defined as $ly = cross(lz,p(:,2)-p(:,1))$. The corresponding normalized unit vectors are nlx, nly , and nlz , respectively.
	The local coordinate system localsys is formed as localsys = $[n(: 1), n]_X$, $n]_X$, $n[X]$,

[p(:,1),nlx,nly,nlz], where p(:,1) is the position of the local origin and nlx, nly, and nlz specifies unit vectors in the direction of the positive local coordinate axes.

Work Plane of Type Explicit

```
[p_wrkpln,localsys] = geomgetwrkpln('explicit',{p_wrkpln1})
```

copies the coordinates p_wrkpln1 and forms the corresponding local coordinate system.

Work Plane of Type Quick

[p_wrkpln,localsys] = geomgetwrkpln('quick', {coordplane,offset})
forms a work plane parallel to the coordinate plane defined by coordplane, which
can have any of the values xy, yz, or zx. The real scalar offset specifies the signed
offset from the coordinate plane.

Work Plane of Type FaceParallel

```
[p_wrkpln,localsys] =
    geomgetwrkpln('faceparallel',{g,fn,dir,offset})
```

creates a work plane, parallel to face fn in 3D geometry object g. The direction dir takes the values +1 or -1 and specifies if the local z-axis, localsys(:,4), should be in the direction of the face's normal, or reversed normal, respectively. The scalar offset specifies the displacement along the local z-axis for the work plane with respect to the face. The face, with number fn, must be planar. If the face is not planar, within the system tolerance, an error message occurs.

Work Plane of Type EdgeAngle

```
[p_wrkpln,localsys] =
    geomgetwrkpln('edgeangle',{g,en,angle,fn,dir})
```

creates a work plane rotated angle radians about the edge en in the 3D geometry object g. The zero-angle is defined by the tangent plane of the face with face number fn. The face fn must be adjacent to the edge en and the face must have a single tangent plane common to all points of the edge en. The direction dir takes the values +1 or -1 and specifies if the rotation should be in positive or negative direction, with respect to the direction of the edge en, respectively. The matrix p_wrkpln, and the system localsys, referred to above, are formed based on the coordinate system induced by the selected edge, where the edge becomes the positive *x*-axis.

Work Plane of Type Vertices

```
[p_wrkpln,localsys] =
    geomgetwrkpln('vertices',{gl,vn,dir,offset})
```

creates a work plane spanned by the three vertices vn in the 3D geometry objects gl. vn is an index vector of length 3, corresponding to the entries in the cell array gl. The direction dir takes the values +1 or -1 and specifies if the local z-axis, localsys(:,4), should be in the direction of the positive normal, or reversed normal, respectively. The positive normal is defined as the cross product of the vectors in the direction from vn(1) to vn(2), and from vn(1) to vn(3), respectively. The scalar offset specifies the displacement along the local z-axis for the work plane with respect to the plane containing the vertices vn. The matrix p_wrkpln, and the system localsys, referred to above, are formed based on the chosen vertices such that, for offset = 0, the vector from vn(1) to vn(2) forms the local x-axis. The local y-axis is formed based on this vector and the local z-axis, as to produce a right-handed local coordinate system.

See Also

geomposition

Purpose	Groups geometry objects into an assembly.		
Syntax	<pre>[g,] = geomgroup(g1,) [g,] = geomgroup(draw,) [g,] = geomgroup(fem,)</pre>		
Description	[g,] = geomgroup(gl,) creates an assembly object g from the geometry objects in the cell array gl.		
	[g,] = geomgroup(draw,) creates an assembly object g from the geometry objects in the draw struct draw.		
	[g,] = geomgroup(fem,) creates an assembly object g from the geometry objects in the draw struct fem.draw.		
	Note that the parts of assembly g are not identical to gl. They are canonized and may have additional domains due to imprints.		
	The function supports the following properties:		
	TABLE I-69: VALID PROPERTY/VALUE PAIRS		

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Imprint	on off	on	Make imprints, when creating pair information
Ns	cell array of strings	{}	Name space
Out	cell array of strings	{'g'}	Cell array of strings: g, gt, st, ft, ct, pt, stx, ftx, ctx, ptx, pairs
Paircand	all none cell array of strings	all	Specifies the geometries which are used to create the pair information
Repairtol	positive scalar	0.0	Tolerance used for repairing gaps

Note: If the first syntax from above is used and the property paircand is a cell array of strings, also the property ns has to be specified.

The output pairs is a cell array of sparse matrices containing the pair information of the operation. Element i contains information for pairs of domains of dimension i-1. The column refers to the source and the row to the destination.

The output gt is a sparse matrix that relates the parts in the assembly to the original geometry objects. If an object has not been modified during the operation the value in gt is 1, otherwise 2.

Examples	<pre>[g pairs] = geomgroup({rect2 move(rect2,[1 0.5])},'out', {'g' 'pairs'}); [gg,stx,ctx,ptx] = getparts(g,'out',{'stx','ctx','ptx'})</pre>
See Also	geomcsg, geomanalyze, getparts

Purpose

Syntax

Import geometry objects from file.

gl = geomimport(filename,...)

Description

gl = geomimport(filename,...) reads the geometry file filename and translates the geometry data using the specified properties into a cell array of geometries gl.

filename can be any of the following formats:

FILE FORMAT	NOTE	FILE EXTENSIONS
Autodesk Inventor	2	.ipt
CATIA V4	2	.model
CATIA V5	2	.CATPart, .CATProduct
COMSOL Multiphysics Binary		.mphbin
COMSOL Multiphysics Text		.mphtxt
DXF		.dxf
GDS		.gds
IGES	I	.igs,.igse
Parasolid	I	.x_t, .x_b
Pro/ENGINEER	2	.prt,.asm
SAT	I	.sat,.sab
STEP	I	.step, .stp
STL		.stl
VDA	2	.vda
VRML		.wrl,.vrml

TABLE I-70: SUPPORTED FILE FORMATS

Note 1: Format requires a license of the CAD Import Module.

Note 2: Format requires a license of a format-specific module.

The following properties are supported

TABLE I-71: VALID PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Coercion	solid face off	solid	Coerce the imported geometry
Keepbnd	on off	on	Keep boundary entities

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Keepfree	on off	off	Keep free edge/point entities
Keepsolid	on off	on	Keep solid entities
Repair	on off	on	Repair imported data
Repairtol	positive scalar	1e-4	Repair tolerance
Importtol	positive scalar	1e-5	Absolute tolerance used when importing CAD Import Module formats.
Report	on off	on	Display a progress window

TABLE I-71: VALID PROPERTY/VALUE PAIRS

When importing COMSOL Multiphysics files and those in the GDS format, the command ignores these properties. See gdsread for GDS specific properties.

For DXF import, the default of Coercion is curve.

For STL and VRML import all properties in meshenrich are also supported.

Coercion can force the import process to knit boundary segments together and possibly try to form solid entities.

Keepbnd, Keepfree, and Keepsolid indicate which type of entities to be considered in the imported data.

Repair determines if the program should process the imported data to improve the quality during the import. These operations include snapping of points, removal of small entities, and improvement of geometric data.

Repairtol is a relative tolerance. It indicates the size of entities to remove, which points to snap together, and similar features.

Importtol is an absolute tolerance. It indicates the size of entities to remove, which points to snap together, and similar features. It is used when importing a file using the CAD Import module, replacing Repairtol.

Report determines if a progress window should appear during the call.

Examples filepath = which('demo1.dxf');
% Importing the DXF file demo1.dxf as a curve2 object.
g = geomimport(filepath);
geomplot(g{1})

Diagnostics	geomimport replaces the functionality of the COMSOL Multiphysics 3.0 functions
	dxfread, igesread, stlread, and vrmlread. We no longer support those
	functions and their properties and thus do not document them.

See Also geom0, geom1, geom2, geom3, meshimport, geomexport

Purpose	Retrieve geometry information.					
Syntax	[xx,]	<pre>[xx,] = geominfo(geom,'Out', {'xx'},)</pre>				
Description	<pre>[xx,] = geominfo(geom, 'Out', {'xx',},) retrieves geometry information specified in property Out from the analyzed geometry geom.</pre>					
	geom is an analyzed geometry, which is a geometry object, a mesh object or a Geometry M-file. The two latter formats are not supported in 3D. The Decomposed Geometry matrix of the PDE Toolbox is supported as well, but this alternative may be eliminated in future releases. For details on analyzed geometries, see the chapter "Geometry Modeling and CAD Tools" on page 23 in the <i>COMSOL</i> <i>Multiphysics User's Guide</i> or the entries geomesg, geomfile, and meshinit in this manual.					
	In the following description, a <i>geometric entity</i> refers to a <i>vertex</i> , an <i>edge segment</i> , a <i>face segment</i> , or a <i>subdomain</i> .					
	• If geom is a 1D analyzed geometry, the geometric entities are vertices and 1D subdomains that are bounded by vertices.					
	 If geom is a 2D analyzed geometry, the geometric entities are vertices, 1D edg segments bounded by vertices, that are assumed to be smooth in the interior, the is, sufficiently differentiable, and 2D subdomains bounded by edge segments. If geom is a 3D analyzed geometry, the geometric entities are vertices, 1D edg segments bounded by vertices, 2D smooth face segments bounded by edges, an 3D subdomains bounded by face segments. 					
	The functio	n geom	info a	ccepts	the following property/values:	
	TABLE I-72: VA	LID PROP	ERTY/VA	LUE PAIR	S.	
	PROPERTY	ID	2D	3D	DESCRIPTION	
	Od $\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{$					
	Odp	\checkmark	\checkmark	\checkmark	Matrix with columns that contain geometric entity dimension number pairs	
	Out	\checkmark	\checkmark	\checkmark	Output arguments, cell array containing strings specifying the output arguments	
	Par	\checkmark	\checkmark	\checkmark	Cell array, where each element is a cell array containing two matrices defining geometric	

Out specifies the geometry information to retrieve and return as output arguments. It is a cell array that can contain string equal to the entries given in Table 1-73.

entity number and parameter values

Par is a cell array containing parameter values and corresponding numbers of geometric entities. Par{m} is a cell containing the matrix Bm and the matrix Sm. Bm is of size nm1-by-nm2, and gives the numbers of entities of dimension d for which the parameters, in the nm1-by-nm2-by-d array Sm, are valid. Bm can also be a scalar or a vector, and as such it is expanded to the size of Sm. Note that the size of the third dimension in Sm, that is, d, defines if the entity number refers to a vertex (d = 0 or Par{m} = {Bm}), an edge (d = 1), or a face (d = 2).

The following table lists the valid outputs to geominfo.

TABLE I-73: OUTPUT ARGUMENTS

ουτ	ID	2D	3D	DESCRIPTION	INPUT PROPERTY
gd	\checkmark	\checkmark	\checkmark	Geometry dimension	
no	\checkmark	\checkmark	\checkmark	Number of objects of the dimensions specified in Od	Od
adj	\checkmark	\checkmark	\checkmark	Adjacency relations of entities in Odp	Odp
хх	\checkmark	\checkmark	\checkmark	Coordinate information	Par
dx		\checkmark	\checkmark	First-order derivative information	Par
ddx		\checkmark	\checkmark	Second-order derivative information	Par
nor		\checkmark	\checkmark	Normal vector information	Par
ff1			\checkmark	First fundamental matrices	Par
ff2			\checkmark	Second fundamental matrices	Par
crv		\checkmark	\checkmark	Curvature information	Par
rng		\checkmark	\checkmark	Parameter range of geometric entities	Od
ud	\checkmark	\checkmark	\checkmark	Up and down subdomains	
se	\checkmark	\checkmark	\checkmark	Start and end vertices of all ID primitive objects	
nmr	\checkmark	\checkmark	\checkmark	Number of subdomains	
nbs	\checkmark	\checkmark	\checkmark	Number of boundary segments	
mp		\checkmark		Coordinates of vertices	
sd				Vertex subdomain numbers	

no is a vector of the same size as Od, containing the number of primitive objects of the dimension as specified in Od.

adj is a cell array of adjacency matrices, where $adj\{k\}$ corresponds to Odp(:,k), and is a sparse matrix where $abs(sign(adj\{k\}(i,j))) = 1$ iff object i of dimension Odp(1,k) is adjacent to object j of dimension Odp(2,k). If the relation Odp(1,k) and Odp(2,k) can be given an orientation, the matrix entries +1 and -1 denotes positive or negative orientation, respectively. If both oriented and non orientable relations exist, -1, +1, and +2 are used, where +2 indicates a non oriented relation. If Odp is a vector of length 2, then adj is a sparse matrix. For subdomain information, the 0-domain is represented as output domain number 1. Thus, there is always an offset of 1 for subdomains.

xx is a cell array of same size as Par containing coordinate information, where $xx\{m\}$ is an nm1-by-nm2-by-gd array, where gd is the geometry dimension, and nm1 and nm2 are given from the size of Par{m}{2}. If the outer curly brackets in Par are not present, then xx is an n1-by-n2-by-gd array.

dx is a cell array of same size as Par containing first order derivative information for edges or faces. For edges, the dx{m} has the same format as $xx{m}$ above. For faces dx{m} is a nm1-by-nm2-by-3-by-2 array, where the last dimension refers to the two vectors, formed by the derivatives of u and v respectively, spanning the tangent plane.

ddx is a cell array of same size as Par containing second order derivative information for edges or faces. For edges, ddx{m} has the same format as $xx{m}$. For faces ddx is a nm1-by-nm2-by-3-by-2-by-2 array, where the last two dimensions refer to the 2-by-2 matrix of second order derivatives in the parameters u and v.

nor is a cell array of same size as Par, where the contents are the normalized normal vectors. They are given on the same format as the contents in xx.

ff1 is a cell array of same size as Par containing the first fundamental matrices of faces, where ff1{m} is an array of size nm1-by-nm2-by-2-by-2. For a parameter point given by the indices im1 and im2, the first fundamental matrix is given by GG = reshape(ff1{m}(im1,im2,:,:),2,2) and the corresponding Jacobian is given by J = reshape(dx{m}(im1,im2,:,:),3,2). It then holds that GG = J'*J.

ff2 is a cell array of same size as Par containing the second fundamental matrices
of faces, where ff2{m} is an array of size nm1-by-nm2-by-2-by-2. For a parameter
point given by the indices im1 and im2, the second fundamental matrix is given by
DD = reshape(ff2{m}(im1,im2,:,:),2,2). If the corresponding normal
derivative DNN and Jacobian J are obtained as above, then DD = -DNN'*J.

crv is a cell array of same size as Par containing curvature information of edges and faces. crv{m} is of size nm1-by-nm2-by-2 in 3D, where for a parameter point defined by the indices im1 and im2, crv{m}(im1,im2,1) is the curvature and crv{m}(im1,im2,2) is the torsion, when referring to an edge. The corresponding values obtained for a face is the Gaussian curvature and the mean curvature, respectively. In 2D, crv{m} is of size nm1-by-nm2 where crv{m}(im1,im2) contains the curvature of an edge for a given parameter.

rng is a cell array of same length as 0d, containing parameter range information for edges or faces. For edges, the first row in a matrix corresponds to the starting parameter value at the starting point, and the second row corresponds to the end parameter value at the end point. For faces, the first and third row contains the lower bounds on parameter values for the u and v parameters respectively. The second and fourth row contains the upper bounds on parameter values for the u and v parameter values for the u and v parameters respectively. The second and fourth row contains the upper bounds on parameter values for the u and v parameters respectively. The range for geometry edges is from zero to the arc-length of each edge. If no 0d is specified, rng is a matrix of range information for all edge curves, in 2D, or all faces, in 3D.

ud is a matrix containing up (left) and down (right) subdomain numbering for boundary segments, in the first and second row, respectively. One column of ud corresponds to one boundary segment.

sd is a vector containing the subdomain numbering of the vertices of mp. If a vertex is adjacent to more than one subdomain, the contents are NaN.

There is a family of low-level geometry functions used by geominfo, for obtaining the geometric data described above. These can be called directly, which in some cases can be preferred. Their names and descriptions are given in the table below.

FUNCTION	DESCRIPTION
flgeomadj	Get geometry adjacency matrices
flgeomec	Get curvature information from curve derivatives
flgeomed	Get coordinates and derivatives for geometry edges
flgeomes	Get parameter space size of geometry edge
flgeomfc	Get curvature from fundamental forms
flgeomfd	Get coordinates and derivatives for geometry faces
flgeomff1	Get first fundamental form from derivatives
flgeomff2	Get second fundamental form from derivatives
flgeomfn	Get normals from face derivatives

TABLE I-74: LOW-LEVEL GEOMETRY FUNCTIONS

FUNCTION	DESCRIPTION
flgeomfs	Get parameter space size of geometry face
flgeomnbs	Get number of geometry boundary segments
flgeomnes	Get number of geometry edge segments
flgeomnmr	Get number of subdomains
flgeomnv	Get number of vertices
flgeomsdim	Get space-dimension of geometry object
flgeomse	Get end-point indices of geometry edges
flgeomud	Get up-down subdomain numbering of geometry faces
flgeomvtx	Get coordinates for geometry vertices

TABLE I-74: LOW-LEVEL GEOMETRY FUNCTIONS

For details on the syntaxes for calling these functions, write help followed by the function name on the command line.

Examples

3D Geometries

To demonstrate the geominfo command, create a solid block object with a circular curve object on top, using the following commands.

```
g3 = geomcsg({block3},{},...
{move(embed(circ1(0.3,'pos',[0.5 0.5])),[0 0 1])})
geomplot(g3,'facelabels','on')
```

The generated object g3 is a solid 3D object consisting of 1 subdomain, 7 faces, 16 edges and 12 vertices. These can be obtained using geominfo with the arguments given below.

From the arguments gd and no, it is clear that g3 is a 3D object with the number of entities as above. The number of faces is also given in nbs, that is, the number of boundary segments. The parameter range of both faces and edges are given in rng. These are of importance when setting up parameter arrays for edge/face information evaluation below.

The following commands set up parameter matrices in two different formats, for faces 4, 5, and 7. The parameter range of these faces is 0 < u < 0.5, 0 < v < 0.5, as given by rng{3}(:,[4 5 7]).

[u,v] = meshgrid(0:0.1:0.5,0:0.1:0.5); S1 = reshape([u(:) v(:)],1,36,2); B1 = 7; S2(1,:,:) = deal([u(:) v(:)]); S2(2,:,:) = deal([u(:) v(:)]); B2 = [4;5];

Appropriate parameter values for the bounding edges of face 7, can be obtained by first creating the face-edge adjacency matrix with geominfo, and then using this information together with the argument rng to set up the parameter vectors. This is done with the following commands.

```
adj = geominfo(g3,'out','adj','odp',[1;2]);
B3 = find(adj{1}(:,7));
for i=1:length(B3)
    S3(i,:,1) = linspace(rng{2}(1,B3(i)),rng{2}(2,B3(i)),10);
end
```

Now, coordinate values of the faces and edges given above, together with coordinates for vertex 3, are obtained as follows.

```
[xx] = geominfo(g3,'out',{'xx'},...
'par',{{B1 S1} {B2 S2} {B3 S3} {3}})
```

To see the obtained results, simply give the following commands.

```
hold on
plot3(xx{1}(:,:,1),xx{1}(:,:,2),xx{1}(:,:,3),'r.')
plot3(xx{3}(:,:,1),xx{3}(:,:,2),xx{3}(:,:,3),'b.')
```

Finally, derivatives and curvatures, for both faces and edges are with the command below. Note that, both curvature measures for all points at face 7 are 0, as is the torsion for the surrounding curves. The curvature of these curves is however nonzero.

```
[dx,crv] = geominfo(g3,'out',{'dx' 'crv'},...
'par',{{B1 S1} {B3 S3}})
```

2D Geometries

Create a solid ellipse, and retrieve coordinates and curvatures for all four edge segments, by the following commands.

```
e = ellip2(0,0,1,2)
[xy,c] = geominfo(e,'out',{'xx','crv'},'par',...
{ones(11,1)*[1 2 3 4],(0:0.1:1)'*ones(1,4)})
```

Plot the obtained coordinates of the ellipse by the command.

plot(xy(:,:,1),xy(:,:,2),'b-')

The curvature of one of the edges are obtained via

figure,plot(0:0.1:1,c(:,1))

The command below retrieves the number of primitive objects (vertices, edges and subdomains) from geometry file (geomfile) cardg:

```
no = geominfo('cardg','out',{'no'},'od',[0 1 2])
```

ID Geometries

A 1D geometry consisting of two subdomains, is created by the following command.

```
g1 = solid1([-1 \ 0.2 \ 1])
```

Since no parameter domain exist only coordinates of vertices can be retrieved. The up and down subdomain of every vertex is given in ud, and the vertex-subdomain adjacency information is given in adj.

Note that the same information is given in ud and adj. The matrix adj is directly obtained from ud via the command:

```
adj = sparse(repmat(1:3,2,1),ud+1,[ones(1,3);-1*ones(1,3)])
```

Compatibility The FEMLAB 2.3 function flgeomepol is obsolete.

See Also geomcsg, geomedit, meshinit

Purpose	Create geometry object.
Syntax	obj = geomobject(input)
Description	<pre>obj = geomobject(input) creates a geometry object from input.</pre>
	input can be any of the following
	• A geometry object. See geom0, geom1, geom2, geom3.
	• A mesh object. See femmesh.
	• A geometry M-file name. See geomfile.
	• A PDE Toolbox geometry description matrix. See pde2geom.
	Note that in 3D, input cannot be a Geometry M-file.
See Also	geomO, geom1, geom2, geom3, femmesh, geomfile, pde2geom

Purpose	Plot geometry.
Syntax	<pre>geomplot(fem,) geomplot(geom,) h = geomplot(fem,) h = geomplot(geom,)</pre>
Description	<pre>geomplot(fem) plots the analyzed geometry fem.geom. For an extended FEM structure, xfem.fem{geomnum}.geom is plotted, where geomnum is 1 by default.</pre>
	geomplot(geom) plots the analyzed geometry geom.
	h = geomplot() additionally returns handles to the plotted axes objects.
	The <i>analyzed geometry</i> can be any of the following geometry representations: a geometry object, a Geometry M-file, or a mesh. The geometry object and Geometry M-file are described in the entries geomesg and geomfile, respectively. The mesh data structure is described in the entry meshinit.
	In 3D, the default plot is a patch plot of the faces with the edge segments and isolated vertices plotted as lines and markers respectively. The face, edge segment, and vertex parts of the plot can be controlled by the property/values starting with face, edge, and point respectively. Subdomains cannot be plotted directly, only indirectly through their adjacent faces.
	In 2D, the default plot is a patch plot of the subdomains with the edge segments and vertices plotted as lines and markers, respectively. The subdomain, edge segment, and vertex parts of the plot can be controlled by the properties starting with sub, edge, and point, respectively. You can turn on indication of curve parameter direction by using the property edgearrows.
	In 1D, the default plot is a line plot of the subdomains with vertices plotted as markers. The subdomain and vertex parts of the plot can be controlled by the properties starting with sub and point, respectively.
	The following table shows the property/value pairs for the geomplot command. The interpretation of the properties in 1D, 2D and 3D varies with dimension. The

design philosophy has been to keep property interpretation constant over space dimension, but to plot these properties as plot objects of different types.

TABLE I-75: VALID PROPERTY/VALUE PAIRS

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Boxcolor		\checkmark		color	k	Control polygon color
Boxstyle		\checkmark		line style		Control polygon line style
Ctrlmarker		\checkmark		marker symbol	0	Control polygon marker style
Ctrlmode		\checkmark		on off	off	Show control polygon
Detail		\checkmark	\checkmark	fine normal coarse	normal	Geometry resolution
Edgearrows		\checkmark		on off	off	Show edge directions with arrows
Edgecolor			\checkmark	color	k	Edge color data
Edgelabels		\checkmark	\checkmark	on off list of strings	off	Edge label list
Edgemode			\checkmark	on off	on	Show edges
Edgestyle			\checkmark	line style	-	Edge line style
Facelabels			\checkmark	on off list of strings	off	Face label list
Facemode			\checkmark	on off	on	Show faces
Labelcolor	\checkmark	\checkmark	\checkmark	color	k	Label color data
Linewidth		\checkmark	\checkmark	numeric	1	Line width
Linewidth	\checkmark			numeric	2	Line width
Markersize	\checkmark	\checkmark	\checkmark	numeric	6	Marker size
Mesh		\checkmark	\checkmark	mesh	new special mesh	Mesh used to render geometry
Pointcolor				color	b	Point color data

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Pointlabels	\checkmark	\checkmark	\checkmark	off on list of strings	off	Point label list
Pointmarker	\checkmark	\checkmark	\checkmark	marker symbol	0	Point marker
Pointmode	\checkmark	\checkmark	\checkmark	on off isolated	on	Show points
Sublabels	\checkmark	\checkmark	\checkmark	on off list of strings	off	Subdomain label list
Submode	\checkmark	\checkmark		on off	on	Show subdomains

TABLE 1-75: VALID PROPERTY/VALUE PAIRS

In addition, the common plotting properties listed under femplot are available.

The properties sublabels, facelabels, edgelabels, and pointlabels control the display of subdomain labels, face labels, edge segment labels, and point labels, respectively.

The properties that control marker type or coloring can handle any standard marker or color type in COMSOL Script. See for example the plot command in the *COMSOL Script Reference Guide*.

Examples 3D Example

Create a simple 3D geometry:

```
c1 = cylinder3(0.5,2,[-1,0,0],[1,0,0]);
c2 = cylinder3(0.2,2,[0,-1,0],[0,1,0]);
g = c1-c2;
```

Plot edges and face labels.

```
geomplot(g,'facemode','off','facelabels','on')
axis equal
```

Plot faces with lighting and without edges and axis in high quality.

```
geomplot(g,'edgemode','off','detail','fine')
light, lighting phong
axis equal, axis off
```

Both faces and edges are plotted by default.

2D Example

Start by creating a simple geometry.

```
clear fem
c1 = circ2;
l1 = curve2([-1,-1,1,1],[-1,1,-1,1]);
p1 = point2(0,0.5);
fem.draw.s.objs = {c1};
fem.draw.c.objs = {l1};
fem.draw.p.objs = {p1};
fem.geom = geomcsg(fem);
```

Plot the standard geometry plot with subdomains indicated as patches, and edge segments and vertices indicated by lines and markers, respectively.

```
geomplot(fem), axis equal
```

Remove patch plot of subdomains, add parameter direction for curves, subdomain numbers, and control polygons.

```
geomplot(fem,'submode','off','edgearrow','on','pointmode',...
'isolated','sublabels','on','ctrlmode','on')
```

```
ID Example
```

Start by creating a simple geometry.

```
clear fem
s1 = solid1([0 0.1 1]);
p1 = point1(2);
fem.draw.s.objs = {s1};
fem.draw.p.objs = {p1};
fem.geom = geomcsg(fem);
```

The standard geometry plot with subdomains indicated as lines, and vertices indicated by markers.

geomplot(fem), axis equal

Change the color of the vertices to red, add vertex labeling, and change the vertex markers to diamonds.

Compatibility The properties pt, ct, ft, and st have been removed in FEMLAB 3.1.

CautionaryThe value numeric of the sublabels, edgelabels, and pointlabels properties
was replaced by on in FEMLAB 1.1. The value numeric is still supported however,
and is equivalent to on.

The default for the ctrlmode property was changed to off in FEMLAB 1.1.

See Also

geomcsg, geomedit

Purpose	Position 3D geometry object in space using work plane info.
Syntax	g3 = geomposition(g32,p_wrkpln)
Description	g3 = geomposition(g32,p_wrkpln) positions the 3D geometry object g32 in space by transforming the point matrix according to the work plane information in p_wrkpln. The geometry g32 is thus assumed to be defined in the local coordinate system of the work plane.
	See geomgetwrkpln for more information on work planes and the work plane points representation p_wrkpln.
See Also	geomgetwrkpln

Purpose	Spline interpolati	Spline interpolation.			
Syntax	c = geomspline	<pre>c = geomspline(p,)</pre>			
Description	- ·	c = geomspline(p,) creates a curve2 or curve3 object from point data p by spline interpolation. The object generated is a closed or open, C^1 or C^2 continuous, spline.			
		omspline accepts th	,	specifies interpolation points. erty/value pairs:	
	PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION	
	Closed	auto on off	auto	Closed or open curve	
	SplineDir	2-by-np matrix 3-by-np matrix		Tangent vectors for the corresponding points in p	
	SplineMethod	uniform chordlength centripetal foley	chordlength	Method for global parameterization	

The property SplineDir is used to specify a tangent vector for the corresponding point in p. This means that the first control point is given and the curve thus generated is only guaranteed to be C^1 (continuous first derivatives). If this property is not given, however, the curve generated is guaranteed to be C^2 (continuous second derivatives). The SplineMethod property does not affect the curve if the SplineDir property is used.

The property SplineMethod controls the method for how to compute the global parameterization of the curve. The global parameterization is a parameter that varies from 0 to 1, from the first interpolated point to the last. For a closed curve the last point is equivalent to the first. The value uniform means that the global parameterization is [0, 1,..., np]/np. The default value chordlength means that the global parameterization is [0, norm(p(:,2)-p(:,1)), norm(p(:,3)-p(:,2)),..., norm(p(:,np)-p(:,np-1))]/ sum(sqrt(sum((diff(p')').^2)')), where the denominator is the total chord length. The values centripetal and foley are two additional methods that handle irregular point sets p more effectively.

The property Closed controls the closure of the spline. If Closed is on the first point is regarded as the last point. The value auto for the property Closed generates a closed curve whenever the first and last points in a scaled version of the point set

	p agree to within 1000*eps in Euclidean distance. Otherwise, an open curve is generated.
	On success, c is a curve2, or curve3 object that passes through the points defined by p. If p does not define a spline curve properly, either an error occurs or a line1, curve2, curve3, or circ1 object is created that meets the requirements in some way.
Example	<pre>% Interpolate irregularly distributed point on a circle. % First create circle data phi=0:0.2:2*pi; phi(end)=[]; % Remove some of the points. phi([1 3 6 7 10 20 21 25 28])=[]; p=[cos(phi);sin(phi)]; % Add some noise. randn('state',17) p=p+0.05*randn(size(p)); plot(p(1,:),p(2,:),'r.') % Interpolate using uniform parameterization. c=geomspline(p,'splinemethod','uniform','closed','on') hold on geomplot(c,'pointmode','off') % Interpolate using centripetal parameterization. c=geomspline(p,'splinemethod','centripetal','closed','on') hold on geomplot(c,'pointmode','off','edgecolor','b') axis equal</pre>
See Also	curve2, curve3

Purpose	Create 3D geometry surface using height data defined on a grid.
Syntax	<pre>f = geomsurf(x,y,z) f = geomsurf(z) s = geomsurf(x,y)</pre>
Description	f = geomsurf(x,y,z) creates a 3D face object f on the grid defined by x and y, using z as height data. The surface created is identical in shape to the surface created with the scripting command surf(x,y,z).
	geomsurf uses piecewise bilinear interpolation, as does surf.
	f = geomsurf(z) creates a 3D face object f , corresponding to the call $surf(z)$.
	s = geomsurf(x,y) creates a 2D solid object s corresponding to the syntax $f = geomsurf(x,y,z)$ with z all zeros.
Example	<pre>% Create randomly generated surface % Create rectangular grid [x,y]=meshgrid(-0.1:0.2:1.1,-0.4:0.2:0.4); % Initialize random generator randn('state',1); % Create random height data z=0.1*randn(size(x)); % Create 3D surface f=geomsurf(x,y,z); % Plot the surface geomplot(f)</pre>
	<pre>% Create approximation to a catenoidal surface % Create grid in spherical coordinates [theta,phi]=meshgrid(pi/8:pi/32:3*pi/8,pi/4:pi/32:pi/2); % The conical surface is expressed in spherical coordinates r=1; x=r.*cos(theta)./sin(phi); y=r.*sin(theta)./sin(phi); z=r.*log((1+sin(phi))./sin(phi)); % Now, create the piecewise bilinear % approximative surface catenoid=geomsurf(x,y,z); % Plot the surface geomplot(catenoid) axis equal</pre>
See Also	face3, meshgrid

Purpose	Extract parts from an assembly object.					
Syntax	[gl,] =	[gl,] = getparts(g,)				
Description	$[gl, \ldots] = getparts(g, \ldots)$ returns a cell array where each element contains a part.					
	The function supports the following properties:					
	PROPERTY PROPERTY VALUE DEFAULT DESCRIPTION					
	Out	cell array of strings	{}	Cell array of strings: stx, ftx, ctx, ptx		
	Part	all none vector of integers	all	Specifies which parts to extract		
Example	g = geomgroup({rect2 move(rect2,[1 0])}); [gg,stx,ctx,ptx] = getparts(g,'out',{'stx','ctx','ptx'});					
See Also	geomgroup, geomcsg					

Purpose	Create helix geometry object.
Syntax	h1 = helix1(r,dh,h) h2 = helix2(dr,r,dh,h,n) h3 = helix3(dr,r,dh,h,n)
Description	h1 = helix1(r,dh,h) creates a helix-shaped curve3 object with radius r, distance between consecutive turns dh, and total height h. The helix is centered at the origin with main axis in the z direction. All arguments are optional; when arguments are omitted, the following default values are used: $r = 1.0$, $dh = 1.0$, and $h = 1.0$.
	h2 = helix2(dr, r, dh, h, n) creates a helix-shaped face3 object with cross-section radius dr, radius r, distance between consecutive turns dh, total height h, and resolution n. The resolution n is an integer that specifies the number of curved sections for every turn; a higher resolution yields a smoother-looking helix. The helix is centered at the origin with main axis in the z direction. All arguments are optional; when arguments are omitted, the following default values are used: dr = 0.1, r = 1.0, dh = 1.0, h = 1.0, and n = 12.
	h3 = helix3(dr,r,dh,h,n) creates a helix-shaped solid3 object with cross-section radius dr, radius r, distance between turns dh, total height h, and resolution n. The resolution n is an integer that specifies the number of curved sections for every turn; a higher resolution yields a smoother-looking helix. The helix is centered at the origin with main axis in the z direction. All arguments are optional; when arguments are omitted, the following default values are used: dr = 0.1, r = 1.0, dh = 1.0, h = 1.0, and n = 12.
Example	The following command generates a solid helix-shaped object:
	h3 = helix3(1,5,1,5,12);
See Also	extrude, loft, revolve

Purpose	Create bilinear hexahedron geometry object.			
Syntax	h2 = hexahedron2(p) h3 = hexahedron3(p)			
Description	h3 = hexahedron3(p) creates a solid hexahedron object with corners in the 3D coordinates given by the eight columns of p. hexahedron3 is a subclass of solid3.			
	h2 = hexahedron2(p) creates a surface hexahedron object with corners in the 3D coordinates given by the eight columns of p. hexahedron3 is a subclass of face3.			
	For a hexahedron approximately aligned to the coordinate planes, the points in p are ordered as follows. The first four points and the last four points projected down to the (x,y) -plane defines two negatively oriented quadrangles. The corresponding plane for the second quadrangle must lie above the plane of the first quadrant in the z direction. Generally oriented hexahedra have the points of p ordered in a similar way, except for a rigid transformation of the defining point set.			
	The default value of p is			
	p=[0 0 1 1 0 0 1 1; 0 1 1 0 0 1 1 0; 0 0 0 0 1 1 1]			
	The 3D geometry object properties are available. The properties can be accessed using the syntax get(object,property). See geom3 for details.			
Example	The following command generates a solid hexahedron object.			
	h3 = hexahedron3([0 0 1 1 0 0 1 1; 0 0.8 1 0 0 1 1.2 0; 0 0.1 0 0.2 1 1 2 1]);			
See Also	face3,geom0, geom1, geom2, geom3			

Purpose	Create polygons.
Syntax	<pre>c = line1(x,y) s = line2(x,y)</pre>
Description	s = line2(x, y) creates a 2D solid object s in the form of a solid polygon with vertices given by the vectors x and y .
	c = line1(x, y) creates a 2D curve object c in the form of an open polygon with vertices given by the vectors x and y .
Examples	<pre>The commands below create an open regular n-gon (n=11) and plot it. n = 11 xy = exp(i*2*pi*linspace(0,1-1/n,n)); l = line1(real(xy),imag(xy)); geomplot(1)</pre>
See Also	arc1, arc2, circ1, circ2, ellip1, ellip2, geomcsg, poly1, poly2

Purpose	Loft 2D geometry sections to 3D geometry.
Syntax	g3 = loft(gl,)
Description	g3 = loft(g1,) lofts the 2D geometry sections in g1 to a 3D geometry object g3.
	gl is a cell array of size 1-by-ng of 2D geometry objects that belongs to one of the subclasses solid2 or curve2. That is, gl{i} contains the geometry object of

The function loft accepts the following property/value pairs:

TABLE I-78:	VALID	PROPERTY/VALUE PAIRS

section number i.

PROPERTY	VALUE	DESCRIPTION
LoftEdge	I-by-ng cell array of integer vectors	Permutation vectors for edges
LoftSgnEdge	I-by-ng cell array of integer vectors	Signed permutation vector for edges
LoftVtxPair	I-by-ng cell array of integer matrices with two rows	Permutation vector for vertex pairs
LoftSecPos	I-by-3 cell array	Positioning for 2D geometry sections
Wrkpln	I-by- <i>ng</i> cell array of 3-by-3 matrices	Work planes for 2D geometry sections
LoftWeights	Matrix of size 2-by-(ng-1)	Cubic lofting weights
LoftMethod	linear cubic	Lofting method

The properties LoftEdge, LoftSgnEdge, or LoftVtxPair are needed to make the connection between edges and vertices in different sections unique.

The property LoftEdge with the value $\{e1, e2, ...\}$ means that the edge with number e1(1) in $g1\{1\}$ should be lofted to match the edge with number e2(1) in $g1\{2\}$ and so on for all elements in e1 and e2.

Likewise, the property LoftSgnEdge with the value {e1,e2,...} means the same thing, except that edges with different directions is indicated by using negative signs. This is often more reliable than LoftEdge above.

The property LoftVtxPair is used in the same way, but uses pairs of vertices instead. Thus, LoftVtxPair with the value $\{v1, v2, ...\}$ means that the vertex with number v1(1,1) in gl{1} is to be matched with vertex number v2(1,1) in

gl{2} and so on. It is required that v1([1 2],1) are the end points on the same curve.

Only one of these properties is allowed. If, however, none is specified then the property/value-pair LoftSgnEdge with the value {[1:nbs],[1:nbs],...} is used as default, where nbs is the number of edges in gl{1}. This means that the edges are considered in order, and is useful for lofting between sections that are simple or similar to each other.

The properties LoftSecPos or Wrkpln are used to specify the geometrical data of each section.

The property LoftSecPos with the value {D,V,R} has the following meaning:

D is either a 1-by-(ng-1) vector or a 3-by-ng matrix that specifies the position for each geometry section. If D is a vector, it contains real numbers that specifies the relative displacement in the local z direction between each pair of consecutive sections in g1, where it is assumed that g1{1} is positioned at z = 0. If D is a matrix, then each column specifies the 3D displacements for each of the sections in g1. Rows 1, 2, and 3 specifies the displacements in the x, y, and z direction, respectively.

V is either a 2-by-ng or a 3-by-ng matrix specifying the tilt-rotations of the geometry objects. If V is a 2-by-ng matrix, then each column specifies rotational angles in spherical coordinates. V(1,:) are the polar angles, that is, the angles between directional normals of each object and the positive *z*-axis, and V(2,:) are the azimuthal angles of the directional normals. If V is a 3-by-ng matrix, then each column specifies a directional normal vector for each section.

R is a 1-by-ng vector that specifies the intrinsic rotation of the geometry sections. Every element of R is a rotational angle, in radians, with respect to the local *z*-axis.

The alternative syntax is the property Wrkpln with the value {T1,T2,...}, where Ti is a matrix of size 3-by-3. Here Ti is understood to specify the work plane for section gl{i}. See geomgetwrkpln for more information on work planes.

Only one of these properties is allowed. If none is specified then the property LoftSecPos with the value {ones(1,ng-1),zeros(2,ng),zeros(1,ng)} is used as default. Moreover, if any of the cells D, V, or R is left empty, the default value is used for that cell.

The property LoftWeights specifies the relative significance of the geometry sections with respect to tangential continuity. This argument has no meaning for linear lofting and is then ignored.

The property LoftMethod can have the values linear or cubic, specifying if the lofting should be linear/ruled or bicubic, respectively. The default method is cubic, with the LoftWeights property set to the value [0.3*ones(1,ng-1); 0.7*ones(1,ng-1)].

Examples

Create a loft between a circle and a square.

gl1 = cell(1,2); gl1{1} = circ2; gl1{2} = rect2(2,2,'pos',[-1 -1]);

Let the edge between vertices number one and two in the circle correspond to the edge between vertex number one and two in the square.

```
tl1 = cell(1,2); tl1{1} = [1;2]; tl1{2} = [1;2];
g1 = loft(gl1,'LoftVtxPair',tl1);
figure, geomplot(g1)
```

Create a loft between edge number 1 in the circle and edge number 1 in the square.

tl2 = cell(1,2); tl2{1} = 1; tl2{2} = 1;

Also, rotate the square.

Create a more complicated example.

gl2 = cell(1,3); gl2{1} = arc2(0,0,2,0,pi/2)-circ2; gl2{2} = rect2(2,2,'pos',[-1 -1]); gl2{3} = gl2{1};

Specify reversed direction of edges by using negative signs.

tl3 = cell(1,3); tl3{1} = 4; tl3{2} = -2; tl3{3} = -3;

Also, rotate the last two sections.

```
g3 = loft(gl2,'LoftSgnEdge',tl3,'LoftSecPos',...
{[-1 -1 0; 0 0 1; 1 1 2]',zeros(2,3),[0 pi/4 pi]});
figure, geomplot(g3)
```

In the latter case, since there are no ambiguities, you could also use unsigned edge numbers by specifying the property LoftEdge.

See Also

extrude, geomgetwrkpln, revolve

Purpose Create an analyzed geometry and/or a draw object from a (deformed) mesh.

Syntax

Description

[xfem,g] = mesh2geom(xfem,args)

[xfem,g] = mesh2geom(xfem,args) returns a new extended fem structure xfem with the fields specified in the destfield property filled, generated from the source specified in the srcdata property. If draw is specified in destfield, the created draw object g is also returned.

The source can be either the deformed geometry from solving an ALE or parameterized geometry problem (deformed), or a mesh (mesh).

The destination can be any nonempty subset of {'draw', 'geom', 'mesh'}, indicating that a mesh, an analyzed geometry, and/or a draw object should be created. If draw is specified, you can use the drawtag property to specify the tag of the new Draw-mode object.

The destination geometry destfem can be an existing geometry or the next undefined geometry, in which case a new geometry is created.

TABLE I-79:	VALID	PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Srcdata	deformed mesh	mesh	Source data: deformed geometry or mesh
Destfield	cell array of strings: mesh geom draw	{'geom', 'mesh'}	Destination: mesh, analyzed geometry, or draw object
Srcfem	positive integer	1	Source geometry
Destfem	positive integer	1	Destination geometry
MCase	integer	0	Source and/or destination mesh case
Drawtag	string		Draw tag to use when creating draw object
Frame	string		Frame to use when retrieving deformed geometry
Solnum	positive integer	last solution	Solution to use when retrieving deformed geometry

Examples

Creating an Analyzed Geometry From a Mesh Create a mesh.

clear fem;

fem.mesh=meshinit(rect2);

Create an analyzed geometry from the mesh, into the new geometry Geom2.

```
xfem=mesh2geom(fem, 'destfem',2);
```

Creating an Analyzed Geometry From a Deformed Mesh Draw two rectangles, one inside the other, and mesh

```
clear fem
g1=rect2(1.8,1.2,'base','corner','pos',[-0.8,-0.8]);
g2=rect2(0.2,0.4,'base','corner','pos',[-0.2,-0.4]);
fem.geom=geomcsg({g1,g2});
fem.mesh=meshinit(fem);
```

Set the inner rectangle to move along the *x*-axis.

```
clear appl
appl.mode.class = 'MovingMesh';
appl.sdim = {'X', 'Y', 'Z'};
appl.assignsuffix = ' ale';
appl.prop.analysis='transient';
appl.prop.weakconstr.value = 'off';
appl.bnd.defflag = \{\{1;1\}\};
appl.bnd.deform = {{0;0},{'t';0}};
appl.bnd.ind = [1,1,1,2,2,2,2,1];
appl.equ.type = {'free','pres'};
appl.equ.presexpr = {{0;0},{'t';0}};
appl.equ.ind = [1,2];
fem.appl{1} = appl;
fem.sdim = {{'X', 'Y'},{'x', 'y'}};
fem.frame = {'ref', 'ale'};
fem=multiphysics(fem);
fem.xmesh=meshextend(fem);
```

Solve the problem:

fem.sol=femtime(fem, 'tlist',[0:0.01:0.1]);

Create an analyzed geometry from the deformed mesh.

```
fem=mesh2geom(fem,'srcdata','deformed','frame','ale');
```

Remesh the created geometry and continue solving.

```
fem.mesh=meshinit(fem);
fem.xmesh=meshextend(fem);
fem.sol=femtime(fem,'tlist',[0:0.01:0.1]);
```

Purpose	Create boundary layer mesh
Syntax	fem.mesh = meshbndlayer(fem,) fem.mesh = meshbndlayer(fem.geom,) fem = meshbndlayer(fem,'out',{'fem'},)
Description	fem.mesh = meshbndlayer(fem,) returns a boundary layer mesh derived from the geometry in fem.geom.
	fem.mesh = meshbndlayer(geom,) returns a boundary layer mesh derived from the geometry geom.
	fem = meshbndlayer(fem, 'Out', 'fem',) modifies the fem structure to include a boundary layer mesh in fem.mesh.
	A boundary layer mesh is a mesh with dense element distribution in the normal direction along specific boundaries. This type of mesh is typically used for fluid flow problems to resolve the thin boundary layers along the no-slip boundaries. In 2D, a layered quadrilateral mesh is used along the specified no-slip boundaries. In 3D, a layered prism mesh or a layered hexahedral mesh is used depending on if the corresponding boundary layer boundaries contain a triangular mesh or a quadrilateral mesh.

The boundary layer mesher inserts boundary layer elements into an existing mesh. If the starting mesh is empty the free mesher is automatically used to create a starting mesh.

Boundary layers are not allowed on isolated boundaries, that is, boundaries with the same subdomain on each side of the boundary.

The function meshbndlayer accepts the following property/value pairs.

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Blbnd	array	all exterior boundaries	Boundary layer boundaries
Blhmin	numeric cell array		Initial boundary layer thickness
Blhminfact	numeric cell array	1	Factor that the default Blhmin is multiplied by
Blnlayers	numeric cell array	8	Number of boundary layers

TABLE I-80: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Blstretch	numeric cell array	1.2	Boundary layer stretching factor
Hauto	numeric	5	Predefined mesh element size
Mcase	numeric	0	Mesh case number
Meshstart	mesh object	empty	Starting mesh
Out	fem mesh	mesh	Output variables
Report	on off	on	Display progress
Subdomain	numericarray auto all none	auto	Specifies the subdomains that are meshed

TABLE 1-80: VALID PROPERTY/VALUE PAIRS	TABLE I-80:	VALID PROPERTY/VALUE PAIRS
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meshbndlayer accepts all property/values that meshinit does. The meshinit command is used to create the starting mesh for the subdomains to be processed by the boundary layer mesher if these are not already meshed.

The property blbnd is an array specifying the boundaries for which boundary layers are created. By default boundary layers are created for all exterior boundaries.

Use the properties blhmin, blstretch, and blnlayers to specify the distribution of the boundary layers. The value of each of these properties is a scalar value for all boundaries or an even numbered cell array where the odd entries contain boundary indices, either as scalar values, or as vectors with boundary indices, and the even entries contain the corresponding parameters. blhmin specifies the thickness of the initial boundary layer, blstretch a stretching factor, and blnlayers the number of boundary layers. This means that the thickness of the *m*th boundary layer (m=1 to blnlayers) is blstretch^(m-1) blhmin. The default value of blhmin is 1/50 of the size of the elements for the corresponding boundary layer boundaries. Note that the number of boundary layers might be automatically reduced in thin regions.

It is also possible to specify the thickness of the initial layer by using the blhminfact property. If you use this property the initial layer thickness is defined as *blhminfact* * *blhmindef*, where *blhmindef* is the default value of the property blhmin. The value of this property is a scalar value for all boundaries or an even numbered cell array where the odd entries contain boundary indices, either as scalar values, or as vectors with boundary indices, and the even entries contain the corresponding parameters.

hauto is an integer between 1 and 9 that controls the element size in the starting mesh. The default value is 5. For more information on this property see meshinit.

The meshstart property is used when meshing a geometry interactively. The value of this property is the starting mesh of the meshing operation. If meshstart does not contain a mesh of the subdomains to be processed a starting mesh is automatically created using the meshinit command before inserting the boundary layer elements.

Use the property subdomain to specify the subdomains to be meshed. If you use this property together with the meshstart property, the value auto means that all subdomains that are not meshed in the starting mesh are meshed, none means that no further subdomains are meshed, and all means that all subdomains are meshed. It is also possible to specify the subdomains to be meshed using a vector of subdomain indices.

Examples

Specify the boundary layer boundaries and the number of boundary layers

```
fem.geom = rect2(10,5) - circ2(1,'pos',[3 2.5]);
fem.mesh = meshbndlayer(fem,'blbnd',[2:3 5:8],...
'blnlayers',{5:8 8});
figure, meshplot(fem)
```

Insert boundary layers to an existing mesh containing both quadrilateral elements and triangular elements.

```
fem.geom = rect2 + rect2(1,1,'pos',[1 0]) - circ2(1.5,0.5,0.2);
fem.mesh = meshmap(fem,'subdomain',1);
fem.mesh = meshinit(fem,'meshstart',fem.mesh);
figure, meshplot(fem)
fem.mesh = meshbndlayer(fem,'blbnd',[2:3 5:6 8:11],...
'meshstart',fem.mesh,...
'subdomain','all');
figure, meshplot(fem)
```

Create a boundary layer mesh consisting of prism elements along the boundary layer boundaries and tetrahedral elements in the interior

```
fem.geom = block3(10,5,5) - sphere3(1,'pos',[3 2.5 2.5]);
fem.mesh = meshbndlayer(fem,'blbnd',[2:13]);
figure, meshplot(fem)
figure, meshplot(fem,'ellogic','x<3')</pre>
```

See Also

meshinit, meshmap, meshsweep

Purpose	Add new mesh cases
Syntax	<pre>fem = meshcaseadd(fem,)</pre>
Description	fem = meshcaseadd(fem) adds one or several new mesh cases to the FEM structure FEM. The mesh cases are typically used as hierarchy in the Geometric multigrid solver. The new mesh cases are constructed by coarsening or refining the mesh (or keeping the same mesh), and possibly changing the order of the shape functions. If the order is changed, the integration point order and the constraint point order is changed accordingly.

The function meshcaseadd accepts the following property/value pairs:

PROPERTY	VALUES	DEFAULT	DESCRIPTION
Mcasekeep	array of nonnegative integers all	Mcaseorig	Mesh cases to keep
Mcaseorig	array of nonnegative integers	lowest existing mesh case	Original mesh case(s)
Meshscale	array of positive numbers	2	Scale factors for mesh size h
Mgauto	meshscale shape anyshape both meshrefine explicit		Method for generating mesh cases
Mggeom	array of positive integers	all	Geometry numbers
Nmcases	positive integer	1	Number of new mesh cases to generate
Report	on off	on	Display progress
Rmethod	regular longest	regular	Mesh refinement method
Shapechg	array of integers	-1	Change in shape function orders

TABLE I-81: VALID PROPERTY/VALUE PAIRS

The function meshcaseadd operates on the FEM structures corresponding to the geometries Mggeom. The FEM structures for other geometries are left unaffected.

Before creating new mesh cases, all existing mesh cases except Mcasekeep are deleted.

The mesh case generation method is determined by the property Mgauto. The new mesh cases will be given the numbers mcmax+1, ..., mcmax+n, where mcmax is the current highest mesh case number.

- If Mgauto=both, shape, anyshape, or meshscale, then the new mesh cases are constructed starting from the mesh case given in the property Mcaseorig (this should be a single nonnegative integer). This process is described in the section "Constructing a Multigrid Hierarchy" on page 521, where the methods are called Coarse mesh and lower order (both), Lower element order first (all) (shape), Lower element order first (any) (anyshape), and Coarse mesh (meshscale). The mesh coarsening factor is given in the scalar Meshscale, the shape function order change amount is given in the scalar Shapechg, and the number of new mesh cases to create is given by the property Nmcases.
- If Mgauto=explicit, then new mesh cases are constructed starting from the mesh case(s) given in the property Mcaseorig. The properties Mcaseorig, Meshscale, and Shapechg should be vectors of the same length n (however if one is scalar, it is expanded to the same length as the other). Mesh case mcmax+i will have a mesh that is coarsened with the factor Meshscale(i), and shape function orders incremented with Shapechg(i) relative to mesh case Mcaseorig(i).
- If Mgauto=meshrefine, then the new mesh cases are constructed by refining the mesh in mesh case Mcaseorig (this should be a single nonnegative integer) repeatedly. The number of new mesh cases to create is given by the property Nmcases. The refinement method can be specified using the property Rmethod, see meshrefine.

The default value of Mgauto is as follows. Let n be the length of the longest among the vectors Meshscale, Mcaseorig, and Shapechg. The default for Mgauto is shape if n = 1, and explicit if n > 1.

The following fields in the FEM structure are affected by meshcaseadd:

mesh, shape, gporder, ***.gporder, cporder, and ***.gporder, where *** is equ, bnd, edg, or pnt. Also, the corresponding fields in the app1 field are affected.

See Also femsolver, meshcasedel

Purpose	Delete mesh cases
Syntax	fem = meshcasedel(fem) fem = meshcasedel(fem,mcases)
Description	<pre>fem = meshcasedel(fem) deletes all mesh cases except 0 from the FEM structure fem.</pre>
	fem = meshcasedel(fem, mcases) deletes the mesh cases in the integer vector mcases from the FEM structure fem.
	The following fields in the FEM structure are affected by meshcasedel:
	mesh, shape, gporder, ***.gporder, cporder, and ***.gporder, where *** is equ, bnd, edg, or pnt. Also, the corresponding fields in the app1 field are affected.
See Also	meshcaseadd

Purpose	Copy mesh between boundaries
Syntax	<pre>fem.mesh = meshcopy(fem,) fem = meshcopy(fem,'out',{'fem'})</pre>
Description	fem.mesh = meshcopy(fem,) copies the mesh between boundaries in the mesh object fem.mesh.
	<pre>fem = meshcopy(fem, 'out', { 'fem'}) modifies the FEM structure to include the new mesh object in fem.mesh.</pre>
	Copy the mesh from one or several source boundaries to one target boundary. The

Copy the mesh from one or several source boundaries to one target boundary. The source boundary (or in the case of several source boundaries, the combined source boundaries) and the target boundary must be of the exact same shape. However, a scaling factor between the boundaries is allowed.

In 3D, the edges around the source and target boundaries are allowed to be partitioned differently, but only in such a way that several edges of the source boundary map to one edge of the target boundary, not the other way around.

The function meshcopy	accepts the following	property/value pairs:

PROPERTY	2D	3D	VALUE	DEFAULT	DESCRIPTION
Direction	\checkmark	V	auto same opposite	auto	Direction between edges
Mcase	\checkmark	\checkmark	integer	0	Mesh case number
Source	\checkmark	\checkmark	numeric array		Source boundaries
Sourceedg		\checkmark	numeric array		Source edges
Target	\checkmark	\checkmark	integer		Target boundary
Targetedg		\checkmark	integer		Target edge

Use the properties source and target to specify the source and target boundaries. Note that source can be either a scalar value or a vector. The property target is always a scalar value. This means that copying from several boundaries is allowed, but you can only copy the mesh to a single boundary.

In 3D, use the properties sourceedg, targetedg, and direction to specify the edge mapping from the source to the target boundary. The property sourceedg can be either a single edge index or a vector of edge indices. The property targetedg is always a single edge index. The property direction specifies the direction

	between sourceedg and targetedg. The possible values are same, opposite, and auto, where the last option means that the direction between the edges is automatically determined by the algorithm. If sourceedg is a vector, then direction refers to the direction between targetedg and the edge with the lowest edge index in sourceedg.
	In 2D, use the property direction to specify the direction between the edges given in the properties source and target. The properties sourceedg and targetedg are not used in 2D.
	If you do not specify how to orient the source mesh on the target boundary through the sourceedg, targetedg, and direction properties, the algorithm attempts to determine the orientation automatically.
	Copying a mesh is only possible if the target boundary is not adjacent to any meshed subdomain. If the target boundary is already meshed, the current mesh is first deleted and the source mesh is then copied to the target boundary.
	In 3D, copying a mesh to a target boundary that is adjacent to a meshed boundary is allowed if the edge between these boundaries has the same number of elements as the corresponding source edges. In this case, the mesh on the target edge is kept, and the copied boundary elements are modified to fit with this edge mesh.
Examples	Mesh Face 1 of a block and copy the mesh to the opposite Face 4.
	<pre>fem.geom = block3; fem.mesh = meshinit(fem,'point',[],'edge',[], 'face',1,'subdomain',[]); fem.mesh = meshcopy(fem,'source',1,'target',4);</pre>
	Mesh Boundaries 1 and 3 of a rectangle and then copy the mesh to Boundary 5.
	<pre>g1 = rect2; g2 = point2(0,0.3); fem.geom = geomcsg({g1 g2}); fem.mesh = meshinit(fem, 'hnumedg', {1 8 3 4}, 'point',[], 'edge',[1,3], 'subdomain',[]); fem.mesh = meshcopy(fem, 'source',[1 3], 'target',5);</pre>
See also	femmesh, meshinit, meshplot

Purpose	Delete elements in mesh.
Syntax	<pre>fem.mesh = meshdel(fem,) mesh = meshdel(mesh,) fem = meshdel(fem,'Out',{'fem'},)</pre>
Description	fem.mesh = meshdel(fem,) deletes elements from the mesh object fem.mesh belonging to domains according to the specified properties.
	<pre>mesh = meshdel(mesh,) deletes elements from the mesh object mesh.</pre>

fem = meshdel(fem, 'out', {'fem'},...) modifies the fem structure to include the new mesh object in fem.mesh.

The function meshdel accepts the following property/values:

TABLE I-82: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Deladj	on off	on	Specifies if elements belonging to adjacent domains of lower dimensions are deleted as well
Edge	integer vector all none	none	Specifies the edge domains for which the elements are deleted
Face	integer vector all none	none	Specifies the face domains for which the elements are deleted
Out	fem mesh	mesh	Output variables
Point	integer vector all none	none	Specifies the vertex domains for which the elements are deleted
Subdomain	integer vector all none	none	Specifies the subdomains for which the elements are deleted

Deleting elements corresponding to a specific domain, all elements on adjacent domains of higher dimension are deleted as well.

Examples

Create a mesh of a 2D geometry with 3 subdomains.

```
fem.geom = rect2+circ2;
geomplot(fem,'sublabels','on','edgelabels','on')
fem.mesh = meshinit(fem);
```

Delete the elements belonging to subdomain 3 only.

```
fem.mesh = meshdel(fem.mesh,'subdomain',3,'deladj','off');
figure, meshplot(fem)
```

Delete the elements belonging to subdomain 1 and all adjacent domains of lower dimensions that can be deleted.

```
fem.mesh = meshdel(fem.mesh,'subdomain',1,'deladj','on');
figure, meshplot(fem)
```

Delete the edge elements belonging to edge 1. Note that the elements belonging to the adjacent subdomain (subdomain 2) are deleted as well.

```
fem.mesh = meshdel(fem.mesh,'edge',1);
figure, meshplot(fem)
```

femmesh, meshenrich, meshinit

See Also

Purpose	Embed a 2D mesh object as a 3D mesh object.						
Syntax	fem1 = meshembed(fem0,) [mesh,geom]= meshembed(fem,'Out',{'mesh','geom'},)						
Description	fem1 = meshembed(fem0,) embeds the 2D geometry object in fem0.geom and the 2D mesh object in fem0.mesh, as a 3D geometry object and a 3D mesh object stored in fem1.geom and fem1.mesh, respectively.						
	[geom, mesh] = meshembed (fem, 'Out', {'geom', 'mesh'},) returns the embedded 3D geometry object in geom and the embedded 3D mesh object in mesh. Valid property/value pairs for the meshembed function are given in the following table. In addition, all embed parameters are supported and are passed to embed for						
	creating the embedded 3D geometry object.						
	PROPERTY VALUES DEFAULT DESCRIPTION						
	Mcase integer 0 Mesh case number						
	Out fem mesh geom fem Output variables						
	Embedding a 2D mesh object as a 3D mesh object, the 2D vertex elements, the 2 boundary elements, the 2D triangular elements, and the 2D quadrilateral element are embedded as 3D vertex elements, 3D edge elements, 3D triangular boundar elements, and 3D quadrilateral boundary elements, respectively.						
See also	embed, meshext	trude,meshrevolve	,femmesh				

Purpose	Make mesh object complete.
Syntax	<pre>fem.mesh = meshenrich(fem,) mesh = meshenrich(mesh,) fem = meshenrich(fem,'Out',{'fem'},)</pre>
Description	fem.mesh = meshenrich(fem,) completes the mesh object fem.mesh wit element information necessary for using the mesh object in a simulation or for converting into a geometry object.
	<pre>mesh = meshenrich(mesh,) completes the mesh object mesh.</pre>
	for - machannich (for lout) (Ifam))) madifies the for structure to

fem = meshenrich(fem, 'out', {'fem'},...) modifies the fem structure to
include the new mesh object in fem.mesh.

with

The function meshenrich accepts the following property/value pairs.

TABLE I-83: VALID PROPERTY/VALUE PAIRS

PROPERTY	2D	3D	VALUE	DEFAULT	DESCRIPTION
Extrangle		\checkmark	numeric	0.01	Maximum angle between boundary element normal and extrusion plane that will cause the element to be a part the extruded face if possible
Faceangle		\checkmark	numeric	1.8	Maximum angle between any two boundary elements in the same face
Facecleanup		\checkmark	numeric	0.01	Avoid creating small faces. Faces with an area less than Facecleanup * the mean face area, are merged with adjacent faces
Facecurv		V	numeric	0.2	Maximum relative curvature deviation between any two boundary elements in the same face
Faceparam		\checkmark	on off	on	Specifies if faces are parameterized
Minareaecurv		\checkmark	numeric	1	Minimum relative area of face to be considered as a face with constant curvature
Minareaextr		\checkmark	numeric	0.05	Minimum relative area of face to be considered extruded

PROPERTY	2D	3D	VALUE	DEFAULT	DESCRIPTION
Minareaeplane		\checkmark	numeric	0.005	Minimum relative area of face to be considered planar
Neighangle	V	V	numeric	0.35	Maximum angle between a boundary element and a neighbor that will cause the elements to be part of the same boundary domain if possible
Out	V	V	fem mesh	mesh	Output variables
Planarangle		V	numeric	0.01	Maximum angle between boundary element normal and a neighbor that will cause the element to be a part the planar face if possible

TABLE I-83: VALID PROPERTY/VALUE PAIRS

Algorithm

These are the main steps of the 3D algorithm:

- I If the domain information (dom field) for the subdomain elements is missing, all subdomain elements are assigned the same domain label.
- **2** Missing boundary elements are added. Boundary elements are required at the boundaries of the subdomains.
- **3** The up-down subdomain information (ud field) for the boundary elements is made complete.
- **4** If the domain information (dom field) for the boundary elements is missing, the face domain partitioning is determined according to the following steps.
 - Search for planar faces according to Planarangle and Minareaplane.
 - Search for extruded faces according to Extrangle and Minareaextr.
 - Search for faces with constant curvature according to Facecurv and Minareacurv. This search is only done for second order elements.
 - The remaining boundary elements are divided into face domains according to Neighangle and Faceangle.
- **5** Exceedingly small faces are merged with neighboring faces according to Facecleanup.
- **6** The faces are parameterized (param field).

	7 Missing edge elements are added. Edge elements are required at the boundaries of the faces. Domain and parameter information (dom and param fields) for the edge elements is also added.
	8 Missing vertex elements are added. Vertex elements are required on the boundaries of the edges.
	The 1D and 2D algorithms work in a similar way.
Example	Create an initial mesh object from mesh point coordinates and tetrahedral element information.
	<pre>load coord.txt; load tet.txt; el = cell(1,0); tet = tet+1; % Lowest mesh point index is zero in tet.txt el{1} = struct('type','tet','elem',tet'); m = femmesh(coord',el);</pre>
	Use meshenrich to create a complete mesh object, that is, a mesh object with boundary elements, edge elements and vertex elements with necessary geometry information.
	<pre>m = meshenrich(m); meshplot(m)</pre>
See Also	geominfo, meshinit, femmesh

Purpose	Export meshes to file.		
Syntax	<pre>meshexport(filename,ml,)</pre>		
Description	<pre>meshexport(filename,ml,) exports the meshes in the cell array ml to a file. ml can also be one single mesh object. The function meshexport supports the following mesh formats</pre>		
	FORMAT	FILE EXTENSION	
	COMSOL Multiphysics text file	.mphtxt	
	COMSOL Multiphysics binary file	.mphbin	
Example	Create a 3D mesh and export in m m = meshinit(block3+cone: meshexport('meshfile.mph	3,'hauto',9)	
See Also	femmesh, meshimport		

Purpose	Extend a mesh to the desired finite element types.			
Syntax	<pre>fem.xmesh = meshextend(fem,) [fem.xmesh, cv] = meshextend(fem,)</pre>			
Description	fem.xmesh = meshextend(fem) extends the (possibly extended) FEM structure fem with the xmesh field. The xmesh object contains the full finite element mesh for the model, and also the full description of the model using an internal syntax (the <i>element syntax</i>).			
	- ,	cv] = meshextend(f les that were multiply		a cell array cv containing
	using this solut			and generates the extended lform is weak meaning that
	The function meshextend accepts the following property/values:.			
	PROPERTY	VALUE	DEFAULT	DESCRIPTION
	Blocksize	positive integer	5000	Block size
	Eqvars	on off cell array	on	Generate equation variables
	Cplbndeq	on off cell array	on	Generate boundary-coupled equation variables
	Cplbndsh	on off cell array	off	Generate boundary-coupled shape variables
	Extend the mesh on these geometries			
	Linshape	integer vector	all meshed geometries	Use linear geometry shape order for inverted elements on these geometries
	Linshapetol	scalar or vector	0.1	Use linear geometry shape order for inverted elements on these geometries
	Mcase	integer vector	all mesh cases	Extend the mesh for these mesh cases

TABLE I-84: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Report	on off	on	Show progress window
Standard	on off	on	Convert standard syntax to element syntax

Use the properties linshape and linshapetol to avoid problems with inverted elements in the extended mesh. linshape is an integer array specifying the geometries where the software avoids inverted elements by using linear geometry shape order for the corresponding elements. linshapetol is the tolerance or a vector of tolerances of the same length as linshape. The tolerance values specify the minimum allowed value of the variable reldetjacmin for elements not being considered inverted.

CompatibilityIf the FEM structure has a version field fem.version and the version is older than
COMSOL Multiphysics 3.2, then the default for fem.solform is equal to
fem.form.

See Also xmeshinfo

Purpose	Extrude a 2D mesh object into a 3D mesh object.			
Syntax	fem1 = meshextrude(fem0,) [mesh,geom]= meshextrude(fem,'Out',{'mesh','geom'},)			
Description	and the 2D mes	h object in fem0.mes	sh, into a 31	2D geometry object in fem0.geom D geometry object and a 3D mesh pectively, according to the given
		meshextrude(fem, pject in geom and the		eom','mesh'},) returns the object in mesh.
	table. In addition	1	eters are sup	function are given in the following oported and are passed to extrude
	PROPERTY	VALUES	DEFAULT	DESCRIPTION
	Elextlavers	I-by-nd cell array		Distribution of mesh element

	TALOLS	DEIAGET	DESCRIPTION
Elextlayers	I-by-nd cell array		Distribution of mesh element layers in extruded mesh
Mcase	integer	0	Mesh case number
Out	fem mesh geom	fem	Output variables

The property Elextlayers defines the distribution of mesh element layers in the extruded mesh. The value of Elextlayers is a cell array where each entry corresponds to a section in the extruded geometry object. If a cell entry is a scalar, it defines the number of equally distributed mesh element layers that is generated for the corresponding extruded section. Alternatively, if a cell entry is a vector, it defines the distribution of the mesh element layers for the corresponding extruded section. The values in the vector, that are sorted and starts with 0, specify the placements, in relative arc length, of the mesh element layers. Note that more element layers might be introduced due to the division of the revolved geometry into sections. By default, the number of element layers is determined such that the distance of each layer is equal to the mean element size in the original 2D mesh.

Extruding a 2D mesh object into a 3D mesh object, the 2D vertex elements, the 2D boundary elements, the 2D triangular elements, and the 2D quadrilateral elements, are extruded into 3D edge elements, 3D quadrilateral boundary elements, 3D prism elements, and 3D hexahedral elements, respectively.

Examples

Create an extruded prism mesh on a cylinder of height 1.3.

```
fem.geom = circ2;
fem.mesh = meshinit(fem);
fem1 = meshextrude(fem, 'distance',1.3);
Create a hexahedral mesh by extruding a quad mesh on a rectangle.
fem.geom = rect2(1,2, 'pos',[0 0]);
fem.mesh = meshmap(fem);
fem1 = meshextrude(fem, 'distance',[1.3 2],...
'displ',[0.4 0;0 -0.2],...
'scale',[2 1;2 1.5],...
'elextlayers',{5 [0 0.2 0.8 1]});
meshplot(fem1);
```

See also extrude, meshembed, meshrevolve, femmesh

Purpose	Convert hexahedral elements to tetrahedral elements			
Syntax	fem.mesh = meshhex2tet(fem,) fem.mesh = meshhex2tet(mesh,)			
Description	fem.mesh = meshhex2tet(fem,) converts hexahedral elements in the mesh object stored in fem.mesh to tetrahedral elements.			
	fem.mesh = meshhex2tet(mesh,) converts hexahedral elements in the mesh object mesh to tetrahedral elements.			
	Each hexahedral element can be split into 5 or 28 tetrahedral elements.			
	The function meshhex2tet accepts the following property/value pairs.			
	TABLE I-85: VALID PROPERTY/VALUE PAIRS			
	PROPERTY VALUE DEFAULT DESCRIPTION			
	Out	fem mesh	mesh	Output properties
	Splithex	diagonal center	diagonal	Specifies if each hexahedral element is split into 5 or 28 tetrahedral elements
	Subdomain	numeric array all none	all	Specifies the subdomains where the hexahedral elements are converted to tetrahedral elements

Use the property subdomain to specify the subdomains where the hexahedral elements are converted to tetrahedral elements. Note that it is not possible to convert hexahedral elements in subdomains that are adjacent to subdomains that should not be converted.

Use the property splithex to specify the technique used for splitting the hexahedral elements. Use the diagonal option to split each hexahedral into 5 tetrahedral elements, and use the center option to split each hexahedral into 28 tetrahedral elements. Note that the quadrilateral elements on the boundaries of the specified subdomains will also be converted, either into two (when the option diagonal is used) or into four triangular elements (when the option center is used).

Examples Create a hex mesh and split each hexahedral element into 5 tetrahedral elements.

```
fem.geom = block3;
fem.mesh = meshmap(fem, 'face', 1);
fem.mesh = meshsweep(fem, 'meshstart', fem.mesh);
fem.mesh = meshhex2tet(fem);
meshplot(fem);
```

Create a hex mesh and split each hexahedral element into 28 tetrahedral elements.

```
fem.geom = block3;
fem.mesh = meshmap(fem, 'face', 1);
fem.mesh = meshsweep(fem, 'meshstart', fem.mesh);
fem.mesh = meshhex2tet(fem, 'splithex', 'center');
meshplot(fem);
```

See Also

femmesh, meshmap, meshquad2tri

Purpose	Import meshes from file.
Syntax	<pre>meshes = meshimport(filename,)</pre>
Description	<pre>meshes = meshimport(filename,) reads the file with name filename using the specified properties and returns a cell array of meshes.</pre>

The function meshimport supports the following mesh formats

FORMAT	FILE EXTENSION
COMSOL Multiphysics text file	.mphtxt
COMSOL Multiphysics binary file	.mphbin
NASTRAN file	.nas .bdf

Valid property/value pairs for the NASTRAN format include.

TABLE I-86: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
elemsplit	on off	off	Specifies if mesh elements of different element forms get different subdomain labels.
enrichmesh	on off	on	Specifies if the imported meshes are enriched.
linearelem	on off	off	Specifies if extended node points are ignored.
materialsplit	on off	on	Specifies if material data in the file is used to determine the domain partitioning of the subdomain elements.
report	on off	on	Determines if a progress window is displayed.

meshimport accepts all property/values that meshenrich does.

elemsplit specifies if mesh elements of different element forms—that is, tetrahedral, pentahedral, or hexahedral—get different subdomain labels. The default value is off.

enrichmesh specifies if the meshes are enriched with domain information—that is, boundary elements, edge elements, and vertex elements. The domain partitioning is controlled by the properties of meshenrich. If the value is off the output meshes are not complete meshes. The default value is on. linearelem determines if extended node points are ignored. If the value is on all imported elements are linear. Otherwise, the order of the imported elements is determined from the order of the elements in the file. The default value is on.

materialsplit determines if material data in the file is used (if available) to determine the domain partitioning of the subdomain elements. If the value is off all subdomain elements in the imported mesh belongs to the same subdomain if possible. The default value is on.

report specifies if a progress window is displayed. The default value is on.

BULK DATA ENTRY
CBAR
CHEXA
CORDIC
CORDIR
CORDIS CORD2C CORD2R CORD2S CPENTA
CQUAD4 CQUAD8
CTETRA
CTRIA3
CTRIA6
GRID

The table below specifies the NASTRAN bulk data entries that are parsed in meshimport.

The NASTRAN bulk data format uses reduced second order elements, that is, the center node on quadrilateral mesh faces (quadNode) and the center node of hexahedral elements (hexNode) are missing. Importing a NASTRAN mesh with second order elements, the coordinates of these missing node points are interpolated from the surrounding node points according to, quadNode = 0.5*quadEdgeNodes-0.25*quadCornerNodes, where quadEdgeNodes is the sum of the coordinates of the surrounding 4 edge nodes and quadCornerNodes is the sum of the coordinates of the surrounding 4 corner nodes, and hexNode = 0.25*hexEdgeNodes-0.25*hexCornerNodes, where hexEdgeNodes is the sum of

	the coordinates of the surrounding 12 edge nodes and hexCornerNodes is the sum of the coordinates of the surrounding 8 corner nodes.
Cautionary	meshimport does not handle NASTRAN files in free field format.
See Also	femmesh, meshenrich, meshexport

Purpose	Create free mesh
Syntax	fem.mesh = meshinit(fem,) fem.mesh = meshinit(geom,) fem = meshinit(fem,'out',{'fem'},)
Description	fem.mesh = meshinit(fem,) returns a mesh object derived from the geometry object fem.geom. It uses a Delaunay algorithm. The mesh size is determined from the shape of the geometry object and various property/value pairs.
	<pre>fem.mesh = meshinit(geom,) returns a mesh object derived from the geometry object geom.</pre>
	The mesh object fem.mesh is the data structure for the mesh. See femmesh for a full description of the mesh object.

fem = meshinit(fem, 'Out', {'fem'},...) modifies the FEM structure to
include the mesh object fem.mesh.

The function meshinit accepts the following property/value pairs:

TABLE I-87:	VALID	PROPERTY/VALUE PAIRS

PROPERTY	0D	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION	
edge			\checkmark	V	numeric array auto all none	auto	Specifies the edges that are meshed	
edgelem		\checkmark	\checkmark	\checkmark	cell array		Edge element distribution	
face				V	numeric array auto all none	auto	Specifies the faces that are meshed	
hauto			\checkmark	\checkmark	numeric	5	Automatic setting o several mesh parameters	
hcurve					numeric	0.3	Curvature mesh size	
hcurve				\checkmark	numeric	0.6	Curvature mesh size	
hcurveedg			\checkmark	\checkmark	numeric array	hcurve	Curvature mesh size for edges	

PROPERTY	0D	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
hcurvefac				\checkmark	numeric array	hcurve	Curvature mesh size for faces
hcutoff			\checkmark		numeric	0.001	Curvature resolution cutoff
hcutoff				\checkmark	numeric	0.03	Curvature resolution cutoff
hcutoffedg			\checkmark	\checkmark	numeric array	hcutoff	Curvature resolution cutoff for edges
hcutofffac				\checkmark	numeric array	hcutoff	Curvature resolution cutoff for faces
hgrad		\checkmark			numeric	1.3	Element growth rate
hgrad				\checkmark	numeric	1.5	Element growth rate
hgradvtx		\checkmark	\checkmark	\checkmark	numeric array	hgrad	Element growth rate for vertices
hgradedg			\checkmark	\checkmark	numeric array	hgrad	Element growth rate for edges
hgradfac				\checkmark	numeric array	hgrad	Element growth rate for faces
hgradsub		\checkmark	\checkmark	\checkmark	numeric array	hgrad	Element growth rate for subdomains
hmax		\checkmark	\checkmark	\checkmark	numeric	estimate	Global maximum element size
hmaxvtx		\checkmark	\checkmark	\checkmark	numeric array	hmax	Maximum element for vertices
hmaxedg			\checkmark	\checkmark	numeric array	hmax	Maximum element for edges
hmaxfac				\checkmark	numeric array	hmax	Maximum element for faces
hmaxsub		\checkmark	\checkmark	\checkmark	numeric array	hmax	Maximum element for subdomains
hmaxfact		\checkmark	\checkmark	V	numeric	1	A factor that the default hmax is multiplied by

TABLE I-87:	VALID PROPERTY/VALUE PA	IRS
-------------	-------------------------	-----

PROPERTY	0D	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
hcurvefac				V	numeric array	hcurve	Curvature mesh size for faces
hcutoff			\checkmark		numeric	0.001	Curvature resolution cutoff
hcutoff				V	numeric	0.03	Curvature resolution cutoff
hcutoffedg			\checkmark	\checkmark	numeric array	hcutoff	Curvature resolution cutoff for edges
hcutofffac				\checkmark	numeric array	hcutoff	Curvature resolution cutoff for faces
hgrad		\checkmark	\checkmark		numeric	1.3	Element growth rate
hgrad				\checkmark	numeric	1.5	Element growth rate
hgradvtx		\checkmark	\checkmark	\checkmark	numeric array	hgrad	Element growth rate for vertices
hgradedg			\checkmark	\checkmark	numeric array	hgrad	Element growth rate for edges
hgradfac				\checkmark	numeric array	hgrad	Element growth rate for faces
hgradsub		\checkmark	\checkmark	\checkmark	numeric array	hgrad	Element growth rate for subdomains
hmax		\checkmark	\checkmark	\checkmark	numeric	estimate	Global maximum element size
hmaxvtx		\checkmark	\checkmark	\checkmark	numeric array	hmax	Maximum element for vertices
hmaxedg			\checkmark	\checkmark	numeric array	hmax	Maximum element for edges
hmaxfac				\checkmark	numeric array	hmax	Maximum element for faces
hmaxsub		\checkmark	\checkmark	\checkmark	numeric array	hmax	Maximum element for subdomains
hmaxfact		\checkmark	\checkmark	V	numeric	1	A factor that the default hmax is multiplied by

PROPERTY	0D	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
hmesh		\checkmark	\checkmark	\checkmark	numeric		Element size given on mesh
hnarrow			\checkmark		numeric	1	Resolution of narrow regions
hnarrow				\checkmark	numeric	0.5	Resolution of narrow regions
hnumedg			\checkmark	\checkmark	cell array		Number of elements for edges
hnumsub		\checkmark			cell array		Number of elements for subdomains
hpnt			\checkmark		numeric	10	Global number of resolution points
hpnt				\checkmark	numeric	20	Global number of resolution points
hpntedg			\checkmark		numeric array	Hpnt	Number of resolution points for edges
hpntfac				\checkmark	numeric array	Hpnt	Number of resolution points for faces
jiggle			\checkmark	\checkmark	on off	on	Improve mesh quality
mcase	\checkmark		\checkmark	\checkmark	integer	0	Mesh case number
mesh		\checkmark	\checkmark	\checkmark	mesh object		Mesh for hmesh
meshstart	\checkmark		\checkmark	\checkmark	mesh object		Starting mesh
methodsub			\checkmark		cell array tri quad	tri	Specify free triangle or free quad mesh
minit			\checkmark	\checkmark	on off	off	Boundary triangulation
mlevel					vtx sub	sub	Meshing level
mlevel			\checkmark		vtx edg sub	sub	Meshing level
mlevel				\checkmark	vtx edg fac sub	sub	Meshing level

TABLE I-87: VALID PROPERTY/VALUE PAIRS

PROPERTY	0D	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
point		\checkmark	V	V	numeric array auto all none	auto	Specifies the vertices that are meshed
out	\checkmark	V	V	\checkmark	fem mesh p e t vg eg vg	mesh	Output variables
report	\checkmark	\checkmark	\checkmark	\checkmark	on off	on	Display progress
subdomain	\checkmark	\checkmark	\checkmark	V	numeric array auto all none	auto	Specifies the subdomains that are meshed
xscale			\checkmark	V	numeric	1	Scale geometry in x direction before meshing
yscale			\checkmark	V	numeric	1	Scale geometry in y direction before meshing
zscale				V	numeric	1	Scale geometry in z direction before meshing

TABLE I-87: VALID PROPERTY/VALUE PAIRS

Use the properties point, edge, face, and subdomain to specify the domains to be meshed. If you use these properties together with the meshstart property, the value auto means that all domains that are not meshed in the starting mesh are meshed and none that no further domains are meshed. all means that all domains not already meshed in the starting mesh and all meshed domains that are not adjacent to a meshed domain of higher dimension are meshed (or remeshed). It is also possible to specify the domains to be meshed (or remeshed) using a vector of domain indices.

The meshstart property is used when meshing a geometry interactively. The value of this property is the starting mesh of the meshing operation.

Use the property methodsub in 2D to specify if subdomains should be meshed with triangles or quads. The property is set to tri or quad, to mesh all subdomains with triangles or quads respectively. The value of the property can also be an even

numbered cell array, where the odd entries contain subdomain indices, either as scalar values, or as vectors with subdomain indices, and the even entries are either quad or tri, specifying triangle or quad elements for each subdomain. Default is tri for all subdomains.

Use the property methodfac in 3D to specify if faces should be meshed with triangles or quads. The property methodfac in 3D is equivalent to the property methodsub in 2D, and is specified in the same way, but for faces instead of subdomains.

The property edgelem is used to explicitly control the distribution of the edge elements in the mesh. The value of this property is an even numbered cell array where the odd entries contain edge indices, either as scalar values, or as a vectors with edge indices, and the even entries contain scalar values or vectors specifying the edge element distribution on the corresponding edge(s). If the edge element distribution is specified by a scalar, the edge elements on the corresponding edge(s) are equally distributed in arc length and the number of edge elements equals the value of the scalar. To get full control over the edge element distribution on an edge, the vector form is used. The values in the vector, that are sorted and starts with 0, specify the relative placement of the mesh vertices along the direction of the corresponding edge(s). The edgelem property can also be used on subdomains in 1D to control the element distribution.

The minit property is related to the way the mesh algorithm works. By turning on minit you can see the initial discretization of the boundaries. This property is only valid for mlevel sub.

Hauto is available in 2D and 3D and is an integer between 1 and 9. This integer is used to set several mesh parameters in order to get a mesh of desired size. Smaller values of hauto generate finer meshes with more elements.

ΗΑυτο	HMAXFACT	HCURVE	HGRAD	HCUTOFF
I	0.15	0.2	1.1	0.0001
2	0.3	0.25	1.2	0.0003
3	0.55	0.25	1.25	0.0005
4	0.8	0.3	1.3	0.001
5	1	0.3	1.3	0.001
6	1.5	0.4	1.4	0.005
7	1.9	0.6	1.5	0.01

TABLE I-88: MESH PARAMETERS SET BY THE PROPERTY HAUTO IN 2D

TABLE I-88: MESH PARAMETERS SET BY THE PROPERTY HAUTO IN 2D

ΗΑυτο	HMAXFACT	HCURVE	HGRAD	HCUTOFF
8	3	0.8	1.8	0.02
9	5	1	2	0.05

TABLE I-89: MESH PARAMETERS SET BY THE PROPERTY HAUTO IN 3D

ΗΑυτο	HMAXFACT	HCURVE	HGRAD	HCUTOFF	HNARROW
I	0.2	0.2	1.3	0.001	1
2	0.35	0.3	1.35	0.005	0.85
3	0.55	0.4	1.4	0.01	0.7
4	0.8	0.5	1.45	0.02	0.6
5	I	0.6	1.5	0.03	0.5
6	1.5	0.7	1.6	0.04	0.4
7	1.9	0.8	1.7	0.05	0.3
8	3	0.9	1.85	0.06	0.2
9	5	I	2	0.07	0.1

Hcurve is a scalar numeric value that relates the mesh size to the curvature of the geometry boundaries. The radius of curvature is multiplied by the hcurve factor to obtain the mesh size along the boundary.

hcurveedg and hcurvefac are matrices with two rows where the first row contains edge indices and face indices respectively, and the second row contains corresponding values of hcurve. If several faces are represented in one patch, the value of hcurvefac for the faces in the patch is set to the minimum value of the hcurvefac values for the corresponding faces.

hcutoff is used to prevent the generation of many elements around small curved parts of the geometry. The interpretation is that when the radius of curvature is smaller than hcutoff*maxdist the radius of curvature is taken as hcutoff*maxdist, where maxdist is the longest axis parallel distance in the geometry.

hcutoffedg and hcutofffac are matrices with two rows where the first row contains edge indices and face indices respectively, and the second row contains corresponding values of hcutoff. If several faces are represented in one patch, the value of hcutofffac for the faces in the patch is set to the minimum value of the hcutofffac values for the corresponding faces.

The property hgrad tells how fast the element size—measured as the length of the longest edge of the element—can grow from a region with small elements to a region with larger elements. If two elements lie one unit length apart, the difference in element size can be at most hgrad.

hgradvtx, hgradedg, hgradfac, and hgradsub are matrices with two rows where the first row contains vertex indices, edge indices, face indices, and subdomain indices respectively, and the second row contains corresponding values of hgrad. If several faces are represented in one patch, the value of hgradfac for the faces in the patch is set to the minimum value of the hgradfac values for the corresponding faces.

The hmax parameter controls the size of the elements in the mesh.meshinit creates a mesh where no element size exceeds hmax. The default hmax value is one fifteenth of the longest axis parallel distance in the geometry in 1D and 2D and one tenth of the longest axis parallel distance in the geometry in 3D.

hmaxvtx, hmaxedg, hmaxfac, and hmaxsub are matrices with two rows where the first row contains vertex indices, edge indices, face indices, and subdomain indices respectively, and the second row contains corresponding values of hmax.

The hmaxfact property is used to scale the defaulted hmax value.

hmesh is a vector with one entry for every mesh vertex or element in the mesh given in the mesh property. This can be used to specify the size of the elements using the mesh provided with the property mesh.

The hnarrow parameter controls the size of the elements in narrow regions. Increasing values of this property decrease the size of the elements in narrow regions. If the value of hnarrow is less than one, elements that are anisotropic in size might be generated in narrow regions.

hnumedg and hnumsub are cell arrays where the odd entries contain edge indices and subdomain indices respectively, and even entries contain number of elements.

The hpnt property controls the number of points that are placed on each edge in 2D and in each parametric direction on each geometry patch in 3D to resolve the geometry.

hpntedg is a matrix with two rows where the first row contains edge indices and the second row contains corresponding values of hpnt.

hpntfac is a matrix with two rows where the first row contains face indices and the second row contains corresponding number of resolution points in each parametric

direction of the underlying geometry patch. If several faces are represented in one patch the value of hpntfac for the faces in the patch is set to the maximum value of the hpntfac values for the corresponding faces.

The jiggle property determines if the quality of the mesh is improved before the mesh is returned.

Use the mesh property to specify a mesh for the property hmesh.

The property mlevel determines to which level the mesh is generated. If it is vtx, only the vertices in the geometry are returned. If it is edg, the edges in the geometry are resolved. If it is fac the edges and faces in the geometry are resolved. If it is sub elements in the subdomains are generated as well.

The properties xscale, yscale, and zscale specify scalar factors in each axis direction that the geometry is scaled by before meshing. The resulting mesh is then scaled back to fit the original geometry. The values of other properties correspond to the scaled geometry. By default, no scaling is done.

Examples

3D Example

Create a 3D mesh of a cylinder.

```
clear fem
fem.geom=cylinder3;
fem.mesh=meshinit(fem);
meshplot(fem)
```

2D Example

Make a simple mesh of a unit square.

```
clear fem
fem.geom = geomcsg({square2(0,1,1)});
fem.mesh = meshinit(fem);
meshplot(fem), axis equal
```

Make the mesh finer than the default.

```
fem.mesh = meshinit(fem, 'hmax', 0.02);
meshplot(fem), axis equal
```

Now, make the mesh denser only near the edge segment to the left.

fem.mesh = meshinit(fem, 'hmaxedg',[1; 0.02]);
meshplot(fem), axis equal

Make a free quad mesh of a circle

```
clear fem
fem.geom = geomcsg({circ2});
```

	fem.mesh = meshinit(fem,'methodsub','quad'); meshplot(fem), axis equal
	ID Example
	Create a mesh on the interval $[0,1]$ that is finer near the point 0 and grows toward 1.
	fem.geom = geomcsg({solid1([0 1])}); fem.mesh = meshinit(fem,'hmax',0.1,'hmaxvtx',[1; 0.001]); meshplot(fem)
	2D Example Dealing with Interactive Meshing
	Create a boundary mesh of a geometry
	clear fem; fem.geom = rect2+circ2; fem.mesh = meshinit(fem,'subdomain','none'); meshplot(fem)
	Mesh subdomain 2 using the boundary mesh as starting mesh
	fem.mesh = meshinit(fem,'subdomain',2,'meshstart',fem.mesh); meshplot(fem)
	Mesh the remaining subdomains using the previous mesh as starting mesh
	fem.mesh = meshinit(fem,'meshstart',fem.mesh); meshplot(fem)
Compatibility	The second and third row in the vg field as well as the second row in the v field will be removed in future versions.
Cautionary	To achieve compatibility with FEMLAB 2.3, the geometry input is automatically converted to a geometry object using the function geomobject. The geometry input can be any analyzed geometry. See geomobject for details.
	If you create a mesh with methodsub set to quad in 2D, or methodfac set to quad in 3D, the generated mesh is not guaranteed to contain only quadrilateral elements. If the algorithm for some reason fails to mesh the entire domain with quad elements, or if the quality of a quad element is very low, some triangular elements are generated instead.
See Also	femmesh, geomcsg, meshplot, meshrefine
Reference	[1] George, P. L., Automatic Mesh Generation—Application to Finite Element Methods, Wiley, 1991.

Purpose	Integrate over arbitrary cross section						
Syntax	I = meshintegrate(p,t,d) I = meshintegrate(p,d) I = meshintegrate(p)						
Description	<pre>I = meshintegrate(p,t,d) computes the integral I over the mesh given by p and t, with values (for each point) in d. d is of size 1-by-np, where np is the number of points in p (=size(p,2)). The elements are considered to be linear.</pre>						
	I = meshintegrate(p,d) assumes t=[1,2,3, (np-1) ; 2,3,4, np], (where np=size(p,2)), i.e., that the mesh is a line and that the points in p are sorted.						
	<pre>I = meshintegrate(p) calls meshintegrate(p(1,:), p(2,:)).</pre>						
	This function is useful for computing integrals along cross sections plotted with postcrossplot, in which case p, t, and d are extracted from the output when the property outtype is set to postdata.						
Examples	Line integral in 2D:						
	<pre>% Just set up a problem: clear fem fem.geom = circ2+rect2; fem.mesh = meshinit(fem); fem.shape = 2; fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = 1; fem.xmesh = meshextend(fem); fem.sol = femstatic(fem); % Make a cross-section plot, with output being a postdata</pre>						
	% structure pd = postcrossplot(fem,1,[0 1;0 1],'lindata','u', 'npoints',100,'outtype','postdata');						
	% Call meshintegrate: I = meshintegrate(pd.p);						
	Line integral in 3D:						
	<pre>% Just set up a problem: clear fem, fem.geom = block3; fem.mesh = meshinit(fem,'hmax',0.15); fem.shape = 2; fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = {1 1 0 0 1 1}; fem.xmesh = meshextend(fem); fem.sol = femstatic(fem);</pre>						

	% Make cross-section plot: pd = postcrossplot(fem,1,[0 1;0 1;0 1],'lindata','u', 'npoints',100,'outtype','postdata');
	% Call meshintegrate: I = meshintegrate(pd.p)
	Surface integral in 3D using the same problem as above:
	<pre>pd = postcrossplot(fem,2,[0 0 0;0 1 0;1 0 1]','surfdata','u', 'outtype','postdata'); I = meshintegrate(pd.p, pd.t, pd.d)</pre>
	This function only works for lines and surfaces actually intersecting the geometry. For plots along geometry boundaries or edges (or 1-D subdomains), better results are achieved using postint.
Cautionary	This function is not implemented for 3-D elements, i.e., when T has four rows.
See also	postcrossplot, postint

Purpose	Create mapped quad mesh.								
Syntax	fem.mesh = meshmap(fem,) fem.mesh = meshmap(geom,) fem = meshmap(fem,'out',{'fem'},)								
Description	fem.mesh = meshmap(fem,) returns a mapped quad mesh derived from the geometry fem.geom. For a 3D geometry, only the faces are meshed.								
	fem.mesh = geometry geo) returr	ns a mapped quad mesh derived from the					
	 fem = meshmap(fem, 'out', {'fem'},) modifies the fem structure to include a mesh in fem.mesh. The quad mesh is generated by a mapping technique. For each subdomain in 2D and face in 3D, a logical mesh is generated on a square geometry and is then mapped onto the real geometry by transfinite interpolation. The following criteria must be met by the input geometry object for the mapping technique to work: Each subdomain/face to be meshed must be bounded by one connected boundary component only. 								
	• Each subdo	main/face to be	meshed m	ust be bounded by at least four edges.					
	• The subdor	nains/faces to be	e meshed m	ust not contain isolated vertices or edges.					
	• The shape of	of each subdomai	in/face to b	e meshed must not differ too much from					
	rectangular	shape.							
	The function	meshmap accepts	the followi	ng property/values:					
	PROPERTY	VALUE	DEFAULT	DESCRIPTION					
	·								

PROPERTY	VALUE	DEFAULT	DESCRIPTION
edgegroups	cell array of size I-by-ns		Determines the grouping of the edges, per subdomain/face, into four edge groups, corresponding to the edges of the logical square
edgelem	cell array		Edge element distribution
face	numeric array auto all none	auto	Specifies the faces in 3D that are meshed
hauto	numeric	5	Predefined mesh element size
mcase	integer	0	Mesh case number

PROPERTY	VALUE	DEFAULT	DESCRIPTION
meshstart	mesh object	empty	Starting mesh
report	on off	on	Display progress
out	fem mesh	mesh	Output variables
subdomain	numeric array auto all none	auto	Specifies the subdomains in 2D that are meshed

The property edgegroups is a cell array where each cell entry, corresponding to each subdomain/face, determines the relation between the edges defining the boundary of the corresponding subdomain/face, and the four edges of the logical square. If a cell entry is left empty, the meshing algorithm splits the edges bounding the subdomain/face into the four edge groups at the vertices corresponding to the four sharpest corners. The relation between the edges of each subdomain/face and the edges of the logical square is specified as a cell array, where each cell entry contains the indices to the edges in the real geometry that correspond to one edge of the logical square.

The property edgelem determines the distribution of the edge elements in the mesh. The value of this property is an even numbered cell array where the odd entries contain edge indices, either as scalar values, or as a vectors with edge indices, and the even entries contain scalar values or vectors specifying the edge element distribution on the corresponding edge(s). If the edge element distribution is specified by a scalar, the edge elements on the corresponding edge(s) are equally distributed in arc length and the number of edge elements equals the value of the scalar. To get full control over the edge element distribution on an edge, the vector form is used. The values in the vector, that are sorted and starts with 0, specify the placements, in arc length, of the mesh vertices along the direction of the corresponding edge(s).

hauto is an integer between 1 and 9 that controls the element size in the generated mesh. The default value is 5 which means that the element size is set to 1/15 in 2D and 1/10 in 3D of the size of the geometry for the elements not affected by the edgelem property. By changing the value of this property, the default element size is multiplied by the following factors.

ΗΑυτο	SCALE FACTOR
I	0.2
2	0.35

ΗΑυτο	SCALE FACTOR
3	0.55
4	0.8
5	I
6	1.5
7	1.9
8	3
9	5

Use the property subdomain in 2D and face in 3D to specify the subdomains/faces to be meshed. If you use this property together with the meshstart property, the value auto means that all subdomains/faces that are not meshed in the starting mesh are meshed, none means that no further subdomains/faces are meshed, and all means that all subdomains/faces are meshed (or remeshed). It is also possible to specify the subdomains/faces to be meshed (or remeshed) using a vector of subdomain/face indices.

The meshstart property is used when meshing a geometry interactively. The value of this property is the starting mesh of the meshing operation.

Note that for the mapping technique to work, opposite edges require the same number of edge elements. If this requirement is not met by the specified values in edgelem, an error is generated.

Examples

Create a mapped quad mesh on a geometry where all subdomains are topologically equivalent with a rectangle.

Create a mapped quad mesh on a geometry with two subdomains.

Create a mesh with both triangle and quad elements

```
fem.geom = geomcomp({circ2(0.5, 'pos',[0 0.5]),rect2,...
                     circ2(0.5, 'pos', [1 0.5])}, 'edge', 7:10);
figure, geomplot(fem)
fem.mesh = meshmap(fem, 'subdomain',2, 'hauto',3);
fem.mesh = meshinit(fem, 'meshstart', fem.mesh, 'hauto', 3);
figure, meshplot(fem);
```

See also

meshdel, meshinit, meshplot

Purpose	Plot mesh.
Syntax	<pre>meshplot(fem,) meshplot(mesh,) h = meshplot()</pre>
Description	<pre>meshplot(fem,) plots the mesh object fem.mesh.</pre>
	meshplot(mesh,) plots the mesh object mesh.
	h = meshplot() additionally returns handles to the plotted axes objects.
	The mesh of the PDE problem is specified by the mesh object. Details on the representation of the mesh can be found in the entry femmesh.
	There is a multitude of options that enables you to plot the mesh in virtually any conceivable way. For 2D and 3D meshes, there are basically two types of mesh plots; the <i>wireframe plot</i> and the <i>patch plot</i> .
	In 3D, the default type is a <i>patch plot</i> , where both the triangular faces of the elements and the boundary elements are rendered. Only the visible faces of the elements are included in the plot, and the boundary elements and other mesh faces are plotted in different colors. The characteristics of this plot type are controlled by the properties that start with edge and bound. The edge properties control the plot of the element faces, and the bound properties control the boundary element plot (the name edge has historic reasons; it was first used in 2D). The color of the edges of the element faces is determined by the property eledgecolor.
	The alternative type of 3D plot is a <i>wireframe plot</i> , consisting of the (1D) edges of the elements and the boundary elements. The properties that start with dedge and dbound control the plot characteristics of this plot type. The dedge properties control the plot of the element edges, and the dbound properties control the plot of the boundary elements. See the 3D example below for how to obtain a patch plot and a wireframe plot of a mesh. You can also plot the mesh edges that lie on geometry edges (curves) with a special color. The characteristics of this plot is controlled by the properties that begin with curve.
	In 2D, there are basically two types of mesh plots. The default type is a <i>wireframe plot</i> of the edges of the elements, where the boundary elements are plotted in a different color. The properties that start with edge and bound control the plot characteristics of this plot type. The edge properties control the plot of the element edges, and the bound properties control the boundary element plot.

The alternative type of 2D plot is a *patch plot* of the triangular elements, where the edges of the elements can have a different color. The properties that start with el control the characteristics of this plot type. See the 2D example below for how to obtain the two plot types. The two plot types can be combined, but doing this is not always useful.

In 1D, the default is to combine the above two types of plots into one plot type. Thus, by default you can see both the elements, the boundary elements, and the intermediate mesh vertices. The plot of the elements is controlled by the properties that begin with el, and the plot of the boundary elements is controlled by the bound properties. The plot of the mesh vertices is controlled by the edge properties.

In all dimensions, you may plot the mesh vertices (sometimes called node points) in a special color. This plot is controlled by the properties that start with node.

The table shows the valid property/value pairs for the meshplot command. The design philosophy has been to keep property interpretation constant over space dimension, but to plot these properties as plot objects of different types. The mode properties that turn the different visualization types on and off, have been marked with the type of the plot produced in the different space dimensions: m for marker plot, 1 for line/wireframe plot, and p for patch plot.

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
bdl		\checkmark	\checkmark	integer vector	all	Boundary list
boundcolor	\checkmark	\checkmark		color	r	Boundary color
boundcolor			\checkmark	color qual	r	Boundary color
boundmarker	\checkmark			marker	0	Boundary marker
boundmode	m	I	Ρ	on off isolated(ID)	on off (3D)	Show boundary elements
curvecolor			\checkmark	color	g	Curve (edge) coloring data
curvemode			I	on off	off	Show mesh edges on curves (edges)
dboundcolor			\checkmark	color	r	Boundary wireframe color
dboundmode			I	on off	off	Show boundary elements as wireframe

TABLE I-90: VALID PROPERTY/VALUE PAIRS

TABLE I-90: VALID PROPERTY/VALUE PAIRS

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
dedgecolor			\checkmark	color	b	Element wireframe color
dedgemode			I	on off	off	Show elements as wireframe
edgecolor	\checkmark			color	b	Color of mesh vertices
edgecolor		\checkmark		color	b	Color of mesh edges
edgecolor			\checkmark	color qual	b	Color of mesh faces
edgemarker	\checkmark			marker	х	Mesh vertex marker
edgemode	m			on off	on	Show mesh vertices
edgemode		I		on off	on	Show wireframe plot of mesh edges
edgemode			Ρ	on off	on	Show triangular mesh faces as patch plot
edl			\checkmark	integer vector	all	Edge list
elcolor	\checkmark			color	k	Element color
elcolor		\checkmark		color	gray	Element color
eledgecolor		\checkmark	\checkmark	color	k	Mesh edge color in patch plot
elkeep		\checkmark	\checkmark	number between 0 and 1	1	Fraction of elements to keep
elkeeptype		\checkmark	\checkmark	min max random	random	Which elements to keep
ellabels	\checkmark	\checkmark		on off	off	Mesh element labels
ellogic			\checkmark	logical expression	1	Select elements using a logical expression
ellogictype			\checkmark	all any xor	all	Interpretation of the logical expression
elmode	I			on off	on	Show elements
elmode		Ρ		on off	off	Show elements
markersize	\checkmark	\checkmark	\checkmark	scalar	6	Marker size
nodecolor		\checkmark	\checkmark	color	k	Mesh vertex (node) color

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
nodelabels	\checkmark	\checkmark	\checkmark	on off	off	Mesh vertex (node) labels
nodemarker	\checkmark	\checkmark	\checkmark	marker		Mesh vertex (node) marker
nodemode	m	m	m	on off	See below	Show mesh vertices (node points)
parent	\checkmark	\checkmark	\checkmark	axes handle		Handle to axes object
pointcolor	\checkmark	\checkmark	\checkmark	color	b	Point color data
pointlabels	\checkmark	\checkmark	\checkmark	off on list of strings	off	Point label list
pointmarker	\checkmark	\checkmark	\checkmark	marker symbol	0	Point marker
pointmode	\checkmark	\checkmark	\checkmark	on off isolated	on	Show points
qualbar		\checkmark	\checkmark	on off	on	Show color bar
qualdlim		\checkmark	\checkmark	I-by-2 numeric	[0,1]	Color limits
qualmap			\checkmark	colormap	jet	Colormap
sdl		\checkmark	\checkmark	integer vector	all	Subdomain list

TABLE I-90: VALID PROPERTY/VALUE PAIRS

In 3D, the properties Sdl, Bdl, Elkeep, Elkeeptype, Ellogic, and Ellogictype determine what part of the mesh is displayed. Only elements lying in subdomains in the list Sdl, and boundary elements lying on boundaries in the list Bdl, are shown. The set of elements is restricted further by the properties Elkeep, Elkeeptype, Ellogic, and Ellogictype. These affect the patch and wireframe plots of the elements (controlled by the Edge and Dedge properties). Only the mesh elements whose corners satisfy the logical expression Ellogic are shown. Ellogictype determines whether all, some, or some but not all of the corners are required to satisfy the condition. Only a fraction Elkeep of the mesh elements are shown. The property Elkeeptype determines whether this fraction is the elements of worst quality, or if it is a random set of elements.

The E1 property group controls the display of the actual element. This is only possible in 1D and 2D. The Edge property group controls the display of the boundaries of the elements. This visualization type is available in all space dimensions. In 1D it is done by displaying marker symbols, in 2D it is done by displaying a wireframe plot of the element edges, and in 3D it is done by displaying

the element edges as patches. In 3D, the property dedge makes it possible to obtain wireframe plots of the elements.

The bound property group controls the display of the boundary elements. In 1D it displays marker symbols, in 2D it displays the boundary elements as a wireframe, and in 3D it displays the boundary elements as a patch plot. In 3D, the property dbound makes it possible to get wireframe plots of the boundary elements.

In 3D, the curve property group controls the display of mesh elements on geometry edges.

The point property group displays the mesh vertex elements as marker symbols in all space dimensions.

The node property group displays the mesh vertices (node points) as marker symbols in all space dimensions.

The ellabels property is available in 1D and 2D and controls the display of mesh element labels. If there are more than one type of mesh elements, for example both triangular and quadrilateral elements in a 2D mesh, the different types are labeled individually. This means that the triangles will be labeled from 1 to the number of triangles, and the quads will be labeled from 1 to the number of quads.

The properties that control marker type or coloring can handle any standard scripting marker or color type (see the plot command). In 3D, the patch coloring can be made according to the element quality, by specifying the color as 'qual'.

meshplot can display meshes where there are no elements on a certain element dimension. In these cases, the default values for curvemode and nodemode change to make the best possible mesh visualization.

Examples

3D Example

Start by creating a 3D geometry and a mesh.

```
c1 = cylinder3(0.2,1,[0.5,0.5,0]);
b1 = block3;
geom = b1-c1;
mesh = meshinit(geom);
```

Plot the mesh as a quality patch plot with parts of the elements excluded by a logical expression. These types of options make it easy to study the mesh inside the geometry.

```
'qualbar','on')
```

You can get a wireframe plot of the same mesh with only a fraction of the tetrahedrons visible, by the command

The plot shows only a small fraction of the elements. It is not possible to get a mesh quality plot by using only wireframes.

2D Example

Start by creating the geometry and a coarse mesh.

```
clear fem
sq1 = square2;
sq2 = move(sq1,0,-1);
sq3 = move(sq1,-1,-1);
fem.geom = sq1+sq2+sq3;
fem.mesh = meshinit(fem,'hmax',0.4);
```

Then plot the mesh as a line plot of the edges of the elements. The element edges are blue except on the boundary elements, where they are red. This is the default mesh plot type.

```
meshplot(fem)
```

You can change the colors of the element edges to yellow and green.

```
meshplot(fem,'edgecolor','y','boundcolor','g')
```

Now, plot the mesh as a patch plot. You need to disable both the element edge and boundary element plots, and enable the element plot.

```
meshplot(fem,'edgemode','off','boundmode','off','elmode','on')
```

You can change the color of the elements to red, with white edges, and add mesh vertex labels by

ID Example

Start by creating the geometry and mesh.

```
clear fem
s1 = solid1([0 0.1 4]);
s2 = solid1([3 4]);
fem.geom = s1+s2;
fem.mesh = meshinit(fem, 'hmax',0.4);
```

 The standard mesh plot in 1D is the following plot.

 meshplot(fem)

 You can turn on node labeling, change the element color to red, and change the element edge boundary coloring to green by

 meshplot(fem, 'elcolor', 'r', 'boundcolor', 'g')

 Compatibility

 meshplot no longer supports the properties boundlabels, curvelabels, ellabels, and labelcolor from FEMLAB 2.3.

 See Also
 femmesh, geomplot, postplot

Purpose	Make a regular mesh on a rectangular geometry.						
Syntax	fem.mesh = meshpoi(fem,nx,ny) fem.mesh = meshpoi(fem,n)						
Description	fem.mesh = meshpoi(fem,nx,ny) constructs a regular mesh on the rectangular geometry specified by fem.geom, by dividing the "x edge" into nx pieces and the "y edge" into ny pieces, and placing $(nx+1)(ny+1)$ points at the intersections. The "x edge" is the one that makes the smallest angle with the x-axis.						
	fem.mesh =	meshpoi(fem,n,) uses nx = ny	= n.			
	The triangular mesh is described by the FEM mesh object mesh. Details on the mesh object can be found in the entry on femmesh.						
	For best performance with poisson, the larger of nx and ny should be a power of 2.						
	The function meshpoi accepts the following property/values:						
	TABLE I-91: VAL	ID PROPERTY/VALUE PAIRS					
	PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION			
	out fem mesh mesh Output variables						
Diagnostics	If fem.geom does not seem to describe a rectangle an error is signalled.						
Cautionary	This function only works in 2D.						
See Also	femmesh, poisson						

Purpose Convert quadrilateral elements to triangular elements

```
Syntax fem.mesh = meshquad2tri(fem,...)
fem.mesh = meshquad2tri(mesh,...)
```

Description

fem.mesh = meshquad2tri(fem,...) converts quadrilateral elements in the mesh object stored in fem.mesh to triangular elements.

fem.mesh = meshquad2tri(mesh,...) converts quadrilateral elements in the mesh object mesh to triangular elements.

Each quadrilateral element can be split into two or four triangular elements.

The function meshquad2tri accepts the following property/value pairs.

PROPERTY	2D	3D	VALUE	DEFAULT	DESCRIPTION
Face		\checkmark	numeric array all none	all	Specifies the faces where the quadrilateral elements are converted to triangular elements
Out	\checkmark	\checkmark	fem mesh	mesh	Output properties
Splitquad	\checkmark	\checkmark	diagonal center	diagonal	Specifies if each quadrilateral element is split into two or four triangular elements
Subdomain	\checkmark		numeric array all none	all	Specifies the subdomains where the quadrilateral elements are converted to triangular elements

TABLE I-92: VALID PROPERTY/VALUE PAIRS

Use the property subdomain in 2D and face in 3D to specify the subdomains and faces, respectively, where the quadrilateral elements are converted to triangular elements. Note that it is not possible to convert quadrilateral elements on faces in 3D that are adjacent to subdomain elements.

Use the property splitquad to specify if each quadrilateral element is split into two or four triangular elements. Use the diagonal option to split each quadrilateral along a diagonal into two triangles and use the center option to split each quadrilateral into four triangles by introducing a new mesh vertex in the centroid of each quadrilateral. Note that to be able to use the center option on faces in 3D you have to provide a geometry object corresponding to the mesh object.

Examples Create a mapped quad mesh on a unit rectangle and split each quadrilateral element into two triangular elements.

```
fem.geom = rect2;
fem.mesh = meshmap(fem);
fem.mesh = meshquad2tri(fem);
meshplot(fem);
```

Create a mapped quad mesh on a unit rectangle and split each quadrilateral element into four triangular elements.

```
fem.geom = rect2;
fem.mesh = meshmap(fem);
fem.mesh = meshquad2tri(fem,'splitquad','center');
meshplot(fem);
```

See Also

femmesh, meshhex2tet, meshmap

Purpose Mesh quality measure.

Syntax

q = meshqual(mesh)

Description

q = meshqual(mesh) returns the mesh element quality for all elements in the mesh object mesh.

The mesh element quality is a number between 0 and 1. The quality is 1 for a perfect element.

Details on the mesh object can be found in the entry on femmesh.

The triangle quality is given by the formula

$$q = \frac{4\sqrt{3}a}{h_1^2 + h_2^2 + h_3^2},$$

where *a* is the area and h_1 , h_2 , and h_3 the side lengths of the triangle. If q > 0.3 the mesh quality should not affect the quality of the solution. q = 1 when $h_1 = h_2 = h_3$. For a quadrilateral,

$$q = \frac{4A}{h_1^2 + h_2^2 + h_3^2 + h_4^2}$$

where h_1, h_2, h_3 , and h_4 are the side lengths. q=1 for a square.

The tetrahedron mesh quality measure is given by

$$q = \frac{72\sqrt{3}V}{(h_1^2 + h_2^2 + h_3^2 + h_4^2 + h_5^2 + h_6^2)^{3/2}}$$

where *V* is the volume, and h_1 , h_2 , h_3 , h_4 , h_5 , and h_6 are the side lengths of the tetrahedron. If q > 0.1 the mesh quality should not affect the quality of the solution.

For a hexahedron,

$$q = \frac{24\sqrt{3}V}{\left(\sum_{i=1}^{12} h_i^2\right)^{3/2}},$$

where h_i are the edge lengths. q=1 for a cube. For a prism,

$$q = \frac{36\sqrt{3}V}{\left(\sum_{i=1}^{9}h_{i}^{2}\right)^{3/2}},$$

where h_i are the edge lengths. q=1 for a right-angled prism where all edge lengths are equal.

The element quality is always 1 in 1D.

See Also meshrefine, femmesh, meshsmooth

ReferenceBank, Randolph E., PLTMG: A Software Package for Solving Elliptic Partial
Differential Equations, User's Guide 6.0, Society for Industrial and Applied
Mathematics, Philadelphia, PA, 1990.

Purpose	Refine a mesh.		
Syntax	<pre>fem.mesh = meshrefine(fem,); fem = meshrefine(fem,'Out',{'fem'},);</pre>		
Description	<pre>fem.mesh = meshrefine(fem,) returns a refined version of the triangular mesh specified by the geometry, fem.geom, and the mesh, fem.mesh.</pre>		
	<pre>fem = meshrefine(fem,'Out',{'fem'},) modifies the FEM structure to include the refined mesh in fem.mesh.</pre>		

meshrefine is supported for meshes containing simplex elements only, that is, lines, triangles, and tetrahedrons.

The function meshrefine accepts the following property/values:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
mcase	integer	0	Mesh case number
out	fem mesh	mesh	Output variables
rmethod	longest regular	see below	Refinement method
subdomain	integer vector all none	all	Specifies the subdomains that are refined
tri	one row vector or two row matrix	all elements once	List of elements to refine (second row is number of refinements)

TABLE I-93: VALID PROPERTY/VALUE PAIRS

The default refinement method in 2D is regular refinement, where all of the specified triangles are divided into four triangles of the same shape. Longest edge refinement, where the longest edge of each specified triangle is bisected, can be demanded by giving longest as Rmethod. Using regular as Rmethod results in regular refinement. Some triangles outside of the specified set may also be refined, in order to preserve the triangulation and its quality.

In 3D, the default refinement method is longest. The regular refinement method is only implemented for uniform refinements.

In 1D, regular refinement, where each element is divided into two elements of the same shape, is always used.

Examples Refine the mesh of the L-shaped membrane several times and plot the mesh for the geometry of the L-shaped membrane:

	<pre>clear fem fem.geom = square2 + move(square2,0,-1) + move(square2,-1,-1); fem.mesh = meshinit(fem, 'hmax',0.4); subplot(2,2,1), meshplot(fem) fem.mesh = meshrefine(fem, 'tri',[1:50;ones(1,50)]); subplot(2,2,2), meshplot(fem) fem.mesh = meshrefine(fem, 'subdomain', [1 2], 'rmethod', 'longest'); subplot(2,2,3), meshplot(fem) fem.mesh = meshrefine(fem, 'subdomain',1); subplot(2,2,4), meshplot(fem)</pre>				
Algorithm	The 2D algorithm is described by the steps below:				
	Pick the initial set of elements to be refined.				
	2 Either divide all edges of the selected elements in half (regular refinement), or divide the longest edge in half (longest edge refinement).				
	3 Divide the longest edge of any element that has a divided edge.				
	4 Repeat step 3 until no further edges are divided.				
	5 Introduce new points on all divided edges, and replace all divided entries in e by two new entries.				
	6 Form the new elements. If all three sides are divided, new elements are formed by joining the side midpoints. If two sides are divided, the midpoint of the longest edge is joined with the opposing corner and with the other midpoint. If only the longest edge is divided, its midpoint is joined with the opposing corner.				
Compatibility	To achieve compatibility with FEMLAB 2.3, the geometry input is automatically converted to a geometry object using the function geomobject. The geometry input can be any analyzed geometry. See geomobjectfor details.				
	The output u is obsolete from FEMLAB 2.2. Use asseminit to interpolate the solution to a new mesh.				
See Also	femmesh, geomcsg, meshinit				

Purpose	Revolve a 2D mesh object into a 3D mesh object.
Syntax	fem1 = meshrevolve(fem0,) [mesh,geom]= meshrevolve(fem,'Out',{'mesh','geom'},)
Description	<pre>fem1 = meshrevolve(fem0,) revolves the 2D geometry object in fem0.geom and the 2D mesh object in fem0.mesh, into a 3D geometry object and a 3D mesh object stored in fem1.geom and fem1.mesh, respectively, according to the given parameters.</pre>

[geom,mesh] = meshrevolve(fem, 'Out', {'geom', 'mesh'},...) returns the 3D geometry object in geom and the 3D mesh object in mesh.

Valid property/value pairs for the meshrevolve function are given in the following table. In addition, all revolve parameters are supported and are passed to revolve for creating the revolved 3D geometry object.

PROPERTY	VALUES	DEFAULT	DESCRIPTION
elrevlayers	scalar vector		Distribution of mesh element layers in revolved mesh
mcase	integer	0	Mesh case number
out	fem mesh geom	fem	Output variables

The property elrevlayers defines the distribution of the mesh element layers in the revolved mesh. If the value of elrevlayers is a scalar, it defines the number of equally distributed mesh element layers in the revolved mesh. Alternatively, if elrevlayers is a vector, it defines the distribution of the mesh element layers in the revolved mesh. The values in the vector, that are sorted and starts with 0, specifies the placements, in relative arc length, of the mesh element layers. By default, the number of element layers is determined such that the distance of each layer is equal to the mean size of the elements in the original 2D mesh.

Revolving a 2D mesh object into a 3D mesh object, the 2D vertex elements, the 2D boundary elements, the 2D triangular elements, and the 2D quadrilateral elements, are revolved into 3D edge elements, 3D quadrilateral boundary elements, 3D prism elements, and 3D hexahedral elements, respectively.

```
Create a revolved prism mesh on a torus:
Examples
```

```
fem.geom = circ2(1, 'pos', [2 0]);
fem.mesh = meshinit(fem);
fem1 = meshrevolve(fem);
```

Create a revolved hex mesh from the *zx* plane:

```
p_wrkpln = geomgetwrkpln('quick', {'zx',10});<br/>ax = [0 1;0.5 2]';<br/>fem.geom = rect2(1.5,1,'pos',[0.5 0]);<br/>fem.mesh = meshmap(fem);<br/>fem1 = meshrevolve(fem, 'angles', [-pi/3 pi/3],...<br/>'revaxis',ax, 'wrkpln',p_wrkpln);<br/>meshplot(fem1);CautionaryRevolving a triangular mesh adjacent to the revolution axis or a mesh containing a<br/>quadrilateral element with only one corner adjacent to the revolution axis is not<br/>supported.See alsomeshembed, meshextrude, femmesh, revolve
```

Purpose Smooth interior mesh vertices and improve quality of a mesh.

```
fem.mesh = meshsmooth(fem,...)
Syntax
                     mesh = meshsmooth(mesh,...)
                     fem = meshsmooth(fem, 'out', {'fem'},...)
```

Description

mesh = meshsmooth(fem, ...) improves the quality of the elements in the mesh fem.mesh by adjusting the mesh vertex positions and by swapping mesh edges.

mesh = meshsmooth(mesh,...) improves the quality of the elements in the mesh object mesh.

```
fem = meshsmooth(fem, 'out', { 'fem' }, ...) modifies the fem structure to
include the new mesh object in fem.mesh.
```

In 3D, meshsmooth is supported for meshes containing tetrahedral elements only.

The function meshsmooth accepts the following property/values:

TABLE 1-94: VALID PROPERTY/VALUE PAIRS

PROPERTY	2D	3D	VALUE	DEFAULT	DESCRIPTION
Jiggleiter	\checkmark		numeric	5	Maximum number of jiggling iterations
Qualoptim	\checkmark		off mean min optim	min	Optimization method
Out	\checkmark	\checkmark	fem mesh	mesh	Output parameters
Subdomain	\checkmark	\checkmark	integer vector all none	all	Specifies the subdomains that are smoothed

Algorithm

2D

Each mesh vertex that is not located on the boundary is moved such that the quality of the surrounding element increases. This process is controlled via the properties Jiggleiter and Qualoptim.

Jiggleiter specifies the maximum number of jiggling iterations. The default value is 5.

Qualoptim specifies the technique that is used to improve the quality of the mesh.

- Off means that no improvement operations are performed.
- Mean means that jiggling is repeated until the mean element quality does not significantly increase, or until the bound Jiggleiter is reached. Furthermore, after every third jiggling iteration, edge swapping operations are performed.

	 Min means that jiggling is repeated until the minimum element quality does not significantly increase, or until the bound Jiggleiter is reached. Furthermore, after every third jiggling iteration, edge swapping operations are performed. Optim means that a mesh quality optimizer is used. This tries to increase the minimum quality to at least 0.8. The Jiggleiter parameter has no effect in this case. 				
	3D				
	Relocation of points similar to the 2D case is combined with edge swapping operations to improve the quality of the tetrahedra.				
Examples	Create a triangular mesh of the L-shaped membrane without quality improvement and improve the quality by calling meshsmooth:				
	<pre>clear fem sq1 = square2(0,0,1); sq2 = move(sq1,0,-1); sq3 = move(sq1,-1,-1); fem.geom = sq1+sq2+sq3; fem.mesh = meshinit(fem,'jiggle','off'); q = meshqual(fem.mesh); minQual1 = min(q) fem.mesh = meshsmooth(fem); q = meshqual(fem.mesh); minQual2 = min(q)</pre>				
See Also	meshqual, femmesh, meshinit				

Purpose	Create swept mesh.
Syntax	<pre>fem.mesh = meshsweep(fem,) fem.mesh = meshsweep(fem.geom,)</pre>
Description	fem.mesh = meshsweep(fem,) returns a swept mesh derived from the 3D geometry in fem.geom.
	fem.mesh = meshsweep(geom,) returns a swept mesh derived from the 3D geometry geom.
	A swept mesh is created for each subdomain by meshing the corresponding source face, if this face is not already meshed, and sweeping the resulting face mesh along the subdomain to the opposite target face.

If the source face for a subdomain is not meshed prior to the meshsweep operation, the source face is automatically meshed with triangles using the free triangle mesher (meshinit). Then, the swept mesh consists of prism elements. To create a swept mesh with hexahedral elements, you need to mesh the source face with quadrilateral elements using either the mapped quad mesher (meshmap), or the free quad mesher (meshinit), prior to the meshsweep operation.

The function meshsweep accepts the following property/value pairs.

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Elsweeplayers	cell array		Distribution of mesh element layers in the sweep direction
Hauto	numeric	5	Predefined mesh element size
Mcase	integer	0	Mesh case number
Meshstart	mesh object	empty	Starting mesh
Out	fem mesh	mesh	Output variables
Report	on off	on	Display progress
Sourceface	cell array		Source faces
Subdomain	numeric array auto all none	auto	Specifies the subdomains that are meshed
Targetface	cell array		Target faces

TABLE I-95: VALID PROPERTY/VALUE PAIRS

The property elsweeplayers determines the distribution of the mesh element layers in the sweep direction. The value of this property is an even numbered cell array where the odd entries contain subdomain indices, either as scalar values, or as vectors with subdomain indices, and the even entries contain scalar values or vectors specifying the element layer distribution for the corresponding subdomain(s). If the element layer distribution is specified by a scalar, the element layers for the corresponding subdomain(s) are equally distributed and the number of element layers equals the value of the scalar. To get full control over the element layer distribution for a subdomain, the vector form is used. The values in the vector, that are sorted and starts with 0, specifies the distances of the element layers along the direction of the sweep for the corresponding subdomain(s).

Hauto is an integer between 1 and 9 that controls the element size in the generated mesh. The default value is 5 which means that the element size is set to 1/10 of the size of the geometry for the elements not affected by the elsweeplayers property.

Use the property subdomain to specify the subdomains to be meshed. If you use this property together with the meshstart property, the value auto means that all subdomains that are not meshed in the starting mesh are meshed, none means that no further subdomains are meshed, and all means that all subdomains are meshed (or remeshed). It is also possible to specify the subdomains to be meshed (or remeshed) using a vector of subdomain indices.

The meshstart property is used when meshing a geometry interactively. The value of this property is the starting mesh of the meshing operation.

The property sourceface is a cell array specifying the source faces for the subdomains. The value of this property is an even numbered cell array where the odd entries contain subdomain indices, either as scalar values, or as vectors with subdomain indices, and the even entries contain the source face indices for the corresponding subdomain(s).

The property targetface defines the target faces for the subdomains. The syntax of this property is the same as for the sourceface property.

If the source and/or target face is not specified for a subdomain, the software automatically tries to determine these faces.

The following criteria must be met by the input geometry object for the sweeping technique to work.

• Each subdomain must be bounded by one shell, that is, a subdomain must not contain holes that do not penetrate the source and target face.

	• There can only be one source face and one target face per subdomain.
	• The source face and target face for a subdomain must be opposite one another in the subdomain's topology.
	• The cross section along the direction of the sweep for a subdomain must be topologically constant.
	Each face about a subdomain to be swept is classified as either a source face, a target face, or a boundary face. The boundary faces are the faces linking the source and target face. The sweep algorithm can handle subdomains with multiple boundary faces in the sweep direction.
	If any of the faces about a subdomain is meshed prior to the meshsweep operation, the following must be fulfilled.
	• If the source and target faces are meshed, these meshes must match.
	• Mapped quad meshes must be applied to the boundary faces.
Algorithm	The subdomains to be meshed are processed in the following order.
	I The subdomains where the source and/or target faces are specified are swept first. These subdomains are swept in the order of increasing subdomain number.
	2 The remaining subdomains, with no specified source face or target face but with one adjacent meshed face, are swept in the order of increasing subdomain number.
	3 Finally, the remaining subdomains, with no specified source face or target face and with none or several adjacent meshed faces, are swept in the order of increasing subdomain number.
	When a swept has been generated for a subdomain, the source face of the next subdomain, that is, the subdomain adjacent to the target face of the current subdomain, is set to the target face of the current subdomain if the source and target faces for the next subdomain are not specified.
	For a subdomain with no specified source and target face, the source face is determined according to the following.
	I If not any face about the subdomain is meshed, the software determines the

I If not any face about the subdomain is meshed, the software determines the opposite face pairs for the subdomain. An opposite face pair is a pair of faces about the subdomain that are not adjacent to each other but to all other faces about the subdomain. The face in these face pairs with lowest geometric degree and face index is used as source face and the opposite face is used as target face. If no

	opposite face pairs exist for the subdomain, an error is thrown and the user is asked to specify the source face.
	2 If there is at least one meshed face about the subdomain, the source face is determined according to the following.
	a The face with lowest geometric degree and face index of the meshed faces about the subdomain that is not a boundary face of another subdomain.
	b The face with lowest geometric degree and face index of the unmeshed faces.
	c The face with lowest geometric degree and lowest face index of all faces about the subdomain.
Examples	Create a swept mesh on a cylinder geometry specifying the element layer distribution in the sweep direction.
	<pre>fem.geom = cylinder3; fem.mesh = meshsweep(fem,'sourceface',{1 3}, 'elsweeplayers',{1 logspace(0,1,11)-1}); meshplot(fem);</pre>
	Create a swept mesh on a helix shaped geometry.
	fem.geom = helix3; fem.mesh = meshsweep(fem,'sourceface',{1 1}); meshplot(fem);
	Create a mesh with both tetrahedrons and prisms using meshinit and meshsweep,
	respectively.
	<pre>fem.geom = block3 + cone3(0.3,1,pi/20,'pos',[1 0.5 0.5],</pre>
	Create a swept mesh with hexahedrons on a block geometry by meshing a source
	face prior to the meshsweep operation using the mapped quad mesher.
	<pre>fem.geom = block3; fem.mesh = meshmap(fem,'face',1); fem.mesh = meshsweep(fem,'meshstart',fem.mesh); meshplot(fem);</pre>
See Also	meshdel, meshinit, meshmap

Purpose	Reflect geometry.			
Syntax	[gm,] = mirror(g,pt,vec,)			
Description	<pre>[gm,] = mirror(g,pt,vec,) creates a mirrored copy of the geometry object g, as reflected in the plane with normal vector vec, centered at pt.</pre>			
	Property value	list for mirror.		
	TABLE I-96: VALID	PROPERTY/VALUE PA	AIRS	
	PROPERTY	VALUE	DEFAULT	DESCRIPTION
	Out	stx ftx ctx ptx	{}	Cell array of output names.
Examples	<pre>In 2D: g = rect2; gm = mirror(g,[1 1],[1 1]); figure,geomplot(g),hold on,geomplot(gm),axis equal In 3D: g = block3;</pre>			
	<pre>gm = mirror(g,[1 1 1],[1 1 1]); figure, geomplot(g), hold on, geomplot(gm), axis equal</pre>			
See Also	geomO, geom1, geom2, geom3,scale			

Purpose	Move geometry object.			
Syntax	[g,] = move(g3,x,y,z,) [g,] = move(g2,x,y,) [g,] = move(gn,x,)			
Description	[g,] = move(g3,x,y,z,) moves a 3D geometry object by the vector (x,y,z) .			
	$[g, \ldots] = move(g2, x, y, \ldots)$ moves a 2D geometry object by the vector (x, y)			
	$[g, \ldots] = move(gn, x, \ldots)$ moves an <i>n</i> D geometry object by the vector x of length <i>n</i> .			
	The function move accepts the following property/values:			
	PROPERTY	ID PROPERTY/VALUE PAIR	DEFAULT	DESCRIPTION
	Out	stx ftx ctx ptx		Output parameters
Examples	The commands below move the circle from the origin to (2,3) and plot the result. c1 = circ2; c2 = move(c1,2,3); geomplot(c2)			
See Also	geomcsg			

Purpose	Multiphysics function.
Syntax	fem1 = multiphysics(fem,) xfem1 = multiphysics(xfem,)
Description	<pre>fem1 = multiphysics(fem) combines the application modes in fem.appl to the composite system fem1.</pre>
	<pre>xfem1 = multiphysics(xfem), where xfem is a structure with a field fem,</pre>

performs the above call for all of the structures xfem.fem{:}. The results are placed in xfem1.fem. In addition the fields elemmph and eleminitmph created for each structure in xfem.fem are concatenated and placed as fields in xfem1.

The function multiphysics accepts the following property/values:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Bdl	cell array of integers integer vector NaN	all boundaries	Only affect the indicated boundaries in fem. bnd
Defaults	on off	off	Return default fields
Diff	cell array of ga, f, r, or g 'on ' 'off'	on for general output form, otherwise of f	Differentiate these coefficients
Edl	cell array of integers integer vector NaN	all edges	Only affect the indicated edges in fem.edg
Outform	'coefficient' 'general' 'weak'	the most general output form of the application modes in fem.app1	Output form
Outsshape	positive integer NaN	generated from the application modes in fem.appl	Output sshape. NaN is the same as not providing Outsshape. Can also be given for each structure in xfem.fem, by giving a cell array of values
Pdl	cell array of integers integer vector NaN	all points	Only affect the indicated points in fem.pnt
Sdl	cell array of integers integer vector NaN	all subdomains	Only affect the indicated subdomains in fem.equ
Shrink	on off	on	Shrink coefficients to most compact form
Simplify	on off	on	Simplify expressions

TABLE I-98: VALID PROPERTY/VALUE PAIRS

The properties Diff, Outform, Outsshape, Rules, and Simplify can also be specified by using fields in the fem structure. The default value for Diff is 'off' if the resulting model has coefficient or weak form and 'on' if it has general form.

For calls of the form xfem1 = multiphysics(xfem,...), The properties Sdl/ Bdl/Edl/Pdl/Outsshape/Outform/Diff/Simplify can be given as cell arrays indicating the properties' values for the different elements of xfem.fem. In this case the number of elements of the cell arrays giving the values must be the same as the number of elements of xfem.fem. For Sdl/Bdl/Edl/Pdl/Outsshape, the value NaN may be given if no value is actually intended to be passed.

For calls of the form xfem1 = multiphysics(xfem,...) Where the property Out is specified, the value 'fem' means xfem1 as described above, but 'equ'/'bnd'/ 'edg'/'pnt'/'dim'/'form'/'shape'/'sshape'/'border'/'var' are returned as cell arrays containing these properties for each of the elements of xfem.fem, and 'elemmph'/'eleminitmph' are concatenations of the individual resulting properties.

AlgorithmThe description below relates to the call fem1 = multiphysics(fem) unless
otherwise stated. The applications are specified as a cell array in the field appl in the
fem structure: fem.appl={a1 a2 ...}.

The notations ***, XXX and xxx used below means the fields equ, bnd, edg, and/ or pnt. *** denotes any of these fields, XXX is used to denote the field which corresponds to the largest dimension (usually equ), while xxx denotes the fields corresponding to lower dimensions (usually bnd, edg, and pnt). Not all of them are always present, rather their presence is dependent on the geometry of the domain and the type of problem being solved.

The table below describes the fields in the appl structure.

TABLE I-99: APPL STRUCTURE FIELDS

FIELD INTERPRETATION			
appl.assign	Application mode variable name assignments		
appl.bnd	Boundary coefficients/application data		
appl.border	Cell array with strings on/off or a corresponding logical vector, one for each solution component		
appl.dim	Cell array with names of the solution components or an integer specifying the number of solution components		
appl.edg	Edge coefficients/application data		

FIELD	INTERPRETATION		
appl.elemdefault	A string indicating what kind of element is the default in this application		
appl.equ	Subdomain coefficients/application data		
appl.form	Problem form: coefficient, general, or weak		
appl.mode	Application mode		
appl.pnt	Point coefficients/application data		
appl.shape	Shape functions. A cell array of shape function objects.		
appl.sshape	Element geometry order. An integer.		
appl.***.shape	Pointers to appl.shape		
appl.XXX.usage	Activate/deactivate domain		
appl.***.gporder	Order of numerical quadrature		
appl.***.cporder	Constraints discretization order		
appl.XXX.init	Initial conditions		
appl.var	Application mode specific variables		

appl.assign contains the application mode variable name assignments. It is a cell array of alternating fixed names and assigned names for the application mode variables. The default is an empty cell array.

appl.bnd is a structure with fields describing the boundary data. The structure contains one field for each of the application-specific boundary parameters, with the field name equal to the parameter name. Each field should be a cell array containing data for the boundaries. appl.bnd also contains a field type, which is a cell array with strings describing the boundary type for each boundary. If only one boundary type is available in the application mode, the type field may be omitted.

appl.dim provides the dependent variable names. The default is obtained from the application mode for physics modes. For PDE modes with no mode field and with a numeric appl.dim, default variable names are substituted according to the global position in the system.

appl.elemdefault contains a string indicating what kind of element is the default in this application. This method is used to generate the defaults for appl.XXX.gporder, appl.XXX.cporder, appl.shape, appl.XXX.shape, and appl.sshape. See elemdefault for a list of valid strings. appl.edg is a structure with fields describing the edge data. The structure contains one field for each of the application-specific edge parameters, with the field name equal to the parameter name. Each field should be a cell array containing data for the edges.

appl.equ is a structure with fields describing the PDE data on subdomains. The structure contains one field for each of the application-specific PDE parameters, with the field name equal to the parameter name. Each field should be a cell array containing data for the subdomains.

appl.mode is a string with the name of the application mode or an application mode object. If the appl.mode field is omitted, the application structure can be used to describe ordinary coefficient/general/weak form PDE problem. This is useful in the definition of multiphysics problems, where coefficient/general/weak form models can be combined with application mode models. See the *COMSOL Modeling Guide* for a description of the application-specific data for each application mode.

appl.pnt is a structure with fields describing the point data. The structure contains one field for each of the application-specific point parameters, with the field name equal to the parameter name. Each field should be a cell array containing data for the points.

appl.shape is a cell array of shape function objects.

appl.sshape is an integer giving the order of geometry approximation.

The fields fem.XXX.shape and fem.xxx.shape are ind-based cell arrays of vectors pointing to elements of appl.shape. Indicates which shape functions to use in each domain group. An empty vector indicates that no shape functions are used in this domain group. Zero indicates that the affected domain group should use defaults or inherit shape functions. Appl.XXX.shape takes defaults, whereas appl.xxx.shape inherits from appl.XXX.shape. Where there is a conflict over which domain group in appl.XXX.shape to inherit from, the first appropriate group is used.

The fields fem.appl.xxx.usage are ind-based cell arrays of ones and zeros indicating domain group usage. Wherever a zero entry exists, the information in appl.xxx.shape is ignored when forming fem.xxx.shape.

The fields app1.XXX.gporder and app1.xxx.gporder indicate the order of quadrature formula to use in the different domain groups. In fully expanded form it is a cell array where each element is a cell array (of positive integers) of length

equal to the number of dependent variables (excluding submode variables) in this mode. Defaulting and inheritance can be induced by using 0. The inherited order is the maximum order used in the objects from the XXX level in contact with the group at the xxx level. Where different elements within a domain group would inherit different orders, some domain group splitting takes place. This field is not present in the appl.pnt field.

The fields appl.XXX.cporder and appl.xxx.cporder indicate domain group constraints discretization order. Behaves exactly as gporder.

The fields appl.XXX.init indicate domain group initial conditions. For format see asseminit.

appl.var contains the application scalar variables. The default is obtained from the application mode for physics modes.

The composite system is created by appending the subsystems in the order they are specified in fem.appl. The affected fields in the fem1 structure are dim, form, equ, bnd, edg, pnt, var, elemmph, eleminitmph, shape, sshape, and border. All the other fields in fem are copied to fem1. In the description below the notation *** is used to represent any or all of the fields equ, bnd, edg, and/or pnt.

The dim field of the composite system, fem1.dim, is a cell array of the dependent variable names:

```
fem1.dim={fem.appl{1}.dim{:}, fem.appl{2}.dim{:}, ...}
```

The default names ('u1', 'u2', ...) Are used for subsystems with integer dim fields and no mode field.

The form of each subsystem may be converted by using flform, in the direction 'coefficient' -> 'general' -> 'weak'. The default form of the composite system, fem1.form, is the first form which all the subsystems may attain, possibly applying flform to some subsystems. The output form can be forced by using the property outform.

First, each application structure is converted to an FEM structure with all the above-mentioned fields. Then the fields in fem1.*** are computed from the corresponding fields in the subsystems according to the table below. The numbers

QUANTITY	COMPOSITE SYSTEM
f, γ, g, r, init weak, dweak, constr gporder, cporder	$\begin{bmatrix} f_1 \\ f_2 \\ \dots \end{bmatrix}$
$c, d_a, e_a, \alpha, \beta, a, q, h$	$\begin{bmatrix} c_1 & 0 & 0 \\ 0 & c_2 & 0 \\ 0 & 0 & \dots \end{bmatrix}$
var, varu, vart	$\left[\mathrm{var}_1 \ \mathrm{var}_2 \ \ldots \right]$

after the coefficient names refer to the subsystems in the order they are specified in fem.appl.

The fields fem.***.expr are kept unchanged as far as possible; the only difference being the permutation of ind-groups to suit a new ind vector if one is generated. For ind groups where a particular expr variable is undefined, the entry [] is used.

Note that the coefficients in the second row of the table are *weakly* coupled in the sense that the corresponding coefficients in the composite system are block diagonal. This puts some limitations on the coupling between the subsystems. By using general form, however, there are no limitations on the composite system except for the da coefficient. The resulting stronger couplings are obtained by using a call to femdiff with the full system resulting from the composition of the subsystems. The properties Diff, Rules, and Simplify control or supplement the call to femdiff. Further couplings between applications can be introduced by the method global_compute which is called for each application before femdiff.

Elements of fem1.***.(f/ga/g/r/weak/dweak/constr/c/da/ea/al/be/a/ q/h) which correspond to subsystems for which the corresponding field does not exist are '0'. Elements of fem1.***.init in such cases are empty strings. Elements in fem1.***.(gporder/cporder) in such cases are 1. The fields fem1.elemmph and fem1.eleminitmph are obtained by concatenating the contents of the results of calling elem_compute for each application. The field fem1.shape is obtained by concatenating the contents of the fields fem.appl{:}.shape. Duplicate shape functions are removed and the fields fem.***.shape are adjusted to take account of this. The field fem1.sshape is obtained by taking the maximum of the fields fem.appl{:}.sshape. The resulting value is overridden if the property Outsshape is given. The fem1.border field is always 1 because coefficients for boundary conditions that are not used due to **border** being on or off in the application mode are set to zero and can be "applied".

Suppose multiphysics has been called previously, and fem is the result of such a call. If changes are made to fem.equ and it is wished to keep them (that is not allow changes to be written over when multiphysics is called again), it is possible to restrict the set of subdomains for which multiphysics writes over fem.equ. Thus giving the property Sdl the value [1 2] in a call to multiphysics results in the coefficients for subdomains l and 2 being "refreshed" from the applications, but all the coefficients in fem.equ relating to other subdomains being kept and copied into fem1.equ. The same principle holds for bnd/edg/pnt using the properties Bdl/Edl/Pdl.

The model "Resistive Heating" uses the multiphysics function. Note that the structure fem contains the data that is *common* for the subsystems, that is, the geometry and the mesh. The electrical subsystem and the heat transfer subsystem are specified in the application structures a1 and a2, and the multiphysics function is used to combine them.

```
clear fem a1 a2
fem.geom = geomcsg({square2(0,0,1)});
fem.mesh = meshinit(fem);
a1.mode = 'ConductiveMediaDC';
a1.assignsuffix = ' dc';
a1.bnd.V0 = \{0.1 \ 0 \ 0\};
a1.bnd.type = {'V' 'nJ0' 'V0'};
a1.bnd.ind = [1 2 2 3];
a1.equ.init = '0.1*(1-x)';
a1.equ.T0 = 293;
a1.equ.T = 'T';
a1.equ.alpha = 0.0039;
a1.equ.res0 = 1.754e-8
a1.equ.sigtype = 'heat';
a2.mode = 'HeatTransfer';
a2.assignsuffix = '_ht';
a2.bnd.T0 = \{300 \ 0\};
a2.bnd.type = {'T' 'q0'};
a2.bnd.ind = [1 2 2 1];
a2.equ.rho = 8930;
a2.equ.C = 340;
a2.equ.k = 384;
a2.equ.Q = 'Q dc';
a2.equ.init = 300;
fem.appl = \{a1 \ a2\};
fem = multiphysics(fem);
fem.xmesh=meshextend(fem);
```

Example

	<pre>fem.sol = femtime(fem,'tlist',linspace(0,3000,41),</pre>			
Diagnostics	All variables in the appl{:}.dim fields must be unique. If any appl{:}.dim has not been provided, no other appl{:}.dim may collide with the defaults. If they do, an error message is generated.			
	If any form appl{:}.form is more general than Outform, an error message is generated.			
Compatibility	To simplify the output of the multiphysics functions in FEMLAB 3.0, its defaults has been changed by the introduction of the properties Shrink and Defaults. The compatibility problems typically occur when you perform changes to the data generated by multiphysics. For example, modifying the α coefficient			
	fem=multiphysics(fem); fem.equ.al{1}{2,1}=; fem.equ.al{1}{3,1}=;			
	may not work as in FEMLAB 2.3 because alpha is often just empty. The code above assumes that fem.equ.al{1} is a 3-by-3 cell array. To obtain fully backward compatible output, use			
	fem=multiphysics(fem,'shrink','off','defaults','on');			
	which makes the above example work.			
	The properties out and rules are obsolete from FEMLAB 3.0.			
	The property Idl is obsolete in FEMLAB 2.2 and later versions.			
	The fields fem.init and fem.usage are no longer constructed. They are superseded by the fields fem.***.init and fem.***.shape, respectively. Fem.***.init is constructed for all fields fem.***, but many entries contain just empty strings.			
See also	femdiff, flform, multiphysics			

Purpose	Convert a PDE Toolbox geometry description to geometry model.			
Syntax	draw = pde2draw(gd,ns,sf)			
Description	fem = pde2draw(gd,ns,sf) converts a PDE geometry description matrix gd, name space matrix ns, and set formula sf to a geometry model draw. The two last arguments are optional.			
See Also	geom0, geom1, geom2, geom3,pde2geom,pde2fem			

Purpose	Convert a PDE Toolbox geometry to a geometry object.			
Syntax	g = pde2geom(dl)			
Description	g = pde2geom(d1) converts a PDE Decomposed Geometry matrix d1 to a geometry object.			
See Also	geom0, geom1, geom2, geom3,pde2draw,pde2fem			

Purpose	Convert a PDE Toolbox problem description to an FEM structure.		
Syntax	fem = pde2fem(g,b,p,e,t,c,a,f,d)		
Description	<pre>fem = pde2fem(g,b,p,e,t,c,a,f,d) converts a PDE problem described by the Decomposed Geometry matrix g, Boundary Condition matrix b, Mesh data p, e, t, and PDE Coefficients c, a, f, d to an FEM structure.</pre>		
Compatibility	PDE Coefficient and Boundary Coefficient M-file functions are not supported.		
See Also	pde2draw, pde2geom, femstruct		

Purpose	Constructor functions for point objects.				
Syntax	<pre>p3 = point3(x,y,z) p3 = point3(vtx,vtxpre,edg,edgpre,fac,mfdpre,mfd) [p3,] = point3(g3,) p2 = point2(x,y) p2 = point2(vtx,edg,mfd) [p2,] = point2(g,) p1 = point1(x) p1 = point1(vtx) [p1,] = point1(g,)</pre>				
Description	p3 = point3(x,y,z) creates	a 3D single	point object with coordinate (x, y, z) .	
	p3 = point3(vtx,vtxpre,edg,edgpre,fac,mfdpre,mfd) creates a 3D point geometry object p3 from the arguments vtx, vtxpre, edg, edgpre, fac, mfdpre, mfd. The arguments must define a valid 3D point object. See geom3 for a description of the arguments.				
	p3 = point3(g3) coerces the	e 3D geomet	ry object g3 to a 3D point object p3.	
	<pre>p2 = point2(x,y) creates a 2D point object consisting of a single point with coordinates (x,y). p2=geom2(vtx,edg,mfd,) creates a 2D point object from the properties v edg, and mfd. The arguments must define a valid 2D point object. See geom2 information on vtx, edg, and mfd. p2 = point2(g2) coerces the 2D geometry object to a point object.</pre>				
	p1 = point1(x) creates a 2D point object consisting of a single point with coordinate x.				
	p1 = point1(vtx,) creates a 1D point object from vtx. The arguments must define a valid 1D point object. See geom1 for information on vtx.				
	<pre>[p1,] = point1(g,) coerces the 1D geometry object to a point object.</pre>				
	The coercion functions [p1,] = point1(g1,), [p2,] = point2(g2,), and [p3,] = point3(g3,) accept the following property/values:				
	TABLE I-100: VALID	PROPERTY/VALUE F	PAIRS		
	PROPERTY	VALUE	DEFAULT	DESCRIPTION	
	Out	stx ftx ctx ptx	{}	Cell array of output names.	

	See geomesg for more information on geometry objects.	
	The nD geometry object properties are available. The properties can be accessed using the syntax get(object,property). See geom for details.	
Example	The commands below create a 2D point object with four points and plot the result.	
	<pre>c1 = circ1; p1 = point2(c1); geomplot(p1)</pre>	
Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported.	
See Also	curve2, curve3, face3, geom0, geom1, geom2, geom3, geomcsg	

Purpose	Fast solution of Poisson's equation on a rectangular grid.			
Syntax	<pre>fem.sol = poisson(fem,)</pre>			
Description	fem.sol = poisson(fem) solves Poisson's equation with Dirichlet boundary conditions on a regular rectangular grid. A combination of sine transforms and tri-diagonal solutions is used for increased performance.			
	The boundar	y conditions must specif	fy Dirichlet cond	itions for all boundary points.
	The mesh fem.mesh must be a regular triangular mesh on a rectangular geometry, for example, generated by meshpoi. Details on the mesh representation can be found in the entry on femmesh.			
	The algorithm handles the f and γ coefficients. The c coefficient is set to one. All other coefficients are ignored. Only a one-dimensional system can be handled, that is, only fem.dim = 1 is allowed.			
	The weak solution form must not be used when generating the extended me Therefore set fem.sol='coefficient' before calling meshextend. Apart from round-off errors, the result should be the same as using femlin. The function poisson accepts the following property/value pairs: TABLE 1-101: VALID PROPERTY/VALUE PAIRS			-
				-
	PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
	Const	cell array		Definition of constants
	Nullfun	flnullorth flspnull	flnullorth	Null space function
	Out	fem sol u	sol	Output variables
Diagnostics	If fem.geom	does not seem to descri	be a rectangle, ar	n error is displayed.
Cautionary	The function elements.	only works in 2D. The	function only su	pports linear Lagrange
Compatibility	The propertie	es Context and Tpoin	t are obsolete fro	om FEMLAB 3.0.
	The property	Variables has been re	enamed to Const	in FEMLAB 2.3.
See Also	meshpoi			

Strang, Gilbert, *Introduction to Applied Mathematics*, Wellesley-Cambridge Press, Cambridge, MA, 1986, pp. 453–458.

Purpose	Create polygons.
Syntax	<pre>c = poly1(x,y) s = poly2(x,y)</pre>
Description	s = poly2(x, y) creates a 2D solid object s in the form of an solid polygon with vertices given by the vectors x and y.
	c = poly1(x,y) creates a 2D curve object c in the form of an closed polygon with vertices given by the vectors x and y.
	See geomesg for more information on geometry objects.
Example	The commands below create a regular n -gon (n =11) and plot it.
	<pre>n = 11 xy = exp(i*2*pi*linspace(0,1-1/n,n)); p = poly1(real(xy),imag(xy)); geomplot(p)</pre>
Cautionary	poly1 and poly2 always creates closed polygon objects. To create open polygon curves, use line1 and line2.
See Also	arc1, arc2,circ1, circ2,ellip1, ellip2,geomcsg,line1, line2

Purpose	Shorthand command for animation in 1D, 2D and 3D.
Syntax	postanim(fem,expr,) M = postanim(fem,expr,) % MATLAB only
Description	<pre>postanim(fem,expr,) plots an animation of the expression expr. The function accepts all property/value pairs that postmovie does. In 1D, this command is just shorthand for the call postmovie(fem,'liny',expr, 'linstyle','bginv',)</pre>
	and in 2D, it is shorthand for
	postmovie(fem,'tridata',expr, 'tribar','on', 'geom','on', 'axisequal','on',)
	and in 3D, this command is just shorthand for
	postmovie(fem,'slicedata',expr, 'slicebar','on', 'geom','on', 'axisequal','on',)
	M = postanim(fem,expr,) additionally returns a matrix in MATLAB movie format. Alternatively, in COMSOL Script, store the animation in a file using the postmovie property Filename.
	If you want to have more control over your animation, use postmovie instead of postanim.
Cautionary	When you are replaying a movie that has been stored in a matrix M, you should explicitly provide a figure handle to the movie command.
	M = postanim(fem,expr,) movie(gcf,M)
	Otherwise the animation does not look good.
Compatibility	The syntax of the command is not compatible with its corresponding FEMLAB 2.1 syntax.
See Also	postplot, postsurf, postcont, postlin, postarrow, postarrowbnd, postflow, postslice, postiso, posttet

Purpose	Shorthand command for subdomain arrow plot in 2D and 3D.
Syntax	postarrow(fem,expr,) h = postarrow(fem,expr,)
Description	postarrow(fem, expr,) plots a subdomain arrow plot for the expressions in the cell array expr. In 2D, expr has length 2 or 3, and in 3D, it has length 3. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call
	postplot(fem,'arrowdata',expr, 'geom','on', 'axisequal','on',)
	h = postarrow(fem,expr,) additionally returns handles to the plotted handle graphics objects.
	If you want to have more control over your arrow plot, use postplot instead of postarrow.
See Also	postplot, postanim, postsurf, postcont, postlin, postarrowbnd, postflow, postprinc, postprincbnd, postslice, postiso, posttet

Purpose	Shorthand command for boundary arrow plot in 2D and 3D.
Syntax	<pre>postarrowbnd(fem,expr,) h = postarrowbnd(fem,expr,)</pre>
Description	postarrowbnd (fem, expr,) plots a boundary arrow plot for the expressions in the cell array expr. In 2D, expr has length 2 or 3, and in 3D, it has length 3. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call
	postplot(fem,'arrowbnd',expr, 'geom','on', 'axisequal','on',)
	h = postarrowbnd(fem,expr,) additionally returns handles to the plotted handle graphics objects.
	If you want to have more control over your arrow plot, use postplot instead of postarrowbnd.
See Also	postplot, postanim, postsurf, postcont, postlin, postarrow, postflow, postprinc, postprincbnd, postslice, postiso, posttet

Purpose	Shorthand command for contour plot in 2D.
Syntax	<pre>postcont(fem,expr,) h = postcont(fem,expr,)</pre>
Description	<pre>postcont(fem,expr,) plots a contour plot for the expression expr. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call postplot(fem, 'contdata',expr, 'contbar','on', 'geom','on', 'axisequal','on',)</pre>
	h = postcont(fem,expr,) additionally returns handles to the plotted handle graphics objects.
	If you want to have more control over your contour plot, use postplot instead of postcont.
Example	Plot the contours of the solution to the equation $-\Delta u = 1$ over a unit circle. Use Dirichlet boundary conditions $u = 0$ on $\partial \Omega$. clear fem
	<pre>fem.geom = circ2; fem.mesh = meshinit(fem); fem.mesh = meshrefine(fem); fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = 1; fem.xmesh = meshextend(fem); fem.sol = femstatic(fem); postcont(fem,'u')</pre>
Compatibility	The syntax of the command is not compatible with its corresponding FEMLAB 2.1 syntax.
See Also	postplot, postanim, postsurf, postlin, postarrow, postarrowbnd, postflow, postprinc, postprincbnd, postslice, postiso, posttet

Purpose	Get coordinates in a model.
Syntax	<pre>coord = postcoord(fem,)</pre>
Description	<pre>coord = postcoord(fem,) returns global coordinates in a model by specifying, for example, a boundary and the number of points on the boundary.</pre>
	To specify start points for particle tracing in postplot, property/values to

postcoord can be specified in the postplot property partstart.

Valid property/value pairs for postcoord are given in the following table.

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
coord	cell array of double vectors		Coordinates where scalar expansion is used
dl	integer vector	all domains	Domain list
edim	-1 1	- 1	Element dimension
frame	string	spatial frame	Coordinate frame
geomnum	positive integer	1	Geometry number
grid	positive integer or vector of domain parameters		Domain parameters
mcase	non-negative integer		Mesh case
npoints	non-negative integer		Number of subdomain points, picked from mesh vertices (edim=-1)
solnum	integer vector all end	See below	Solution number
т	double vector	See below	Time for evaluation
U	solution object solution vector scalar	fem.sol or O	Solution(s) for evaluation

TABLE I-102: VALID PROPERTY/VALUE PAIRS

The properties Frame, Solnum, T, and U are only used when the model fem contains moving meshes. The default behavior of Solnum and T are described in postinterp.

Examples

% Set up a 2D geometry with a mesh and an extended mesh clear fem fem.geom = circ2+rect2; fem.mesh = meshinit(fem);

	<pre>fem.xmesh = meshextend(fem);</pre>
	% Get the coordinates of 17 evenly spaced points on % boundaries 1,2,4 coord = postcoord(fem,'edim',1,'grid',17,'dl',[1,2,4]);
Compatibility	This function was introduced in COMSOL Multiphysics 3.3.
See Also	postinterp, postplot

Purpose	Cross-section plot.
Syntax	postcrossplot(fem,cdim,dom,) h = postcrossplot(fem,cdim,dom,) [h,data] = postcrossplot(fem,cdim,dom,)
Description	postcrossplot(fem,cdim,dom,) displays a plot of an expression, including an FEM solution, in one or several cross sections of the geometry, with space dimension cdim and defined by dom (coordinates) or on mesh elements of space dimension cdim, specified by dom (domain list). The argument dom is therefore either a number of points to specify a cross section or a list of domains. To specify a cross section, dom must be one of the following, where sdim denotes the space dimension of the geometry:
	• An sdim-by-2 matrix to specify a cross-section line (the line between the two points in the columns of dom) in 2D and 3D. Used with properties lindata and surfdata. Here, cdim must be 1.
	• A 3-by-3 matrix to specify a cross-section plane (the plane containing all three points in the columns of dom) in 3D. Used with property surfdata. Here, cdim must be 2.
	• An sdim-by-np matrix to specify np points for point plots. Used with property pointdata. Here, cdim must be 0.
	To specify a list of domains (geometry vertices, geometry edges, geometry boundaries, or geometry subdomains), dom must be:
	• A 1-by-nd integer matrix, where nd is the number of domains with space dimension cdim (vertices, edges, boundaries, or subdomains) to plot on. When used with surfdata in 3D, nd must be one.
	In 1D, if all entries in dom are integers greater than or equal to 1, they are interpreted as indices to geometry vertices. Otherwise, they are interpreted as coordinates. To specify a coordinate which is an integer greater than or equal to 1, use the property pointtype set to coord.
	The expressions that are plotted can be COMSOL Multiphysics expressions involving variables, in particular <i>application mode variables</i> .
	The following plot types can be made:
	Point plots (ID, 2D, 3D) Plot of an expression on any geometry vertex or arbitrary point in the subdomains of the geometry. This is most useful when there are several

solutions, in which case this plot shows the value of the expression in the selected points for the different solutions. If there is only one solution, the values in the specified points are displayed in the x-axis range [0 1].

Line plots on domains (1D, 2D, 3D) Plot of an expression on a set of connected 1D domains (edges in 3D, boundaries in 2D and subdomains in 1D). If Linxdata is not specified, this is done by folding out the arc length of the 1D domain to the x-axis of the resulting plot, and letting the value of the expression be set on the x-axis. If dom contains more than one domain, the different arc lengths are just added to each other on the x-axis. In this case, the domains have to be connected and so that not more than two selected domains meet in the same vertex.

If Linxdata is specified, this is the quantity on the x-axis in the resulting plot. Using Linxdata, you can project cross sections to, e.g., the x-axis by setting Linxdata to x.

The direction of the path is so that the start point is the point along the path with lowest geometry vertex number. If the selected domains form a closed curve, so that this point is also the end point, the direction is in the direction of the domain with lowest number.

If there are several solutions (that is, Solnum or T is a vector), the curves for the different solutions can be either plotted in the same *x*-*y*-plot or can be extruded along the third axis to generate a surface. This is controlled by specifying either the property Lindata or Surfdata.

Line plots on cross sections (2D, 3D) In 2D and 3D, plot of an expression along a straight line, defined between the two points in dom, in the geometry. The points in dom are regarded as the end points of the cross-section line and are the x-axis limits in the resulting plot, hence if the line between the two specified points do not intersect the geometry, the resulting plot will be empty.

Surface plots on domains (2D, 3D) Plot of an expression on a boundary in 3D or on a set of subdomains in 2D. In 3D, if Surfxdata and Surfydata are not specified, the boundary is plotted in an xy-plane where x and y correspond to the (s,t)-parameters of the boundary. If Surfxdata and Surfydata are specified, these represent the quantities on the x- and y-axis in the resulting plot. Using Surfxdata and Surfydata, you can project a cross section to, for example, the xy-plane of the geometry by setting Surfxdata to x and Surfydata to y. If there are several solutions, all plots for the different solutions are displayed along the third axis, as a slice plot. In 3D, dom must be a single integer.

Surface plots on cross sections (3D) Plot of an expression on one or more 2D cross-section planes, defined by the three points in dom, of the 3D geometry. If more than one cross section is selected, or only one cross section is selected but there are several solutions, all plots for the different cross sections/solutions are displayed along the third axis, as a slice plot. The cross-section plane is the plane containing the three points in the columns of dom.

For line plots, if more than one curve is plotted (either one cross section and several solutions or vice versa), the different curves can be either plotted in the same x-y-plot or can be extruded along the third axis to generate a surface. This is controlled by specifying either the property Lindata or Surfdata.

h = postcrossplot(fem,cdim,dom,...) additionally returns handles or a
postdata structure (depending on the value of the property Outtype) to the plotted
objects.

Valid property/value pairs for postcrossplot are given in the following table, where the columns **S**, **L**, and **P** denotes if the property has effect on surface, line and point plots, respectively.

PROPERTY NAME	s	L	Р	PROPERTY VALUE	DEFAULT	DESCRIPTION
Axistype	\checkmark	\checkmark	\checkmark	cell array of strings lin or log		X-, Y- and Z-axis types
Const		\checkmark	\checkmark	cell array		Definition of constants
Cont	\checkmark	\checkmark	\checkmark	off on	off	Make output continuous
Crosslicecs	\checkmark			local global	global	Coordinate system to plot cross-section slices in
Geom	\checkmark			off on		Show geometry contour
Frame	\checkmark	\checkmark	\checkmark	string	spatial frame	Coordinate frame
Geomnum	\checkmark	\checkmark	\checkmark	integer (or vector of integers when cdim=2 and dom is 3-by-3)	I	Geometry number
Lincolor		\checkmark	\checkmark	colorspec cell array of colorspec	cycle	Line color
Lindata		\checkmark		string		Expression to plot
Linewidth			\checkmark	numeric	0.5	Line width

PROPERTY NAME	s	L	Р	PROPERTY VALUE	DEFAULT	DESCRIPTION
Linlegend		\checkmark	\checkmark	off on	off	Color legend
Linmarker		\checkmark	\checkmark	marker specifier cell array of marker specifiers	none	Line marker
Linstyle		\checkmark	\checkmark	symbol cell array of symbols	'_'	Line style
Linxdata		\checkmark		string		Line x-axis expression
Markersize		\checkmark	\checkmark	integer	6	Size of markers
Npoints		\checkmark		integer	200	Number of points on each line, when dom is a cross section
Outtype	\checkmark	\checkmark	\checkmark	handle postdata	handle	Output type
Phase	\checkmark	\checkmark	\checkmark	scalar	0	Phase angle
Pointdata			\checkmark	string		Expression to plot
Pointtype			\checkmark	coord vertex	vertex	Point plot type when cdim=0 and dom is integer(s).
Pointxdata			\checkmark	string		Point plot x-axis expression
Refine	\checkmark	\checkmark		integer auto	See posteval	Refinement of element in evaluation
Sdl	V			integer vector or cell array of integer vectors	all	Subdomain list for cross-section slice plots
Solnum	\checkmark	\checkmark	\checkmark	integer vector all end	all	Solution numbers
Spacing	\checkmark	\checkmark		integer	I	Number of planes/lines or vector with distances, when dom is a cross section
Surfbar	\checkmark			off on	off	Color bar
Surfdata	\checkmark	\checkmark		string		Expression to plot
Surfdlim	\checkmark			[min max]	full range	Surface plot color limits
Surfedgestyle	V			flat interp none bg bginv colorspec	none	Triangle edge style

PROPERTY NAME	s	L	Ρ	PROPERTY VALUE	DEFAULT	DESCRIPTION
Surffacestyle	\checkmark			flat interp none bg bginv colorspec	interp	Triangle face style
Surfmap	\checkmark			colormap		Colormap
Surfxdata	\checkmark			string		Surface x-axis expression
Surfydata	\checkmark			string		Surface y-axis expression
т	\checkmark	\checkmark	\checkmark	vector		Times for evaluation
U	\checkmark	\checkmark	\checkmark	solution object or vector	fem.sol or O	Solution for evaluation

In addition, the common plotting properties listed under femplot are available.

If the field fem.sol.u does not exist and the property U is not specified, expressions not depending on the solution can still be plotted.

The notation colorspec in the value column denotes a *color specification*. See postplot for a description of this.

The property Phase is described in posteval.

Examples

```
3D Example
clear fem
fem.geom = geomcsg({cylinder3});
fem.mesh = meshinit(fem);
fem.equ.c = 1; fem.equ.f = 1; fem.equ.da = 1;
fem.bnd.h = 1;
fem.shape = 2;
fem.xmesh = meshextend(fem);
fem.sol = femtime(fem, 'tlist',0:0.01:0.1);
```

Plot solutions on a cross section:

Plot fourth solution on five cross sections with geometry boundaries:

Plot on a boundary 6 for the last time value:

```
postcrossplot(fem,2,6,'surfdata','ux','cont','on','solnum',11)
```

Compare this with

```
postplot(fem,'tridata','ux','cont','on','bdl',6,'geom','on',...
'solnum',11)
```

Plot on a boundary 6 for some time values with overlaid mesh:

postcrossplot(fem,2,6,'surfdata','ux','cont','on',... 'surfedgestyle','k','refine',1,... 'solnum',[1,4,7,11])

Plot along a line intersecting the geometry:

linpts = [-1 1;-1 1;0 1];
postcrossplot(fem,1,linpts,'lindata','u','npoints',100)

Same but with time extrusion:

```
postcrossplot(fem,1,linpts,'surfdata','u','npoints',100,...
'camlight','on')
```

Plot along some connected edges:

```
postcrossplot(fem,1,[4 5 8 11],'lindata','t1x')
% Compare this with the postplot call
postplot(fem,'lindata','t1x','edl',[4 5 8 11],'linbar','on',...
'geom','off')
```

Point plot on n points in geometry:

```
n = 30;
pts = [linspace(-1,1,n);linspace(0,1,n);linspace(0,1,n)];
postcrossplot(fem,0,pts,'pointdata','u')
```

2D Examples

Time-dependent problem (Heat equation)

```
clear fem
fem.geom = geomcsg({rect2});
fem.mesh = meshinit(fem);
fem.equ.c = 1; fem.equ.f = 1; fem.equ.da = 1;
fem.bnd.h = 1;
fem.shape = 2;
fem.xmesh = meshextend(fem);
fem.sol = femtime(fem, 'tlist',0:0.01:0.1);
```

Plot solution along the diagonal for all time-steps:

postcrossplot(fem,1,[0 1;1 0],'lindata','u')

Plot solution at time step 4 in several parallel cross sections:

```
postcrossplot(fem,1,[0 1;1 0],'lindata','u*x','solnum',4,...
```

	'spacing',5,'lincolor','r')
	Plot along three boundaries for the first five time steps:
	postcrossplot(fem,1,[1 2 3],'lindata','ux','cont','on', 'solnum',1:5)
	Same but with time-extrusion:
	postcrossplot(fem,1,[1 2 3],'surfdata','ux','cont','on', 'solnum',1:5)
	Make point plot of square of solution on three points in geometry:
	pts = [0.2 0.3 0.6;0.1 0.7 0.2]; postcrossplot(fem,0,pts,'pointdata','u^2')
Compatibility	In FEMLAB 3.0a, extrusion plots, i.e., when plotting for several solutions (Solnum or T is a vector), cdim is 1, and Surfdata is used, can only be made for plots where the extrusion axis represents the solution. Extrusions cannot be made between parallel lines for cross-section line plots. Also, all line plots are all plotted in the x-y plane, also for several solutions and for several parallel cross-section lines.
	The property Variables has been renamed to Const in FEMLAB 2.3.
See Also	postplot, postinterp

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Purpose	Plot a post data structure.
Syntax	postdataplot(pd,) h = postdataplot(pd,)
Description	postdataplot(pd,) displays a plot of a <i>post data</i>

postdataplot(pd,...) displays a plot of a post data structure, typically returned by posteval. The function supports plotting postdata structures with element dimension 1 or 2, corresponding to the posteval property Edim.

h = postdataplot(pd,...) additionally returns handes to the plotted objects.

The function postdataplot accepts the following property/values:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Colorbar	off on	on	Display a color scale
Colormap	colormap	jet(1024)	The color map
Edgestyle	flat interp none bg bginv colorspec	none	Surface edge style
Facestyle	flat interp none bg bginv colorspec	interp	Surface face style

TABLE I-104: VALID PROPERTY/VALUE PAIRS

In addition, the common plotting properties listed under femplot are available.

The notation colorspec in the value column denotes a *color specification*. See postplot for details.

Compatibility The function postdataplot was introduced in COMSOL Multiphysics 3.4.

See Also

posteval

PurposeEvaluate expressions in subdomains, boundaries, edges or vertices.Syntax[v1,v2,...,vn] = posteval(fem,e1,e2,...,en,...)Description[v1,v2,...,vn] = posteval(fem,e1,e2,...,en,...) returns values
v1,v2,...,vn of the expressions e1,e2,...,en. The expressions can be evaluated
on any domain type: subdomain, boundary, edge, and vertex, using one or several
solutions.

The values vi are *post data*, a structure with fields p, t, q, d, and elind. The field p contains node point coordinate information. The number of rows in p is the number of space dimensions. The field t contains the indices to columns in p of a simplex mesh, each column in t representing a simplex. The field q contains the indices to columns in p of a quadrilateral mesh, each column in q representing a quadrilateral. The field d contains data values. The columns in d correspond to node point coordinates in columns in p. There is one row in d for each solution (see the properties Solnum and T below). The data contains the real part of complex-valued expressions. The field elind contains indices to mesh elements for each point.

The string expressions can be any COMSOL Multiphysics expressions involving variables, in particular *application mode variables*.

The function posteval accepts the following property/values:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Bpoint	double matrix		Local coordinates for quadrilateral and block elements
Const	cell array		Definition of constants
Cont	off on internal	off	Smoothing
Dl	integer vector or cell array of integer vectors	all domains	Domain lists
Edim	integer	full	Element dimension
Frame	integer	spatial frame	Coordinate frame
Geomnum	positive integer	1	Geometry number
Phase	scalar	0	Phase angle
Prpoint	double matrix		Local coordinates for prism elements

TABLE I-105: VALID PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Refine	integer auto	3	Refinement of element for evaluation points
Solnum	integer vector all end	See below	Solution number
Spoint	double matrix		Local coordinates for simplex elements
т	double vector		Time for evaluation
Triangulate	off on	off	Divide quad elements into triangles
U	solution object or vector	fem.sol or O	Solution for evaluation

TABLE I-105: VALID PROPERTY/VALUE PAIRS

The property Refine constructs evaluation points by making a regular refinements of each element. Each mesh edge is divided into Refine equal parts. If auto is used, an automatic refinement value is computed internally and used, which depends on the maximum element order and the number of elements evaluated on. This value is most useful in postplot.

Use the properties Spoint, Bpoint, and Prpoint to specify arbitrary local element evaluation points for simplex elements (triangular, tetrahedral, and edge elements), quadrilateral/block elements, and prism elements, respectively. If you specify any of these properties, the fields t and q in the output postdata structure are empty, and the property Cont is neglected.

The property Edim decides which elements to evaluate on. Evaluation takes place only on elements with space dimension Edim. If not specified, Edim=sdim is used, where sdim is the space dimension of the geometry. For example, in a 3D model, if evaluation is done on edges (1D elements), Edim is 1. Similarly, for boundary evaluation (2D elements), Edim is 2, and for subdomain evaluation (3D elements), Edim is 3 (default in 3D).

The property D1 controls on which domains (subdomains, boundaries, etc.) evaluation should take place. If Geomnum is a vector, D1 must be a cell array of the same length as Geomnum containing domain lists for each geometry.

The property Cont controls if the post data is forced to be continuous on element edges. When Cont is set to internal, only elements not on interior boundaries are made continuous.

	The expressions ei are evaluated for one or several solutions. Each solution generates an additional row in the d field of the post data output structure. The properties Solnum and T control what solutions are used for the evaluations. If Solnum is provided, the solution indicated by the indices provided with the Solnum property are used. It T is provided, solutions are interpolated The property T can only be used for time dependent solutions. If nether Solnum nor T is provided, a single solution is evaluated. For parametric and time-dependent solutions, the final solution is used.					
	For time-dependent problems, the variable t can be used in the expressions ei . The value of t is the interpolation time when the property T is provided, and the time for the solution, when Solnum is used. Similarly, lambda and the parameter are available as eigenvalues for eigenvalue problems and as parameter value for parametric problems, respectively.					
	When the property Phase is used, the solution vector is multiplied with exp(i*phase) before evaluating the expression.					
Example	Solve Poisson's equation on two rectangles and evaluate the solution on one of them and the negative solution on the other.					
	<pre>clear fem fem.geom = square2(1,'pos',[0 -1])+square2; fem.mesh = meshinit(fem); fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = 1; fem.equ.expr = {'uu' {'u','-u'}}; fem.xmesh = meshextend(fem); fem.sol = femstatic(fem); pd = posteval(fem,'uu');</pre>					
Compatibility	The properties Spoint and Bpoint was re-introduced and Prpoint was introduced in COMSOL Multiphysics 3 .2a.					
	The FEMLAB 3.0 output type has been changed to a structure containing data suitable for further postprocessing. The new output format is incompatible with FEMLAB 2.3 and earlier versions.					
	The properties Context, Contorder, Posttype, and Spoint are obsolete from FEMLAB 3.0.					
	The property Variables has been renamed to Const in FEMLAB 2.3.					

The properties Bd1, Epoint, Sd1, and Tpoint, are obsolete from FEMLAB 2.2. Use the D1 property to specify domain lists. The post data format has changed in FEMLAB 2.2 and later versions.

The variable name lambda introduced in FEMLAB 1.2 can introduce a variable name conflict for old models.

See Also postdataplot, postglobaleval, postinterp, postint

Purpose	Shorthand command for streamline plot in 2D and 3D.				
Syntax	<pre>postflow(fem,expr,) h = postflow(fem,expr,)</pre>				
Description	postflow(fem, expr,) plots a streamline plot for the expressions in the cell array expr. In 2D, expr has length 2, and in 3D, it has length 3. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call				
	postplot(fem,'flowdata',expr, 'geom','on', 'axisequal','on',)				
	<pre>h = postflow(fem,expr,) additionally returns handles to the plotted handle graphics objects.</pre>				
	If you want to have more control over your streamline plot, use postplot instead of postflow.				
See Also	postplot, postanim, postsurf, postcont, postlin, postarrow, postarrowbnd, postprinc, postprincbnd, postslice, postiso, posttet				

Purpose	Evaluate globally defined expressions, such as solutions to ODEs.					
Syntax	<pre>data = postglobaleval(fem,) data = postglobaleval(fem,expr,)</pre>					
Description	postglobaleval(fem, expr,) is the evaluation function for globally defir expressions, such as solution variables for ODEs and other space-independent equations.					
		contains the expressions to ressions in fem.ode.dim		e a cell array of strings. If		
	<pre>data = postglobaleval(fem,) returns a structure with fields x, y, and legend. This structure is compatible with the import/export structure used by COMSOL Reaction Engineering Lab. The values can be plotted with plot(data.x, data.y) legend(data.legend) Valid property/value pairs for postglobaleval are given in the following table.</pre>					
	PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION		
	Const	cell array		Definition of constants		
	Phase	scalar	0	Phase angle		
	Solnum	integer vector all end	all	Solution number		
	Т	vector		Times for evaluation		
	U	Solution for evaluation				
	The property Phase is described in posteval.					
Examples	Example: Solve t	he Lotka-Volterra equati	ions for two pop	ulations r and f		

```
clear fem
fem.ode.dim={'r','f'};
fem.ode.f={'r*(1-2*f)-rt','-f*(3-r)-ft'};
fem.ode.init={'10','1'};
fem.ode.dinit={'0','0'};
fem.geom=solid1([0,1]);
fem.mesh = meshinit(fem);
fem.xmesh = meshextend(fem);
fem.sol = femtime(fem,'tlist',[0,1]);
% Evaluate 'r' and 'f' for all time steps
data = postglobaleval(fem);
```

	% Evaluate 'r+f' and 'r*f' and plot the result data = postglobaleval(fem,{'r+f','r*f'}) plot(data.x, data.y) legend(data.legend)
Compatibility	This function was introduced in COMSOL Multiphysics 3.2a.
See Also	postglobalplot, postinterp

Purpose	Plot globally def	ined expressions, such as	solutions to OD	Es.	
Syntax		t(fem,expr,) lplot(fem,expr,)			
Description		t (fem, expr,) is the as solution variables for	-	r globally defined e-independent equations.	
	The input expr of strings.	The input expr contains the expressions to plot. It must be a string or a cell array of strings.			
		h = postglobalplot(fem,expr,) additionally returns handles or a postdata structure (depending on the value of the property Outtype) to the plotted objects.			
	Valid property/v	alue pairs for postgloba	lplot are given	in the following table.	
	TABLE I-107: VALID P	ROPERTY/VALUE PAIRS			
	PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION	
	Const	cell array		Definition of constants	
	Lincolor	colorspec cell array of colorspec	cycle	Line color	
	Linewidth	numeric	0.5	Line width	
	Linlegend	off on	off	Color legend	
	Linmarker	marker specifier cell array of marker specifiers	none	Line marker	
	Linstyle	symbol cell array of symbols	1 _ 1	Line style	
	Linxdata	string		Line x-axis expression	
	Markersize	integer	6	Size of markers	
	Outtype	handle postdata	handle	Output type	
	Phase	scalar	0	Phase angle	
	Solnum	integer vector all end	all	Solution number	
	Т	vector		Times for evaluation	
	U	solution object or vector	fem.sol or O	Solution for evaluation	

In addition, the common plotting properties listed under femplot are available.

	If Outtype is 'handle', postglobalplot returns a vector of handles to the plots. If Outtype is 'postdata', the function returns a post data structure. The post data structure has the same format as the output from posteval.
	The notation colorspec in the value column denotes a <i>color specification</i> . See postplot for a description of this specification.
	The property Phase is described in posteval.
Examples	<pre>Example: Solve the Lotka-Volterra equations for two populations r and f clear fem fem.ode.dim={'r','f'}; fem.ode.dimt={'10','1'}; fem.ode.init={'10','1'}; fem.ode.dinit={'0','0'}; fem.geom=solid1([0,1]); fem.mesh = meshinit(fem); fem.xmesh = meshextend(fem); fem.sol = femtime(fem,'tlist',[0,1]); % Plot the solutions r and f with legend postglobalplot(fem,{'r','f'},'linlegend','on'); % Plot the r population versus the f population postglobalplot(fem,'r','linxdata','f')</pre>
Compatibility	This function was introduced in COMSOL Multiphysics 3.2a.
See Also	postcrossplot, postinterp

Purpose	Extract Gauss points and Gauss point weights.
Syntax	gp = postgp(type,order) [gp,gpw] = postgp(type,order)
Description	postgp(type,order) returns the gauss points of order order for an element of type type.
	[gp,gpw] = postgp(type,order) additionally returns the Gauss point weights.
	The Gauss points and their weights are the ones used in postint when computing integrals.
	The input type must be one of the following: vtx, edg, tri, quad, tet, prism, or hex corresponding to a vertex element, an edge element, a triangular element, a quadrilateral element, a tetrahedral element, a prism element, and a hexahedral element, respectively.
Examples	% The second order Gauss points for a triangular element gp = postgp('tri',2);
	% The third order Gauss points and their weights for a hexahedral % element [gp,gpw] = postgp('hex',3);
Compatibility	This function was introduced in COMSOL Multiphysics 3.2a.
See Also	posteval, postint

Purpose	Integrate expressions in domains with arbitrary space dimension.
Syntax	<pre>[v1,v2,,vn] = postint(fem,e1,e2,,en,)</pre>
Description	<pre>[v1,i2,,vn] = postint(fem,e1,e2,,en,) returns the integrals v1,v2,,vn of the expressions e1,,en. The integrals can be evaluated on any domain type: subdomain, boundary, edge, and vertex, using one or several solutions. When the several solutions are provided, each vi is a vector with values</pre>

corresponding to the solutions.

The expressions that are integrated can be expressions involving variables, in particular *application mode variables*.

postint accepts the following property/value pairs:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Const	cell array		List of assignments of constants
Dl	integer vector	all domains	Domain list
Edim	integer	full	Element dimension
Frame	string	spatial frame	Coordinate frame
Geomnum	positive integer	1	Geometry number
Intorder	positive integer	See below	Integration order
Phase	integer vector	0	Phase angle
Solnum	integer vector all end	See below	Solution numbers
Т	double vector	See below	Time for evaluation
U	solution object or vector	fem.sol or O	Solution(s) for evaluation

TABLE I-108: VALID PROPERTY/VALUE PAIRS

The expressions ei are integrated for one or several solutions. Each solution generates an element in the output vectors vi. The properties Solnum and T control what solutions are used for the evaluations. If Solnum is provided, the solution indicated by the indices provided with the Solnum property are used. It T is provided, solutions are interpolated The property T can only be used for time dependent solutions. If nether Solnum nor T is provided, a single solution is evaluated. For parametric and time-dependent solutions, the final solution is used. For eigenvalue solution the first solution is used.

	For time-dependent problems, the variable t can be used in the expressions ei. The value of t is the interpolation time when the property T is provided, and the time for the solution, when Solnum is used. Similarly, lambda and the parameter are available as eigenvalues for eigenvalue problems and as parameter value for parametric problems, respectively.
Examples	<pre>Compute the integral of the solution to Poisson's equation on the unit disk using weak constraints. Use weak constraint to obtain accurate flux. clear fem fem.dim = {'u' 'lm'}; fem.geom = circ2; fem.mesh = meshinit(fem); fem.shape={'shlag(2,''u'')' 'shlag(2,''lm'')'}; fem.equ.c = {{1 0}}; fem.equ.f = {{1 0}}; % make shape function for u active on subdomain fem.equ.shape={1}; fem.bnd.weak = {{'test(u)*lm' 'test(lm)*(-u)'}}; % make shape functions for u and lm active on boundary fem.bnd.shape={[1 2]}; fem.solform = 'general'; fem.sol = femstatic(fem); postint(fem,'u')</pre>
	Verify that the integral of the source term in Poisson's equation on the unit disk cancels the integral of the flux over the boundary. To have access to the variables f1 and ncu1, you must use the General solution form.
	postint(fem,'f1') postint(fem,'lm','edim',1)
	You can also use the variable neu to compute the flux, but it is much less accurate.
	postint(fem,'-ncu1','edim',1)
Compatibility	The properties Context, Cont, and Contorder are obsolete from FEMLAB 3.0.
	The property Variables has been renamed to Const in FEMLAB 2.3.
See Also	posteval

Purpose	Evaluate expressions in arbitrary points.
Syntax	<pre>[v1,v2,,vn,pe] = postinterp(fem,e1,e2,,en,xx,) [pio,pe] = postinterp(fem,xx,) [v1,v2,,vn] = postinterp(fem,e1,e2,,en,pio,)</pre>
Description	<pre>[v1,v2,,vn,pe] = postinterp(fem,e1,e2,,en,xx,) returns the values v1,v2,,vn of the expressions e1,e2,,en in the points xx.</pre>
	[pio,pe] = postinterp(fem,xx,) computes a <i>PostInterp object</i> pio, which contains information about where the points xx are located.
	<pre>[v1,v2,,vn] = postinterp(fem,e1,e2,,en,pio,) returns the values v1,v2,,vn of the expressions e1,e2,,en in the points given by the PostInterp object pio.</pre>
	The columns of the matrix xx are the coordinates for the evaluation points. If the number of rows in xx equals the space dimension, then xx are global coordinates, and the property Edim determines the dimension in which the expressions are evaluated. For instance, Edim=2 means that the expressions are evaluated on boundaries in a 3D model. If Edim is less than the space dimension, then the points in xx are projected onto the closest point on a domain of dimension Edim. If, in addition, the property Dom is given, then the closest point on domain number Dom in dimension Edim is used.
	If the number of rows in xx is less than the space dimension, then these coordinates are parameter values on a geometry face or edge. In that case, the domain number for that face or edge must be specified with the property Dom.
	The expressions that are evaluated can be expressions involving variables, in particular <i>application mode variables</i> .
	The matrices $v1, v2,, vn$ have size k-by-size(xx,2), where k is the number of solutions for which the evaluation is carried out, see below. The value of expression ei for solution number j in evaluation point xx(:,m) is vi(j,m).
	The vector pe contains the indices m for the evaluation points $xx(:,m)$ that are outside the mesh, or, if a domain is specified, are outside that domain.

postinterp accepts the following property/value pairs:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Blocksize	positive integer	5000	Block size
Const	cell array		List of assignments of constants
Dom	positive integer		Domain number
Edim	0 1 2 3	<pre>size(xx,1)</pre>	Element dimension for evaluation
Ext	number between 0 and 1	0.1	Extrapolation distance
Frame	string	spatial frame	Coordinate frame
Geomnum	positive integer	1	Geometry number
Matherr	off on	off	Error for undefined operations
Mcase	non-negative integer		Mesh case
Phase	scalar	0	Phase angle
Solnum	integer vector all end	See below	Solution numbers
т	double vector	See below	Time for evaluation
U	solution object solution vector scalar	fem.sol or O	Solution(s) for evaluation

The properties Blocksize and Const are described in assemble.

The property Ext determines how far the extrapolation reaches. A positive value Ext means that for points outside the mesh, the evaluation is carried out by extrapolation from the nearest mesh element, provided that the distance to the mesh element is at most ext times the element diameter, roughly. Other (more distant) points outside the mesh give the value NaN in the value matrices vi.

The property Matherr is described in femsolver.

If the property U does not specify the mesh case number, it is given by the property Mcase. The default is the mesh case that has the greatest number of degrees of freedom.

The property Phase is described in posteval.

The property U specifies the solution for which the evaluation is carried out. If U is not specified, then it is taken from fem.sol if it exists; otherwise it is the zero vector.

The expressions ei are interpolated for one or several solutions. The properties Solnum and T control what solutions are used for the evaluations. If Solnum is provided, the solution indicated by the indices provided with the Solnum property are used. It T is provided, solutions are interpolated at the given times. The property T can only be used for time dependent solutions. If neither Solnum nor T is provided, a single solution is evaluated. For parametric and time-dependent solutions, the final solution is used. For eigenvalue solution the first solution is used.

For time-dependent problems, the variable t can be used in the expressions ei. The value of t is the interpolation time when the property T is provided, and the time for the solution, when Solnum is used. Similarly, lambda and the parameter are available as eigenvalues for eigenvalue problems and parameter value for parametric problems, respectively.

A subsequent evaluation with [v1,v2,...,vn] =
postinterp(fem,e1,e2,...,en,pio,...) is faster than using xx instead of pio.
In this form of the call, only the properties Const, Phase, Solnum, T, and U are used.CompatibilityThe properties Context, Cont, and Contorder are obsolete from FEMLAB 3.0.
In FEMLAB 3.0, the interpolation structure is as a Java object.
The property Variables has been renamed to Const in FEMLAB 2.3.
The syntax and capabilities of this function has changed since FEMLAB 2.1.

See Also posteval

Purpose	Shorthand command for isosurface plot in 3D.
Syntax	postiso(fem,expr,) h = postiso(fem,expr,)
Description	postiso(fem, expr,) plots an isosurface plot for the expression expr. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call
	postplot(fem,'isodata',expr, 'isobar','on', 'geom','on', 'axisequal','on',)
	h = postiso(fem,expr,) additionally returns handles to the plotted handle graphics objects.
	If you want to have more control over your isosurface plot, use postplot instead of postiso.
See Also	postplot, postanim, postsurf, postcont, postlin, postarrow, postarrowbnd, postflow, postprinc, postprincbnd, postslice, posttet

Purpose	Shorthand command for line plot in 1D, 2D and 3D.
Syntax	<pre>postlin(fem,expr,) h = postlin(fem,expr,)</pre>
Description	postlin(fem, expr,) generates a line plot for the expression expr. The function accepts all property/value pairs that postplot does. In 1D, this command is just shorthand for the call
	postplot(fem,'liny',expr, 'linstyle','bginv',)
	and in 2D, it is shorthand for
	postplot(fem,'lindata',expr, 'linz',expr, 'linbar','on', 'axisequal','on',)
	and in 3D, it is shorthand for
	postplot(fem,'lindata',expr, 'linbar','on', 'axisequal','on',)
	h = postlin(fem,expr,) additionally returns handles to the plotted handle graphics objects.
	If you want to have more control over your line plot, use postplot instead of postlin.
See Also	postplot, postanim, postsurf, postcont, postarrow, postarrowbnd, postflow, postprinc, postprincbnd, postslice, postiso, posttet

Purpose	Compute ma	Compute maximum value of an expression.						
Syntax		m = postmax(fem,expr,) [m,p] = postmax(fem,expr,)						
Description	The function	<pre>m = postmax(fem,expr,) returns the maximum value of the expression expr. The function accepts all property/value pairs that posteval does, except cont. In addition, the following property/value pairs are accepted:</pre>						
	PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION				
	Out	all sollist	all	Return min over all solutions, or min per solution, specified with solnum or t				
	Useinf	on off	on	Allow infinity to be maximum value				
	containing t	<pre>[m,p] = postmax(fem,expr,) additionally returns the sdim-by-l matrix p containing the coordinate for which the maximum value occurs, where sdim is the space dimension of the geometry.</pre>						
	Note that the property Refine (see posteval) specifies the refinement used for finding the element in which the maximum value occurs. This element is then refined further to find the maximum value within the element. Therefore, the coordinate for which the maximum value of expr is attained, is not necessarily a node in the mesh.							
Cautionary	When expr value calcula	-	lex number	s, the real part is used in the maximum				
See Also	posteval,p	oostmin						

Compute minimum value of an expression.

Purpose Syntax

m = postmin(fem,expr,...)
[m,p] = postmin(fem,expr,...)

Description

m = postmin(fem,expr,...) returns the minimum value of the expression expr. The function accepts all property/value pairs that posteval does, except cont. In addition, the following property/value pairs are accepted:

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Out	all sollist	all	Return min over all solutions, or min per solution, specified with solnum or t
Useinf	on off	on	Allow -infinity to be minimum value

[m,p] = postmin(fem,expr,...) additionally returns the sdim-by-l matrix p containing the coordinate for which the minimum value occurs, where sdim is the space dimension of the geometry.

Note that the property Refine (see posteval) specifies the refinement used for finding the element in which the minimum value occurs. This element is then refined further to find the maximum value within the element. Therefore, the coordinate for which the minimum value of expr is attained, is not necessarily a node in the mesh.

CautionaryWhen expr is evaluated to complex numbers, the real part is used in the minimum
value calculation.

See Also posteval, postmax

Purpose	Postprocessing animation function.
Syntax	postmovie(fem,) postmovie({fem1,fem2,fem3,},) M = postmovie(fem,) % MATLAB only
Description	postmovie(fem,) is the general solution animation function. It supports all property/value pairs that postplot supports, and in addition to that, it supports a set of property/value pairs that is exclusive for animation.
	The input fem must be an FEM structure or a cell array of FEM structures. When it is a cell array, the properties solnum and t must, if specified, be cell arrays of the same size as fem. In this case, the FEM structures must have the same solution type, for example, all time-dependent solutions.
	M = postmovie(fem,) additionally returns a matrix in the MATLAB movie format. Alternatively, in COMSOL Script, you can store the animation in a file using the property Filename.
	The command can generate a sequence of image files containing all images in the movie. In addition, the command can generate an AVI movie file.
	Valid and a start for the sector of a start of the sector

Valid property/value pairs for the postmovie function are given in the following table. In addition, all postplot parameters are supported and are passed to postplot. See the entry on postplot for a description of the post data formats.

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Aviautoplay	on off	on	In Matlab on Windows, try to launch program associated with AVI-extension and play generated AVI-movie
Avicompression	string		Compression used for AVI-movie
Aviquality	integer between 0 and 100	75	Quality of AVI-movie
Filename	string		Output file name
Filetype	avi jpg tiff png	avi	Output file type
Fps	integer	12	Frames per second
Height	integer	480	Height of image/movie files
Repeat	integer	5	Number of repeats

TABLE I-II0: VALID PROPERTY/VALUE PAIRS

	PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION	_
	Reverse	on off	off	Make movie backwards	
	Resol	integer	150	Resolution	
	Solnum	integer vector all end	all	Solution numbers	
	Statfunctype	string	full half linear	Static plot function	
	Statnframes	integer	11	Number of frames in animation of static solution	
	Width	integer	640	Width of image/movie files	-
Example Cautionar	y Who expl	en you are replayin icitly provide a fig	· · · •		
	Oth	erwise the animati	on does not look g	good.	
Compatib	and	Qtcomp on Mac ar	1 1	as well as the properties Qtrate LAB 3.0a. The option avi wor y to avi.	
See Also	pos	teval, postplot			

Purpose	Postprocessing plot function.
Syntax	<pre>postplot(fem,) h = postplot(fem,)</pre>
Description	<pre>postplot(fem,) is the general solution plot function. It can display an FEM solution in several different ways. The command works for both 1D, 2D, and 3D geometries. h = postplot(fem,) additionally returns handles or postdata corresponding</pre>

to the drawn axes objects. See properties Out and Outtype.

The function postplot accepts the following property/value pairs:

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Arrowbnd			\checkmark	vector post spec		Boundary arrow data
Arrowbndz		\checkmark		post spec		Boundary arrow height data
Arrowcolor		\checkmark	\checkmark	colorspec	red	Subdomain arrow color
Arrowcolorbnd		\checkmark	\checkmark	colorspec	blue	Boundary arrow color
Arrowcoloredg			\checkmark	colorspec	black	Edge arrow color
Arrowdata			\checkmark	vector post spec		Arrow data
Arrowedg			\checkmark	vector post spec		Edge arrow data
Arrowscale		\checkmark	\checkmark	numeric	auto	Subdomain arrow scale
Arrowscalebnd		\checkmark	\checkmark	scalar	auto	Boundary arrow scale
Arrowscaleedg			\checkmark	scalar	auto	Edge arrow scale
Arrowstyle		\checkmark	\checkmark	proportional normalized	proportional	Subdomain arrow style
Arrowstylebnd		\checkmark	\checkmark	proportional normalized	proportional	Boundary arrow style
Arrowstyleedg			\checkmark	proportional normalized	proportional	Edge arrow style
Arrowtype			\checkmark	arrow cone arrow3d	cone	Subdomain arrow type
Arrowtypebnd		\checkmark	\checkmark	arrow cone arrow3d	cone	Boundary arrow type

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Arrowtypeedg			\checkmark	arrow cone arrow3d	cone	Edge arrow type
Arrowxspacing		\checkmark	\checkmark	number of arrows or vector specifying x-coordinates	15 (in 2D) 7 (in 3D)	Arrow x-spacing
Arrowyspacing		\checkmark	\checkmark	number of arrows or vector specifying y-coordinates	15 (in 2D) 7 (in 3D)	Arrow y-spacing
Arrowz		\checkmark		post spec		Arrow height data
Arrowzspacing			\checkmark	number of arrows or vector specifying z-coordinates	7 (in 3D)	Arrow z-spacing
Bdl		\checkmark	\checkmark	list of boundary numbers	all	Boundary list
Bndmarker		\checkmark	\checkmark	marker specifier	square	Boundary max/min marker type
Bndmarkersize		\checkmark	\checkmark	integer	6	Size of boundary max min markers
Const	\checkmark	\checkmark	\checkmark	cell array		Definition of constants
Cont	V	\checkmark	\checkmark	off on internal	off	Make output continuous
Contbar		\checkmark		off on	on	Show color bar for contours
Contcolorbar		\checkmark		off on	on	Show color bar for contour colors
Contcolordata		\checkmark		post spec		Contour color data
Contcolordlim		\checkmark		[min max]	full range	Contour color limits
Contdata		\checkmark		post spec		Contour data
Contdlim		\checkmark		[min max]	full range	Contour limits
Contlabel		\checkmark		off on	off	Show contour labels
Contlevels		\checkmark		number of levels or a vector specifying levels	20	Contour levels
Contmap		\checkmark		colormap		Colormap for contour plot

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Arrowtypeedg			\checkmark	arrow cone arrow3d	cone	Edge arrow type
Arrowxspacing		\checkmark	\checkmark	number of arrows or vector specifying x-coordinates	15 (in 2D) 7 (in 3D)	Arrow <i>x</i> -spacing
Arrowyspacing		\checkmark	\checkmark	number of arrows or vector specifying y-coordinates	15 (in 2D) 7 (in 3D)	Arrow y-spacing
Arrowz		\checkmark		post spec		Arrow height data
Arrowzspacing			\checkmark	number of arrows or vector specifying z-coordinates	7 (in 3D)	Arrow z-spacing
Bdl		\checkmark	\checkmark	list of boundary numbers	all	Boundary list
Bndmarker		\checkmark	\checkmark	marker specifier	square	Boundary max/min marker type
Bndmarkersize		\checkmark	\checkmark	integer	6	Size of boundary max/ min markers
Const	\checkmark	\checkmark	\checkmark	cell array		Definition of constants
Cont	\checkmark	\checkmark	\checkmark	off on internal	off	Make output continuous
Contbar		\checkmark		off on	on	Show color bar for contours
Contcolorbar		\checkmark		off on	on	Show color bar for contour colors
Contcolordata		\checkmark		post spec		Contour color data
Contcolordlim		\checkmark		[min max]	full range	Contour color limits
Contdata		\checkmark		post spec		Contour data
Contdlim		\checkmark		[min max]	full range	Contour limits
Contlabel		\checkmark		off on	off	Show contour labels
Contlevels		\checkmark		number of levels or a vector specifying levels	20	Contour levels
Contmap		\checkmark		colormap		Colormap for contour plot

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Contrefine		\checkmark		integer auto	auto	Refinement of elements for contour plots
Contstyle		\checkmark		bg bginv interp cycle	interp	Contour style
Contz		\checkmark		post spec		Contour height data
Deformauto		\checkmark	\checkmark	on off	on	Auto scaling
Deformbnd		\checkmark	\checkmark	vector post spec		Deform data for boundaries
Deformdata		\checkmark	\checkmark	vector post spec		Deformation data
Deformedg			\checkmark	vector post spec		Deform data for edges
Deformscale	\checkmark	\checkmark	\checkmark	numeric		Deformation scale factor for subdomains
Deformscalebnd		\checkmark	\checkmark	numeric		Deformation scale factor for boundaries
Deformscaleedg			\checkmark	numeric		Deformation scale factor for edges
Deformscalesub			\checkmark	numeric		Deformation scale factor for subdomains
Edgmarker			\checkmark	marker specifier	square	Edge max/min marker type
Edgmarkersize			\checkmark	integer	6	Size of edge max/min markers
Edl			\checkmark	integer vector	all	Edge list
Ellogic	\checkmark	\checkmark	\checkmark	logical expression	1	Logical expression for elements to include
Ellogictype	\checkmark	\checkmark	\checkmark	all any xor	all	Interpretation of logical expression
Flowback		\checkmark	\checkmark	off on	on	Integrate streamlines backwards
Flowbar		\checkmark	\checkmark	off on	on	Color bar for streamline color data
Flowcolor		\checkmark	\checkmark	interp bg bginv colorspec	interp	Streamline color
Flowcolordata				post spec		Streamline color data

postplot

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Flowcolordlim		\checkmark	\checkmark	[min max]	full range	Streamline color data limits, works only when flowtype is line
Flowdata		\checkmark	\checkmark	vector post spec		Streamline velocity field
Flowdens		\checkmark	\checkmark	none uniform velocity	none	Type of streamline density
Flowdist		\checkmark		numeric	0.05	Separating distance factor
Flowdist			\checkmark	numeric	0.15 or [0.05,0.15]	Separating distance factor
Flowdistdel		\checkmark	\checkmark	numeric	0.2	Minimum Delaunay distance
Flowdistend		\checkmark	\checkmark	numeric	0.5	Terminating distance factor
Flowinitref		\checkmark	\checkmark	integer	1	Boundary element refinement
Flowdignoredist		\checkmark	\checkmark	numeric	0.5	Fraction of streamline length to ignore
Flowlines		\checkmark	\checkmark	integer	20	Number of streamlines
Flowlooptol		\checkmark	\checkmark	numeric	0.01	Streamline loop tolerance
Flowmap		\checkmark	\checkmark	colormap	jet	Colormap for streamline color data
Flowmaxsteps		\checkmark	\checkmark	integer	400	Maximum number of integration steps
Flowmaxtime		\checkmark	\checkmark	integer	100	Maximum integration time
Flownormal		\checkmark	\checkmark	off on	off	Normalize velocity field
Flowradiusdata		\checkmark	\checkmark	post spec		Streamline radius data
Flowrefine		\checkmark	\checkmark	integer auto	auto	Refinement of elements for streamline plots

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Flowsat		\checkmark	\checkmark	numeric	1.3	Streamline saturation factor
Flowseed		\checkmark	\checkmark	I-by-sdim numeric		First start point for streamlines when flowdens is not none
Flowstart		\checkmark	\checkmark	centers ginput edges cellarray{x y z}	centers	Starting points for streamlines
Flowstattol		\checkmark	\checkmark	positive scalar	1e-2	Streamline stationary point stop tolerance
Flowtol		\checkmark		positive scalar	1e-3	Streamline integration tolerance
Flowtol			\checkmark	positive scalar	1e-2	Streamline integration tolerance
Flowtuberes		\checkmark	\checkmark	numeric	8	Tube resolution for streamlines
Flowtubescale		\checkmark	\checkmark	numeric		Tube radius scale for streamlines
Flowtype		\checkmark	\checkmark	line tube	tube (line in 2D)	Type of streamline
Flowz				post spec		Streamline height data
Frame	\checkmark		\checkmark	string	spatial frame	Coordinate frame
Geom	V	\checkmark	\checkmark	off on	on	Show geometry contours
Geomcolor	V	\checkmark	\checkmark	bg bginv colorspec	bginv	Geometry contours color
Geomnum	\checkmark	\checkmark	\checkmark	integer	1	Geometry number
Isobar			\checkmark	on off	on	Isosurface color bar
Isocolorbar			\checkmark	on off	on	Color bar for isosurfaces color data
Isocolordata			\checkmark	post spec		Isosurface color data
Isocolordlim			\checkmark	[min max]	full range	Isosurface color limits
Isocolormap			\checkmark	colormap	jet	Colormap for isosurface color data
Isodata			\checkmark	Post spec		Isosurface data
Isodlim			\checkmark	[min max]	full range	Isosurface limits

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Isoedgestyle			\checkmark	flat interp none bg bginv colorspec	none	Isosurface edge style
Isofacestyle			\checkmark	flat interp none bg bginv colorspec	interp	Isosurface face style
Isolevels			\checkmark	number of levels or a vector specifying levels	5	Isosurface levels
Isomap			\checkmark	colormap		Colormap for isosurface plot
Isostyle			\checkmark	bg bginv color	color	Isosurface style
Linbar	\checkmark	\checkmark	\checkmark	off on	on	Line color bar
Lindata	\checkmark	\checkmark	\checkmark	post spec		Line data
Lindlim	\checkmark	\checkmark	\checkmark	[min max]	full range	Line limits
Linmap	\checkmark	\checkmark	\checkmark	colormap		Line colormap
Linrefine	V	\checkmark	V	integer auto	auto	Refinement of elements for line plots
Linstyle	\checkmark	\checkmark	\checkmark	flat interp none bg bginv colorspec	interp	Line style
Liny	\checkmark			post spec		Line y data
Linz	\checkmark	\checkmark		post spec		Line <i>z</i> data
Maxminbnd		\checkmark	\checkmark	post spec		Max/min marker on boundaries
Maxminedg			\checkmark	post spec		Max/min marker on edges
Maxminsub	\checkmark	\checkmark	\checkmark	post spec		Max/min marker on subdomains
Out	\checkmark		\checkmark	cell array of strings all	all	Output
Outtype	\checkmark	\checkmark	\checkmark	handle postdata	handle	Output type
Partatol		\checkmark	\checkmark	numeric vector		Absolute tolerances for particle tracing
Partbar			\checkmark	off on	on	Show color bar for particle tracing

TABLE I-III: VALID PROPERTY/VALUE PA	RS
--------------------------------------	----

TABLE I-III: VALID PROPERTY/VALUE PAIRS

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Partbnd		\checkmark	\checkmark	stick disappear	stick	Particle point boundary behavior
Partcolordata		\checkmark	\checkmark	post spec		Particle tracing line color
Partdropfreq		\checkmark	\checkmark	numeric		Time between each particle release
Partdroptimes		\checkmark	\checkmark	numeric vector		Time values to release particles
Partedgetol		\checkmark	\checkmark	numeric auto	0.001	Edge tolerance for particle tracing
Partdata		\checkmark	\checkmark	cell-array of strings		Particle force
Parthmax		\checkmark	\checkmark	numeric auto		Maximum time step for particle tracing
Parthstart		\checkmark	\checkmark	numeric auto		Initial time step for particle tracing
Partlinecolor		\checkmark	\checkmark	colorspec	blue	Particle tracing line color
Partmap		\checkmark	\checkmark	colormap		Colormap for particle tracing
Partmass		\checkmark	\checkmark	string	1	Particle mass
Partmaxsteps		\checkmark	\checkmark	integer auto	1000	Maximum number of steps for particle tracing
Partplotas		\checkmark	\checkmark	lines points both along	lines	Particle tracing plot type
Partpointcolor		\checkmark	\checkmark	colorspec	red	Particle tracing point color
Partpointscale		\checkmark	\checkmark	numeric		Point radius scale
Partradiusdata			\checkmark	post spec		Particle tracing tube radius
Partres		\checkmark	\checkmark	integer	5	Resolution of pathline for particle tracing
Partrtol		\checkmark	\checkmark	numeric		Relative tolerance for particle tracing

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Partstart		\checkmark	V	numeric matrix, cell array of double vectors, or cell array of property/ values to postcoord		Start points for particle tracing
Partstatic		\checkmark	\checkmark	off on	off	Use instantaneous flow field
Partstatictend			\checkmark	numeric auto	auto	End time for stationary flows
Parttstart			\checkmark	numeric auto		Initial time for particle tracing
Parttuberes		\checkmark	\checkmark	numeric	8	Tube resolution
Parttubescale		\checkmark	\checkmark	numeric		Tube radius scale
Parttvar		\checkmark	\checkmark	string		Particle integration time variable name
Partvelstart		\checkmark	\checkmark	cell-array of strings	Zero velocity	Initial velocity for particle tracing
Partvelvar		\checkmark	\checkmark	cell-array of strings		Particle velocity variable names
Phase	\checkmark	\checkmark	\checkmark	scalar	0	Phase angle
Princbnd			\checkmark	vector post spec		Boundary principal plot data
Princcolor		\checkmark	\checkmark	colorspec	blue	Subdomain principal plot color
Princcolorbnd			\checkmark	colorspec	blue	Boundary principal plot color
Princdata			\checkmark	vector post spec		Subdomain principal plot data
Princscale		\checkmark	\checkmark	numeric	auto	Subdomain principal plot scale
Princscalebnd			\checkmark	scalar	auto	Boundary principal plot scale
Princstyle		\checkmark	\checkmark	proportional normalized	proportional	Subdomain principal plot style
Princstylebnd			\checkmark	proportional normalized	proportional	Boundary principal plot style

TABLE I-III: VALID PROPERTY/VALUE PAIRS

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Princtype				arrow cone arrow3d	cone	Subdomain principal plot type
Princtypebnd			\checkmark	arrow cone arrow3d	cone	Boundary principal plot type
Princxspacing		\checkmark	\checkmark	number of arrows or vector specifying x-coordinates	8 (in 2D) 5 (in 3D)	Arrow <i>x</i> -spacing for subdomain principal plot
Princyspacing		\checkmark	\checkmark	number of arrows or vector specifying y-coordinates	8 (in 2D) 5 (in 3D)	Arrow <i>y</i> -spacing for subdomain principal plot
Princz		\checkmark		post spec		Subdomain principal plot height data
Princzspacing			\checkmark	number of arrows or vector specifying z-coordinates	8 (in 3D)	Arrow <i>z</i> -spacing for subdomain principal plot
Refine	\checkmark	\checkmark	\checkmark	integer auto	auto	Refinement of elements for all plot types
Sdl	\checkmark	\checkmark	\checkmark	list of subdomain numbers	all	Subdomain list
Slicebar		\checkmark	V	off on	on	Show color bar for slice plot
Slicedata		\checkmark		post spec		Slice plot data
Slicedlim		\checkmark	\checkmark	[min max]	full range	Slice plot limits
Sliceedgestyle		\checkmark	\checkmark	flat interp none bg bginv colorspec	none	Slice plot edge style
Slicefacestyle		\checkmark	\checkmark	flat interp none bg bginv colorspec	interp	Slice plot face style
Slicemap		\checkmark	\checkmark	colormap		Colormap for slice plot
Slicerefine		\checkmark	\checkmark	integer auto	auto	Refinement of elements for slice plots

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Slicexspacing		\checkmark	\checkmark	number of slices or vector specifying x-coordinates	5	Slice plot <i>x</i> -positions
Sliceyspacing		\checkmark	\checkmark	number of slices or vector specifying y-coordinates	[]	Slice plot <i>y</i> -positions
Slicezspacing			\checkmark	number of slices or vector specifying z-coordinates	[]	Slice plot <i>z</i> -positions
Solnum	\checkmark	\checkmark		integer end	1	Solution number
Submarker	\checkmark	\checkmark		marker specifier	square	Subdomain max/min marker type
Submarkersize	\checkmark	\checkmark	\checkmark	integer	6	Size of subdomain max/min markers
т	\checkmark	\checkmark	\checkmark	scalar		Time for evaluation
Tetbar			\checkmark	off on	on	Show color bar for subdomain plot
Tetdata			\checkmark	Post spec		Subdomain plot data
Tetdlim			\checkmark	[min max]	full range	Subdomain plot limits
Tetedgestyle			\checkmark	flat interp none bg bginv colorspec	none	Subdomain plot edge style
Tetfacestyle			\checkmark	flat interp none bg bginv colorspec	interp	Subdomain plot face style
Tetkeep			\checkmark	number between 0 and I	1	Fraction of elements to keep
Tetkeeptype			\checkmark	min max random	random	Which elements to keep
Tetmap			\checkmark	colormap		Subdomain plot colormap
Tetmaxmin			\checkmark	on off	off	Show subdomain plot max/min markers
Tetrefine			\checkmark	integer auto	auto	Refinement of elements for subdomain plots

TABLE I-III: VALID PROPERTY/VALUE PAIRS

PROPERTY	ID	2D	3D	VALUE	DEFAULT	DESCRIPTION
Tribar		\checkmark	\checkmark	off on	off	Surface color bar
Tridata		\checkmark		post spec		Surface data
Tridlim		\checkmark		[min max]	full range	Surface limits
Triedgestyle		\checkmark	\checkmark	flat interp none bg bginv colorspec	none	Surface edge style
Trifacestyle		\checkmark	\checkmark	flat interp none bg bginv colorspec	interp	Surface face style
Trimap		\checkmark	\checkmark	colormap		Surface colormap
Trimaxmin		\checkmark		on off	off	Show surface max/min markers
Trirefine		\checkmark	\checkmark	integer auto	auto	Refinement of elements for surface plots
Triz		\checkmark		post spec		Triangle height
U	\checkmark	\checkmark		solution vector	fem.sol.u	Solution for evaluation

TABLE I-III: VALID PROPERTY/VALUE PAIRS

The properties Out and Outtype control the format of the output h when the syntax h = postplot(fem,...) is used:

If Out is 'all' (default), output corresponding to all plotted objects are returned. The property Out can also be a cell array containing any of the strings 'geom', 'slice', 'iso', 'tet', 'tri', 'cont', 'lin', 'flow', 'partline', 'partpoint', 'arrow', 'arrowbnd', 'arrowedg', 'maxminsub', 'maxminbnd', 'maxminedg', and 'light'. These correspond to the plots made using the properties 'geom', 'slicedata', 'isodata', 'tetdata', and so on. When Out is a 1-by-*n* cell-array, the output h is a cell array of the same size, matching the strings in Out.

If Outtype is 'handle', handle-graphics handles to the plots (as specified with Out) are returned in a vector of handles if Out is 'all', otherwise in a cell array.

If Outtype is 'postdata', post data structures are returned in a cell array. The post data structures have the same format as the output from posteval. In addition, for particle tracing plots (using 'partdata'), the postdata structure contains the fields parttime and partvel, containing the time and velocity vector, respectively,

associated to each point on the path. Also, for both particle tracing line plots and streamline plots, the fields startpts and endpts are included, containing the coordinates of each plotted line's start and end point, respectively.

If the property Refine is specified, its value is used for all specified plot types; that is, it overrides all other properties ending with refine. See posteval.

The properties Princdata and Princbnd can either be specified as the names of the three principal stress or strain values in a 1-by-3 cell array of strings, for example, { 's1', 's2', 's3'}, or as the expressions for the value and then the vector components for each principal direction, for example,

{'e1','e1x','e1y','e1z','e2','e2x','e2y','e2z','e3','e3x','e3y', 'e3z'}.

The camera properties (Campos, Camtarget, etc.) override the setting of view if both are used.

The notation colorspec in the value column denotes a *color specification*, that is a single letter string: y, m, c, r, g, b, w, and k, meaning yellow, magenta, cyan, red, green, blue, white, and black, respectively (also 'yellow', 'magenta', etc. are acceptable as color specification), or a 1-by-3 numeric array with RGB values.

Post spec is one of the following.

- A string with an expression to be evaluated. It can be a COMSOL Multiphysics expression involving variables, in particular *application mode variables*.
- A cell array, where the first entry is a string with an expression to be evaluated or a cell array of such strings, and the other entries are parameters passed to posteval.

Vector post spec is a cell array of Post specs.

The properties can be grouped in terms of what plot entity it refers to. The table below shows this grouping.

PLOT ENTITY	ID	2D	3D	PROPERTY NAMES STARTING WITH
Arrows		Ω	Ω	arrow
Arrows		$\Omega 6$	$\Omega 6$	arrowbnd
Arrows			$\partial^2 \Omega$	arrowedg
Contours		Ω		cont
Isosurfaces			Ω	iso

TABLE I-112: PROPERTY GROUPING

PLOT ENTITY	ID	2D	3D	PROPERTY NAMES STARTING WITH
Lines	Ω	$\Omega 6$	$\partial^2 \Omega$	lin
Principal stress/strain plots		Ω	Ω	princ
Principal stress/strain plots			$\Omega 6$	princbnd
Slices			Ω	slice
Particle tracing		Ω	Ω	part
Streamlines		Ω	Ω	flow
3D subdomains			Ω	tet
Surfaces		Ω	$\Omega 6$	tri

TABLE I-112: PROPERTY GROUPING

The symbol $\partial \Omega$ indicates the boundary of the domain, and the symbol Ω indicates the domain itself. For the boundary of the domain, post data evaluated on the boundary is plotted. For the domain itself, post data evaluated on the domain is plotted.

Examples

3D Example

Solve the Poisson equation on a unit square:

```
clear fem
fem.geom = block3;
fem.mesh = meshinit(fem, 'hmax',0.15);
fem.equ.c = 1; fem.equ.f = 1;
fem.bnd.h = {1 1 0 0 1 1};
fem.xmesh = meshextend(fem); fem.sol = femstatic(fem);
```

Plot the solution as a slice plot

postplot(fem, 'slicedata', 'u')

Plot the solution using isosurfaces

postplot(fem,'isodata','u','scenelight','on')

Plot lighted cones showing the gradient together with geometry edges

2D Example

Solve the Poisson equation on the unit circle

```
clear fem
fem.geom = circ2; fem.mesh = meshinit(fem);
fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = 1;
fem.xmesh = meshextend(fem); fem.sol = femstatic(fem);
```

	Plot the solution as triangle color and z -height, and $u.*x$ as contour lines
	postplot(fem,'tridata','u','contdata','u*x', 'triz','u','contz','u');
	Plot 30 streamlines for the field (-uy, $x*ux$) with color data u.
	postplot(fem,'flowdata',{'-uy','x*ux'}, 'flowlines',30,'flowcolordata','u')
Cautionary	Some default values have changed from FEMLAB 2.3 resulting in slightly different plots.
Compatibility	The properties contlabel, context, contorder, and tetmarker are no longer supported in FEMLAB 3.0.
	Properties ending with maxmin are no longer supported. To plot max/min markers, use the properties maxminsub, maxminbnd, and maxminedg to plot markers on subdomains, boundaries, and edges, respectively.
	The support for outputs from posteval as Post spec, has only a limited support and is not recommended.
	The properties starting with Princ are added in FEMLAB 3.1.
	The property contlabel is added in COMSOL Multiphysics 3.2a.
See Also	geomplot, meshplot, postanim, postarrow, postarrowbnd, postcont, postcrossplot, posteval, postflow, postiso, postlin, postmovie, postprinc, postprincbnd, postslice, postsurf, posttet

Purpose	Shorthand command for subdomain principal stress/strain plot in 2D and 3D.
Syntax	<pre>postprinc(fem,expr,) h = postprinc(fem,expr,)</pre>
Description	postprinc(fem,expr,) plots a subdomain principal stress/strain plot for the expressions in the cell array expr, which can have length 3 or 12. See property Princdata in postplot. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call
	postplot(fem,'princdata',expr, 'geom','on', 'axisequal','on',)
	h = postprinc(fem,expr,) additionally returns handles to the plotted handle graphics objects.
	If you want to have more control over your principal stress/strain plot, use postplot instead of postprinc.
See Also	postplot, postanim, postsurf, postcont, postlin, postarrowbnd, postflow, postprincbnd, postslice, postiso, posttet
Compatibility	This function was added in FEMLAB 3.1.

Purpose	Shorthand command for boundary principal stress/strain plot in 2D and 3D.
Syntax	<pre>postprincbnd(fem,expr,) h = postprincbnd(fem,expr,)</pre>
Description	postprincbnd(fem,expr,) plots a boundary principal stress/strain plot for the expressions in the cell array expr, which can have length 3 or 12. See property Princdata in postplot. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call
	postplot(fem,'princbnd',expr, 'geom','on', 'axisequal','on',)
	h = postprincbnd(fem,expr,) additionally returns handles to the plotted handle graphics objects.
	If you want to have more control over your principal stress/strain plot, use postplot instead of postprincbnd.
See Also	postplot, postanim, postsurf, postcont, postlin, postarrowbnd, postflow, postprinc, postslice, postiso, posttet
Compatibility	This function was added in FEMLAB 3.1.

Purpose	Shorthand command for slice plot in 3D.
Syntax	<pre>postslice(fem,expr,) h = postslice(fem,expr,)</pre>
Description	postslice(fem, expr,) plots a slice plot for the expression expr. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call
	postplot(fem,'slicedata',expr, 'slicebar','on', 'geom','on', 'axisequal','on',)
	<pre>h = postslice(fem,expr,) additionally returns handles to the plotted handle graphics objects.</pre>
	If you want to have more control over your slice plot, use postplot instead of postslice.
See Also	postplot, postanim, postsurf, postcont, postlin, postarrow, postarrowbnd, postflow, postiso, postprinc, postprincbnd, posttet

Purpose	Shorthand command for surface plot in 2D and 3D.			
Syntax	<pre>postsurf(fem,expr1,) postsurf(fem,expr1,expr2) h = postsurf(fem,)</pre>			
Description	postsurf(fem,expr1,expr2) plots a surface plot on subdomains in 2D colored according to the expression expr1 and with height according to expr2. For a 3D model, it plots a colored surface plot on the boundaries, colored according to expr1. The function accepts all property/value pairs that postplot does. In 2D, this command is just shorthand for the call			
	postplot(fem,'tridata',expr1, 'triz',expr2, 'tribar','on', 'axisequal','on',)			
	and in 3D, it is shorthand for			
	postplot(fem,'tridata',expr1, 'tribar','on', 'geom','on', 'axisequal','on',)			
	h = postsurf(fem,) additionally returns handles to the plotted handle graphics objects.			
	If you want to have more control over your surface plot, use postplot instead of postsurf.			
Example	Surface plot of the solution to the equation $-\Delta u = 1$ over the geometry defined by the L-shaped membrane. Use Dirichlet boundary conditions $u = 0$ on $\partial \Omega$. sq1 = square2(0,0,1); sq2 = move(sq1,0,-1); sq3 = move(sq1,-1,-1); clear fem fem.geom = sq1+sq2+sq3; fem.mesh = meshinit(fem); fem.equ.c = 1; fem.equ.f = 1; fem.bnd.h = 1; fem.xmesh = meshextend(fem); fem.sol = femstatic(fem); postsurf(fem,'u')			
See Also	postplot, postanim, postcont, postlin, postarrow, postarrowbnd, postflow, postprinc, postprincbnd, postslice, postiso, posttet			

Purpose	Shorthand command for subdomain plot in 3D.				
Syntax	<pre>posttet(fem,expr,) h = posttet(fem,expr,)</pre>				
Description	posttet (fem, expr,) plots a subdomain plot for the expression expr. The function accepts all property/value pairs that postplot does. This command is just shorthand for the call				
	postplot(fem,'tetdata',expr, 'tetbar','on', 'geom','on', 'axisequal','on',)				
	<pre>h = postcont(fem,expr,) additionally returns handles to the plotted handle graphics objects.</pre>				
	If you want to have more control over your subdomain plot, use postplot instead of posttet.				
See Also	postplot, postanim, postsurf, postcont, postlin, postarrow, postarrowbnd, postflow, postprinc, postprincbnd, postslice, postiso				

Purpose	Create rectangular pyramid geometry object.					
Syntax	<pre>rp3 = pyramid3 rp2 = pyramid2 rp3 = pyramid3(a,b,h) rp2 = pyramid2(a,b,h) rp3 = pyramid3(a,b,h,rat) rp2 = pyramid3(a,b,h,rat) rp3 = pyramid3(a,b,h,rat,) rp2 = pyramid3(a,b,h,rat,)</pre>					
Description	and side lengths	s of bottom sur	face equal to	r pyramid geometry object with height o one, axis along the coordinate <i>z</i> -axis, origin. pyramid3 is a subclass of		
	ec3 = pyramic side lengths a a			ctangular pyramid geometry object with		
	ec3 = pyramic between the top			yramid with the non-negative ratio rat		
	The functions p			cept the following property/values:		
	PROPERTY	VALUE	DEFAULT	DESCRIPTION		
	Axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object		
	Const	Cell array of strings	{}	Evaluation context for string inputs		
	Displ	2-by- <i>nd</i> matrix	[0;0]	Displacement of extrusion top		
	Pos	Vector of reals or cell array of	[0 0]	Position of the bottom surface		

For more information on input arguments and properties see gency13.

Rotational angle about Axis (radians)

strings

real or string 0

Rot

ec2 = pyramid2(...) creates a surface rectangular pyramid geometry object, without bottom and top faces, according to the arguments described for pyramid3. pyramid2 is a subclass of gency12.

Pyramid objects have the following properties:

TABLE I-II4:	PYRAMID	OBIECT	PROPERTIES

PROPERTY	DESCRIPTION
a, b	Side lengths
n	Height
rat	Ratio
lx, dy	Semi-axes
x, y, z, xyz	Position of the object. Components and vector forms
ix2	Rotational angle of symmetry axis
ax3	Axis of symmetry
ot	Rotational angle

In addition, all 3D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom3 for details.

Compatibility The FEMLAB 2.3 syntax is obsolete but still supported. The numbering of faces, edges and vertices is different from the numbering in objects created in 2.3.

econe2, econe3, gency12, gency13

See Also

Purpose	Create rectangle geometry object.
Syntax	<pre>obj = rect2 obj = rect1 obj = rect2(lx,ly,) obj = rect1(lx,ly,)</pre>
Description	<pre>obj = rect2 creates a solid rectangle geometry object with all side lengths equal to 1, and the lower left corner at the origin. rect2 is a subclass of solid2. obj = rect2(lx,ly,) creates a solid rectangle object with side lengths equal</pre>
	to 1x and 1y, respectively, and the lower left corner at the origin. 1x and 1y are positive real scalars, or strings that evaluate to positive real scalars, given the evaluation context provided by the property Const.
	The function rect1 similarly creates curve rectangle objects.

The functions rect2 and rect1 accept the following property/values:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
base	corner center	corner	Positions the object either centered about pos or with the lower left corner at pos
const	Cell array of strings	{}	Evaluation context for string inputs
pos	Vector of reals or cell array of strings	[0 0]	Position of the object
rot	Real or string	0	Rotational angle about pos (radians)

TABLE I-115: VALID PROPERTY/VALUE PAIRS

obj = rect1(...) creates a curve circle geometry object with properties as given for the rect2 function. rect1 is a subclass of curve2.

Rectangle objects have the following properties:

TABLE I-II6: RECTANGLE OBJECT PROPERTIES

PROPERTY	DESCRIPTION
lx,ly	Side lengths
base	Base point
х, у	Position of the object
rot	Rotational angle

Example	In addition, all 2D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom2 for details. The commands below create a geometry object corresponding to the L-shaped membrane using the union of three rectangles and plot the result.
	<pre>sq1 = rect2(1,1); sq2 = move(sq1,0,-1); sq3 = move(sq1,-1,-1); lshape = sq1+sq2+sq3 geomplot(lshape);</pre>
Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported.
See Also	geomcsg,curve2, curve3,square1, square2

Syntax	g3 = revolve(g2,)						
-	-						
Description		g3 = revolve(g2,) revolves the 2D geometry object g2 into a 3D geometry object g3 according to given parameters.					
	The function	The function revolve accepts the following property/values:					
	TABLE I-117: VALID PROPERTY/VALUE PAIRS						
	PROPERTY	VALUES	DEFAULT	DESCRIPTION			
	angles	I-by-2 vector	[0 2*pi]	Revolution angle			
	polres	scalar	50	Polygon resolution			
	revaxis	2-by-2 matrix	[0 0; 0 1]	Revolution axis			
	wrkpln	3-by-3 matrix	[0 1 0; 0 0 1; 0 0 0]	Work plane for 2D geometric cross section			
	the plane def between the	ined by the property	wrkpln is revolv property angle	ved about the revolution axis es. angles can also be given			
	the plane def between the scalar, in whi The property	ined by the property angles defined by the ch case the first angle revaxis is a 2-by-2	wrkpln is revolv property angle is assumed to b matrix where the	wed about the revolution axis es. angles can also be given be 0. e first column specifies a poin			
	the plane def between the scalar, in whi The property the axis, and	ined by the property angles defined by the ch case the first angle revaxis is a 2-by-2 the second column sp thes the number of para	wrkpln is revolv property angle is assumed to b matrix where the pecifies the direc	wed about the revolution axis es. angles can also be given be 0. e first column specifies a point ction of the revolution axis.			
Examples	the plane def between the scalar, in whi The property the axis, and polres defin of the edges.	ined by the property angles defined by the ch case the first angle revaxis is a 2-by-2 the second column sp thes the number of para	wrkpln is revolv property angle is assumed to b matrix where the pecifies the direc	e first column specifies a point			
Examples	the plane def between the scalar, in whi The property the axis, and polres defin of the edges. Create torus	ined by the property angles defined by the ch case the first angle r revaxis is a 2-by-2 the second column sp tes the number of para	wrkpln is revolv property angle is assumed to b matrix where the pecifies the direc ameter value pai	wed about the revolution axis es. angles can also be given be 0. e first column specifies a point ction of the revolution axis.			
Examples	the plane def between the scalar, in whi The property the axis, and polres defin of the edges. Create torus re = reve	ined by the property angles defined by the ch case the first angle revaxis is a 2-by-2 the second column sp the second column sp the the number of para	wrkpln is revolv property angle is assumed to b matrix where the pecifies the direc ameter value pai	wed about the revolution axis es. angles can also be given be 0. e first column specifies a point ction of the revolution axis.			
Examples	the plane def between the scalar, in whi The property the axis, and polres defin of the edges. Create torus re = rev Create revolv p_wrkpln ax = [0	<pre>ined by the property angles defined by the ch case the first angle revaxis is a 2-by-2 the second column sp tes the number of para about the y-axis: olve(circ2(1, 'pos red object from zx-plate</pre>	wrkpln is revolv property angle is assumed to b matrix where the pecifies the direc ameter value pai ',[2 0])); ane: 'quick',{'zx' os',[1 0]),'a	<pre>ved about the revolution axis es. angles can also be given be 0. e first column specifies a point ction of the revolution axis. rs in the polygon representati ',10}); angles',[-pi/3 pi/3],</pre>			

Purpose	Rotate geon	netry object.				
Syntax	[g3,] = [g3,] = [g3,Q,c] = [g,] =	<pre>[g,] = rotate(g,r,) [g3,] = rotate(g,r,v,c,) [g3,] = rotate(g,r,vx,vy,vz,cx,cy,cz,) [g3,Q,c] = rotate(g,) [g,] = rotate(g,r,c,) [g,] = rotate(g,r,cx,cy,)</pre>				
Description	[g,] = about the <i>z</i> -		otates the 2D or 3	BD geometry object g by r radians		
	[g3,] = rotate(g,r,v,c,) rotates the 3D geometry object g by r rad about the axis v=(vx,vy,vz), with center of rotation c=(cx,cy,cz). v can als a vector of spherical coordinates, where v(1) is the polar angle, that is, the an between the axis of rotation and the positive z-axis, and v(2) is the azimuthal a of the axis of rotation.					
		<pre>- rotate(g,r,vx,v ents of the axis and c</pre>		,) is the same as above, but are explicitly given.		
	[g3,Q,c] = rotate(g,) additionally returns a rotation matrix Q corresponding to rotation given by r and v centered at the origin. The translation vector c is also returned for convenience. This means that a point set p, of size 3-by-n, containing 3D point coordinates, that is to be rotated in the same way as g, is transformed according to prot = $Q*(p-cp)+cp$, where $cp = repmat(c(:),1,size(p,2))$ represents the center of rotation.					
	<pre>[c,] = rotate(g,r,c,) rotates a 2D geometry object about the point c=(cx,cy).</pre>					
	[c,] = coordinates	,) is the sam) is the same as above, but the center			
	The function rotate accepts the following property/values:					
	TABLE I-II8: VALID PROPERTY/VALUE PAIRS					
	PROPERTY VALUE DEFAULT DESCRIPTION					
	Out stx ftx ctx empty Output parameters ptx ptx ptx ptx ptx					

See geomcsg for more information on geometry objects.

Example

The command below rotates the ellipse by 1 radian about (2,3) and plots the result.

```
e1 = ellip2(0,0,1,3);
e2 = rotate(e1,1,2,3);
geomplot(e2)
```

See Also

geomcsg

Purpose	Scale geome	Scale geometry object.				
Syntax	<pre>[g,] = scale(g3,fx,fy,fy,) [g,] = scale(g3,fx,fy,fy,x,y,z,) [g,] = scale(g3,fxyz,xyz,) [g,] = scale(g2,fx,fy,) [g,] = scale(g2,fx,fy,x,y,) [g,] = scale(g1,fx,) [g,] = scale(g1,fx,x,)</pre>					
Description	<pre>[g,] = scale(g3,xscale,yscale,zscale,) scale the 3D geometry object g3 by (xscale, yscale, zscale) about the origin.</pre>					
		scale(g,xscale,y ject g3 by (xscale,		x,y,z,) scale the 3D e) about (x,y,z).		
	[g,] =	scale(g,xyzscale	,xyz,) scale	e the 3D geometry object g3 by		
	the vector f	xyz about the vector	xyz.			
	[g,] = about the or) scale the 2D) geometry object by (fx,fy)		
	[g,] = s about (x,y)		y,) scale the	e 2D geometry object by (fx,fy)		
	[g,] = scale(g2,fx,) scale the 1D geometry object by fx about th origin.					
	[g,] =	scale(g2,fx,x,	.) scale the 1D	geometry object by fx about x.		
	The function scale accepts the following property/values:					
	TABLE I-II9: VA	LID PROPERTY/VALUE PAI	RS			
	PROPERTY	VALUE	DEFAULT	DESCRIPTION		
	Out	stx ftx ctx ptx	empty	Output parameters		
	See geomcsg for more information on geometry objects.					
Examples	The commands below scale the unit circle by $(1,2)$ about $(2,3)$ and plot the					
	c1 = circ2; c2 = scale(c1,1,2,2,3); geomplot(c2)					
See Also	geomcsg					

Purpose	Create an Argyris shape function object.
Syntax	obj = sharg_2_5(basename)
Description	The Argyris shape function object is used to implement the Argyris element of order 5 on triangles in 2D.
	$obj = sharg_2_5(basename)$ basename is a string.
	For more information, see "The Argyris Element" on page 456.
See Also	shbub, shcurl, shdens, shdisc, shdiv, shgp, shherm, shlag, shuwhelm

Purpose	Create a bubble element shape function object.
Syntax	obj = shbub(mdim,basename)
Description	The bubble element shape function object is used to implement finite elements of bubble type of order $mdim+1$ on a simplex.
	mdim is the maximum dimension of the bubble and basename is a string.
	For more information, see "Divergence Elements" on page 467.
See Also	sharg_2_5, shcurl, shdens, shdisc, shdiv, shgp, shherm, shlag, shuwhelm

Purpose	Create a vector shape function object.
Syntax	obj = shcurl(order,fieldname)
Description	The vector shape function object is used to implement finite elements of curl (edge) type of order order on all types of mesh elements (also called Nédélec elements). fieldname is a string with the field name or a cell array containing the names of the field components.
	For more information, see "The Curl Element" on page 461.
Compatibility	COMSOL Multiphysics 3.3: Replaces the shvec vector shape function object. shvec still works for backward compatibility reasons.
See Also	sharg_2_5, shbub, shdens, shdisc, shdiv, shgp, shherm, shlag, shuwhelm

Purpose	Create a density element shape function object.
Syntax	obj = shdens(mdim,order,basename)
Description	The density element shape function object is used to implement finite elements of density type on any mesh element type.
	mdim is the maximum dimension of the element, order is the default element order, and basename is a string.
	For more information, see "Density Elements" on page 465.
See Also	sharg_2_5, shbub, shcurl, shdisc, shdiv, shgp, shherm, shlag, shuwhelm

Purpose	Create a discontinuous element shape function object.
Syntax	obj = shdisc(mdim,order,basename)
Description	The discontinuous element shape function object is used to implement finite elements of discontinuous type on any mesh element type.
	mdim is the maximum dimension of the element, order is the default element order, and basename is a string.
	For more information, see "Discontinuous Elements" on page 464.
Compatibility	Since COMSOL Multiphysics 3.2 the meaning of the degrees of freedom has changed. This means that you have to re-solve models made in earlier versions of COMSOL Multiphysics that include discontinuous elements.
See Also	sharg_2_5, shbub, shcurl, shdens, shdiv, shgp, shherm, shlag, shuwhelm

Purpose	Create a divergence shape function object.
Syntax	obj = shdiv(order,fieldname)
Description	The divergence shape function object is used to implement finite elements of divergence type of order order on any type of mesh element. fieldname is a string with the field name or a cell array containing the names of the field components. For more information, see "Divergence Elements" on page 467.
Compatibility	Since COMSOL Multiphysics 3.2 the meaning of the degrees of freedom has changed. This means that you have to re-solve models made in earlier versions of COMSOL Multiphysics that include divergence elements.
	The syntax obj = shdiv(fieldname) is obsolete but still works in COMSOL Multiphysics 3.3.
See Also	sharg_2_5, shbub, shcurl, shdens, shdisc, shgp, shherm, shlag, shuwhelm

Purpose	Create a Gauss-point shape function object.
Syntax	obj = shgp(mdim, order, basename)
Description	The Gauss-point shape function object is used to implement finite elements of Gauss-point type of any order on any type of mesh element. The shape function have the degrees of freedoms in the points determined by the Gauss-point pattern for the element type. mdim is the maximum dimension of the element, order is a positive integer and determines the number of points used through the order of the Gauss-point pattern. basename is a string. The variable basename is evaluated as the degree of freedom value in the nearest Gauss point.
See Also	sharg_2_5, shbub, shcurl, shdens, shdisc, shdiv, shherm, shlag, shuwhelm

Purpose	Create a Hermite shape function object.
Syntax	obj = shherm(order, basename)
Description	The Hermite shape function object is used to implement finite elements of Hermite type of any order on mesh elements of any type. order is a positive integer greater than 2, and basename is a string. The variable basename is represented as a polynomial of degree (at most) order in the local coordinates. For more information, see "The Hermite Element" on page 458.
See Also	<pre>sharg_2_5, shbub, shcurl, shdens, shdisc, shdiv, shgp, shlag, shuwhelm</pre>

Purpose	Create a Lagrange shape function object.
Syntax	obj = shlag(order, basename)
Description	The Lagrange shape function object is used to implement finite elements of Lagrange type of any order on any type of mesh element. order is a positive integer and basename is a string. The variable basename is represented as a polynomial of degree (at most) order in the local coordinates.
	For more information, see "The Lagrange Element" on page 455.
Examples	The following three sequences set up shape functions for the variables u and v of order 1 and 2, respectively, using the standard syntax:
	fem.dim = {'u' 'v'}; fem.shape = [1 2];
	fem.dim = {'u' 'v'}; fem.shape = {'shlag(1,''v'')' 'shlag(2,''v'')'}};
	fem.dim = {'u' 'v'}; fem.shape = {'shlag(''basename'',''u'',''order'',1)' 'shlag(''basename'',''v'',''order'',2)'}
See Also	sharg_2_5, shbub, shcurl, shdens, shdisc, shdiv, shgp, shherm, shuwhelm

Purpose	Create a scalar plane-wave basis function object.
Syntax	<pre>obj = shuwhelm(ndir,basename,'kvar') obj = shuwhelm(ndir,basename,'kvar',{'xvar','yvar'}) obj = shuwhelm('ndir',ndir,'basename',basename,'kexpr','kvar', 'xexpr',{'xvar','yvar'})</pre>
Description	The scalar plane-wave basis function object, shuwhelm, implements scalar plane-wave basis functions for solving scalar wave equations of Helmholtz type using an <i>ultraweak variational formulation</i> . The plane-wave basis functions are discontinuous between mesh elements. ndir is a positive integer for the number of directions of the plane-wave basis functions and basename is a string. 'kvar' is a variable representing the wave number. You can also add expressions for the transformation of the spatial coordinates. The default values are the global <i>x</i> , <i>y</i> , and <i>z</i> (in 3D) directions, typically 'x', 'y', and 'z'. For PML domains (perfectly matched layers), where the spatial coordinates are mapped to a complex domain, the spatial coordinates in the PML domain provide the coordinate transformation, for example, 'PMLx_acpr' and 'PMLy_acpr' (in 2D), where acpr is the name of the application mode.
See Also	sharg_2_5, shbub, shcurl, shdens, shdisc, shdiv, shgp, shherm, shlag

Purpose	Constructor functions for solid objects.
Syntax	<pre>p3 = solid3(vtx,vtxpre,edg,edgpre,fac,mfdpre,mfd) [s3,] = solid3(g3,) s3 = solid3(g2) p2 = solid2(vtx,edg,mfd) [s2,] = solid2(g,) s1 = solid1(x) s1 = solid1(vtx) [s1,] = solid1(g,) s0 = solid0(full) [s0,] = solid0(p,)</pre>
Description	 s3 = solid3(vtx,vtxpre,edg,edgpre,fac,mfdpre,mfd) creates 3D solid geometry object s3 from the arguments vtx, vtxpre, edg, edgpre, fac, mfdpre, mfd. The arguments must define a valid 3D solid object. See geom3 for a description of these arguments.
	[\$3,] = \$01id3(g3,) coerces the 3D geometry object g3 to a 3D solid object \$3.
	s3 = solid3(g2) coerces the 2D geometry object g2 to a 3D solid object s3, by embedding g2 in the plane, $z = 0$.
	p2 = solid2(vtx,edg,mfd) creates a 2D solid geometry object from the properties vtx, edg, and mfd. The arguments must define a valid 2D solid object. See geom2 for a description of these arguments.
	[s2,] = solid2(g,) coerces the 2D geometry object to a 2D solid object.
	s1 = solid1(x) creates a 1D solid object that spans all the coordinate values in the vector x.
	<pre>s1 = solid1(vtx) creates a lD solid geometry object from vtx. The arguments must define a valid 2D solid object. See geom1 for a description of this argument.</pre>
	[\$1,] = solid1(g,) coerces the 1D geometry object to a 1D solid object.
	g = solid0(full) creates a 0D solid geometry object, where the Boolean full determines if the object is empty or not.
	g = solid0(p) creates a 0D solid geometry object, where p is a matrix of size 0-by-1.
	[s0,] = solid0(g,) coerces the 0D geometry object to a 0D solid object.

The coercion functions $[s0,] = solid0(g1,), [s1,] =$
<pre>solid1(g1,), [s2,] = solid2(g2,), and [s3,] =</pre>
solid3(g3,) accept the following property/values:

TABLE I-120: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Out	stx ftx ctx ptx	{}	Cell array of output names

See geomcsg for more information on geometry objects.

The nD geometry object properties are available. The properties can be accessed using the syntax get(object, property). See geom for details.

Examples The following commands create a unit curve circle object, coerce it to a curve object, and then back to a solid object.

c1 = circ2 c2 = curve2(c1) c3 = solid2(c2)

Compatibility The FEMLAB 2.3 syntax is obsolete but still supported.

See Also curve2, curve3, face3, geomcsg, geom0, geom1, geom2, geom3, point1, point2, point3

Purpose	Get number of solutions in a solution object.	
Syntax	<pre>sz = solsize(fem.sol)</pre>	
Description	<pre>sz = solsize(fem.sol) returns the number of solutions in the femsol object fem.sol.</pre>	

Purpose	Create a spherical geometry object.	
Syntax	<pre>obj = sphere3 obj = sphere2 obj = sphere3(r) obj = sphere2(r) obj = sphere3(r,) obj = sphere2(r,)</pre>	

Description

obj = sphere3 creates a solid sphere geometry object with center at the origin and semi-axes equal to 1. sphere3 is a subclass of ellipsoid3.

obj = sphere3(r, ...) creates a solid sphere object with radius r. r is a positive real scalar, or a string that evaluates to a positive real scalar, given the evaluation context provided by the property const.

The functions sphere3/sphere2 accept the following property/values:

TABLE I-121: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object
const	Cell array of strings	{}	Evaluation context for string inputs
pos	Vector of reals or cell array of strings	[0 0]	Position of the object
rot	real or string	0	Rotational angle about axis (radians)

axis sets the local z-axis, stated either as a directional vector of length 3, or as a 1-by-2 vector of spherical coordinates. axis is a vector of real scalars, or a cell array of strings that evaluate to real scalars, given the evaluation context provided by the property const. See gency13 for more information on axis.

pos sets the position of the center of the object. pos is a vector of real scalars, or a cell array of strings that evaluate to real scalars, given the evaluation context provided by the property const.

rot is an intrinsic rotational angle for the object, about its local z-axis provided by the property axis. pos is a real scalar, or a string that evaluate to a real scalar, given

the evaluation context provided by the property const. The angle is assumed to be in radians if it is numeric, and in degrees if it is a string.

obj = sphere2(...) creates a surface sphere object with the properties as given for the sphere3 function. sphere2 is a subclass of ellipsoid2.

Sphere objects have the following properties:

TABLE I-122: SPHERE OBJECT PROPERTIES

PROPERTY DESCRIPTION	
r	Radius
x, y, z, xyz	Position of the object. Components and vector forms
ax2	Rotational angle of symmetry axis
ax3	Axis of symmetry
rot	Rotational angle

In addition, all 3D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom3 for details.

Examples	The following commands create a surface and solid sphere object, where the position and radius are defined in the two alternative ways.
	s2 = sphere2(1,'pos',[0 0 0],'axis',[0 0 1],'rot',0) s3 = sphere3(4)
Compatibility	The representation of the sphere objects has been changed. The FEMLAB 2.3 syntax is obsolete but still supported. If you use the old syntax or open 2.3 models containing spheres they are converted to general face or solid objects.
See Also	geomO, geom1, geom2, geom3,ellipsoid2, ellipsoid3

Purpose	Split a geometry object.			
Syntax	[gg,] =	split(g,)		
Description	<pre>[gg,] = split(g,) returns a cell array where each cell entry contains a geometry object. When g is solid, face, curve, and point objects, the output gg contains object of the respective type. When g is a geometry object, the output contains a combination of solid, face, curve, and point objects. The function scale accepts the following property/values: TABLE 1-123: VALID PROPERTY/VALUE PAIRS</pre>			
	PROPERTY VALUE DEFAULT DESCRIPTION Out stx ftx ctx empty Output parameters ptx ptx Output parameters			
Examples	<pre>Split union of a solid circle and a solid rectangle. g = solid2(geomcsg({rect2,circ2})); gg = split(g);</pre>			
See Also	geomO, geom1, geom2, geom3			

Purpose	Create square geometry objects.					
Syntax	obj = square2 obj = square1 obj = square2(1,) obj = square1(1,)					
Description	obj = square2 creates a solid square geometry object with all side lengths equal to 1, and the lower left corner at the origin. square2 is a subclass of rect2 and solid2.					
	<pre>obj = square2(1,) creates a solid square object with side lengths equal to 1. 1 is a positive real scalar, or a string that evaluates to a positive real scalar, given the evaluation context provided by the property const.</pre>					
	The function square1 similarly creates curve square objects.					
	The functions square2/square1 accept the following property/values:					
	TABLE I-124: VALID PROPERTY/VALUE PAIRS					
	PROPERTY VALUE DEFAULT DESCRIPTION					

PROPERTY	VALUE	DEFAULT	DESCRIPTION
base	corner center	corner	Positions the object either centered about pos or with the lower left corner in pos
const	Cell array of strings	{}	Evaluation context for string inputs
pos	Vector of reals or cell array of strings	[0 0]	Position of the object
rot	Real or string	0	Rotational angle about pos (radians)

obj = square1(...) creates a curve circle geometry object with properties as given for the rect2 function. square1 is a subclass of rect1 and curve2.

Square objects have the following properties:

TABLE I-125: SQUARE OBJECT PROPERTIES

PROPERTY	DESCRIPTION
1	Side length
base	Base point
х, у	Position of the object
rot	Rotational angle

	In addition, all 2D geometry object properties are available. All properties can be accessed using the syntax get(object,property). See geom2 for details.
Example	The commands below create a unit solid square geometry object and plot it.
	<pre>sq1 = square2(1); geomplot(sq1)</pre>
Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported.
See Also	geomcsg, rect1, rect2

Purpose	Create a tangent to a 2D geometry object.
Syntax	g = tangent(g1,g2,) g = tangent(g1,p1,)
Description	g = tangent(g1,g2,) creates a common tangent line between geometry object g1 and geometry object g2.
	g = tangent(g1,p1,) creates a tangent line from geometry object g1 to a

point p1.

The function tangent accepts the following property/values pairs:

TABLE I-126: VALID PROPERTY/VALUE PAIRS

PROPERTY NAME	PROPERTY VALUE	DEFAULT	DESCRIPTION
Edim1	0 or I	geometry dependent	Starting point element dimension: 0 for vertex, 1 for edge
Edim2	0 or I	geometry dependent	Ending point element dimension: 0 for vertex, 1 for edge
Dom1	positive integer	1	Starting point domain number
Dom2	positive integer	1	Ending point domain number
Start1	number between 0 and 1	0.5	Starting point parameter value on specified edge
Start2	number between 0 and 1	0.5	Ending point parameter value on specified edge
Out	cell array of strings	{}	Additional output data (see Table 1-127)

The following properties are valid in the Out cell array:

TABLE I-127: OUTPUT DATA TYPES

ENTRY IN OUT CELL ARRAY	DESCRIPTION
Doml	Domain number of starting point
Dom2	Domain number of ending point
Paraml	Parameter value of starting point
Param2	Parameter value of ending point

TABLE I-127: OUTPUT DATA TYPES

ENTRY IN OUT CELL ARRAY	DESCRIPTION
CoordI	Coordinate of starting point
Coord2	Coordinate of ending point

Examples

The following commands generate a tangent from the unit circle to the point (2, 0) and plot the result:

```
c=circ2;
t=tangent(c,[2 0]);
geomplot(c); hold on; geomplot(t);
```

The following commands generate a common tangent between two circles and plot the result:

```
c1=circ2;
c2=circ2(1,'pos',[2 2]);
t=tangent(c1,c2,'dom1',4,'dom2',4);
geomplot(c1); hold on; geomplot(c2);
geomplot(t);
```

Purpose	Create a tetrahedron geometry object.
Syntax	t2 = tetrahedron2(p) t3 = tetrahedron3(p)
Description	t3 = tetrahedron3 creates a solid tetrahedron object with the corners at the origin and at the distance 1 from the origin along each positive coordinate axis. tetrahedron3 is a subclass of solid3.
	t3 = tetrahedron3(p) creates a solid tetrahedron object with the corners given by the four columns of p.
	t2 = tetrahedron2() creates a surface tetrahedron object. tetrahedron2 is a subclass of face3.
	The 3D geometry object properties are available. The properties can be accessed using the syntax get(object,property). See geom3 for details
Examples	The following command generates a solid tetrahedron object.
	t3 = tetrahedron3([0 0 1 1; 0 0.8 1 0; 0 0.1 0 0.2]);
See Also	face3,geom0, geom1, geom2, geom3

Create torus geometry object.
<pre>t3 = torus3 t2 = torus2 t3 = torus3(rmaj,rmin) t2 = torus2(rmaj,rmin,phi) t3 = torus3(rmaj,rmin,phi) t3 = torus3(rmaj,rmin,phi,) t2 = torus2(rmaj,rmin,phi,)</pre>
 t3 = torus3 creates a solid torus object with directrix radius 1 and generatrix radius 0.5 about the z-axis. torus3 is a subclass of solid3. t3 = torus3(rmaj,rmin) creates a solid torus with directrix radius rmaj and generatrix radius rmin, where rmaj>rmin. t3 = torus3(rmaj,rmin,phi) additionally sets the revolution angle phi of the torus.

The functions torus3/torus2 accept the following property/ values:

TABLE I-128: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Axis	Vector of reals or cell array of strings	[0 0]	Local z-axis of the object
Const	Cell array of strings	{}	Evaluation context for string inputs
Pos	Vector of reals or cell array of strings	[0 0]	Position of the bottom surface
Rot	real or string	0	Rotational angle about Axis (radians)

t3 = torus2(...) works similarly to above, but creates a surface torus object. torus2 is a subclass of face3.

Torus objects have the following properties:

PROPERTY	DESCRIPTION	
rmaj	Directrix	
rmin	Generatrix	
revang	Revolution angle	
x, y, z, xyz	Position of the object. Components and vector forms	
ax2	Rotational angle of symmetry axis	
ax3	Axis of symmetry	
rot	Rotational angle	
	In addition, all 3D geometry object properties are available. Al properties can be accessed using the syntax get(object,property). See geom3 for details.	
	For more information on geometry objects, see geom and geomcsg.	
Compatibility	The FEMLAB 2.3 syntax is obsolete but still supported. The numbering of faces, edges and vertices is different from the numbering in objects created in 2.3.	
Examples	The following commands generate a surface and solid torus, respectively.	
	t2 = torus2(2,1,pi,'pos',[0 0 0],'axis',[0 0 1]) t3 = torus3(10,2,pi/2,'pos',[1 1 1],'axis',[0 0 -100], 'rot',pi/3);	
See Also	face3,geom0, geom1, geom2, geom3	

TABLE 1-129: TORUS OBJECT PROPERTIES

Purpose	Get extended mesh information.	
Syntax	<pre>out = xmeshinfo(fem,); out = xmeshinfo(xmesh,);</pre>	
Description	The xmeshinfo function provides information about the numbering of elements, nodes, and degrees of freedom (DOFs) in the extended mesh and in the matrices returned by assemble and the solvers.	

TABLE I-130: VALID PROPERTY/VALUE PAIRS

PROPERTY	VALUE	DEFAULT	DESCRIPTION
Dofname	string cell array of strings	all	DOF names
Edim	integer vector	all	Element dimensions
Geomnum	integer vector	all	Geometry numbers
Ldof	cell array	all	Local DOFs
Lnode	real matrix	all	Local coordinates of nodes
Mcase	integer	mesh case with largest number of DOFs	Mesh case
Meshtype	vtx edg tri quad tet hex prism edg2 tri2 quad2 tet2 hex2 prism2 cell array of these strings	all	Mesh element types
Null	sparse matrix	identity	Null-space matrix for constraint elimination
Out	mcases femindex dofs nodes edims types dofnames ndofs elements cell array of these strings	dofs	Output names
Solcomp	string cell array of strings	all	DOFs solved for

The properties Mcase, Geomnum, Edim, and Meshtype determine the part of the extended mesh that information is requested for. The properties Lnode, Dofname, and Ldof determine the local nodes, DOF names, and local DOFs that information is requested for. The property Ldof is a cell array where the first row contains DOF names, and the remaining rows contain local coordinates.

NUMBERING CONVENTIONS

The numbering provided by xmeshinfo corresponds to the numbering in the mesh data structure (see femmesh). The extended mesh uses a different numbering internally. All numberings are 1-based.

- **Elements**. For each mesh element type, the element numbering of femmesh is used.
- Node points. The node points in femmesh have the same numbers in the extended mesh. Additional node points have higher numbers (these are arbitrarily ordered).
- Local node points. The numbering of the local node points within a mesh element is different from the numbering in femmesh. However, the same definition of the local coordinate system is used. In the extended mesh, the local node points are ordered in lexicographical order of their local coordinates. In femmesh, the mesh vertices come first, in lexicographical order, and then come the other node points in lexicographical order (the latter are only present for a second-order mesh).
- **DOFs**. By default, the DOF number is the index in the complete set of degrees of freedom of the model. If the property Solcomp is given, the DOF number is the index in the set of DOFs solved for. If the property Null is given, it is assumed that the Eliminate constraint handling method is used, and the DOF number is the index in the set of unconstrained DOFs. This assumes a simple form of the constraints, where each constraint only constrains one DOF. In other words, each column of the Null matrix has a single nonzero element. If Null does not have this form, an error message is given. The Null matrix is an output from the solvers (see femlin).

OUTPUTS WITH GLOBAL SCOPE

mcases = xmeshinfo(xmesh, 'out', 'mcases') returns an integer vector containing all mesh cases in the extended mesh xmesh.

femindex = xmeshinfo(xmesh, 'out', 'femindex') returns
an integer vector containing indices into xfem.fem for all
geometries in xmesh. That is, geometry geomnum in xmesh is
xfem.fem{femindex(geomnum)}.geom.

dofs = xmeshinfo(xmesh, 'out', 'dofs') returns
information about DOFs in xmesh for the mesh case given by
the property Mcase. The return value dofs is a struct with the
following fields:

FIELD	CONTENTS
mcase	Mesh case number
geomnums	Geometry numbers for all DOFs (vector)
nodes	Node numbers for all DOFs (vector)
coords	Global coordinates for all DOFs. The kth column of this matrix contains the coordinates of DOF number k
names	DOF names. Note that this is a subset of the property Dofname (if given)
nameinds	Indices into names for all DOFs (vector). The value 0 means that the DOF is not present in names
solcompinds	Indices into set of DOFs solved for (determined by property Solcomp) for all DOFs (vector). This field is only present if the Null property is given
alldofinds	Indices into total set of DOFs in the model for all DOFs (vector). This field is only present if the Solcomp property is given

TABLE I-131: DOFS STRUCT

OUTPUTS RELATED TO GEOMETRIES

nodes = xmeshinfo(xmesh, 'out', 'nodes') returns
information about nodes in the part of xmesh determined by the

properties Mcase and Geomnum. The return value nodes is a struct or a cell array of structs with the following fields:

FIELD	CONTENTS	
mcase	Mesh case number	
geomnum	Geometry number	
names	DOF names in this geometry. Note that this is a subset of the property Dofname (if given)	
dofs	DOF numbers for all nodes in this geometry. dofs(k , n) is the DOF number for DOF name names{ k } at node point n . A value 0 means that there is no DOF with this name at the node	
coords	Global coordinates for all nodes. The nth column of the matrix coords contains the coordinates of node point number n	

TABLE I-132: NODES STRUCT CORRESPONDING TO A GEOMETRY

OUTPUTS RELATED TO MESH ELEMENT TYPES

edims = xmeshinfo(xmesh, 'out', 'edims') returns a vector containing the element dimensions in the part of xmesh determined by the properties Mcase, Geomnum, Edim, and Meshtype.

types = xmeshinfo(xmesh, 'out', 'types') returns a cell array of strings containing the mesh element types in the part of xmesh determined by the properties Mcase, Geomnum, Edim, and Meshtype.

dofnames = xmeshinfo(xmesh, 'out', 'dofnames') returns a cell array of strings containing the DOF names in the part of xmesh determined by the properties Mcase, Geomnum, Edim, Meshtype, Lnode, Dofname, and Ldof.

ndofs = xmeshinfo(xmesh, 'out', 'ndofs') returns the number of DOFs in the part of xmesh determined by the properties Mcase, Geomnum, Edim, Meshtype, Lnode, Dofname, and Ldof.

elements = xmeshinfo(xmesh, 'out', 'elements') returns information about mesh elements in the part of xmesh determined by the properties Mcase, Geomnum, Edim, and Meshtype. The return value elements is a struct or a cell array of structs with the following fields:

FIELD	CONTENTS	
mcase	Mesh case number	
geomnum	Geometry number	
edim	Element dimension	
type	Mesh element type	
lnodes	Local coordinates of nodes. The kth column of the matrix 1nodes contains the coordinates of local node point number k. Note that 1nodes is a subset of the property Lnode (if given)	
nodes	Node point indices for all mesh elements of type type. nodes(k,el) is the node point number within geometry geomnum (see the output nodes) for local node point k within mesh element el. A value 0 means that there is no node point at this location	
ldofs	A cell array containing the local DOFs. The first row contains DOF names, and the remaining rows contain local coordinates. If the property Ldof is given, 1dofs is restricted to a subset. Otherwise, 1dofs is restricted by the properties Lnode and Dofname (if given)	
dofs	DOF numbers for all mesh elements of type type. dofs(k,el) is the DOF number for local DOF ldofs(:,k) within mesh element el. A value 0 means that there is no DOF at this location	

TABLE I-133: ELEMENTS STRUCT CORRESPONDING TO A MESH ELEMENT TYPE

Examples

Assume that fem.mesh is an imported NASTRAN mesh with second-order tetrahedral elements, where node point numbering starts at 1. Use second-order Lagrange elements:

```
m= meshimport('nastrandemo1.nas');
fem.mesh = m{1};
fem.dim = 'u';
fem.shape = 2;
fem.equ.c = 1;
fem.bnd.h = 1;
fem.xmesh = meshextend(fem);
```

To get the DOF number corresponding to node point number 22 in the NASTRAN mesh, type

```
nodes = xmeshinfo(fem,'out','nodes');
nodes.dofs(1,22)
```

Compute an eliminated stiffness matrix and a null-space matrix by

```
[Kc,Null]=femstatic(fem,'out',{'Kc' 'Null'});
```

To find the node point number corresponding to column 30 of Kc, and its coordinates, type

```
dofs = xmeshinfo(fem,'out','dofs','null',Null);
n = dofs.nodes(30)
dofs.coords(:,30) % alternatively:
nodes.coords(:,n)
```

To find the six DOF numbers in tetrahedron element 10 of the mesh, type

```
elements = xmeshinfo(fem,'out','elements',...
'meshtype','tet2');
elements.dofs(:,10)
```

To find the total number of DOFs on the boundary, type

```
xmeshinfo(fem,'out','ndofs','edim',2)
```

See Also

femmesh, meshextend

Diagnostics

This chapter contains lists of the most common error messages that may occur in COMSOL Multiphysics. The lists also include an explanation of the error and possible causes and workarounds.

Error Messages

This section summarizes the most common error messages and solver messages generated by COMSOL Multiphysics. All error messages are numbered and sorted in different categories according to the following table.

TABLE 2-1. ERROR HESSAGE CATEGORIES		
NUMBERS	CATEGORY	
1000-1999	Importing Models	
2000–2999	Geometry Modeling	
3000–3999	CAD Import	
4000-4999	Mesh Generation	
5000-5999	Point, Edge, Boundary, and Subdomain Specification	
6000–6999	Assembly and Extended Mesh	
7000–7999	Solvers	
8000-8999	Postprocessing	
9000–9999	General	

TABLE 2-1: ERROR MESSAGE CATEGORIES

For error messages that do not appear in the following lists, contact COMSOL's support team for help.

2000–2999 Geometry Modeling

ERROR NUMBER	ERROR MESSAGE	EXPLANATION
2118	Negative output from empty input	Incorrect Geometry M-file.
2119	Non scalar output from empty input	Incorrect Geometry M-file.
2120	Normal directions are inconsistent	Incorrect input data from STL/VRML import.
2138	Self intersections not supported	Curves resulting in self-intersections are not supported.
2140	Singular extrusions not supported	Error in input parameters.
2141	Singular revolutions not supported	The revolved mesh has a singularity at the z axis. If possible, create the cylinder using a 3D primitive or by revolving the geometry before meshing.

ERROR NUMBER	ERROR MESSAGE	EXPLANATION
2146	Subdomain must bounded at least four boundary segments	Incorrect geometry for mapped mesh.
2147	Subdomain must bound one connected edge component only	Incorrect geometry for mapped mesh.
2190	Invalid radius or distance	Incorrect input parameters to fillet/chamfer
2197	Operation resulted in empty geometry object	Geometry operation resulted in an empty geometry object which is not allowed. Make sure an empty geometry object is not created.
2209	Geometry to revolve may not cross axis of revolution	The axis of revolution and the geometry intersect. Check the dimension of the geometry and the definition of the axis for the revolution.

TABLE 2-2: GEOMETRY MODELING ERROR MESSAGES

4000–4999 Mesh Generation

ERROR NUMBER	ERROR MESSAGE	EXPLANATION
4002	A degenerated tetrahedron was created	The mesh generator ran into numerical difficulties while creating tetrahedrons with a size based on user-controlled parameters. Causes could be too small and narrow subdomains relative to the rest of the geometry or exceedingly short boundary segments. Try to avoid creating small and narrow subdomains and very short boundary segments that are adjacent to longer boundary segments.
4003	A degenerated triangle was created	The mesh generator ran into numerical difficulties while creating triangles with a size based on user-controlled parameters. Causes could be too small and narrow subdomains relative to the rest of the geometry or exceedingly short boundary segments. Try to avoid creating small and narrow subdomains and very short boundary segments that are adjacent to longer boundary segments.

ERROR NUMBER	ERROR MESSAGE	EXPLANATION	
4012	Cannot create mapped mesh for this geometry	The geometry does not fulfill the topological requirements for a mapped mesh. Changes in input parameters or further subdomain division can possibly help this.	
4026	Failed create matching edge discretizations	Cannot make mapped mesh with the given input parameters.	
4029	Failed to insert point	Problems inserting point at given coordinate. Manually inserting a point there may help.	
4031	Failed to respect boundary element on geometry edge	The mesh generator failed in making the elements compatible with the geometry object's edges. The reason for this could be that the face mesh is too coarse or contains adjacent elements with large differences in scale. Another reason can be that some subdomains in the geometry are too narrow with respect to the rest of the geometry.	
4032	Failed to respect boundary element on geometry face	See Error message 4031.	
4044	Internal error boundary respecting	See Error message 4031.	
4054	Invalid topology of geometry	The geometry object cannot be used for creating a mapped mesh. It must be subdivided.	
4055	Isolated entities found	Entities that are not connected to the boundaries of a geometry objects is found. The mapped mesh generator does not support such isolated entities.	
4119	Singular edge detected	The geometry object contains an edge of zero length.	

TABLE 2-3: MESH GENERATION ERROR MESSAGES

6000–6999 Assembly and Extended Mesh

ERROR NUMBER	ERROR MESSAGE	EXPLANATION
6008	Circular variable dependency detected	A variable has been defined in terms of itself, possibly in a circular chain of expression variables. Make sure that variable definitions are sound. Be cautious with equation variables in equations.
6063	Invalid degree of freedom name	The software does not recognize the name of a degree of freedom. Check the names of dependent variables that you have entered for the model. See also Error 7192.
6139	Wrong number of DOFs in initial value	The current solution or the stored solution has for some reason the wrong number of degrees of freedom, sometimes due to a change of the implementation of elements between two versions of the software. To overcome the problem, go to the Initial value area in the Solver Manager , and select Initial value expression . Then the initial value expressions is evaluated without using the current or stored solution.
6140	Wrong number of dofs in linearization poir	The current solution or the stored solution has for some reason the wrong number of degrees of freedom, sometimes due to a change of the implementation of elements between two versions of the software. To overcome the problem, go to the Value of variables not solved for and linearization point area in the Solver Manager, and click the Use setting from Initial value frame button or the Zero button.
6163	Divide by zero	A property in the model contains a divisor that becomes zero. Check to make sure that division by zero does not occur in any expression or coefficient.
6164	Duplicate variable name	A variable name has two different definitions. For instance, the same variable name appears two or more times for a dependent variable, a constant, an expression variable, or a coupling variable. Remove or rename one of the variables.

TABLE 2-4: ASSEMBLY AND EXTENDED MESH ERROR MESSAGES

ERROR NUMBER	ERROR MESSAGE	EXPLANATION	
6170	Failed to evaluate variable	An error occurred when evaluating the variable. The domains in which COMSOL Multiphysics tried to evaluate the variable are indicated. Also, the error message shows the expression that COMSOL Multiphysics was unable to evaluate. Make sure that you have defined the variables correctly in the indicated domains.	
6176	Attempt to evaluate real logarithm of negative number	An expression contains $log(a)$, where a becomes negative or zero. To make the logarithm well-defined, make sure that a>0. Often, a becomes only slightly negative (due to approximations in the solution process). Then, a possible solution is to use $log(a+e)$, where e is a small constant. Another remedy is to use $log(abs(a))$. If you do want to have a complex logarithm, go to the Advanced tab of Solver Parameters and select the Use complex functions with real input check box.	
6177	Matrix has zero on diagonal	When the equations have a structure such that the stiffness matrix (Jacobian matrix) has zeros on the diagonal, it is not possible to use the following linear system solvers/ preconditioners/smoothers: all versions of SOR and Diagonal scaling (Jacobi). Try the Vanka preconditioner/smoother instead.	
6194	Attempt to evaluate non-integral power of negative number	An expression contains a^b , where a becomes negative and b is non an integer. To make the power well-defined, make sure that $a>0$. Often, a becomes only slightly negative (due to approximations in the solution process). Then, a possible solution is to use $(a+e)^b$, where e is a small constant. Another remedy is to use $abs(a)^b$. If you do want to have a complex number a^b , go to the Advanced tab of Solver Parameters and select Use complex functions with real input.	

TABLE 2-4: ASSEMBLY AND EXTENDED MESH ERROR MESSAGES

ERROR NUMBER	ERROR MESSAGE	EXPLANATION
6199	Attempt to evaluate real square root of negative number	The model contains a sqrt (square root) function that takes the square root of a negative number. Either make sure that the square-root argument is nonnegative or select the Use complex functions with real input check box on the Advanced tab in the Solver Parameters dialog box.
6204	Undefined function call	An expression contains an undefined function name. Check that the function name is correct and that the function is in COMSOL Multiphysics' or MATLAB's path.
6206	Internal evaluation error: unexpected NaN encountered	Not-A-Number (NaN) appears unexpectedly. A possible cause is improperly defined coupling variables. As a first step, check that the definitions of the source and destination domains of any coupling variables or periodic boundary conditions are correct.
6245	Unsupported integration order	Integration order is too high. For triangular elements, the order can be up to 10, and for tetrahedral elements, the order can be up to 8. Find more information in the section "Numerical Quadrature" on page 472.

TABLE 2-4: ASSEMBLY AND EXTENDED MESH ERROR MESSAGES

ERROR NUMBER	ERROR MESSAGE	EXPLANATION	
7001	Adaption only implemented for tetrahedral meshes	It is only possible to use adaptive mesh refinement in 3D for models using tetrahedral mesh elements. Either turn off adaptive mesh refinement or switch from brick or prism elements to tetrahedral elements.	
7002	Adaption only implemented for triangular meshes	It is only possible to use adaptive mesh refinement in 2D for models using triangular mesh elements. Either turn off adaptive mesh refinement or switch from quadrilateral elements to triangular elements.	
7022	Segregated solver steps do not involve all of solcomp	The groups for the segregated solver do not include all dependent variables. One reason for this error could be that some boundary conditions (for example, for laminar inflow in fluid-flow models) add dependent variables that are not initially in the model.	
7033	Error in UMFPACK back substitution	The stiffness matrix (Jacobian matrix) is singular or almost singular, which leads to undefined operations in UMFPACK's back substitution. This error could appear for 3D models with a mesh where some elements are close to being two-dimensional. Try to modify or refine the mesh to avoid elements with low quality.	
7043	Initial guess leads to undefined function value	This error message usually appears when you have set up an expression that returns "not a value," that is, it is undefined, for the initial condition you have set. For instance, this happens if an expression contains a divisor that becomes zero or a logarithm of a negative value. To solve the problem, change the expression or the initial value so that the expression is well-defined when substituting the initial value of the variables. Also, watch out for warnings in the Log window.	

TABLE 2-5: SOLVER ERROR MESSAGES

TABLE 2-5: SOLVER ERROR MESSAGES

RROR NUMBER ERROR MESSAGE EXPLANATION		EXPLANATION
7067	System matrix is zero	This error message appears if there are no volume elements in the mesh. In the case that you have a mapped surface mesh, try sweeping or extruding the surface mesh to get a volume mesh.
7069	Maximum number of linear iterations reached	The iterative linear system solver did not converge due to a bad initial guess or a bad preconditioner. Increase the limit on the number of linear iterations or use a better preconditioner. If possible, use a direct linear system solver.
7081	No parameter name given	The parametric solver does not find a name for the parameter. Check the Name of parameter edit field on the General page of the Solver Manager.
7092	Out of memory in Algebraic multigrid	The Algebraic multigrid solver/ preconditioner ran out of memory. See error 7144 regarding general memory-saving tips.
7093	Out of memory during back substitution	The solver ran out of memory during back substitution. See error 7144 regarding general memory-saving tips.
7094	Out of memory during LU factorization	The solver ran out of memory during LU factorization. See error 7144 regarding general memory-saving tips.

TABLE 2-5: SOLVER ERROR MESSAGES

ERROR NUMBER	ERROR MESSAGE	EXPLANATION	
7111	Singular matrix	The system matrix (Jacobian matrix or stiffness matrix) is singular, so the solver cannot invert it. Usually this means that the system is underdetermined. Check that all equations are fully specified and that the boundary conditions are appropriate. For instance, in a stationary model you usually need to have a Dirichlet condition on some boundary. A singular matrix could also occur if mesh elements are of too low quality. If the minimum element quality is less than 0.005 you might be in trouble. Another reason for this error message is that you have different element orders for two variables that are coupled by, for example, a weak constraint. Use the same element order for all variables that are coupled.	
7136	Very ill-conditioned preconditioner. The relative residual is more than 1000 times larger than the relative tolerance	You need to improve the quality of the preconditioner to get an accurate solution. For the Incomplete LU preconditioner, lower the drop tolerance.	
7144	Out of memory in adaptive solver	The adaptive solver ran out of memory. The adaptive mesh refinement has generated a too fine mesh. In general, when you run out of memory, try to use memory-efficient modeling techniques such as utilizing symmetries, solving models sequentially, and selecting memory efficient solvers. See the chapter "Solving the Model" on page 359 in the COMSOL Multiphysics User's Guide for more information. See also the COMSOL Installation and Operations Guide for information about system memory management.	
7145	Out of memory in eigenvalue solver	The eigenvalue solver ran out of memory. See error 7144 regarding general memory-saving tips.	
7146	Out of memory in stationary solver	The stationary solver ran out of memory. See error 7144 regarding general memory-saving tips.	

TABLE 2-5: SOLVER ERROR MESSAGES

ERROR NUMBER	ERROR MESSAGE	EXPLANATION
7147	Out of memory in time-dependent solver	The time-dependent solver ran out of memory. See error 7144 regarding general memory-saving tips.
7165	Incomplete LU back substitution failed	The stiffness matrix (Jacobian matrix) is singular or almost singular, which leads to undefined operations in the Incomplete LU preconditioner's back substitution.
7192	Invalid degree of freedom name in manual scaling	The name of a dependent variable in the Manual scaling edit field on the Advanced page in the Solver Parameters dialog box does not match any of the dependent variables in the model.
7199	Reordering failed	One of the PARDISO reordering algorithms failed. Try a different reordering algorithm or try turning off row preordering.

9000–9999 General Errors

TABLE 2-6:	GENERAL	ERROR	MESSAGES

ERROR NUMBER	ERROR MESSAGE	EXPLANATION
9037	Failed to initialize 3D graphics. OpenGL not fully supported	OpenGL is not available on the computer. This can happen if your graphics card does not support OpenGL or if you have a Unix/ Linux computer where OpenGL has not been configured.
9052	Invalid address/port	You did not enter the correct server name or server port when trying to connect a client to a server.
9084	Server connection error	The client somehow lost the connection to the server. For example, the server crashed unexpectedly, or the power saving mechanism on a laptop shut down the TCP/ IP connection.

TABLE 2-6: GENERAL ERROR MESSAGES

ERROR NUMBER	ERROR MESSAGE	EXPLANATION
9143 License error	License error	The most common reasons for this message:
		The license file license.dat has been removed from the right directory in the COMSOL software installation. The license.dat file must be located in the \$COMSOL34/license directory, where \$COMSOL34 is the COMSOL 3.4 installation directory.
		The license manager has not started properly. Please find the FLEXIm log file (named by the person who started the license manager). Inspect this file to see the server status. Send it to support@comsol.com if you are in doubt about how to interpret this file.
		It is crucial that you use the correct license.dat file on both the server and the clients
9178	Error in callback	An error occurred when calling a COMSOL Script or MATLAB function from COMSOL Multiphysics. Make sure that the M-file that defines the function is correct and exists in the current path. Note that all inputs are vectors of the same size, and that the output should be a vector of the same size.

Solver Error Messages

These error messages can appear during solution and appear on the **Log** tab in the Progress window.

SOLVER ERROR MESSAGE	EXPLANATION
Cannot meet error tolerances. Increase absolute or relative tolerance.	The time-dependent solver cannot solve the model to the specified accuracy.
Error in residual computation Error in Jacobian computation	The evaluation of the residual or the Jacobian generated an error during a time-dependent solution. An additional message states the direct error. Some possible reasons are division by zero, range and overflow errors in mathematica functions, and interpolation failure in coupling variables with time-dependent mesh transformation.
Failed to find a solution	The nonlinear solver failed to converge. An additional error message gives some more details. See the description for that message.
Failed to find a solution for all parameters, even when using the minimum parameter step	During a parametric solution, the nonlinear iteration did not converge despite reducing the parameter step length to the minimum allowed value. The solution may have reached a turning point or bifurcation point.
Failed to find a solution for initial parameter	The nonlinear solver failed to converge for the initial value of the parameter during a parametric solution. An additional error message gives some more details. See the description for that message.
Failed to find consistent initial values	The time-dependent solver could not modify the initial conditions given to a DAE system to satisfy the stationary equations at the initial time. Make sure the initial values satisfy the equations and boundary conditions. In many cases, this can be achieved by solving for only the algebraic variables using a stationary solver before starting the time-dependent solver.
III-conditioned preconditioner. Increase factor in error estimate to X	The preconditioner is ill-conditioned. The error in the solution might not be within tolerances. To be sure to have a correct solution, open the Linear System Solver Settings dialog box from the General tab of Solver Parameters. Select Linear system solver in the tree, and increase Factor in error estimate to the suggested number X. Alternatively use a better preconditioner or tune the settings for the preconditioner.

TABLE 2-7: SOLVER ERROR MESSAGES IN LOG WINDOW

SOLVER ERROR MESSAGE	EXPLANATION
Inf or NaN found, even when using the minimum damping factor	Despite reducing the step size to the minimum value allowed, the solver cannot evaluate the residual or modified Newton direction at the new solution iterate. This essentially means that the current approximation to the solution is close to the boundary of the domain where the equations are well-defined. Check the equations for divisions by zero, powers, and other functions that can become undefined for certain inputs.
Inverted mesh element near coordinates (x, y, z)	In some mesh element near the given coordinates, the (curved) mesh element is (partially) warped inside-out. More precisely, the Jacobian matrix for the mapping from local to global coordinates has a negative determinant at some point. A possible reason is that the linear mesh contains a tetrahedron whose vertices all lie on a boundary. When improving the approximation of the boundary using curved mesh elements, the curved mesh element becomes inverted. To see whether this is the case, you can change Geometry shape order to 1 in the Model Settings dialog box, which means that curved mesh elements will not be used. You can usually avoid such bad tetrahedra by using a finer mesh around the relevant boundary. Another reason for this error message can be that the mesh becomes inverted when using a deformed mesh.
Last time step is not converged.	The last time step returned from the time-dependent solver is not to be trusted. Earlier time steps are within the specified tolerances.
Matrix is singular	When encountered during time-dependent solution: the linear system matrix (which is a linear combination of the mass-, stiffness-, and possibly, damping-matrices) is singular. Usually the problem originates from the algebraic part of a DAE. In particular, the cause can often be found in weak constraints or constraint-like equations like the continuity equation in incompressible flow.
Maximum number of linear iterations reached	The iterative linear system solver failed to compute a Newton direction in the specified maximum number of iterations.

TABLE 2-7: SOLVER ERROR MESSAGES IN LOG WINDOW

SOLVER ERROR MESSAGE	EXPLANATION
Maximum number of Newton iterations reached	The nonlinear solver could not reduce the error below the desired tolerance in the specified maximum number of iterations. This is sometimes a sign that the Jacobian is not complete or badly scaled. It may even be almost singular, if the system is underdetermined. If the returned solution seems reasonable, it might be enough to restart the solver with this solution as the initial guess.
No convergence, even when using the minimum damping factor	The nonlinear solver reduced the damping factor below the minimum value allowed. The solver reduces the damping factor each time a computed step did not lead to a decrease in the error estimate. Make sure the model is well-posed, in particular that there are enough equations and boundary conditions to determine all degrees of freedom. If the model is well-posed, it should have one or more isolated solutions. In that case, the error is probably due to the initial guess being too far from any solution.
Nonlinear solver did not converge	During a time-dependent solution, the nonlinear iteration failed to converge despite reducing the time step to the minimum value allowed. Usually, the error is related to the algebraic part of a DAE. For example, the algebraic equations can have reached a turning point or bifurcation point. The error can also appear when the algebraic equations do not have a unique solution consistent with the given initial conditions. Make sure algebraic equations have consistent initial values and that they have a unique solution for all times and values reached by the other variables.
Not all eigenvalues returned	When the eigenvalue solver terminated (stopped by the user or due to an error), it had not found the requested number of eigenvalues. The eigenvalues returned can be trusted.
Not all parameter steps returned	After premature termination of the parametric solver, only some of the requested solutions have been computed.
Predicted solution guess leads to undefined function value	The solver computes the initial guess for the new parameter value based on the solution for the previous parameter value. This initial guess led to an undefined mathematical operation. Try using another Predictor on the Parametric tab of Solver Parameters .

TABLE 2-7: SOLVER ERROR MESSAGES IN LOG WINDOW

SOLVER ERROR MESSAGE	EXPLANATION
Repeated error test failures. May have reached a singularity.	During a time-dependent solution, the error tolerances could not be met despite reducing the time step to the minimum value allowed.
Returned solution has not converged.	The solution returned by the stationary solver is not to be trusted. It might, however, be useful as initial guess after modifying equations or solver settings.
The elasto-plastic solver failed to find a solution	The Newton iteration loop for the computation of the plastic state at some point in the geometry did not converge.

The Finite Element Method

This chapter contains a theoretical background to the finite element method and an overview of the finite element types in COMSOL Multiphysics. Sections in this chapter also explain how COMSOL Multiphysics forms the system of equations and constraints that it solves and the implications of Dirichlet conditions involving several solution components in a multiphysics model.

Understanding the Finite Element Method

This section describes how the Finite Element Method (FEM) approximates a PDE problem with a problem that has a finite number of unknown parameters, that is, a *discretization* of the original problem. This concept introduces *finite elements*, or *shape functions*, that describe the possible forms of the approximate solution.

Mesh

The starting point for the finite element method is a mesh, a partition of the geometry into small units of a simple shape, *mesh elements*. For more information about the types of elements that are available in 1D, 2D, and 3D, see "Mesh Elements" on page 286.

Sometimes the term "mesh element" means any of the mesh elements—mesh faces, mesh edges, or mesh vertices. When considering a particular d-dimensional domain in the geometry (that is, a subdomain, boundary, edge, or vertex), then by its mesh elements you mean the d-dimensional mesh elements contained in the domain.

Finite Elements

Once you have a mesh, you can introduce approximations to the dependent variables. For this discussion, concentrate on the case of a single variable, u. The idea is to approximate u with a function that you can describe with a finite number of parameters, the so-called *degrees of freedom* (DOF). Inserting this approximation into the weak form of the equation generates a system of equations for the degrees of freedom.

Start with a simple example: linear elements in 1D. Assume that a mesh consists of just two mesh intervals: 0 < x < 1 and 1 < x < 2. Linear elements means that on each mesh interval the continuous function u is linear (affine). Thus, the only thing you need to know in order to characterize u uniquely is its values at the *node points* $x_1 = 0$, $x_2 = 1$, and $x_3 = 2$. Denote these as $U_1 = u(0)$, $U_2 = u(1)$, $U_3 = u(2)$. These are the *degrees of freedom*.

Now you can write

$$u(x) = U_1 \varphi_1(x) + U_2 \varphi_2(x) + U_3 \varphi_3(x)$$

where $\varphi_i(x)$ are certain piecewise linear functions. Namely, $\varphi_i(x)$ is the function that is linear on each mesh interval, equals 1 at the *i*th node point, and equals 0 at the other node points. For example,

$$\varphi_1(x) = \begin{cases} 1-x & \text{if } 0 \le x \le 1\\ 0 & \text{if } 1 \le x \le 2 \end{cases}$$

The $\varphi_i(x)$ are called the *basis functions*. The set of functions u(x) is a linear function space called the *finite element space*.

For better accuracy, consider another finite element space corresponding to quadratic elements. Functions u in this space are second-order polynomials on each mesh interval. To characterize such a function, introduce new node points at the midpoint of each mesh interval: $x_4 = 0.5$ and $x_5 = 1.5$. You must also introduce the corresponding degrees of freedom $U_i = u(x_i)$. Then, on each mesh interval, the second-degree polynomial u(x) is determined by the degrees of freedom at the endpoints and the midpoint. In fact, you get

$$u(x) = U_1 \varphi_1(x) + U_2 \varphi_2(x) + U_3 \varphi_3(x) + U_4 \varphi_4(x) + U_5 \varphi_5(x)$$

where the basis functions $\varphi_i(x)$ now have a different meaning. Specifically, $\varphi_i(x)$ is the function that is quadratic on each mesh interval, equals 1 at the *i*th node point, and equals 0 at the other node points. For example,

$$\varphi_1(x) = \begin{cases} (1-x)(1-2x) & \text{if } 0 \le x \le 1 \\ 0 & \text{if } 1 \le x \le 2 \end{cases}$$

In general, you specify a finite element space by giving a set of basis functions. The description of the basis functions is simplified by the introduction of *local coordinates* (or *element coordinates*). Consider a mesh element of dimension d in an n-dimensional geometry (whose space coordinates are denoted $x_1,...,x_n$). Consider also the *standard d-dimensional simplex*

$$\xi_1 \ge 0, \xi_2 \ge 0, \dots, \xi_d \ge 0, \xi_1 + \dots + \xi_d \le 1$$

which resides in the local coordinate space parametrized by the local coordinates ξ_1 , ..., ξ_d . If d = 1, then this simplex is the unit interval. If d = 2, it is a triangle with two 45 degree angles, and if d = 3 it is a tetrahedron. Now you can consider the mesh element as a linear transformation of the standard simplex. Namely, by letting the

global space coordinates x_i be suitable linear (affine) functions of the local coordinates, you get the mesh element as the image of the standard simplex.

When described in terms of local coordinates, the basis functions assume one of a few basic shapes. These are the *shape functions*. In the example with linear elements in 1D, any basis function on any mesh element is one of the following:

$$\phi = \xi_1, \qquad \phi = 1 - \xi_1, \qquad \phi = 0$$

Thus the first two are the shape functions in this example (0 is not counted as a shape function). In the example with quadratic elements in 1D, the shape functions are

$$\phi = (1 - \xi_1)(1 - 2\xi_1), \qquad \phi = 4\xi_1(1 - \xi_1), \qquad \phi = \xi_1(2\xi_1 - 1)$$

CURVED MESH ELEMENTS

When using higher-order elements (that is, elements of an order > 1), the solution has a smaller error. The error also depends on how well the mesh approximates the true boundary. To keep errors in the finite element approximation and the boundary approximation at the same level, it is wise to use *curved mesh elements*. They are distorted mesh elements that can approximate a boundary better than ordinary straight elements (if the problem's boundary is curved). You can get curved mesh elements by writing the global coordinates x_i as polynomials of order k (the *geometry shape order*) in the local coordinates ξ_j . (The earlier example took k = 1). Then the mesh element is the image of the standard simplex. For mesh elements that do not touch the boundary, there is no reason to make them curved, so they are straight. It is customary to use the same order k here as for the order of the (Lagrange) element. This is referred to as using *isoparametric elements*.

The order k is determined by the geometry shape order for the frame (coordinate system) associated with the finite element. You can control the geometry shape order using the **Model Settings** dialog box. The frame is determined by the property frame to the finite element (the default is the reference frame); see "Shape Function Variables" on page 171. For certain finite elements, the geometry shape order given by the frame can be overridden by the property sorder.

If a curved mesh element becomes too distorted, it can become inverted and cause problems in the solution (see "Avoiding Inverted Mesh Elements" on page 356 in the *COMSOL Multiphysics User's Guide*).

THE LAGRANGE ELEMENT

The preceding examples are special cases of the Lagrange element. Consider a positive integer k, the order of the Lagrange element. The functions u in this finite element space are piecewise polynomials of degree k, that is, on each mesh element u is a polynomial of degree k. To describe such a function it suffices to give its values in the Lagrange points of order k. These are the points whose local (element) coordinates are integer multiples of 1/k. For example, for a triangular mesh in 2D with k = 2, this means that you have node points at the corners and side midpoints of all mesh triangles. For each of these node points p_i , there exists a degree of freedom $U_i = u(p_i)$ and a basis function φ_i . The restriction of the basis function φ_i to a mesh element is a polynomial of degree (at most) k in the local coordinates such that $\varphi_i = 1$ at node i, and $\varphi_i = 0$ at all other nodes. Thus the basis functions are continuous and you have

$$u = \sum_{i} U_i \varphi_i$$

The Lagrange element of order 1 is called the linear element. The Lagrange element of order 2 is called the quadratic element.

The Lagrange elements are available with all types of mesh elements. The order k can be arbitrary, but the available numerical integration formulas usually limits its usefulness to $k \le 5$ ($k \le 4$ for tetrahedral meshes).

Syntax for the Lagrange Element (shlag)

To specify a Lagrange shape function in the **shape** edit field on the **Element** page of **Subdomain** settings, enter a string of the form shlag(k, basename), where k is the order (a positive integer) and *basename* is the name of the variable (a string enclosed in single quotes), for example, shlag(2, 'u'). There is also an alternative syntax shlag(...) based on property names and values. The following properties are allowed:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
basename	variable name		Base variable name
order	positive integer		Basis function order
frame	string	reference frame	Frame
border	positive integer		Alias for order
sorder	positive integer	determined by frame	Geometry shape order

TABLE 3-1: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHLAG SHAPE FUNCTION

It is not possible to abbreviate the property names, and you must write them in lowercase letters enclosed in single quotation marks. For example:

```
shlag('order',2,'basename','u')
```

Note: When using the property name/value syntax for shlag in COMSOL Script or MATLAB, you must enter the command as a string with each string argument enclosed in two single quotes because they become strings within a string: 'shlag(''order'',2,''basename'',''u'')'.

The Lagrange element defines the following variables. Denote **basename** with u, and let x and y denote (not necessarily distinct) names of space coordinates. The variables are (where sdim = space dimension and edim = mesh element dimension):

- u
- ux, meaning the derivative of u with respect to x, defined on edim = sdim
- *uxy*, meaning a second derivative, defined on edim = sdim
- *uTx*, the tangential derivative variable, meaning the *x*-component of the tangential projection of the gradient, defined on edim < sdim
- *uTxy*, meaning *xy*-component of the tangential projection of the second derivative, defined when edim < sdim

When computing the derivatives, the global space coordinates are expressed as polynomials of degree (at most) sorder in the local coordinates.

Note: The use of isoparametric elements means that u is not a polynomial in the global coordinates (if k > 1), only in the local coordinates.

THE ARGYRIS ELEMENT

For a function represented with Lagrange elements, the first derivatives between mesh elements can be discontinuous. In certain equations (for example, the biharmonic equation) this can be a problem. The *Argyris element* has basis functions with continuous derivatives between mesh triangles (it is defined in 2D). The second order derivative is continuous in the triangle corners. On each triangle, a function u in the Argyris finite element space is a polynomial of degree 5 in the local coordinates.

The Argyris element is available with triangular meshes only.

When setting Dirichlet boundary conditions on a variable that has Argyris shape functions, a locking effect can occur if the boundary is curved and constraint order (cporder) 5 is used. Use cporder=4 if the boundary is curved and cporder=5 for straight boundaries.

Syntax for the Argyris Element (sharg_2_5)

To specify Argyris shape functions in the **shape** edit field on the **Element** tab in the **Subdomain Settings** dialog box, enter a string of the form sharg_2_5(*basename*), where *basename* is the name of the variable (a string enclosed in single quotes), for example, sharg_2_5('u'). There is also an alternative syntax sharg_2_5(...) based on property names and values. The following properties are allowed:

TABLE 3-2: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHARG SHAPE FUNCTION

PROPERTY	VALUE	DEFAULT	DESCRIPTION
basename	variable name		Base variable name
frame	string	reference frame	Frame

The property names cannot be abbreviated and must be written in lowercase letters enclosed in single quotation marks.

```
Example: sharg_2_5('basename', 'u').
```

Note: When using the property name/value syntax for sharg in COMSOL Script or MATLAB, you must enter the command as a string with each string argument enclosed in two single quotes because they become strings within a string: 'sharg_2_5(''basename'', ''u''')'.

The Argyris element defines the following degrees of freedom (where u is the base name and x and y are the space coordinate names):

- *u* at corners
- *ux* and *uy* at corners, meaning derivatives of *u*
- uxx, uxy, and uyy at corners, meaning second derivatives
- *u*n at side midpoints, meaning a normal derivative. The direction of the normal is to the right if moving along an edge from a corner with lower mesh vertex number to a corner with higher number

The Argyris element defines the following field variables (where sdim = space dimension = 2 and edim = mesh element dimension):

- u
- *ux*, meaning the derivative of *u* with respect to *x*
- *uxy*, meaning a second derivative, defined for edim = sdim and edim = 0
- *uxTy*, the tangential derivative variable, meaning the *y*-component of the tangential projection of the gradient of *ux*, defined for 0 < edim < sdim

When computing the derivatives, the global space coordinates are always expressed with shape order 1 in the Argyris element.

THE HERMITE ELEMENT

On each mesh element, the functions in the Hermite finite element space are the same as for the Lagrange element, namely, all polynomials of degree (at most) k in the local coordinates. The difference lies in which DOFs are used. For the Hermite element, a DOF u exists at each Lagrange point of order k, except at those points adjacent to a corner of the mesh element. These DOFs are the values of the function. In addition, other DOFs exist for the first derivatives of the function (with respect to the global coordinates) at the corners (ux and uy in 2D). Together, these DOFs determine the polynomials completely. Note that the functions in the Hermite finite element space have continuous derivatives between mesh elements at the mesh vertices. However, at other common points for two mesh elements, these derivatives are not continuous. Thus, you can think of the Hermite element as lying between the Lagrange and Argyris elements.

The Hermite element is available with all types of mesh elements. The order $k \ge 3$ can be arbitrary, but the available numerical integration formulas usually limits its usefulness to $k \le 5$ ($k \le 4$ for tetrahedral meshes).

When setting Dirichlet boundary conditions on a variable that has Hermite shape functions, a locking effect can occur if the boundary is curved and the constraint order cporder is the same as the order of the Hermite element. This means that the derivative becomes over constrained at mesh vertices at the boundary, due to the implementation of the boundary conditions. To prevent this locking, you can specify cporder to be the element order minus 1.

Syntax for the Hermite Element (shherm)

To specify Hermite shape functions in the **shape** edit field on the **Element** tab in the **Subdomain Settings** dialog box, enter a string of the form shherm(k, basename),

where k is the order (an integer > 2), and *basename* is the name of the variable (a string enclosed in single quotes), for example shherm(3, 'u'). There is also an alternative syntax shherm(...) based on property names and values. The following properties are allowed:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
basename	variable name		Base variable name
order	integer >= 3		Basis function order
frame	string	reference frame	Frame
border	integer		Alias for order
sorder	positive integer	determined by frame	Geometry shape order

TABLE 3-3: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHHERM SHAPE FUNCTION

The property names cannot be abbreviated and must be written in lowercase letters enclosed in single quotation marks.

Example: shherm('order',3,'basename','u').

Note: When using the property name/value syntax for shherm in COMSOL Script or MATLAB, you must enter the command as a string with each string argument enclosed in two single quotes because they become strings within a string: 'shherm(''order'',3,''basename'',''u'')'.

The Hermite element defines the following degrees of freedom:

- The value of the variable basename at each Lagrange node point that is not adjacent to a corner of the mesh element.
- The values of the first derivatives of basename with respect to the global space coordinates at each corner of the mesh element. The names of these derivatives are formed by appending the space coordinate names to basename.

The Hermite element defines the following field variables. Denote basename with u, and let x and y denote (not necessarily distinct) names of space coordinates. The variables are (where sdim = space dimension and edim = mesh element dimension):

• u

- *ux*, meaning the derivative of *u* with respect to *x*, defined when edim = sdim or edim=0
- *uxy*, meaning a second derivative, defined when edim = sdim
- *uTx*, the tangential derivative variable, meaning the *x*-component of the tangential projection of the gradient, defined when 0 < edim < sdim
- *uTxy*, meaning *xy*-component of the tangential projection of the second derivative, defined when edim < sdim

When computing the derivatives, the global space coordinates are expressed as polynomials of degree (at most) sorder in the local coordinates.

BUBBLE ELEMENTS

Bubble elements have shape functions that are zero on the boundaries of the mesh element and have a maximum in the middle of the mesh element. The shape function (there is only one for each mesh element) is defined by a lowest-order polynomial that is zero on the boundary of the element.

The bubble element are available with all types of mesh elements.

Syntax for Bubble Elements (shbub)

To specify discontinuous shape functions in the **shape** edit field on the **Element** tab in the **Subdomain Settings** dialog box, enter a string of the form shbub(*mdim*, *basename*), where *mdim* is the dimension of the mesh elements for which the shape functions exist, and *basename* is the name of the variable (a string enclosed in single quotes), for example shbub(3, 'u'). There is also an alternate syntax shbub(...) based on property names and values. The following properties are allowed:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
basename	variable name		Base variable name
mdim	nonnegative integer	sdim	Dimension of the mesh elements on which the bubble exist
frame	string	reference frame	Frame
sorder	positive integer	determined by frame	Geometry shape order

TABLE 3-4: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHBUB SHAPE FUNCTION

The property names cannot be abbreviated and must be written in lowercase letters enclosed in single quotation marks.

Example: shbub('mdim',3,'basename','u').

Note: When using the property name/value syntax for shbub in COMSOL Script or MATLAB, you must enter the command as a string with each string argument enclosed in two single quotes because they become strings within a string: 'shbub(''mdim'',3,''basename'',''u'')'.

The bubble element has a single degree of freedom, **basename**, at the midpoint of the mesh element.

The bubble element defines the following field variables. Denote basename with u, and let x and y denote (not necessarily distinct) names of space coordinates. The variables are (where sdim = space dimension and edim = mesh element dimension):

- u, defined when edim \leq mdim, u = 0 if edim < mdim.
- *ux*, meaning the derivative of *u* with respect to *x*, defined when edim = mdim = sdim.
- uTx, the tangential derivative variable, meaning the *x*-component of the tangential projection of the gradient, defined when mdim < sdim and edim \leq mdim . uTx = 0 if edim < mdim.
- *u*T*xy*, meaning the *xy*-component of the tangential projection of the second derivative, defined when mdim < sdim and edim ≤ mdim . *u*T*xy* = 0 if edim < mdim.

THE CURL ELEMENT

In electromagnetics, *curl elements* (also called *vector elements* or *Nédélec's edge elements*) are popular. Each mesh element has DOFs corresponding only to tangential components of the field. For example, in a tetrahedral mesh in 3D each of the three edges in a triangle face element has degrees of freedom that are tangential components of the vector field in the direction of the corresponding edges, and in the interior there are degrees of freedom that correspond to vectors tangential to the triangle itself (if the element order is high enough). Finally, in the interior of the mesh tetrahedron there a degrees of freedom in all coordinate directions (if the element order is high enough). This implies that tangential components of the vector field are continuous across element boundaries, but the normal component is not necessarily continuous. This also implies that the curl of the vector field is an integrable function, so these elements are suitable for equations using the curl of the vector field.

The curl elements are available for all types of mesh elements. The polynomial order of the curl element can be at most 3 in 3D, and at most 4 in 2D and 1D.

Syntax for the Curl Element (shcurl)

To specify curl shape functions in the **shape** edit field on the **Element** page in the **Subdomain Settings** dialog box, enter a string of the form shcurl(*k*, *fieldname*) where *fieldname* is the name of the vector field (a string enclosed in single quotes), and *k* is the order (a positive integer), for example shcurl(3, 'E'). Alternatively, use the syntax shcurl(*k*, *compnames*), where *compnames* is a cell array of strings with the vector components, for example shcurl(3, {'Ex' 'Ey'}). There is also a syntax shcurl(...) based on property names and values. The following properties are allowed:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
fieldname	string		Field name
compnames	cell array of strings	derived from fieldname	Names of vector field components
dofbasename	string	See below	Base name of degrees of freedom
dcompnames	string	See below	Names of the anti-symmetrized components of the gradient of the vector field
order	integer		Basis function order
frame	string	reference frame	Frame
border	positive integer	order	Alias for order
sorder	positive integer	given by frame	Geometry shape order

TABLE 3-5: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHCURL SHAPE FUNCTION

The property names cannot be abbreviated and must be written in lowercase letters enclosed in single quotation marks.

Example: shcurl('compnames', {'Ex' 'Ey'}, 'dofbasename', 'tE').

Note: When using the property name/value syntax for shcurl in COMSOL Script or MATLAB, you must enter the command as a string with each string argument (including arguments within arguments) enclosed in two single quotes because they become strings within a string:

'shcurl(''compnames'',''{''Ex'',''Ey''}'',''dofbasename'',''tE'')'.

The default for compnames is fieldname concatenated with the space coordinate names. The default for dofbasename is tallcomponents, where allcomponents is the concatenation of the names in compnames.

The property dcompnames lists the names of the component of the antisymmetric matrix

$$dA_{ij} = \frac{\partial A_j}{\partial x_i} - \frac{\partial A_i}{\partial x_i}$$

where A_i are the vector field components and x_i are the space coordinates. The components are listed in row order. If a name is the empty string, the field variable corresponding to that component is not defined. If you have provided compnames, the default for the entries in dcompnames is compnames(*j*) sdimnames(*i*) compnames(*i*) sdimnames(*j*) for off-diagonal elements. If only fieldname has been given, the default for the entries are dfieldname sdimnames(*i*)sdimnames(*j*). Diagonal elements are not defined per defaults. For example, shcurl('order',3,'fieldname', 'A', 'dcompnames', $\{'', '', 'curlAy', 'curlAz', '', '', 'curlAx', ''\}$).

The curl element defines the following degrees of freedom: dofbasename dc, where d = 1 for DOFs in the interior of an edge, d = 2 for DOFs in the interior of a surface, etc., and c is a number between 0 and d - 1.

The curl element defines the following field variables (where comp is a component name from compnames, and dcomp is a component from dcompnames, sdim = space dimension and edim = mesh element dimension):

- comp, meaning a component of the vector, defined when edim = sdim.
- tcomp, meaning one component of the tangential projection of the vector onto the mesh element, defined when edim < sdim.

- comp*x*, meaning the derivative of a component of the vector with respect to global space coordinate *x*, defined when edim = sdim.
- tcompTx, the tangential derivative variable, meaning the *x* component of the projection of the gradient of tcomp onto the mesh element, defined when edim < sdim. Here, *x* is the name of a space coordinate.
- dcomp, meaning a component of the anti-symmetrized gradient, defined when edim = sdim.
- tdcomp, meaning one component of the tangential projection of the anti-symmetrized gradient onto the mesh element, defined when edim < sdim.

For performance reasons, prefer using **dcomp** in expressions involving the curl rather than writing it as the difference of two gradient components.

For the computation of components, the global space coordinates are expressed as polynomials of degree (at most) sorder in the local coordinates.

DISCONTINUOUS ELEMENTS

The functions in the discontinuous elements space are the same as for the Lagrange element, with the difference that the basis functions are discontinuous between the mesh elements. All degrees of freedom are located in the element interior.

The discontinuous elements are available with all types of mesh elements. The polynomial order k can be arbitrary, but the available numerical integration formulas usually limits its usefulness to $k \le 5$ ($k \le 4$ for tetrahedral meshes).

Syntax for the Discontinuous Element (shdisc)

To specify discontinuous shape functions in the **shape** edit field on the **Element** tab in the **Subdomain Settings** dialog box, enter a string of the form shdisc(*mdim*, *order*, *basename*), where mdim is the dimension of the mesh elements for which the shape functions exist, *order* is the order (a positive integer) and *basename* is the name of the variable (a string enclosed in single quotes), for example shdisc(3,2, 'u'). There is also an alternative syntax shdisc(...) based on property names and values. The following properties are allowed:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
basename	variable name		Base variable name
order	integer		Basis function order

TABLE 3-6: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHDISC SHAPE FUNCTION

TABLE 3-6: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHDISC SHAPE FUNCTION

PROPERTY	VALUE	DEFAULT	DESCRIPTION
mdim	nonnegative integer	sdim	Dimension of the mesh elements where the discontinuous element exists
frame	string	reference frame	Frame
border	nonnegative integer	order	Alias for order
sorder	positive integer	given by frame	Geometry shape order

The property names cannot be abbreviated and must be written in lowercase letters enclosed in single quotation marks.

Example: shdisc('mdim',3,'order',2,'basename','u').

Note: When using the property name/value syntax for shdisc in COMSOL Script or MATLAB, you must enter the command as a string with each string argument enclosed in two single quotes because they become strings within a string: 'shdisc(''mdim'',3,''order'',2,''basename'',''u'')'.

The discontinuous element defines the following field variables. Denote basename with u, and let x denote names of space coordinates. The variables are (where edim is the mesh element dimension):

- *u*, defined when edim = mdim.
- ux, meaning the derivative of u with respect to x, defined when edim = mdim = sdim.
- *uTx*, the tangential derivative variable, meaning the derivative of *u* with respect to *x*, defined when edim = mdim < sdim.

DENSITY ELEMENTS

The functions in the density elements space are the same as for the discontinuous element, if the mesh element is not curved. If the element is curved, the functions define a density of the given order in *local* coordinates and the value in global coordinates is dependent on the transformation between local coordinates and global coordinates.

The discontinuous elements are available with all types of mesh elements. The order k can be arbitrary, but the available numerical integration formulas usually limits its usefulness to $k \le 5$ ($k \le 4$ for tetrahedral meshes).

Syntax for the Density Element (shdens)

To specify discontinuous shape functions in the **shape** edit field on the **Element** tab in the **Subdomain Settings** dialog box, enter a string of the form shdens(*order*, *basename*), where *order* is the order (a positive integer) and *basename* is the name of the variable (a string enclosed in single quotes), for example shdens(2, 'u'). There is also an alternative syntax shdens(...) based on property names and values. The following properties are allowed:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
basename	variable name		Base variable name
order	integer		Basis function order
frame	string	reference frame	Frame
border	nonnegative integer	order	Alias for order
sorder	positive integer	given by frame	Geometry shape order

TABLE 3-7: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHDENS SHAPE FUNCTION

The property names cannot be abbreviated and must be written in lowercase letters enclosed in single quotation marks.

Example: shdens('order',2,'basename','u').

Note: When using the property name/value syntax for shdens in COMSOL Script or MATLAB, you must enter the command as a string with each string argument enclosed in two single quotes because they become strings within a string: 'shdens(''order'',2,''basename'',''u'')'.

The density element defines the following field variable. Denote basename with u. The variable is (where edim is the mesh element dimension):

• *u*, defined when edim = sdim.

DIVERGENCE ELEMENTS

For modeling the **B** (magnetic flux density) and **D** (electric displacement) fields in electromagnetics, the divergence elements are useful. The DOFs on the boundary of a mesh element correspond to normal components of the field. In addition, there are DOFs corresponding to all vector field components in the interior of the mesh element of dimension sdim (if the order is high enough). This implies that the normal component of the vector field is continuous across element boundaries, but the tangential components are not necessarily continuous. This also implies that the divergence of the vector field is an integrable function, so these elements are suitable for equations using the divergence of the vector field.

The divergence element are available with all types of mesh elements. The polynomial order of the curl element can be at most 3 in 3D, and at most 4 in 2D and 1D.

Syntax for Divergence Elements (shdiv)

To specify divergence shape functions in the **shape** edit field on the **Element** tab in the **Subdomain Settings** dialog box, enter a string of the form shdiv(*fieldname*) where *fieldname* is the name of the vector field (a string enclosed in single quotes), for example shdiv('B'). Alternatively, use the syntax shdiv(*compnames*), where *compnames* is a cell array of strings with the vector components, for example, shdiv({'Bx' 'By'}). There is also a syntax shdiv(...) based on property names and values. The following properties are allowed:

PROPERTY	VALUE	DEFAULT	DESCRIPTION
fieldname	variable name		Name of vector field
compnames	cell array of strings	derived from fieldname	Names of vector field components
dofbasename	string	see below	Base name of degrees of freedom
divname	string	see below	Name of divergence field
order	integer	1	Basis function order
frame	string	reference frame	Frame
border	positive integer	order	Alias for order
sorder	positive integer	given by frame	Geometry shape order

TABLE 3-8: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHDIV SHAPE FUNCTION

The property names cannot be abbreviated and must be written in lowercase letters enclosed in single quotation marks.

Example: shdiv('compnames', {'Bx' 'By'}, 'dofbasename', 'nB').

Note: When using the property name/value syntax for shdiv in COMSOL Script or MATLAB, you must enter the command as a string with each string argument (including arguments within arguments) enclosed in two single quotes because they become strings within a string:

'shdiv(''compnames'',''{''Bx'',''By''}'',''dofbasename'',''nB'')'.

The default for compnames is fieldname concatenated with the space coordinate names. The default for dofbasename is nallcomponents, where allcomponents is the concatenation of the names in compnames.

The vector element defines the following degrees of freedom: dofbasename on element boundaries, and dofbasename sdim c, c = 0, ..., sdim - 1 for DOFs in the interior.

The divergence element defines the following field variables (where comp is a component name from compnames, divname is the divname, sdim = space dimension and edim = mesh element dimension):

- comp, meaning a component of the vector, defined when edim = sdim.
- ncomp, meaning one component of the projection of the vector onto the normal of mesh element, defined when edim = sdim-1.
- comp*x*, meaning the derivative of a component of the vector with respect to global space coordinate *x*, defined when edim = sdim.
- ncompTx, the tangential derivative variable, meaning the *x* component of the projection of the gradient of ncomp onto the mesh element, defined when edim < sdim. Here, *x* is the name of a space coordinate. ncompTx = 0.
- divname, means the divergence of the vector field.

For performance reasons, prefer using divname in expressions involving the divergence rather than writing it as the sum of sdim gradient components.

For the computation of components, the global space coordinates are expressed as polynomials of degree (at most) sorder in the local coordinates.

SCALAR PLANE WAVE BASIS FUNCTION

The scalar plane wave basis function, shuwhelm, is used to implement scalar plane wave basis functions for solving scalar wave equations of Helmholtz type using an *ultraweak*

variational formulation (UWVF). These basis functions are discontinuous in between mesh elements.

Syntax for the Scalar Plane Wave Basis Function (shuwhelm) To specify scalar plane wave basis functions in the **shape** edit field on the **Element** page in the **Subdomain Settings** dialog box, enter a string of the form shuwhelm(*ndir*, *basename*, *kvar*), where *ndir* is the number of directions for the waves (a positive integer), *basename* is the name of the variable (a string enclosed in single quotes), and *kvar* is the name of a variable for the wave number (a string enclosed in single quotes), for example shuwhelm(1, 'p', 'k'). In addition, you can use the syntax shuwhelm(*ndir*, *basename*, *kvar*, {*xvar*, *yvar*}) to specify the expressions for the spatial coordinate transformation as strings in a cell array, such as {'x', 'y'} (in 2D). In domains that represent perfectly matched layers, the spatial coordinates are mapped to a complex domain, and special spatial coordinate variables provide the transformation of the spatial coordinates. There is also an alternative syntax shuwhelm(...) based on property names and values. The following properties are allowed:

PROPERTY VALUE DESCRIPTION DEFAULT Base variable name basename variable name Number of wave directions ndir integer Variable for the wave number kexpr string {'x','y','z'} xexpr cell array of Expressions for the x, y, and z coordinate transformations strings

TABLE 3-9: VALID PROPERTY NAME/VALUE PAIRS FOR THE SHUWHELM WAVE BASIS FUNCTION

The property names cannot be abbreviated and must be written in lowercase letters enclosed in single quotation marks.

Example: shuwhelm('ndir',2,'basename','u').

Note: When using the property name/value syntax for shuwhelm in COMSOL Script or MATLAB, you must enter the command as a string with each string argument enclosed in two single quotes because they become strings within a string: 'shuwhelm(''ndir'',2, ''basename'', ''u''')'.

The scalar plane wave basis function defines the following field variables. Denote **basename** with u, and let x denote names of a space coordinates. The variables are (where sdim = space dimension):

• u

• ux, meaning the derivative of u with respect to x, defined on edim = sdim

Discretization of the Equations

This section describes how COMSOL Multiphysics forms the discretization of the PDE. Consider a 2D problem for simplicity. The starting point is the weak formulation of the problem. First comes the discretization of the constraints

$$0 = R^{(2)} \text{ on } \Omega$$
$$0 = R^{(1)} \text{ on } B$$
$$0 = R^{(0)} \text{ on } P$$

starting with the constraints on the boundaries, *B*. For each mesh element in *B* (that is, each mesh edge in *B*), consider the Lagrange points of some order *k* (see "The Lagrange Element" on page 455). Denote them by $x_{mj}^{(1)}$, where *m* is the index of the mesh element. Then the discretization of the constraint is

$$0 = R^{(1)}(x_{mj}^{(1)}),$$

that is, the constraints must hold pointwise at the Lagrange points. The Lagrange point order k can be chosen differently for various components of the constraint vector $R^{(1)}$, and it can also vary in space. COMSOL Multiphysics' data structures denote the k as cporder The constraints on subdomains Ω and points P are discretized in the same way. (Nothing needs to be done with the points P.) You can collect all these pointwise constraints in one equation 0 = M, where M is the vector consisting of all the right-hand sides.

COMSOL Multiphysics approximates the dependent variables with functions in the chosen finite element space(s). This means that the dependent variables are expressed in terms of the degrees of freedom as

$$u_l = \sum_i U_i \varphi_i^{(l)}$$

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where $\varphi_i^{(l)}$ are the basis functions for variable u_l . Let U be the vector with the degrees of freedoms U_i as its components. This vector is called the *solution vector* because it is what you want to compute. M depends only on U, so the constraints can be written 0 = M(U).

Now consider the weak equation:

$$0 = \int_{\Omega} W^{(2)} dA + \int_{B} W^{(1)} ds + \sum_{P} W^{(0)}$$
$$-\int_{\Omega} v \cdot h^{(2)T} \mu^{(2)} dA - \int_{B} v \cdot h^{(1)T} \mu^{(1)} ds - \sum_{P} v \cdot h^{(0)T} \mu^{(0)}$$

where $\mu^{(i)}$ are the Lagrange multipliers. To discretize it, express the dependent variables in terms of the DOFs as described earlier. Similarly, approximate the test functions with the same finite elements (this is the Galerkin method):

$$v_l = \sum_i V_i \varphi_i^{(l)}$$

Because the test functions occur linearly in the integrands of the weak equation, it is enough to require that the weak equation holds when you choose the test functions as basis functions:

$$v_l = \varphi_i^{(l)}$$

When substituted into the weak equation, this gives one equation for each i. Now the Lagrange multipliers must be discretized. Let

$$\Lambda_{mj}^{(d)} \,=\, \mu^{(d)}(x_{mj}^{(d)}) w_{mj}^{(d)}$$

where $x_{mj}^{(d)}$ are the Lagrange points defined earlier, and $w_{mj}^{(d)}$ are certain weights (see the following discussion). The term

$$\int_{B} \varphi_i \cdot h^{(1)T} \mu^{(1)} ds$$

is approximated as a sum over all mesh elements in B. The contribution from mesh element number m to this sum is approximated with the Riemann sum

$$\sum_{j} \varphi_{i}(x_{mj}^{(1)}) \cdot h^{(1)T}(x_{mj}^{(1)}) \mu^{(1)}(x_{mj}^{(1)}) w_{mj}^{(1)} = \sum_{j} \varphi_{i}(x_{mj}^{(1)}) \cdot h^{(1)T}(x_{mj}^{(1)}) \Lambda_{mj}^{(1)}$$

where $w_{mj}^{(1)}$ is the length (or integral of ds) over the appropriate part of the mesh element. The integral over Ω and the sum over P is approximated similarly.

All this means that you can write the discretization of the weak equation as

$$0 = L - N_F \Lambda$$

where L is a vector whose *i*th component is

$$\int_{\Omega} W^{(2)} dA + \int_{B} W^{(1)} ds + \sum_{P} W^{(0)}$$

evaluated for $v_l = \varphi_i^{(l)}$. A is the vector containing all the discretized Lagrange multipliers $\Lambda_{mj}^{(d)}$. N_F is a matrix whose *i*th row is a concatenation of the vectors

$$\varphi_i(x_{mj}^{(d)})h^{(d)}(x_{mj}^{(d)})^T$$

For problems using *ideal constraints*, N_F is equal to the *constraint Jacobian matrix* N, which is defined as

$$N = -\frac{\partial M}{\partial U}$$

To sum up, the discretization of the stationary problem is

$$0 = L(U) - N_F(U)\Lambda$$
$$0 = M(U)$$

The objective is to solve this system for the solution vector U and the Lagrange multiplier vector Λ . L is called the *residual vector*, M is the *constraint residual*, and N_F is the *constraint force Jacobian matrix*. Note that M is redundant in the sense that some pointwise constraints occur several times. Similarly, Λ is redundant. Solvers remove this redundancy.

NUMERICAL QUADRATURE

The integrals occurring in the components of the residual vector L (as well as K, as noted later in this discussion) are computed approximately using a *quadrature formula*. Such a formula computes the integral over a mesh element by taking a

weighted sum of the integrand evaluated in a finite number of points in the mesh element. The *order* of a quadrature formula on a 1D, triangular, or tetrahedral element is the maximum number k such that it exactly integrates all polynomials of degree k. For a quadrilateral element, a formula of order k integrates exactly all products $p(\xi_1)q(\xi_2)$, where p and q are polynomials of degree k in the first and second local coordinates, respectively. A similar definition holds for hexahedral and prism elements. Thus the accuracy of the quadrature increases with the order. On the other hand, the number of evaluation points also increases with the order. As a rule of thumb, you can take the order to be twice the order of the finite element being used. COMSOL Multiphysics' data structures refer to the order of the quadrature formula as gporder (gp stands for Gauss points). The maximum available order of the quadrature formula (the gporder value) is:

- 41 for 1D, quadrilateral, and hexahedral meshes
- 30 for triangular and prism meshes
- 8 for tetrahedral meshes

TIME-DEPENDENT PROBLEMS

The discretization of a time-dependent problem is similar to the stationary problem

$$0 = L(U, U, U, t) - N_F(U, t) \Lambda$$
$$0 = M(U, t)$$

where U and Λ now depend on time t.

LINEARIZED PROBLEMS

Consider a linearized stationary problem (see "The Linear or Linearized Model" on page 366). The linearization "point" u_0 corresponds to a solution vector U_0 . The discretization of the linearized problem is

$$\begin{split} K(U_0)(U-U_0) + N_F(U_0)\Lambda &= L(U_0) \\ N(U_0)(U-U_0) &= M(U_0) \end{split}$$

where K is called the *stiffness matrix*, and $L(U_0)$ is the *load vector*. For problems given in general or weak form, K is the Jacobian of -L:

$$K = -\frac{\partial L}{\partial U}$$

The entries in the stiffness matrix are computed in a similar way to the load vector, namely by integrating certain expressions numerically. This computation is called the *assembling* the stiffness matrix.

If the original problem is linear, then its discretization can be written

$$KU + N_F \Lambda = L(0)$$
$$NU = M(0)$$

Similarly, for a time-dependent model the linearization involves the *damping matrix*

$$D = -\frac{\partial L}{\partial \dot{U}}$$

and the mass matrix

$$E = -\frac{\partial L}{\partial \ddot{U}}$$

When E = 0, the matrix D is often called the mass matrix instead of the damping matrix.

EIGENVALUE PROBLEMS

The discretization of the eigenvalue problem is

$$\lambda^{2} E(U_{0})U - \lambda D(U_{0})U + K(U_{0})U + N_{F}(U_{0})\Lambda = 0$$
$$N(U_{0})U = 0$$

where U_0 is the solution vector corresponding to the linearization "point." If the underlying problem is linear, then D, K, and N do not depend on U_0 , and you can write

$$KU + N_F \Lambda = \lambda DU - \lambda^2 EU$$
$$NU = 0$$

WEAK CONSTRAINTS

Weak constraints present an alternative way to discretize the Dirichlet conditions, as opposed to the pointwise constraints described earlier. The idea is to regard the Lagrange multipliers $\mu^{(d)}$ as field variables and thus approximate them with finite

elements. This concept also introduces corresponding test functions $v^{(d)}$. Multiply the Dirichlet conditions with these test functions and integrate to end up with the following system in the case of a stationary problem in 2D:

$$0 = \int_{\Omega} W^{(2)} dA + \int_{B} W^{(1)} ds + \sum_{P} W^{(0)}$$

- $\int_{\Omega} v \cdot h^{(2)T} \mu^{(2)} dA - \int_{B} v \cdot h^{(1)T} \mu^{(1)} ds - \sum_{P} v \cdot h^{(0)T} \mu^{(0)}$
$$0 = \int_{\Omega} v^{(2)} \cdot R^{(2)} dA$$

$$0 = \int_{B} v^{(1)} \cdot R^{(1)} ds$$

$$0 = \sum_{P} v^{(0)} \cdot R^{(0)}$$

You could add these weak equations to form a single equation. This treatment of the Lagrange multipliers as ordinary variables has thus produced a weak equation without constraints. This can be useful if the Lagrange multipliers are of interest in their own right.

Take care when combining pointwise and weak constraints. For instance, if you have both types of constraints for some variable and the constraints are in adjacent domains, the resulting discretization does not work. Note that you can obtain pointwise constraints from the weak constraints formulation by using the basis functions

$$\delta(x-x_{mj}^{(d)})$$

for the Lagrange multipliers and their test functions, that is, let

$$\mu^{(d)} = \sum_{m,j} \Lambda^{(d)}_{mj} \delta(x - x^{(d)}_{mj})$$

where δ is Dirac's delta function.

What Equations Does COMSOL Multiphysics Solve?

This section explains how COMSOL Multiphysics forms the system of equations and constraints that it solves. It also discusses the implications of Dirichlet conditions involving several solution components in a multiphysics model.

You specify material parameters and boundary conditions in a number of *application modes*. Enter these settings in the **Subdomain Settings**, **Boundary Settings**, **Edge Settings**, and **Point Settings** dialog boxes, which you open from the **Physics** menu. Each application mode forms one or several PDEs and boundary conditions from these settings. If you use one of the PDE modes, you specify the equation coefficients and terms directly.

The software collects all the equations and boundary conditions formulated by the application modes into one large system of PDEs and boundary conditions. This process also includes converting equations and boundary conditions to the selected *solution form*, which can be *coefficient form*, *general form*, or *weak form*. You can select the solution form on the **Advanced** page in the **Solver Parameters** dialog box. This dialog box also provides the option *automatic* (the default setting), which means that you let the software select the solution form. COMSOL Multiphysics uses the weak solution form unless you have chosen to use the adaptive solver. The software then selects the general solution form because the adaptive solver does not work with the weak form. If any of the equation system forms is weak, the solution form is also the weak form even if you use the adaptive solver because it is not possible to convert the equations from weak form to general form.

In the **Model Settings** dialog box you can specify the *equation system form*—the form of the system of equations and boundary conditions that you can see in the **Equation System** dialog boxes. This form can differ from the solution form. If it does, COMSOL Multiphysics first converts the equations to the solution form before solving. If you use a PDE mode, notice the difference between the form of the PDE in the application mode, the equation system form, and the solution form.

Occasionally you might want to change the PDEs generated by the application modes. You can do this by editing the settings in **Equation System>Subdomain Settings** dialog box (see "Viewing and Modifying the Full Equation System" on page 213 in the *COMSOL Multiphysics User's Guide*). Similarly, you can change the boundary conditions generated by the application modes in the **Equation System>Boundary Settings** dialog box (see "Modifying Boundary Settings for the Equation System" on page 237 in the *COMSOL Multiphysics User's Guide*). If you have PDEs or constraints on edges or points, you can also modify the equations that the application modes generate in the **Equation System>Edge Settings** and **Equation System>Point Settings** dialog boxes.

The Equation System/Solution Forms

COEFFICIENT FORM

In the coefficient equation system form, the PDEs and boundary conditions are written in following form (for a time-dependent model):

$$e_{a}\frac{\partial^{2} u}{\partial t^{2}} + d_{a}\frac{\partial u}{\partial t} + \nabla \cdot (-c\nabla u - \alpha u + \gamma) + \beta \cdot \nabla u + \alpha u = f \quad \text{in } \Omega$$
$$\mathbf{n} \cdot (c\nabla u + \alpha u - \gamma) + qu = g - h^{T}\mu \qquad \text{on } \partial\Omega$$

$$hu = r$$
 on $\partial\Omega$

In addition to these PDEs, there can be weak-form contributions; see the **weak** and **dweak** edit fields on the **Weak** page of the **Equation System>Subdomain Settings** dialog box. If these edit fields are nonzero, COMSOL Multiphysics modifies the above PDE by:

- Converting the PDE to the weak form by multiplying it by a test function, integrating, and integrating the flux term by parts.
- Adding the **dweak** term to the left side of the resulting weak equation and adding the **weak** term to the right side.
- Adding weak-form contributions from the Equation System>Boundary Settings, Equation System>Edge Settings, and Equation System>Point Settings dialog boxes to the resulting weak equation.

In addition to the above Dirichlet boundary condition, hu = r, there can be additional constraints in the **constr** edit field on the **Weak** page of the **Equation System>Boundary Settings** dialog box. The expressions in the **constr** edit field are constrained to be equal to zero. Similarly, the constraints in the **constr** edit fields in **Equation System>Subdomain Settings, Equation System>Edge Settings**, and **Equation System>Point Settings** dialog boxes are enforced.

GENERAL FORM

In the general equation system form, the PDEs and boundary conditions are written in following form (for a time-dependent model):

$$\begin{cases} e_a \ddot{u} + d_a \dot{u} + \nabla \cdot \Gamma = F & \text{in } \Omega \\ -\mathbf{n} \cdot \Gamma = G - h^T \mu & \text{on } \partial \Omega \\ 0 = R & \text{on } \partial \Omega \end{cases}$$

Just like for the coefficient form, you can modify this PDE by adding weak-form contributions in the **weak** and **dweak** edit field (see earlier discussion). Similarly, there can be additional constraints in the **constr** edit fields.

WEAK FORM

In the weak equation system form, the PDEs are written solely in the weak formulation (see the weak and dweak edit fields on the Weak tab of Equation System>Subdomain Settings, Equation System>Boundary Settings, Equation System>Edge Settings, and Equation System>Point Settings dialog boxes). The dweak fields contribute to the left side of the equations, and the weak fields contribute to the right side of the equations.

You specify constraints in the constr edit fields in the Equation System>Subdomain Settings, Equation System>Boundary Settings, Equation System>Edge Settings, and Equation System>Point Settings dialog boxes. These expressions are constrained to be equal to zero.

The Full Equation System

In addition to the PDEs and boundary conditions that you can view in the dialog boxes from the **Equation System** submenu on the **Physics** menu, there can sometimes be extra contributions, which are generated by the application modes or by periodic conditions and identity conditions. You cannot view these contributions directly in the user interface but only in the Model M-file or by exporting the FEM structure to the command window. The extra contributions show up in the fields elemmph and elemcpl of the FEM structure. They occur in the following cases:

- · Periodic conditions generate extra constraints.
- Identity conditions generate extra constraints.

- Dirichlet boundary conditions for the tangential component of a vector field discretized using vector elements generate extra constraints.
- The Shell application mode in the Structural Mechanics Module generates extra contributions to the equations.

The full system of equations and constraints is approximated using the finite element method; see "Discretization of the Equations" on page 470.

Notes on Constraints in Multiphysics Models

In a multiphysics model, if a Dirichlet boundary condition involves two different dependent variables and there is also a Neumann boundary condition, that Neumann boundary condition is not the one displayed in the application mode. The displayed Neumann boundary condition is modified by adding an extra Lagrange multiplier term on the right-hand side.

To explain this, assume that you want to solve the system of PDEs

$$\begin{cases} -u_{xx} = 1\\ -v_{xx} = 1 \end{cases}$$

on the interval 0 < x < 1 with the Dirichlet boundary conditions u = 0 and v = 0 at x = 0, and u = v at x = 1, and the Neumann boundary condition $v_x = 0$ at x = 1. Use two PDE, Coefficient Form application modes (one for u and one for v) and the general equation system form.

The general form boundary conditions at x = 1 (which you can inspect in the **Equation System>Boundary Settings** dialog box) read

$$\begin{cases} -\mathbf{n} \cdot \Gamma = G - h^T \mu \\ 0 = R \end{cases}$$

where

$$\mathbf{n} = 1, \quad \Gamma = \begin{bmatrix} -u_x \\ -v_x \end{bmatrix}, \quad G = \begin{bmatrix} 0 \\ 0 \end{bmatrix}, \quad R = \begin{bmatrix} v - u \\ 0 \end{bmatrix}$$

The matrix h is

$$h = -\begin{bmatrix} \frac{\partial R_1}{\partial u} & \frac{\partial R_1}{\partial v} \\ \frac{\partial R_2}{\partial u} & \frac{\partial R_2}{\partial v} \end{bmatrix} = \begin{bmatrix} 1 & -1 \\ 0 & 0 \end{bmatrix}$$

Thus, the resulting boundary conditions are

$$\begin{cases} \begin{bmatrix} u_x \\ v_x \end{bmatrix} = \begin{bmatrix} -1 & 0 \\ 1 & 0 \end{bmatrix} \begin{bmatrix} \mu_1 \\ \mu_2 \end{bmatrix}$$
$$\begin{bmatrix} 0 \\ 0 \end{bmatrix} = \begin{bmatrix} v - u \\ 0 \end{bmatrix}$$

The boundary condition $u_x = -\mu_1$ is expected. It just says that $-u_x$ is equal to the Lagrange multiplier μ_1 , but because μ_1 is an unknown this condition can be eliminated. However, the condition $v_x = \mu_1$ is not expected and it invalidates the argument. The resulting boundary condition is $v_x = -u_x$. Note that the boundary-condition description in the PDE application mode for v incorrectly states that the Neumann condition is $v_x = 0$.

The reason for this unexpected result is the Lagrange multiplier term μ_1 in the right-hand side of the Neumann boundary condition for *v*. Such Lagrange multiplier terms are often called *constraint forces* in a structural mechanics model. The moral is that if you have a Dirichlet boundary condition involving both *u* and *v*, you get constraint forces in the Neumann boundary conditions for *both* variables *u* and *v*. This means that the Neumann boundary-condition description in the application modes must be modified if you use such Dirichlet boundary conditions.

The term $h^{T}\mu$ in the right-hand side of the Neumann boundary condition is what characterizes *ideal constraints*. In the above example, you would like to have a *non-ideal constraint* where the term $h^{T}\mu$ is changed to

$$\begin{bmatrix} -1 & 0 \\ 0 & 0 \end{bmatrix} \begin{bmatrix} \mu_1 \\ \mu_2 \end{bmatrix}$$

⊢ ¬

To accomplish this, you can remove the constraint specification on the **Coefficients** tab of the **Boundary Settings** dialog box. Instead select **User defined** from the **Constraint type** list on the **Weak** page. Then enter the constraint and the constraint force independently in the **constr** and **constrf** edit fields, respectively.

Advanced Geometry Topics

4

This chapter describes some advanced geometry topics that are part of the solid modeling tools in COMSOL Multiphysics.

Advanced Geometry Topics

Rational Bézier Curves

A rational Bézier curve is a parameterized curve of the form

$$\mathbf{b}(t) = \frac{\displaystyle\sum_{i=0}^{p} \mathbf{b}_{i} w_{i} B_{i}^{p}(t)}{\displaystyle\sum_{i=0}^{p} w_{i} B_{i}^{p}(t)} \quad , 0 \leq t \leq 1$$

where the functions

$$B_i^p(t) = \binom{p}{i} t^i (1-t)^{p-i}$$

are the *Bernstein basis* functions of *degree* p; $\mathbf{b}_i = (x_1, ..., x_n)$ are the control vertices of the *n*-dimensional space; and w_i are the weights, which should always be positive real numbers. A rational Bézier curve has a direction defined by the parameter *t*. This direction is used to uniquely determine subdomain numbers to the left and to the right of a curve in 2D.

Note: The parameter t used in this section is named s or s1 when used as a variable. It does not represent the arc length of a curve but is equivalent to the Bézier parameter as described above.

The end-point interpolation property corresponds to $\mathbf{b}(0) = \mathbf{b}_0$ and $\mathbf{b}(1) = \mathbf{b}_p$. Another useful property of the rational Bézier curves is that the direction of the tangent vector at t = 0 and t = 1 is determined by the vectors $\mathbf{b}_1 - \mathbf{b}_0$ and $\mathbf{b}_p - \mathbf{b}_{p-1}$, respectively. That is, the curve will always be tangent to the line connecting the control vertices \mathbf{b}_0 and \mathbf{b}_1 and also to the line connecting \mathbf{b}_{p-1} and \mathbf{b}_p . When joining curves at end points, aligning the (nonzero) tangent vectors assures tangential continuity. This technique produces visually smooth transitions between adjacent curves.

There is also an interaction between the control polygon and the curve. For instance, the curve is always contained in the convex hull of its control polygon,

 $\{\mathbf{b}_0, \mathbf{b}_1, ..., \mathbf{b}_p\}$. A useful property is that of invariance under translation, rotation, and scaling. Translating, rotating, or scaling the control polygon by a certain amount, translates, rotates, or scales the curve that the polygon defines by exactly the same amount. In formal terms, this property of rational Bézier curves is called *affine invariance*.

A rational Bézier curve is equivalent to a polynomial Bézier curve (or simply a Bézier curve) if the control weights w_0, \ldots, w_p are all equal. In this case the denominator equals the binomial expansion of $(t + (1 - t))^p$, in which each term is one of the Bernstein basis functions. This implies that the polynomial Bézier curves are a subset of the rational Bézier curves.

Note that a line could be viewed as a rational Bézier curve of degree 1.

Conic Sections

Rational Bézier curves of degree 2 can represent all conic sections: circles, ellipses, parabolas, and hyperbolas. Elliptical or circular curve segments are often called arcs. The conic sections are also called quadric curves or quadrics. Because the parameter *t* is constrained to be in the interval [0, 1], only a segment of the conic section is represented. A 2nd degree curve consists of three control vertices and three weights. There is a simple rule for classifying a 2nd degree curve if the end point weights are set to 1, only allowing the central weight w_1 to vary: if $w_0 = w_2 = 1$, then $0 < w_1 < 1$ gives ellipses, $w_1 = 1$ gives parabolas, and $w_1 > 1$ gives hyperbolas. For a fixed control polygon, at most one value of w_1 (among the ellipses generated by letting $0 < w_1 < 1$) gives a circle segment. For example, a quarter of a full circle is generated by a control polygon with a right angle and with a central weight of $1 / \sqrt{2}$.

RELATION TO CURVES ON IMPLICIT FORM.

A rational Bézier curve of degree 2 is a *rational parameterization* of an algebraic curve of degree 2, that is, a curve on the familiar implicit form for quadrics

$$ax^2 + by^2 + cxy + dx + ey + f = 0$$

The unit circle, for example, has a=b=1, f=-1 and c=d=e=0. The set of rational Bézier curves of degree 2 is essentially equivalent to the set of algebraic curves of degree 2.

Cubic Curves

Rational Bézier curves of degree 3 (cubic curves) have more dynamic properties than conic section curves. A cubic curve has four control vertices and four weights, making it possible, for example, to create a self-intersecting control polygon or a zigzag control polygon. A self-intersecting polygon may give rise to a self-intersecting curve, a loop.

A zigzag control polygon generates an S-shaped curve containing a point of inflection where the tangent line lies on both sides of the curve.

A curve with a cusp is a limiting case of a curve with a loop. A cusp occurs when a loop shrinks so that the area enclosed in the loop approaches zero. At the cusp the tangent vector of the curve vanishes. That is, the curve has no well-defined tangent line at the cusp.

RELATION TO CURVES ON IMPLICIT FORM

The set of rational Bézier curves of degree 3 is a strict subset of the set of algebraic curves of degree 3, that is, curves that contain terms of the type x^3 , x^2y , xy^2 , y^3 , x^2 , and so on in their implicit form. This is because some algebraic curves of degree 3 do not have a rational parameterization.

Rational Bézier Surfaces

When you create a 3D geometry object with a curved boundary, COMSOL Multiphysics represents it by rational Bézier surfaces. The software supports two types of Bézier surfaces: rectangular and triangular. A rectangular Bézier surface has a mixed degree (m, n), which represents the degree of the surface in terms of two parameters, often named *s* and *t*. A triangular Bézier surface has a single degree, *m*, just as a Bézier curve.

A rectangular rational Bézier surface of degree p-by-q is described by

$$\mathbf{S}(s,t) = \frac{\sum_{i=0}^{p} \sum_{j=0}^{q} \mathbf{b}_{i,j} w_{i,j} B_{i}^{p}(s) B_{j}^{q}(t)}{\sum_{i=0}^{p} \sum_{j=0}^{q} w_{i,j} B_{i}^{p}(s) B_{j}^{q}(t)}, \ 0 \le s, t \le 1 \ ,$$

where B_i^p and B_j^q are the Bernstein basis functions of degree p and q, respectively, as described in the previous section. This surface description is called rectangular because

the parameter domain is rectangular, that is, the two parameters s and t can vary freely in given intervals.

Another form of surface description is the triangular surface, also called a *Bézier triangle*. A triangular rational Bézier surface is defined as

$$\mathbf{S}(s,t) = \frac{\displaystyle\sum_{i+j \leq p} \mathbf{b}_{i,j} w_{i,j} B_{i,j}^p(s,t)}{\displaystyle\sum_{i+j \leq p} w_{i,j} B_{i,j}^p(s,t)} \quad , 0 \leq s,t \leq 1$$

which differs from the Bézier curve description only by the use of *bivariate* Bernstein polynomials instead of *univariate*, for the curve case. The bivariate Bernstein polynomials of degree *p* are defined as

$$B^{p}_{i,j}(s,t) = \frac{p!}{i!j!(p-i-j)!}s^{i}t^{j}(1-s-t)^{p-i-j}, \quad i+j \le p$$

where the parameters s and t must fulfill the conditions

$$\left\{ \begin{array}{l} 0 \le s, t \\ s+t \le 1 \end{array} \right.$$

which form a triangular domain in the parameter space, therefore the name of this surface description.

The normal vector, $\mathbf{n}(s, t)$, for a point, $\mathbf{S}(s, t)$, at a surface that is defined as

$$\mathbf{n}(s,t) = \frac{\partial \mathbf{S}}{\partial s}(s,t) \times \frac{\partial \mathbf{S}}{\partial t}(s,t)$$

determines the direction of the surface. This direction is used to define the up- and down subdomains of a surface.

The Bézier surfaces are contained in the convex hull of their control points. Bézier surfaces also have the affine invariance property: invariance of surface under translation, rotation, and scaling. Boundary curves of a Bézier surface are Bézier curves, and the corners in the parameter grid that define the control points all lie on the surface.

The simplest form of surface is a *plane*. A Bézier triangle of degree 1 can define a plane spanned by three distinct control points. A rectangular Bézier surface of degree (1,1), on the other hand, forms a bilinear surface where the boundary curves are lines.

COMSOL Multiphysics supports rectangular surfaces of mixed degree at most (3,3) and triangular surfaces of degree 1 to represent planar surfaces. Rectangular rational Bézier surfaces of mixed degree up to (2,2) can represent all common CAD surfaces, including bilinear surfaces, cylinders, cones, spheres, ellipsoids, and tori. The (3,3)-degree rational Bézier curves assist in the creation of additional free-form surfaces. To model a cone or a cylinder you need a rectangular surface of degree (2,1). Modeling a sphere or a torus requires rectangular surfaces of degree (2,2).

CONTROL VERTICES AND WEIGHTS

A rectangular rational Bézier surface of degree (m,n) is defined by a control net consisting of (m+1)-by-(n+1) control vertices assigned a positive weight. The surface always interpolates the four corner points of the control net. A change in the net's shape produces a change in the surface's appearance. Its shape mimics that of the control net. The higher the surface degree the more complicated the shapes you can create. Increasing the weight pulls the curve toward the corresponding control vertex. This interaction between the control net and the surface makes the rational Bézier surface representation useful.

TRIMMED SURFACES

The 3D geometry objects in COMSOL Multiphysics are formed by a set of trimmed rational Bézier surfaces. A cylinder consists of four trimmed rectangular degree (2,1) surfaces and two trimmed triangular planar surfaces. The planar surfaces are trimmed by boundary curves in the parameter space so only a circular portion of each planar surface is used. For the curved surfaces, the boundary curves in the parameter space are lying on the rectangular boundary of the surface.

When using geometry modeling operations, the Bézier surfaces are trimmed by the intersection curves between surfaces. By trimming surfaces, surface boundaries can take virtually any shape. The connected surfaces of a 3D geometry object are called faces. A surface can be divided into any number of faces, which are curved areas bounded by trimming (intersection) curves.

Note: The parameters s and t used in this section are equivalent to the variables s1 and s2.

RATIONAL BÉZIER REPRESENTATIONS AND NURBS

The *NURBS* representation (*nonuniform rational B-spline*) is another popular curve and surface representation scheme. It is usually possible to split a curve having a NURBS representation into a sequence of rational Bézier curves.

Parameterization of Curves and Surfaces

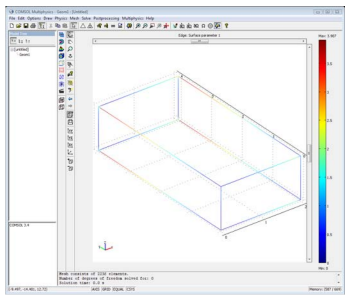
The curves and surfaces of a geometry object can have several mathematical representations. Thus, a local parameter s_1 is defined for curves, and two local parameters s_1 and s_2 are defined for faces. These parameters prove helpful when setting up a model or postprocessing the solution. More precisely, for each value of the curve parameter s_1 within its domain of definition, there is a unique point on the curve, while each pair of values (s_1 , s_2) corresponds to a unique point on a face.

The faces and edges in the COMSOL geometry representation consist of trimmed surfaces and curves, respectively. Thus there is a well-defined boundary in the parameter domain that determines the valid values of s_1 and s_2 . In 2D, the possible values of the curve parameter s_1 often lie in the interval [0, 1], but in 3D the parameter domain is more complicated for surfaces as well as for curves.

The best way to determine the parameterization is to plot the parameter values. For a block, you can do this in the way described here:

- I Open the Model Navigator, select 3D in the Space dimension list and click OK.
- 2 Draw a **Block** object with side lengths 2, 4, and 1.
- **3** To enter Postprocessing mode, where the software displays the parameterization of the curves and faces, click the **Solve Problem** button on the Main toolbar.
- **4** Open the **Plot Parameters** dialog box, for example by clicking the **Plot Parameters** button on the Main toolbar. Clear the **Slice** plot type check box and select **Edge** plot instead.
- 5 Find the Edge page in the Plot Parameters dialog box and set the Expression to s1.

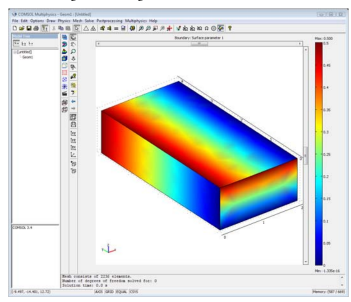




The different geometric variables available for plotting appear in the tables in "Geometric Variables" on page 165.

7 To visualize the surface parameters on the faces, select Boundary plot in the PlotParameters dialog box, and enter either s1 or s2 as the Expression on the Boundary

page. Make sure to clear the **Smooth** check box before plotting to avoid incorrect smoothing over the edges.



Geometric Variables

Note: In the following table of geometric variables, replace all letters in italic font with the actual names for the independent variables (spatial coordinates) in the model. Replace x, y, and z with the first, second, and third spatial coordinate variable, respectively. If the model contains a deformed mesh, you can replace the symbols x, y, z with either the spatial coordinates (x, y, z by default) or the reference coordinates (X, Y, Z by default).

The geometric variables in the following table characterize geometric properties.

DOMAIN \ SPACE DIM	ID	2D	3D	ND
POINT				
EDGE			s1,t1x,t1y, t1z	

DOMAIN \ SPACE DIM	ID	2D	3D	ND
BOUNDARY	dnx,nx,unx	dnx,dny,nx, ny,s,tx,ty, unx,uny	dnx, dny, dnz, nx, ny, nz, s1, s2, t1x, t1y, t1z, t2x, t2y, t2z, unx, uny, unz	
SUBDOMAIN				h,sd, reldetjac, reldetjacmin
ALL	x,xg,dvol	x,y,xg,yg, dvol	x,y,z,xg,yg, zg,dvol	dvol,dom, meshtype, meshelement

In this table the space dimension refers to the number of independent variables. Most geometric variables of interest are defined on boundaries.

The variables xg, yg, and zg contain the spatial coordinate values of the original geometry as opposed to the standard spatial coordinate variables x, y, and z, which are based on polynomial shape functions. It is the standard spatial coordinate variables' values that COMSOL Multiphysics uses to compute the solution. The difference between these two sets of spatial coordinate variables is normally very small. If a deformed mesh is used, the variables xg, yg, zg are only available when x, y, z are the reference coordinates (X, Y, Z by default).

Advanced Solver Topics

This chapter describes some advanced solver settings in COMSOL Multiphysics settings that you for most simulations need not worry about. It also examines the various solvers in COMSOL Multiphysics in some detail.

Advanced Solver Settings

The **Advanced** page in the **Solver Parameters** dialog box controls solver settings you normally do not need to change.

Analysis:	General Parametric Stationary	Adaptive Advanced
Auto select solver	Constraint handling method: Null-space function:	Elimination + Automatic +
Stationary Time dependent Eigenvalue Parametric	Assembly block size:	5000 Instraint matrix and in symmetry detection al input
Stationary segregated Parametric segregated	Stop if error due to undefined Store solution on file Solution form:	Automatic +
*	Scaling of variables Type of scaling:	Automatic
Adaptive mesh refinement	and the second	
Adaptive mesh refinement	Manual scaling: Row equilibration:	On v
Adaptive mesh refinement	Manual scaling: Row equilibration:	0n •
Adaptive mesh refinement	Manual scaling: Row equilibration:	On 🔹

The Advanced page of the Solver Parameters dialog box.

The following sections describe the settings on this page.

Constraint Handling, Null-Space Functions, and Assembly Block Size

The **Constraint handling method** list controls how COMSOL Multiphysics handles constraints. The default elimination method is always preferable, but see "Constraint Handling" on page 499 for details.

The default selection in the **Null-space function** list is **Automatic**, which means that COMSOL Multiphysics chooses the most appropriate of the orthonormal and sparse null-space functions. To override this choice, select **Orthonormal** or **Sparse** in the **Null-space function** list. The orthonormal null-space function computes an orthonormal basis for the null space of the constraint Jacobian *N*. For models that involve constraints (Dirichlet boundary conditions) that couple values at several nodes,

this method typically runs out of memory. For example, this can happens if the model contains:

- · Periodic boundary conditions
- Identity conditions
- · Constraints involving coupling variables, for instance integral constraints
- Constraints involving derivatives (but you should always rewrite constraints on normal derivatives as Neumann boundary conditions).

In these cases, the sparse null-space function performs better than the orthonormal. On the other hand, the sparse null-space function has the following drawbacks:

- It does not always work well together with the Geometric multigrid solver/ preconditioner.
- The computation of initial values is less efficient.
- When two boundaries with different Dirichlet boundary conditions meet, the value at the joining node is less predictable. In the case of the orthonormal null-space function, the average value is obtained.

The **Assembly block size** edit field determines the number of mesh elements that the solver processes together during the assembly process (default = 5000). A lower value results in lower memory usage, but it can also make the assembly slower.

Settings Related to Complex-Valued Data and Undefined Operations

TAKING THE COMPLEX CONJUGATE FOR COMPLEX-VALUED MODELS

For a complex-valued model, the Use Hermitian transpose of constraint matrix and in symmetry detection check box affects the meaning of the Lagrange multiplier term $h^T \mu$ in the Neumann boundary condition of the general or coefficient form, and in general the term $N_F \Lambda$ in the discretized model. If you select this check box, the complex conjugate is taken for the matrix N_F (that is $N_F \rightarrow N_F^*$). This check box also affects the automatic symmetry detection. By default, the complex conjugate is not taken. This check box is only active when Nonsymmetric or Automatic is selected in the Matrix symmetry list on the General tab. Otherwise, the setting of the check box is determined by the choice Symmetric/Hermitian. That is, if Symmetric is selected, the conjugate -is taken.

USING COMPLEX FUNCTIONS WITH REAL INPUT

If a function takes real inputs, you can assume the output is real by default. For instance, sqrt(-1) generates an error message. To change this behavior, select the **Use complex functions with real input** check box.

STOPPING IF ERROR DUE TO UNDEFINED OPERATIONS OCCURS

By default, the solver stops with an error message when it encounters an undefined mathematical operation in an expression that appears in the model settings, for instance, division by zero or square root of a negative number. To change this behavior, clear the **Stop if error due to undefined operation** check box. Then the solver treats the result of the operation as Inf (infinity) or NaN (not a number). This feature can be useful in a nonlinear problem where the steps in the iterative solution process lead to variable values for which an expression is undefined. The solver then reduces the step size in the Newton iteration when it encounters Inf or NaN so that it can find a solution.

Storing Solutions on File

By default, COMSOL Multiphysics stores the solution in memory. If you select the **Store solution on file** check box, the solution is primarily stored in a file. This option is useful if you do a time-dependent or parametric simulation with a large number of time steps or parameter steps. The large amount of solution data would otherwise fill up the computer's memory. The software deletes the file that it creates when the solution is no longer needed (for instance, when you exit COMSOL Multiphysics). This file is located in the default directory for temporary files provided by the operating system. You can override this location with an option when starting COMSOL Multiphysics (see the chapter "Running COMSOL" in the *COMSOL Installation and Operations Guide*).

Solution Form

The *solution form* determines the form into which COMSOL Multiphysics converts a PDE and its boundary conditions before solving it. For a description of how the software forms the PDE system, see "What Equations Does COMSOL Multiphysics Solve?" on page 476.

The solution form does not have to be the same as the *equation system form*. The equation system form is the form in which the **Equation System** dialog boxes display the equations. You select the equation system form in the **Model Settings** dialog box. If the

solution form is different from the equation system form, COMSOL Multiphysics transforms the equations to the solution form before solving.

When using a PDE mode, be sure not to confuse the PDE form with the solution form; you can, for instance, solve a PDE, Coefficient Form model using the general solution form.

When selecting the solution form, you have (at most) four options: *automatic* (the default), *coefficient*, *general*, and *weak*. With the automatic option, COMSOL Multiphysics selects the solution form according to the following rules:

- If you use the adaptive solver with **Residual method** set to **Coefficient** and the equation system form is *not* the weak form, COMSOL Multiphysics selects the general solution form.
- In all other cases, COMSOL Multiphysics selects the weak solution form

The reason the software selects the general solution form when using the adaptive solver with **Residual method** set to **Coefficient** is that it does not work with the weak solution form (see "The Adaptive Solver Algorithm" on page 505).

The only situation when you need to manually select the solution form is when you want to use *equation variables* (for instance, cux and ncu), because these are not available for the weak solution form (see "Special Variables" on page 175 of the *COMSOL Multiphysics User's Guide*). Consider the following aspects when selecting the solution form:

- The weak solution form is usually the best choice because you normally get a correct Jacobian (see "The Importance of a Correct Jacobian Matrix" on page 363 of the *COMSOL Multiphysics User's Guide*) and the assembly is somewhat faster than for the coefficient and general forms.
- The general solution form generates an incorrect Jacobian if the model has derivatives in the boundary conditions or some terms in the PDE depend on a second-order spatial derivative. It also generates an incorrect Jacobian if some coefficient or term in the PDE or boundary conditions depends on a coupling variable.
- The general solution form generates an incorrect Jacobian if the model contains time derivatives in other places than multiplying the d_a or e_a coefficients, or in the **weak** or **dweak** edit fields.
- The coefficient solution form is more restricted than the general form. In addition to the disadvantages of the general form, this solution form results in an

incorrect Jacobian if some of the coefficients depend on the solution. Therefore use the coefficient form only for linear single-physics or uncoupled models.

Note that not all solution forms might be available, depending on the formulation of the equations in the application modes in use. For instance, it is not possible to solve the nonlinear Navier-Stokes equations in coefficient form.

Manual Control of Reassembly

It is important for the efficiency of the time-stepping algorithm to assemble the timeindependent matrices only once. The solver automatically detects the coefficients in your equations that are time dependent and reassembles only those quantities. The nonlinear and parametric solvers also follow this logic (with the parameter playing the role of time).

If the Jacobian is incorrect (see "The Importance of a Correct Jacobian Matrix" on page 363 of the *COMSOL Multiphysics User's Guide*), the automatic detection can fail, which means that you might get an incorrect solution. In this case you must either manually control the reassembly (see below) or, better, use the weak solution form, which you specify on the **General** tab. If you use periodic boundary conditions, identity conditions, or coupling variables, the automatic detection is too sensitive, which means that the solution you get is correct but the reassembly process might take an unnecessarily long time. For such models, you can speed up time-stepping, by manually specifying which matrices are constant. To do so, first select the **Manual control of reassembly** check box and then select the check boxes in this area according to the following guidelines:

- Select the **Load constant** check box if the PDE and Neumann boundary conditions are linear with time-independent coefficients and right-hand sides. For the discretized model, this means that the residual vector L depends linearly on U $(L = L_0 KU)$ and that L_0 , K, and the mass matrix D are constant. It is assumed that the Jacobian matrix K is correct.
- Select the **Jacobian constant** check box if the Jacobian matrix *K* is time-independent. You can also choose this option if you want to use the same Jacobian throughout the time-dependent or nonlinear solver. This choice cuts down linear-system factorization/preconditioning time but causes more iterations because the Newton iteration is degraded into a fixed-point iteration.
- Select the **Damping (mass) constant** check box if the coefficients of the first-order time-derivative terms are time independent (often the case). In the discretized model, this means that the damping (sometimes called mass) matrix *D* is constant.

- Select the **Mass constant** check box if the coefficients of the second-order time-derivative terms are time independent (often the case). In the discretized model, this means that the mass matrix *E* is constant.
- Select the **Constraint constant** check box if the Dirichlet boundary conditions (constraints) are linear and time-independent. For the discretized model, this means that the constraint residual M depends linearly on $U (M = M_0 NU)$ and that M_0 and N are constant. It is also assumed that the constraint Jacobian N is correct.
- Select the **Constraint Jacobian constant** check box if the Dirichlet boundary conditions are linear with time-independent coefficients (not right-hand side). For the discretized model this means that *N* is constant.

Scaling of Variables and Equations

If the dependent variables in your model have widely different magnitudes, the solver might have problems with the resulting ill-conditioned matrix. For instance, in a structural-mechanics problem the displacements can be of the order of 0.0001 m while the stresses are 1,000,000 Pa (1 MPa). To remedy this situation, COMSOL Multiphysics internally rescales the variables so that a well-scaled system results.

The default is *automatic scaling*, which works well for most models. It is based on the magnitudes of the elements in the Jacobian and mass matrices. If you know the order of magnitudes of the variables in advance, you can opt for *manual scaling*. For instance, suppose that a problem involves two degrees of freedom, u and sigma, and that the values of u are on the order of 10^{-4} and the values of sigma are approximately 10^{6} . To take this knowledge into account, in the Scaling of variables area, select Manual from the Type of scaling list and type u 1e-4 sigma 1e6 in the Manual scaling edit field. When you start the solvers, they will internally use the rescaled degrees of freedom U = u/1e-4 and Sigma = sigma/1e6, which both are of the order 1. If you provide an initial value that gives a good estimate of the scales of your variables, another choice is to use *initial-value based scaling* by selecting Initial value based in the Type of scaling list.

Note: The automatic scaling in COMSOL Multiphysics does not work when using the nonlinear stationary solver and one solution component is identically zero in the solution (the solver does not converge). In this case use **Manual** or **None**.

Even if variables are well scaled, equations can have very different scales. To overcome this problem you can equilibrate the equations by selecting **On** in the **Row equilibration** list (the default). To turn off equation scaling, choose **Off** in the **Row equilibration** list. To preserve the possible symmetry of the matrix, COMSOL Multiphysics does not use row equilibration in the following cases:

- · Automatic matrix symmetry detection is used and the system matrices are symmetric
- Symmetric or Hermitian is selected in the Matrix symmetry list
- The Conjugate gradients or Geometric multigrid solver is used
- The eigenvalue solver is used.

THE RESCALED LINEAR SYSTEM

The rescaling of the discretized linear system (see "The Discretized Linearized Model" on page 367 of the *COMSOL Multiphysics User's Guide*) occurs before constraint handling. Assume that the degrees of freedom U_i are expressed terms of rescaled degrees of freedom \tilde{U}_i according to the formula

$$U_i = s_i U_i$$

where s_i are positive scale factors. Using a diagonal matrix S, the relation between U and U is

$$U = SU$$
,

and you can write the rescaled linear system as

$$\begin{bmatrix} \tilde{K} & \tilde{N}F \\ \tilde{N} & 0 \end{bmatrix} \begin{bmatrix} \tilde{U} \\ \tilde{\Lambda} \end{bmatrix} = \begin{bmatrix} \tilde{L} \\ \tilde{M} \end{bmatrix}$$

where

$$\Lambda = R\Lambda \qquad NF = SN_FR \qquad K = SKS \qquad N = RNS$$

and

$$L = SL, M = RM$$
.

Here, R_{i} is a diagonal matrix of positive scale factors chosen such that the rows in the matrix N are of magnitude 1.

Consider the linear (scaled) system

$$\begin{bmatrix} K & N_F \\ N & 0 \end{bmatrix} \begin{bmatrix} U \\ \Lambda \end{bmatrix} = \begin{bmatrix} L \\ M \end{bmatrix}$$

The Lagrange multiplier vector Λ is typically underdetermined, and COMSOL Multiphysics does not solve for it. Similarly, the constraint NU = M often contains the same equation several times. To handle this problem, COMSOL Multiphysics turns to a *constraint-handling method* using elimination, Lagrange multipliers, or stiff springs. Select the desired constraint-handling method on the **Advanced** page of the **Solver Parameters** dialog box.

ELIMINATION CONSTRAINT HANDLING

This is the default constraint-handling method. The solver computes a solution U_d to the constraint NU = M as well as a matrix Null, whose columns form a basis for the null space of N. For non-ideal constraints $(N_F \neq N^T)$ then a matrix Nullf is also computed, whose column forms a basis for the null space of N_F^T . Then it obtains the solution as $U = \text{Null } U_n + U_d$, where U_n is the solution of $K_e U_n = L_e$, where

$$\begin{cases} K_e = \text{Nullf}^T K \text{Null} \\ L_e = \text{Nullf}^T (L - KU_d) \end{cases}$$

For the ideal constraint case Nullf = Null, the corresponding eliminated D and E matrices are

$$D_e = \text{Nullf}^T D \text{Null}, \quad E_e = \text{Nullf}^T E \text{Null}$$

LAGRANGE MULTIPLIERS CONSTRAINT HANDLING

The linear solver computes a matrix Range, whose columns form a basis for the range of N, and a matrix Rangef, whose columns form a basis for the range of N_F^T . Then it constrains Λ to be of the form $\Lambda = s$ Rangef Λ where s is a scaling factor. This transforms the system to

- -

$$K_l \begin{vmatrix} U \\ \hat{\Lambda} \end{vmatrix} = L_l$$

where

$$K_{l} = \begin{bmatrix} K & sN_{F} \text{Rangef} \\ s\text{Range}^{T}N & 0 \end{bmatrix} \qquad \qquad L_{l} = \begin{bmatrix} L \\ s\text{Range}^{T}M \end{bmatrix}.$$

COMSOL Multiphysics solves this system for both U and Λ , but it returns only U. The corresponding D matrix is

$$D_l = \begin{bmatrix} D & 0 \\ 0 & 0 \end{bmatrix}$$

For the ideal constraint case Rangef = Range.

STIFF SPRINGS CONSTRAINT HANDLING

This method approximates the constraint NU = M by

$$NU = M + \frac{1}{\kappa}\Lambda$$

where κ is a suitably large constant. Eliminating Λ gives the system $K_s U = L_s$, where

$$\left\{ \begin{array}{l} K_{s}=K+\kappa N_{F}N\\ L_{s}=L+\kappa N_{F}M \end{array} \right.$$

•

The corresponding D matrix is $D_s = D$.

Solver Algorithms

The Nonlinear Solver Algorithm

The nonlinear solver uses an affine invariant form of the damped Newton method as described in Ref. 1. You can write the discrete form of the equations as f(U) = 0, where f(U) is the residual vector and U is the solution vector. Starting with the initial guess U_0 , the software forms the linearized model using U_0 as the linearization point. It solves the discretized form of the linearized model $f'(U_0)\delta U = -f(U_0)$ for the Newton step δU using the selected linear system solver ($f'(U_0)$ is the Jacobian matrix). It then computes the new iteration $U_1 = U_0 + \lambda \delta U$, where λ ($0 \le \lambda \le 1$) is the damping factor. Next the modified Newton correction estimates the error E for the new iteration U_1 by solving $f'(U_0)E = -f(U_1)$. If the relative error corresponding to E is larger than the relative error in the previous iteration, the code reduces the damping factor λ and recomputes U_1 . This algorithm repeats the damping-factor reduction until the relative error is less than in the previous iteration or until the damping factor underflows the minimum damping factor. When it has taken a successful step U_1 , the algorithm proceeds with the next Newton iteration.

A value of $\lambda = 1$ results in Newton's method, which converges quadratically if the initial guess U_0 is sufficiently close to a solution. In order to enlarge the domain of attraction, the solver chooses the damping factors judiciously. Nevertheless, the success of a nonlinear solver depends heavily on a carefully selected initial guess. Thus you should spend considerable time providing the best value for U_0 , giving at least an order of magnitude guess for different solution components.

CONVERGENCE CRITERION

The nonlinear iterations terminate when the following convergence criterion is satisfied: Let U be the current approximation to the true solution vector, and let E be the estimated error in this vector. The software stops the iterations when the relative tolerance exceeds the relative error computed as the weighted Euclidean norm

err =
$$\left(\frac{1}{N}\sum_{i=1}^{N} (|E_i|/W_i)^2\right)^{1/2}$$

Here N is the number of degrees of freedom and $W_i = \max(|U_i|, S_i)$, where S_i is a scale factor that the solver determines on the basis of the **Type of scaling** option selected

in the Scaling of variables area on the Advanced page according to the following rules:

- For Automatic, S_i is the average of $|U_j|$ for all DOFs *j* having the same name as DOF *i* times a factor equal to 10^{-5} if the Highly nonlinear problem check box is selected or 0.1 otherwise.
- For Manual, S_i is the value for DOF *i* given in the Manual scaling edit field.
- For **Initial value based**, S_i is the factor s (see below) times the average of $|U_{0j}|$ for all DOFs j having the same name as DOF i, where U_0 is the solution vector corresponding to the initial value.
- For None, $W_i = 1$. In this case, err is an estimate for the absolute error.

The Augmented Lagrangian Solver Algorithm

Denoting the main components in step i of the algorithm U_i and the corresponding augmented-Lagrangian components V_i , the following steps describe the algorithm:

- I Initialize U_0 and V_0 , and set i = 0.
- **2** Solve the nonlinear problem for $U = U_{i+1}$ with U_i as initial data and with $V = V_i$ held fixed.
- **3** Solve the linear problem for $V = V_{i+1}$ with $U = U_{i+1}$ held fix.
- **4** If $|V_{i+1} V_i| / |V_{i+1}| \le \delta_V$ and $|U_{i+1} U_i| / |U_{i+1}| \le \delta_U$, or $i > i_{\text{max}}$, then terminate, else set i = i+1 and go to Step 2.

This procedure is called Uzawa iterations (or segregated iterations). The value in the **Tolerance** edit field controls the tolerance δ_V in the convergence criterion, and you control the other tolerance δ_U using the **Relative tolerance** edit field in the **Nonlinear settings** area. The value in the **Maximum number of iterations** edit field controls the i_{max} parameter in Step 4. You can choose the linear system solver used for Step 3 in the **Solver** list. The Structural Mechanics Module uses this process to solve contact problems with the augmented-Lagrangian technique. See the *Structural Mechanics Module User's Guide* for more information.

The Time-Dependent Solver Algorithm

The finite element discretization of the time-dependent PDE problem is

$$0 = L(U, U, U, t) - N_F(U, t)\Lambda$$
$$0 = M(U, t)$$

which is often referred to as the *method of lines*. Before solving this system, the algorithm eliminates the Lagrange multipliers Λ . If the constraints 0 = M are linear and time independent and if the constraint force Jacobian N_F is constant then the algorithm also eliminates the constraints from the system. Otherwise it keeps the constraints, leading to a differential-algebraic system.

To solve the above ODE or DAE system, COMSOL Multiphysics uses a version of the DAE solver DASPK created by Linda Petzold at the University of California, Santa Barbara (see Ref. 3 and Ref. 4). The DASPK solver, in turn, is based on the older code DASSL (see Ref. 2), which uses variable-order variable-step-size backward differentiation formulas (BDF). Thus the solver is an implicit time-stepping scheme, which implies that it must solve a possibly nonlinear system of equations at each time step. It solves the nonlinear system using a Newton iteration, and it then solves the resulting systems with an arbitrary COMSOL Multiphysics linear system solver. The linearization of the above system used in the Newton iteration is

$$EV + DV + KV = L - N_F \Lambda$$

 $NV = M$

where $K = -\partial L/\partial U$ is the stiffness matrix, $D = -\partial L/\partial U$ is the damping matrix, and $E = -\partial L/\partial U$ is the mass matrix. When E = 0, D is often called the mass matrix.

For problems with second order time-derivatives ($E \neq 0$), extra variables are internally introduced so that a first order time-derivative system can be formed. The vector of extra variables, here U_v , comes with the extra equation

$$\dot{U} = U_v$$

where U denotes the vector of original variables. The original ODE or DAE system is by this procedure expanded to double size but the linearized system is reduced to the original size with the matrix $E + \sigma D + \sigma^2 K$, where σ is a scalar inversely proportional to the time-step. By the added equation the original variable U is therefore always a differential variable (index-0). The variable U_v is excluded from the error test, unless **Consistent initialization of DAE systems** is set to **On**, in which case the differential U_v -variables are included in the error test and the **Error estimation strategy** applies to the algebraic U_v -variables.

The Eigenvalue Solver Algorithm

Finite element discretization leads to the generalized eigenvalue system

$$(\lambda - \lambda_0)^2 E U - (\lambda - \lambda_0) D U + K U + N_F \Lambda = 0$$
$$N U = 0$$

where the solver evaluates E, D, K, N and N_F for the solution vector U_0 , λ denotes the eigenvalue, and λ_0 is the linearization point. If E = 0, it is a linear eigenvalue problem; if E is nonzero, it is a quadratic eigenvalue problem. To solve the quadratic eigenvalue problem, COMSOL Multiphysics reformulates it as a linear eigenvalue problem. After constraint handling, it is possible to write the system in the form $Ax = \lambda Bx$.

More general eigenvalue problems sometimes arise when boundary conditions or material properties are nonlinear functions of the eigenvalue. These cases can be handled as a series of quadratic eigenvalue problems. COMSOL Multiphysics treats general dependences on the eigenvalue by assembling a quadratic approximation around the eigenvalue linearization point λ_0 . Normally, iteratively updating the linearization point leads to rapid convergence.

Finding the eigenvalues closest to the shift σ is equivalent to computing the largest eigenvalues of the matrix $C = (A - \sigma B)^{-1}B$. To do this, the solver uses the ARPACK FORTRAN routines for large-scale eigenvalue problems (Ref. 5). This code is based on a variant of the Arnoldi algorithm called the *implicitly restarted Arnoldi method* (IRAM). The ARPACK routines must perform several matrix-vector multiplications Cv, which they accomplish by solving the linear system $(A - \sigma B)x = Bv$ using one of the linear system solvers.

The Parametric Solver Algorithm

The parametric solver performs a loop around the usual stationary solver in which it estimates the initial guess using the solution for the previous parameter value. If the nonlinear solver does not converge, it tries a smaller parameter step; the size of this step it determined on the basis of the convergence speed of the Newton iteration. These criteria for step size selection are based on work in Ref. 6.

The Stationary Segregated Solver Algorithm

CONVERGENCE CRITERION

The segregated solver terminates if, for all the groups j, the error estimate is smaller than the corresponding tolerance,

$$\operatorname{err}_{i,k} < \operatorname{tol}_i$$
,

where the error estimate in segregated iteration k is

$$\operatorname{err}_{j,k} = \max(\operatorname{e}_{j,k}^{N}, \operatorname{e}_{j,k}^{S}).$$

The number tol_j is taken from the **Relative tolerance** edit field for the corresponding group settings for the **Stationary segregated** solver on the **General** page of the **Solver Parameters** dialog box. Furthermore,

$$e_{j,k}^{N} = \max_{l} (1 - \alpha_{l}) \left[\frac{1}{N_{j}} \sum_{i=1}^{N_{j}} \left(\frac{\left| (\Delta U^{l,j,k})_{i} \right|}{W^{j}_{i}} \right)^{2} \right]^{1/2}$$

is an estimate of the largest damped Newton error. Here l is taken for all iterations in all substeps solving for the group j, α_l is the damping factor, $\Delta U^{l,j,k}$ is the Newton increment vector, and N_j is the number of DOFs. The weight factor W^j_i is described below. Moreover,

$$\mathbf{e}_{j,k}^{S} = \left[\frac{1}{N_{j}} \sum_{i=1}^{N_{j}} \left(\frac{\left|(U^{j,k} - U^{j,k-1})_{i}\right|}{W^{j}_{i}}\right)^{2}\right]^{1/2}$$

is the relative increment over one complete iteration k. In this expression, $U^{j,k}$ is the segregated solution vector for the group j, and $W^{j}_{i} = \max(|U^{j}_{i}|, S_{i})$, where S_{i} is a scale factor that the solver determines from the settings in the **Scaling of variables** area on the **Advanced** page.

Now examine the possible choices in the Type of scaling list:

- For Automatic, S_i is the factor 0.1 times the average of $|U_m|$ for all DOFs *m* having the same name as DOF *i*.
- For Manual, S_i is the value for DOF i given in the Manual scaling edit field.
- For **Initial value based**, S_i is the factor 0.1 times the average of $|U_{0m}|$ for all DOFs m having the same name as DOF i, where U_0 is the solution vector corresponding to the initial value.
- For None, $W_i = 1$.

The Adaptive Solver Algorithm

The L2 norm error estimate relies on an assumption of a strong stability estimate for the PDE problem (satisfied, for example, for Poisson's equation over a domain with a

smooth boundary). From such an assumption, it is possible to show that there is a constant C, such that the L2 norm of the error, e_l , in the *l*th equation satisfies

$$|e_l| \leq C |h^{q_l} \rho_l|$$

where ρ_l is the residual in the *l*th equation and q_l is the stability estimate derivative order. The adaptive solver algorithm uses the following L2-norm error indicator:

$$\left(\int\limits_{\Omega}\sum\limits_{l}s_{l}^{-2}h^{2q_{l}}\left|
ho_{l}
ight|^{2}dA
ight)^{rac{1}{2}}$$

with the default value $q_l = 2$. This formula also introduces the scaling factors s_l for the residual with the default value $s_l = 1$. The local error indicator for a mesh element is

$${\sum_l} {s_l^{-2} h^{2q_l}} { au_l^2} { au_l^2} A$$

where *A* is the area (volume, length) of the mesh element, and τ_l is the absolute value of the *l*th equation residual (one number per mesh element). The residual methods *Weak* and *Coefficient* compute τ_l in rather different ways.

The Coefficient residual method uses an explicit so-called strong form of the PDE to compute the equation residual τ_l on each mesh element. This method evaluates the PDE residual at the center of each element takes normal flux jumps to neighboring elements into account. However, it does not take residual contributions from equations formulated with the Weak solution form into account. Neither does it add terms in the Weak edit fields, and constraint forces do not contribute to the residual. Because there is no compelling reason to use the Coefficient residual method, you should therefore avoid selecting it. It is provided for backward compatibility only.

The Weak residual method (the default) uses an auxiliary mesh case to estimate the residual τ_l . This method automatically generates the mesh case by increasing the order of the shape functions used (by one) for the problem considered, while using the same mesh. The solution is mapped to this auxiliary mesh case and the discrete residual vector L is assembled. The equation residual τ_l for a mesh element is computed by:

- Finding how the *l*th field variable (dependent variable) is coupled to the degrees of freedom's
- Taking an average per element for the corresponding components of *L*.

Due to the extra residual assembly work, the Weak residual method is somewhat slower than the Coefficient residual method. On the other hand, the Weak method is more general; for example, it supports vector elements. It also takes boundary fluxes into account. Degrees of freedom that are constrained do not contribute to the residual.

The adaptive solver performs the following iterative algorithm (Ref. 7):

- I Solve the problem on the existing mesh using the stationary or eigenvalue solver.
- 2 Evaluate the residual of the PDE on all mesh elements.
- **3** Estimate the error in the solution on all mesh elements. The computed error estimate is really just an *error indicator* because the estimate involves an unknown constant (*C* above).
- **4** Terminate execution if it has made the requested number of refinements or if it has exceeded the maximum number of elements.
- 5 Refine a subset of the elements based on the sizes of the local error indicators.
- **6** Repeat from Step 1.

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Linear System Solvers

The UMFPACK Direct Solver

UMFPACK is the default linear system solver. It solves general systems of the form Ax = b using the nonsymmetric-pattern multifrontal method and direct LU factorization of the sparse matrix A. It employs the COLAMD and AMD approximate minimum degree preordering algorithms to permute the columns so that the fill-in is minimized. The code, written in C, uses level-3 BLAS (Basic Linear Algebra Subprograms) for optimal performance. COMSOL Multiphysics uses UMFPACK version 4.2 written by Timothy A. Davis (Ref. 1).

In the Linear System Solver Settings dialog box you can set the *memory-allocation factor*, a positive number (default = 0.7). The solver makes a rough estimate of the required memory before performing the calculations. The memory-allocation factor dictates how much memory COMSOL Multiphysics should allocate. A value of 0.7 results in using 70% of the estimate. A low allocation factor saves memory, but the simulation might then run much slower.

If you select the **Check tolerances** check box, COMSOL Multiphysics estimates and checks the error after the solution phase. For information about the error estimate see the section "Convergence Criteria" on page 513. By default the error estimate is turned off for UMFPACK.

You can also set the *pivot threshold*, a number between 0 and 1 (default = 0.1). The solver permutes rows for stability. In any given column the algorithm accepts an entry as a pivot element if its absolute value is greater than or equal to the pivot threshold times the largest absolute value in the column. A low pivot threshold means less fill-in and thus saves memory. If the default setting leads to poor accuracy in the linear solution, try to increase the pivot threshold from the default to, for example, 1 (which means that the linear solver always applies partial pivoting). This action can lead to a more stable solution process and a more accurate solution of the linear systems.

When using UMFPACK as a preconditioner, you can also provide a *drop tolerance* in the range 0 to 1. A value of 0.01 means that it drops matrix entries smaller than 1% of the maximum value in each column of the LU factors. Doing so reduces the size of the factors and reduces memory requirements. However, the dropping occurs only when writing the LU factors, and it does not affect the rest of the factorization process. In

contrast, in the Incomplete LU preconditioner the element dropping affects the rest of the factorization process, which leads to a more memory-efficient preconditioner.

The SPOOLES Direct Solver

The SPOOLES solver works on general systems of the form Ax = b using the multifrontal method and direct LU factorization of the sparse matrix A. When the matrix A is symmetric or Hermitian, the solver uses an LDLT version of the algorithm, which saves half the memory. SPOOLES uses several preordering algorithms to permute the columns and thereby minimize the fill-in. The code is written in C. COMSOL Multiphysics uses SPOOLES version 2.2 developed by Cleve Ashcraft and collaborators (Ref. 2).

In the **Linear System Solver Settings** dialog box you choose among the following preordering algorithms:

- Minimum degree (the default algorithm)
- Nested dissection
- Multisection
- The best of nested dissection and multisection

If you select the **Check tolerances** check box, COMSOL Multiphysics estimates and checks the error after the solution phase. For information about the error estimate see the section "Convergence Criteria" on page 513. By default the error estimate is turned off for SPOOLES.

You can also specify a pivot threshold in the range of 0 to 1 (default = 0.1). When using SPOOLES as a preconditioner, you can provide a drop tolerance in the range of 0 to 1 (see "The UMFPACK Direct Solver" on page 508).

The PARDISO Direct Solver

The parallel sparse direct linear solver PARDISO works on general systems of the form Ax = b. In order to improve sequential and parallel sparse numerical factorization performance, the solver algorithms are based on a Level-3 BLAS update, and they exploit pipelining parallelism with a combination of left-looking and right-looking supernode techniques. The code is written in C and Fortran. COMSOL Multiphysics uses the PARDISO version developed by Olaf Schenk and collaborators (Ref. 3), which is included with Intel MKL (Intel Math Kernel Library).

In the **Linear System Solver Settings** dialog box you choose among the following preordering algorithms:

- · Minimum degree
- Nested dissection (the default algorithm)

You can also specify if the solver should use a maximum weight matching strategy by choosing row preordering on (default) or off.

In order to avoid pivoting, PARDISO uses a pivot perturbation strategy that tests the magnitude of the potential pivot against a constant threshold of

 $\varepsilon = \alpha |PP_{\text{MPS}}D_rAD_cP|_{\infty}$, where P and P_{MPS} are permutation matrices, D_r and D_c are diagonal scaling matrices, and $|.|_{\infty}$ is the infinity norm. If the solver encounters a tiny pivot during elimination, it sets it to $\operatorname{sign}(l_{ii})\varepsilon |PP_{\text{MPS}}D_rAD_cP|_{\infty}$. You can specify the pivot threshold ε . The perturbation strategy is not as robust as ordinary pivoting. In order to improve the solution PARDISO uses iterative refinements.

COMSOL Multiphysics can optionally estimate and check the error after the solution phase. You control this option through the **Check tolerances** list. For the **Automatic** selection, error checking is at least done for problems where pivot perturbation or iterative improvement has been used. For information about the error estimate, see the section "Convergence Criteria" on page 513. By default the error checking is enabled **(On)**. You can disable the check by instead selecting **Off**.

For information about running COMSOL Multiphysics using parallelism, see "Parallel COMSOL" on page 63 in the COMSOL Installation and Operations Guide.

Note: PARDISO is available on Linux and Windows only. On Sun and Mac, COMSOL Multiphysics switches to UMFPACK instead.

The TAUCS Cholesky Direct Solver

The TAUCS Cholesky direct solver handles systems of the form Ax = b, where A is a positive definite symmetric sparse matrix (see "Which Models Are Positive Definite?" on page 399 of the *COMSOL Multiphysics User's Guide*) using a multifrontal supernodal Cholesky factorization. It employs the multiple minimum degree reordering algorithm to permute the rows and columns and thus minimize the fill-in. Written in C, it uses level-3 BLAS for maximum performance. COMSOL Multiphysics uses TAUCS version 2.2 written by Sivan Toledo and collaborators (Ref. 4).

Due to the algorithm's recursive nature, it can run out of stack space for large models. If this happens, you can increase the value of the STACKSIZE parameter in the appropriate COMSOL startup script. For PC/Windows set the STACKSIZE parameter in the \bin\comsol.opts file, located in the COMSOL installation directory; for UNIX/Linux/Mac OS X set the STACKSIZE parameter in the /bin/comsol command, located in the COMSOL installation directory. The default value is 2m (2 MB). For example, if you edit the script and double the value of STACKSIZE to 4m (4 MB), the maximum recursion depth for the algorithm also doubles. Continue doubling the value of STACKSIZE until the TAUCS algorithm is successful.

Note: To use the TAUCS Cholesky and LDLT solvers, you must select **Symmetric** or **Hermitian** in the **Matrix symmetry** list in the **Solver Parameters** dialog box.

The TAUCS LDLT Direct Solver

This linear system solver handles real symmetric or Hermitian matrices. It has no parameters to set. You can use the direct LDLT (TAUCS) solver as an alternative to the SPOOLES symmetric solver.

The GMRES Iterative Solver

This linear system solver uses the restarted GMRES (generalized minimum residual) method (see Ref. 5 and Ref. 6). This is an iterative method for general linear systems of the form Ax = b. For fast convergence it is important to use an appropriate *preconditioner* (see "Selecting a Preconditioner" on page 395 of the *COMSOL Multiphysics User's Guide*).

The value in the **Number of iterations before restart** edit field in the **Linear System Solver Settings** dialog box specifies the number of iterations the solver performs until it restarts (the default is 50). There is no guarantee that a restarted GMRES will converge for a small restart value. A larger restart value increases the robustness of the interactive procedure, but it also increases memory use and computational time. For large problems, the computational cost is often very large to produce a preconditioner of such a high quality that the termination criteria are fulfilled for a small number of iterations and for a small restart value. For those problems it is often advantageous to set up a preconditioner with a somewhat lesser quality and instead increase the restart value or iterate more steps. Doing so typically increases the condition number for the preconditioned system, so an increase in the error-estimate factor might be needed as well.

Two slightly different versions of GMRES are available in COMSOL Multiphysics. The difference between these two versions is whether left or right preconditioning is used (see "The Preconditioned Linear System" on page 400 of the *COMSOL Multiphysics User's Guide*). Select the preconditioning type from the **Preconditioning** list. The default choice is left preconditioning. Normally, the two versions of GMRES have similar convergence behavior (see Ref. 7). If the preconditioner is ill-conditioned there could, however, be differences in the behavior.

For information about the convergence criterion used by GMRES and the **Relative** tolerance and Factor in error estimate edit fields, see "Convergence Criteria" on page 513.

If the solver does not converge, it terminates with an error message when it reaches the value in the Maximum number of iterations edit field (default = 10,000).

The FGMRES Iterative Solver

This solver uses the restarted FGMRES (flexible generalized minimum residual) method (see Ref. 8). The solver is a variant of the GMRES solver that can handle a wider class of preconditioners in a robust way. You can, for example, use any iterative solver as preconditioner for FGMRES. The downside with the method is that it uses twice as much memory as GMRES for the same value in the **Number of iterations before restart** edit field. FGMRES uses right preconditioning and therefore has the same convergence criterion as right-preconditioned GMRES. If FGMRES is used together with a constant preconditioner such as, for example, the Incomplete LU preconditioner, then the FGMRES solver is identical to the right preconditioned GMRES solver.

For information about the convergence criterion used by FGMRES and the **Relative** tolerance and Factor in error estimate edit fields, see "Convergence Criteria" on page 513.

The Conjugate Gradients Iterative Solver

This solver uses the conjugate gradients iterative method (see Ref. 9, Ref. 10, and Ref. 11). It is an iterative method for linear systems of the form Ax = b where the matrix A is positive definite and (Hermitian) symmetric (see "Which Models Are Positive Definite?" on page 399 of the *COMSOL Multiphysics User's Guide*). Sometimes the

solver also works when the matrix is not positive definite, especially if it is close to positive definite. This solver uses less memory and is often faster than the GMRES solver, but it applies to a restricted set of models.

For fast convergence it is important to use an appropriate *preconditioner* (see "Selecting a Preconditioner" on page 395 of the *COMSOL Multiphysics User's Guide*), which should be positive definite and (Hermitian) symmetric.

Select the preconditioning type from the **Preconditioning** list. The default choice is left preconditioning. For the Conjugate gradient method this choice only affects the convergence criterion and not the algorithm itself. For information about the convergence criterion and the **Relative tolerance** and **Factor in error estimate** edit fields, see "Convergence Criteria" on page 513

Convergence Criteria

When you use an iterative solver COMSOL Multiphysics estimates the error of the solution while solving. Once the error estimate is small enough, as determined by the convergence criterion

$$\left| \mathbf{M}^{-1}(b - Ax) \right| < \operatorname{tol} \cdot \left| \mathbf{M}^{-1}b \right|$$
(5-1)

the software terminates the computations and returns a solution. When you use a direct solver COMSOL Multiphysics can optionally make a check (**Error check**), to determine if the above convergence criterion is fulfilled after the solution step. If the error criterion is not met, the solution process is stopped an error message is given.

The definitions of M for the various solvers are:

- For UMFPACK, PARDISO and SPOOLES, M = LU, where L and U are the LU factors computed by the solver.
- For left-preconditioned GMRES and left-preconditioned Conjugate Gradients, *M* is the preconditioner matrix.
- For the remaining iterative solvers, *M* is the identity matrix.

The convergence criterion 5-1, states that the iterations terminate when the relative (preconditioned) residual times the factor ρ is less than a tolerance tol. You can set the factor ρ in the **Factor in error estimate** edit field (default = 400). For solvers where M is equal to the identity matrix, the iterations can sometimes terminate too early with an incorrect solution if the system matrix A is ill-conditioned. For solvers where M is not equal to the identity matrix, the iterations can sometimes terminate too early if M

is a poor preconditioner. If the iterations terminate too early due to an ill-conditioned system matrix or a poor preconditioner, increase the factor ρ to a number of the order of the condition number for the matrix $M^{-1}A$. Note that if ρ is greater than the condition number for the matrix $M^{-1}A$, the convergence criterion implies that the relative error is less than tol: $|x - A^{-1}b| < \operatorname{tol} \cdot |A^{-1}b|$.

LINEAR SYSTEM TOLERANCE

For the **Stationary** linear solver and the **Stationary segregated** solver, the tolerance tol in the convergence criterion 5-1 is the value specified in the **Relative tolerance** edit field in the **Linear System Solver Settings** dialog box.

For the **Stationary** nonlinear solver, tol is adaptive and based on the maximum of the number entered in the **Relative tolerance** edit field in the **Linear System Solver Settings** dialog box and the number entered in the **Relative tolerance** edit field on the **Stationary** page of the **Solver Parameters** dialog box. During the nonlinear iterations tol can, however, be larger or smaller than this number.

For the **Parametric** solvers, tol is used in the same way as for the corresponding **Stationary** solver.

When using the **Time dependent** solver, tol is the maximum of the number in the **Relative tolerance** edit field in the **Linear System Solver Settings** dialog box and the number in the **Relative tolerance** edit field on the **General** page of the **Solver Parameters** dialog box.

When using the **Eigenvalue** solver together with an iterative method, tol is the number in the **Relative tolerance** edit field in the **Linear System Solver Settings** dialog box times a safety factor of 10^{-4} . When using a direct method, tol is the number in the **Relative tolerance** edit field in the **Linear System Solver Settings** dialog box without any safety factor.

For the main components of the **Augmented Lagrangian** solver, tol is used in the same way as for the **Stationary** solver. For the **Augmentation components** the error check for the direct solvers is disabled.

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Preconditioners for the Iterative Solvers

The Incomplete LU Preconditioner

The Incomplete LU preconditioner performs an incomplete LU factorization of the system matrix A. That is, it drops small elements during the column-oriented Gaussian elimination (see Ref. 1 and Ref. 2). Thus it saves memory, and the resulting factors L and U are approximate. The resulting preconditioner is an approximation to A. The preconditioner supports threshold drop, fill-ratio drop, and threshold pivoting. It can optionally respect the nonzero pattern in the original matrix. The preconditioner accepts matrices in symmetric and Hermitian format but expands these to full storage before factorization.

In the Linear System Solver Settings dialog you can select a drop rule (see the following section) from the Drop using list. Depending on the selected drop rule, you can specify a Drop tolerance or a Fill ratio. You can also control the drop tolerance on the General tab of the Solver Parameters dialog box either numerically (by supplying a number between 0 and 1) or using the Memory efficiency/Precond. quality slider. A smaller drop tolerance means that the preconditioner drops fewer elements and so the preconditioner becomes more accurate. This leads to fewer iterations in the iterative solver, but on the other hand memory requirements and preconditioning time increases. A larger drop tolerance means that the preconditioning time decrease. In this case, however, the preconditioner becomes less accurate, which leads to more iterations in the iterative solver, or, if the drop tolerance is too high, to no convergence at all. Often it is most efficient to use as high a drop tolerance as possible, that is, choose it so that the iterative solver barely converges.

You can also set the **Pivot threshold**, a number between 0 and 1 (default = 1). The solver permutes rows for stability. In any given column, if the absolute value of the diagonal element is less than the pivot threshold times the largest absolute value in the column, it permutes rows such that the largest element is on the diagonal. Thus the default value 1 means that it uses partial pivoting.

Once the approximate factors L and U have been computed, you can use the incomplete LU factorization as an iterative preconditioner/smoother (see "The SSOR, SOR, SORU, and Diagonal Scaling (Jacobi) Algorithms" on page 527). Here,

 $M = (LU)/\omega$, where ω is a relaxation factor, and L and U are the approximate factors. When using incomplete LU factorization as a preconditioner or smoother, it performs a fixed number of sweeps as dictated by the value in the **Number of iterations** edit field in the **Linear System Solver Settings** dialog box (default = 1).

Specify ω in the **Relaxation factor (omega)** edit field (default = 1).

SELECTING A DROP RULE

The Incomplete LU preconditioner uses either the *threshold drop rule* (the default) or the *fill-ratio drop rule*. The preconditioner drops (neglects) an element during the elimination phase if its absolute value is smaller than the Euclidean norm of the entire column times a *drop tolerance*. In contrast, the fill-ratio drop rule limits the number of nonzeros in the incomplete factors L and U, and it keeps the largest absolute values. The number of values it keeps depends on the number of nonzeros in the corresponding column of the original matrix times a fill-ratio factor. There are two exceptions to these drop rules:

- The preconditioner never drops diagonal elements.
- The preconditioner optionally drops nonzeros in positions where the original matrix is nonzero. The default is to keep these nonzeros. To make the preconditioner drop them, clear the **Respect pattern** check box in the settings for the Incomplete LU preconditioner.

The primary problem with setting up a preconditioner is the tradeoff between resources (computer time and memory) and the preconditioner's quality. The computational cost of setting up a preconditioner with the Incomplete LU preconditioner is at least proportional to the number of nonzeros in the produced factors L and U. An advantage of using the fill-ratio drop rule is that you can estimate and limit the cost beforehand; the main disadvantage is that the quality of the preconditioner is typically not as good as using the threshold drop rule with a drop tolerance resulting in the same number of nonzeros. However, with the threshold drop rule there is no good way of estimating resource requirements beforehand. Furthermore, there is no general formula for these drop rules that gives a drop tolerance or fill ratio that guarantees fast convergence for a certain iterative method. Therefore, it is often necessary to rely on experiments and experience for this difficult and, from a performance point of view, crucial choice.

The TAUCS Incomplete Cholesky Preconditioner

This TAUCS incomplete Cholesky preconditioner is applicable to models where the system matrix is (Hermitian) symmetric positive definite (see "Which Models Are Positive Definite?" on page 399 of the *COMSOL Multiphysics User's Guide*). It performs an incomplete Cholesky factorization LL^T of the system matrix A. The resulting preconditioner $M = LL^T$ is an approximation to A. The code, written in C, uses a column-based left-looking approach with row lists. COMSOL Multiphysics uses TAUCS version 2.2 written by Sivan Toledo and collaborators (see http://www.tau.ac.il/~stoledo/taucs).

In the **Linear System Solver Settings** dialog box you can specify a value for the drop tolerance and select an option for a modified Cholesky factor. You can also control the drop tolerance on the **General** tab of **Solver Parameters** dialog box either numerically (giving a number between 0 and 1) or using the **Memory efficiency/Precond. quality** slider. The preconditioner drops elements from the *L* matrix if they are smaller than the drop tolerance times the norm of the corresponding column of *L*, provided that they are not on the diagonal and not in the nonzero pattern of *A* (see "The Incomplete LU Preconditioner" on page 516 for details on the implications of changing the drop tolerance). If you select the **Modified Cholesky** check box (which is not the default state) the preconditioner modifies the factor *L* so that the row sums of LL^T are equal to the row sums of the input matrix.

Note: To use the incomplete Cholesky preconditioner you must select **Symmetric** or **Hermitian** in the **Matrix symmetry** list in the **Solver Parameters** dialog box.

The Geometric Multigrid Solver/Preconditioner

The Geometric multigrid solver or preconditioner is a fast and memory-efficient iterative method for elliptic and parabolic models (see "Elliptic and Parabolic Models" on page 400 of the *COMSOL Multiphysics User's Guide*). It performs one or several cycles of the geometric multigrid method. The classical multigrid algorithm uses one or several auxiliary meshes that are coarser than the original (fine) mesh. The idea is to perform just a fraction of the computations on the fine mesh. Instead, it performs computations on the coarser meshes when possible, which leads to fewer operations. The size of the extra memory used for the coarser meshes and associated matrices is

comparable to the size of the original data. This leads to an iterative algorithm that is both fast and memory efficient. See Ref. 3 for more information.

COMSOL Multiphysics uses a hierarchy of *multigrid levels* where each corresponds to a mesh and a choice of shape functions. Thus, in addition to coarsening the mesh it is possible to construct a new "coarser" level by lowering the order of the shape functions. The number of degrees of freedom decreases when you go to a coarser multigrid level. The meshes for the different levels can be constructed either "manually" or automatically. The automatic version applies a coarsening algorithm to the fine mesh, which leads to meshes that are not aligned to each other. There is also an option to generate the finer meshes from the coarsest mesh by successive mesh refinements, which leads to aligned (nested) meshes. The manual option can be useful when you have a quadrilateral, hexahedral, or prism mesh, or when you for some other reason wish to control details in the meshes.

To describe the multigrid algorithm, assume that you have N + 1 multigrid levels numbered from 0 to N, where 0 is the finest level (the level for which you seek the solution). To solve the linear system $A_0x = b$ (corresponding to level 0), the algorithm must reform the system matrices $A_1, ..., A_N$ for the coarse multigrid levels. It must also compute the *prolongation matrices* P_i that map a solution x vector on level i to the corresponding solution vector P_ix on the next finer level i-1.

The prolongation matrices are constructed using plain interpolation from one multigrid level to the other. The system matrices for the coarse levels can be constructed in two ways:

- By assembling A_i on the mesh of level i (the default method).
- By projection from the finer level: $A_i = P_i^T A_{i-1} P_i$. This is also called the Galerkin method. It typically leads to more nonzero elements in the system matrix A_i , but the convergence can be faster than in the default method.

The following algorithm describes one multigrid cycle:

- I The input to the algorithm is some initial guess x_0 to the solution of the system $A_0 x = b$.
- 2 Starting with x_0 , apply a few iterations of a *presmoother* to the linear system $A_0x = b$, yielding a more accurate iterate x_{0s} . Typically the presmoother is some simple iterative algorithm such as SOR, but you also chose an arbitrary iterative solver.
- 3 Compute the residual $r_0 = b A_0 x_{0s}$. The presmoother "smooths" the residual so the oscillations in *r* have such a long wavelength that they are well resolved on the

next coarser level (1). Therefore, project the residual onto level 1 by applying the transpose of the prolongation: $r_1 = P_1^T r_0$.

- 4 If N = 1 use the *coarse solver* to solve the system $A_1x_1 = r_1$. The coarse solver is typically a direct solver such as UMFPACK. The number of degrees of freedom on level 1 is less than for level 0, which means that solving $A_1x_1 = r_1$ is less expensive. If instead N > 1, solve the system $A_1x_1 = r_1$ (approximately) by recursively applying one cycle of the multigrid algorithm for levels 1, 2, ..., N. In both cases the obtained solution x_1 is called the *coarse grid correction*.
- 5 Map the coarse grid correction to level 0 using the prolongation matrix: $x_{0c} = x_{0s} + P_1 x_1$.
- **6** Starting with x_{0c} , apply a few iterations of a *postsmoother* to the linear system $A_0x = b$, yielding a more accurate iterate x_{0mg} . The default postsmoother is SORU (the version of SOR using the upper triangle of the matrix). The iterate x_{0mg} is the output of the multigrid cycle.

The cycle just described is called the V-cycle. The recursive call in Step 4 (when N > 1) is a also a V-cycle. For the W-cycle and the F-cycle, the Steps 1–6 above are the same but with the twist that the recursive call in Step 4 is substituted with two multigrid calls for the coarser levels. For the W-cycle these two calls are recursive calls, they are W-cycle calls. For the F-cycle the first call is a W-cycle and the second a V-cycle.

For only two multigrid levels (N = 1) these cycles are the same because the algorithm uses the coarse solver in Step 4. Also note that the amount of work on the finest level is the same for the different cycles. Normally the V-cycle is sufficient, but the W-cycle and the F-cycle can be useful for more difficult problems.

When using multigrid as a preconditioner, the action of this preconditioner is obtained by applying a fixed number of multigrid cycles. When using multigrid as a solver, the multigrid cycle repeats until it reaches convergence.

When using multigrid as a preconditioner for the conjugate gradients method for a symmetric matrix A, the preconditioning matrix M should also be symmetric. This requirement is fulfilled if the matrices M (see "The SSOR, SOR, SORU, and Diagonal Scaling (Jacobi) Algorithms" on page 527) associated with the presmoother and the postsmoother are transposes of each other. For instance, this is the case if the presmoother is SOR and the postsmoother is SORU, and if the same number of smoothing steps is used. This combination with two smoothing steps is the default.

NOTES ON THE EFFICIENCY OF SMOOTHERS

COMSOL Multiphysics performs smoothing on all but the coarsest multigrid level. A smoother should be computationally cheap and be effective at reducing the part of the error that has a high spatial frequency on the mesh to which it is applied. Therefore, the applying a smoother on several meshes with a hierarchy of mesh sizes results in a more efficient solver than if the smoother were applied only on the finest mesh.

The efficiency of the multigrid method with simple iterations as a smoother (such as the Jacobi and SOR iteration) hinges on the ellipticity of the underlying mathematical problem. For Helmholtz problems originating from an equation

$$-\nabla \cdot \left(\frac{1}{a}\nabla u\right) - \omega^2 u = f$$

or

$$\nabla \times \left(\frac{1}{a} \nabla \times \mathbf{E}\right) - \omega^2 \mathbf{E} = \mathbf{F}$$

the obtained linear problem is indefinite for large frequencies ω . For these problems, a simple iteration amplifies smooth eigenmodes if the mesh is too coarse and makes these methods unsuitable as smoothers. To determine when to use a simple iteration, apply the *Nyquist criterion*

$$h_{\max} < \frac{\lambda}{2} = \frac{\pi}{\omega \sqrt{a}}$$

which says that the mesh must have at least two mesh elements per wavelength. Thus, when using the Geometric multigrid solver for these type of problems, you should ensure that this criterion is fulfilled on the coarsest mesh if simple iterations is used as a smoother. In situations where the criterion is not fulfilled on coarse meshes GMRES can be a suitable smoother (Ref. 4).

CONSTRUCTING A MULTIGRID HIERARCHY

The multigrid hierarchy can be constructed either automatically or manually. To select which method to use, go to the **General** tab of the **Solver Parameters** dialog box and click the **Settings** button. This action opens the **Linear System Solver Settings** dialog box.

If you use multigrid as a preconditioner, select **Preconditioner** in the list to the left, otherwise select **Linear system solver**.

Preconditioner:	Geometric multigrid
Preconditioner	2
Multigrid cycle:	V-cycle
Hierarchy generation method:	Lower element order first (all)
Automatic Manual	
Number of levels:	2
Mesh coarsening factor:	2
Refinement method:	Regular 👻
Use hierarchy in geometries:	Geom1
	Number of iterations: Multigrid cycle: Hierarchy generation method: Automatic <u>Manual</u> Number of levels: Mesh coarsening factor: Refinement method:

Preconditioner settings for the Geometric multigrid solver.

In the Hierarchy generation method list you can select among the following methods:

- Lower element order first (all) (default). In this method, the coarse levels are constructed automatically from the finest level by lowering the shape-function orders in steps of one. When a given shape function order cannot be decreased more, the mesh is instead coarsened by the factor given in the Mesh coarsening factor edit field (default = 2). The coarsened mesh is constructed by generating a new mesh of the geometry such that the element edge size at any location is approximately equal to the mesh coarsening factor times the element edge size in the old mesh. This procedure repeats until the number of multigrid levels (including the finest level) equals the Number of levels (default = 2).
- Lower element order first (any). In this method, the coarse levels are constructed automatically from the finest level by lowering the shape-function orders in steps of one. When none of the given shape function orders can be decreased more, the mesh is instead coarsened by the factor given in the Mesh coarsening factor edit field (default = 2). The coarsened mesh is constructed by generating a new mesh of the geometry such that the element edge size at any location is approximately equal to the mesh coarsening factor times the element edge size in the old mesh. This

procedure repeats until the number of multigrid levels (including the finest level) equals the **Number of levels** (default = 2).

- **Coarse mesh and lower order**. The coarse levels are constructed automatically from the finest level by coarsening the mesh and lowering the order of the shape functions at the same time. More precisely, the first coarse level is constructed by coarsening the mesh by a factor given in the **Mesh coarsening factor** edit field (default = 2) and lowering the shape function orders by 1. If some of the shape function orders cannot be decreased, only the mesh coarsening is done. This procedure repeats until the number of multigrid levels (including the finest level) equals the **Number of levels** (default = 2).
- **Coarse mesh**. The coarse levels are constructed automatically from the finest level by coarsening the mesh by the factor given in the **Mesh coarsening factor** edit field (default = 2). This procedure repeats until the number of multigrid levels (including the finest level) equals the **Number of levels** (default = 2).
- **Refine mesh**. In this method the current mesh is refined multiple times until the number of multigrid levels (including the finest level) equals the **Number of levels** (default = 2). Thus the given mesh becomes the coarsest multigrid level, and the solution is delivered for a refined mesh. When you use the **Refine mesh** method, the software automatically selects the **Keep generated mesh cases** check box because the refined mesh is needed in postprocessing. You can also make a selection from the **Refinement method** list. Selecting **Regular** (the default) refines the elements in a regular pattern, whereas **Longest** refines only the longest element edge. For 3D models, we recommend the **Longest** method because it produces fewer mesh elements. The **Longest** method is not available for 1D models.
- Manual. When you select this method the Manual tab shows a list of all available *mesh* cases where a mesh case is a mesh together with the choice of shape functions (and corresponding integration orders and constraint point orders). You can construct new ones by going to the Mesh menu and choosing Mesh Cases, which opens the Mesh Case Settings dialog box where you add and delete them. Each mesh case is identified by a nonnegative number. Existing mesh cases appear at the bottom of the Mesh menu, where it also indicates the *current mesh case*. The current mesh case also appears in the status bar. The mesh, shape functions, integration orders, and constraint point orders are specific to the mesh case. To change those settings for a mesh case, first make that mesh case current by selecting it in the list on the Mesh menu. Then modify any desired settings, for instance changing mesh parameters and

generating a new mesh, or changing the shape functions in the **Subdomain Settings** dialog box. For more information, see "Mesh Cases" on page 355.

When you have defined some mesh cases, go to the settings for **Geometric multigrid** in the **Linear System Solver Settings** dialog box. On the **Manual** tab you can choose for each mesh case whether it should be used in the multigrid hierarchy (select the **Use** check box) and whether the system matrix should be assembled on this level (select the **Assemble** check box). By default the hierarchy includes all mesh cases, and matrices are not assembled on the coarse levels; that is, the software uses Galerkin projection. The solver sorts the multigrid levels according to decreasing number of degrees of freedom. The solution is delivered for the finest of the selected mesh cases, and that mesh case is made current when the solver returns.

If you select the **Assemble on all levels** check box on the **Automatic** tab, the solver assembles the system matrices on the coarse levels (the default). Otherwise the coarse level matrices are formed using the Galerkin projection method.

When using the automatic hierarchy generation methods, the default behavior is that the solver deletes the coarse levels when it finishes. If you want to inspect them, select the **Keep generated mesh cases** check box, which makes the generated levels appear as new mesh cases. When this happens, the hierarchy generation method changes to the manual method. This means that the solver reuses the generated mesh cases the next time you solve, which saves some work.

The automatic hierarchy generation methods operate only on the geometries specified in the **Use hierarchy in geometries** edit field, where you provide a space-separated list of positive numbers. The mesh coarsening and shape function changes are applied only to these geometries.

Note: The automatic hierarchy generation methods construct coarsened meshes consisting of isotropic triangles or tetrahedra. If the original mesh contains quadrilaterals, hexahedrons, or prisms, or if it is anisotropic, you get better results by constructing the coarse meshes manually.

SETTINGS FOR THE MULTIGRID SOLVER/PRECONDITIONER

Apart from settings controlling the multigrid hierarchy, you can specify the following settings in the **Linear System Solver Settings** dialog box. If multigrid is used as a preconditioner, you can specify the *number of iterations* (default = 2). This gives the

number of times the multigrid cycle is performed each time the preconditioner is applied.

If you use multigrid as a linear system solver, you can instead specify a *relative tolerance*, a *factor in error estimate*, and a *maximum number of iterations*. For information about the convergence criterion used by multigrid and the **Relative tolerance** and **Factor in error estimate** edit fields, see "Convergence Criteria" on page 513. The tolerance in the convergence criterion is determined by the nonlinear stationary solver or the time-dependent solver. When using the linear stationary solver or the eigenvalue solver, you can adjust the tolerance in the **Relative tolerance** edit field (default = 10^{-6}).

If the solver does not converge, it terminates with an error message when it reaches the value in the **Maximum number of iterations** edit field (default = 10,000).

You can also select the type of multigrid cycle: V-cycle, W-cycle, or F-cycle.

SETTINGS FOR THE SMOOTHERS

To control the settings for the presmoother, select one in the list on the left side of the **Linear System Solver Settings** dialog box. In the **Presmoother** list you can select among the following smoothers: SOR (default), SORU, SSOR, SOR vector, SORU vector, SSOR vector, SOR gauge, SORU gauge, SSOR gauge, Jacobi, Vanka, Incomplete LU, GMRES, FGMRES, Conjugate gradients, and Algebraic multigrid. Change settings for the selection in the **Presmoother** area (see the sections on the specific smoothers in the following sections). For instance, it is possible to control the number of smoothing iterations here.

You control settings for the postsmoother in a similar fashion. The default postsmoother is SORU (the version of SOR using the upper triangle of the matrix).

When solving an electromagnetics model using vector elements for a PDE involving the curl-curl operator, you should select the SOR vector presmoother and the SORU vector postsmoother.

When solving fluid-dynamics problems using the incompressible Navier-Stokes equations or when using weak constraints, an algebraic so-called saddle-point problem is the result. These problems often have zeros on the diagonal of the system matrix, which makes the standard smoothers fail. Use the Vanka smoother (or Incomplete LU) in that case.

SETTINGS FOR THE COARSE SOLVER

To control the settings for the coarse solver, select its name in the list to the left in the Linear System Solver Settings area. In the Coarse solver list you can choose from the following: UMFPACK, SPOOLES, PARDISO, TAUCS Cholesky (if Symmetric is selected), GMRES, FGMRES, Conjugate gradients, Algebraic multigrid, SSOR, SOR, SORU, SSOR vector, SSOR gauge, and Jacobi. Normally choose a direct solver (UMFPACK, SPOOLES, PARDISO, or TAUCS Cholesky). Make any desired modifications to the settings in the Coarse solver area (refer to the sections on the specific linear system solvers).

When an iterative solver is used as coarse solver you can choose whether to solve using a tolerance (default) or to perform a fixed number of iterations. Choose either **Use tolerance** or **Fixed number of iterations** in the **Termination** list. Note that some default values for an iterative solver, when used as a coarse solver, are different from the default values when the solver is used as a linear system solver, preconditioner, or smoother. The edit fields that have different default values are: **Relative tolerance** (default = 0.1), **Factor in error estimate** (default = 1), **Maximum number of iterations** (default = 500), and **Number of iterations** (default = 10).

The Algebraic Multigrid Solver/Preconditioner

The algebraic multigrid solver or preconditioner performs one or several cycles of the algebraic multigrid method. This is similar to the geometric multigrid algorithm (see "The Geometric Multigrid Solver/Preconditioner" on page 518), the difference being that it constructs the multigrid levels directly from the finest-level system matrix A_0 . That is, it constructs the prolongations P_i from A_0 without using auxiliary meshes. It constructs the coarse level matrices A_i from A_0 with the Galerkin projection method. The advantage is that you need not bother about the coarse multigrid levels. The disadvantages are twofold:

- Algebraic multigrid does not work well for vector-valued PDEs in COMSOL's implementation. That is, it handles only scalar PDEs.
- COMSOL's implementation does not support complex-valued system matrices.

In the Linear System Solver Settings dialog box you can control the automatic construction of the multigrid hierarchy with the Maximum number of levels, Max DOFs at coarsest level, and Quality of multigrid hierarchy edit fields. Coarse levels are added until the number of DOFs at the coarsest level is less than the max DOFs at coarsest level (default = 5000) or until it has reached the maximum number of levels, including the finest level (default = 6). In the Quality of multigrid hierarchy edit field specify an

integer value between 1 and 10 (default = 3) to make a tradeoff between memory requirements and preconditioner quality. For instance, if the linear solver does not converge or if it uses too many iterations, try a higher value to increase the accuracy in each iteration, meaning fewer iterations. In contrast, if the algebraic multigrid algorithm runs into memory problems, try a lower value to use less memory. When using algebraic multigrid as a preconditioner, it is also possible to set the value for the quality of the multigrid hierarchy in the **Linear system solver** area on the **General** tab of **Solver Parameters** dialog box either numerically or by using the **Memory efficiency**/**Preconditioner quality** slider.

The remaining settings for the algebraic multigrid solver/preconditioner and its smoothers and coarse solver are identical to those for the Geometric multigrid solver (see "The Geometric Multigrid Solver/Preconditioner" on page 518).

The SSOR, SOR, SORU, and Diagonal Scaling (Jacobi) Algorithms

These simple and memory-efficient solvers/preconditioners/smoothers are based on classical iteration methods for solving a linear system of the form Ax = b. Given a relaxation factor ω (usually between 0 and 2), a sweep of the Jacobi (diagonal scaling) method transforms an initial guess x_0 to an improved approximation $x_1 = x_0 + M^{-1}(b - Ax_0)$, where $M = D/\omega$, and D is the diagonal part of A.

The SOR (successive over-relaxation) method uses the same formula with $M = L + D/\omega$, where *L* is the strictly lower triangular part of *A*. When $\omega = 1$ (the default), the Gauss-Seidel method is obtained. In the SORU method, $M = U + D/\omega$, where *U* is the strictly upper triangular part of *A*. The SOR and SORU methods use a more accurate approximation of the matrix, which leads to fewer iterations but slightly more work per iteration than in the Jacobi method.

The SSOR (symmetric successive over-relaxation) method is one SOR sweep followed by a SORU sweep. The output x_1 for an input x_0 also comes from the above formula but with

$$M = \frac{\omega}{2-\omega} \left(L + \frac{D}{\omega} \right) D^{-1} \left(U + \frac{D}{\omega} \right)$$

When A is symmetric, the SSOR method has the advantage that M is symmetric. Symmetry of the preconditioner matrix is necessary when using the conjugate gradients iterative method. In such cases, the SSOR preconditioner is preferable to the SOR preconditioner. A blocked version of the SOR algorithms is available. It is optimized for parallel computations. In this case M is constructed from a column permuted version of A.

When these algorithms run as linear system solvers, they perform sweeps until they have established convergence or they have reached the maximal number of iteration. You control this aspect with the parameters in the **Relative tolerance**, **Factor in error estimate**, and **Maximum number of iterations** edit fields in the same way as for the other iterative solvers (see for instance "Settings for the Multigrid Solver/Preconditioner" on page 524 of the *COMSOL Multiphysics User's Guide* and "Convergence Criteria" on page 513 of this *Reference Guide*).

When the algorithms run as preconditioners or smoothers, they perform a fixed number of sweeps as dictated by the value in the **Number of iterations** edit field in the **Linear System Solver Settings** dialog box (default = 2).

Specify ω in the **Relaxation factor (omega)** edit field (default = 1).

The SSOR Vector, SOR Vector, and SORU Vector Algorithms

These preconditioners/smoothers are intended for problems involving the $\nabla \times (a \nabla \times .)$ curl-curl operator and where you use so-called *vector elements*. The vector elements are available primarily for electromagnetic-wave simulations in the RF Module. The algorithm is an implementation of the concepts in Ref. 7 and Ref. 8. The algorithm applies SOR iterations on the main linear equation Ax = b but also makes SOR iterations on a projected linear equation $T^TATy = T^Tb$. Here the projection matrix, T, is the discrete gradient operator, which takes values of a scalar field in the mesh vertices and computes the vector-element representation of its gradient. Loosely speaking, the argument for using this projection is the following: For example, let the linear equation Ax = b represent the discretization of a PDE problem originating from the vector Helmholtz equation

$$\nabla \times (a \nabla \times \mathbf{E}) + c \mathbf{E} = \mathbf{F}$$

for the unknown vector field **E**, where *a* and *c* are scalars, and **F** is some right-hand side vector. Standard preconditioners/smoothers cannot smooth the error in the null space of the operator $\nabla \times (a \nabla \times .)$. This null space is the range of the gradient operator. This algorithm adds a correction $\mathbf{E} \rightarrow \mathbf{E} + \nabla \phi$ to the standard SOR smoothed solution (or residual), where it computes ϕ from SOR iterations on a projected problem. The projected problem is obtained by taking the divergence (or discretely $-T^T$) of the Helmholtz equation and plugging in the correction. You then obtain (for clarity, boundary constraints are disregarded)

$$-\nabla \cdot (c\nabla \phi) = -\nabla \cdot \mathbf{F},$$

which, if *c* is definite (strictly positive or strictly negative), is a standard elliptic type of equation for the scalar field ϕ .

When using this algorithm as a smoother for the multigrid solver/preconditioner, it is important—for the correct discrete properties of the projected problem—to generate nested meshes. Also note that it does assembly on all mesh levels (controlled by the multigrid **Assemble** check box). You can generate nested meshes through manual mesh refinements or do so automatically by going to the **Linear System Solver Settings** dialog box and selecting **Refine mesh** from the **Hierarchy generation method** list.

The projection matrix T is computed in such a way that non-vector shape functions are disregarded, and therefore you can use it in a multiphysics setting. It can also handle contributions from different geometries. Non-vector shape function variables are not affected by the correction from the projected system, and the effects on them are therefore the same as when you apply the standard SOR algorithm (see above).

The parameter in the **Number of iterations** edit field in the **Presmoother** (or **Postsmoother**) area controls the number of main iterations (default = 2). For each main iteration, the algorithm makes a number of SOR iterations for the projected equation system; set that number (default = 1) in the **Number of secondary iterations** edit field.

In more detail, to preserve symmetry as a preconditioner and also when used as symmetric pre- and postsmoother in a multigrid setting, the SOR iterations are done in the following order:

- In each main iteration, the SOR vector version of this algorithm makes one SOR iteration on the main system followed by a number of secondary SOR iterations on the projected system.
- In each main iteration, the SORU vector version first makes a number of secondary SORU iterations on the projected system followed by one SORU iteration on the main system.
- In each main iteration, the SSOR vector version makes one SOR iteration on the main system followed by a number of secondary SSOR iterations on the projected system and then one SORU iteration on the main system.

You specify the relaxation factor ω in the **Relaxation factor (omega)** edit field (default = 1). It applies to all the different types of SOR iterations in this algorithm.

These preconditioners/smoothers are primarily intended for 3D magnetostatic problems in the AC/DC Module discretized with vector elements. The smoothers are basically SOR smoothers with some added functionality.

Magnetostatic problems are often formulated in terms of a magnetic vector potential. The solution of problems formulated with such a potential is in general not unique. Infinitely many vector potentials result in the same magnetic field, which typically is the quantity of interest. A finite element discretization of such a problem results in a singular linear system of equations, Ax = b. Despite being singular, these systems can be solved using iterative solvers, provided that the right hand side of the discretized problem is the range of the matrix A. For discretized magnetostatic problems, the range of A consists of all divergence free vectors. Even if the right side of the mathematical problem is divergence free, the right side of the finite element discretization might not be numerically divergence free. To ensure that b is in the range of A, SOR gauge performs a divergence cleaning of the right side by using the matrices T and T^T ; see "The SSOR Vector, SOR Vector, and SORU Vector Algorithms" on page 451. To this end, the system $T^TT\psi = -T^Tb$ is first solved. Adding $T\psi$ to b will then make the numerical divergence of the right side small.

As in the case of SOR there are blocked versions available that perform better in when running on a parallel machine.

In addition to the initial divergence cleaning, SOR gauge also performs a number of cleaning iterations in each linear solver iteration. You can control the number of such divergence cleaning iterations in the **Number of secondary iterations** edit field in the **Linear System Solver** area. The default number of secondary iterations is 1. In the **Variables** edit field you can specify which vector degrees of freedom to include in the divergence cleaning (this applies both to the initial and secondary cleaning iterations). By default, all vector degrees of freedom are included in the divergence cleaning.

The settings **Number of iterations** and **Relaxation factor (omega)** work in the same way as for the usual SOR smoothers; see "The SSOR, SOR, SORU, and Diagonal Scaling (Jacobi) Algorithms" on page 450.

The Vanka Algorithm

This preconditioner/smoother is intended for, but not restricted to, problems involving the Navier-Stokes equations. Formally it applies to saddle-point problems. A saddle-point problem is a problem where the (equilibrium) solution is neither a maximum nor a minimum. The corresponding linear system matrix is indefinite, and often it has zeros on the diagonal. This is the case for the Navier-Stokes equations but also for problems formulated with weak constraints.

The algorithm is a local smoother/preconditioner of Vanka type. It is based on the ideas in Ref. 9, Ref. 10, and Ref. 11. It is possible to describe it as a block SOR method, where the local coupling of the degrees of freedom (DOFs) determines the blocks. The important idea in this algorithm is to use the Lagrange multiplier variable (or set of variables) to form the blocks. For illustration purposes, consider the Navier-Stokes equations. For these equations the pressure variable plays the role of Lagrange multiplier. The linearized equations on discrete form has the following structure:

$$A\begin{bmatrix} U\\ P\end{bmatrix} = \begin{bmatrix} S & D^T\\ D & 0 \end{bmatrix} \begin{bmatrix} U\\ P\end{bmatrix} = \begin{bmatrix} F\\ G\end{bmatrix}$$

where U and P are the velocity and pressure degrees of freedom, respectively. The algorithm loops over the Lagrange multiplier variable DOFs, here the pressure DOFs P_j , and finds the direct connectivity to this DOF. To do so, the algorithm locates the nonzero entries in the matrix column corresponding to P_j . The row indices of the nonzero entries defines the DOFs U_k , and the software forms a local block matrix based on this connectivity:

$$A_j = \begin{bmatrix} S_j & D_j^T \\ D_j & 0 \end{bmatrix}$$

One Vanka update loops over all P_i and updates

$$\begin{bmatrix} U_j \\ P_j \end{bmatrix} \leftarrow \begin{bmatrix} U_j \\ P_j \end{bmatrix} + \omega A_j^{-1} \left(\begin{bmatrix} F \\ G \end{bmatrix} - A \begin{bmatrix} U \\ P \end{bmatrix} \right)_j$$

where the $(.)_j$ denotes the restriction of a vector to the rows corresponding to the block j. ω is a relaxation parameter. The algorithm does not form the inverses of the block matrices explicitly. Instead, it computes the Vanka update either with a LAPACK direct solver subroutine call or by a GMRES iterative method subroutine call. The GMRES method is the restarted GMRES without preconditioning. The algorithm relies on that it is possible to invert the submatrices A_j . If it is not possible, the algorithm gives an error message. Note that a zero on the diagonal of A or A_j is not necessarily a problem for this updating strategy.

Note: If you use the Vanka algorithm as preconditioner, or as smoother to a multigrid preconditioner when either GMRES or Conjugate gradients is used as the linear system solver, you should use the **Direct** option in the **Solver** list in order to get a stationary preconditioner. The **GMRES** option can be useful if you use the FGMRES method as linear system solver since it can handle preconditioners that are not stationary. The **GMRES** option can also be useful if you use the Vanka algorithm as smoother to a multigrid solver because GMRES can be a bit faster than the direct solver.

In general, the Vanka update does not necessarily update all DOFs. This is, for example, the case for problems with weak constraints, where only a small subset of the problem's DOFs are directly coupled to the Lagrange multipliers for the constraints. Another example is the Navier-Stokes equations (or similar types of equations) coupled to other equations, but where the coupling is not directly through the pressure variable. This is, for example, the case with the k- ϵ turbulence model. The Vanka algorithm automatically detects DOFs that are not updated by the above Vanka updating procedure and performs, for each Vanka update, a number of SSOR sweeps for these DOFs. This part of the algorithm is denoted the *SSOR update*. The SSOR update only works for a submatrix that has a nonzero diagonal. Just as the SOR and Jacobi preconditioner algorithms, this algorithm gives an error message if it finds zeros on the diagonal for the DOFs in the SSOR update.

As in the case of SOR there is a blocked version that works on a permuted version of the system matrix. It is especially suited for parallel computations.

Control the number of Vanka updates and consecutive SSOR updates by the parameter in the **Number of iterations** edit field in the **Linear System Settings** dialog box. For the Vanka update, control the Lagrange variables used for the local block definitions by the **Variables** edit field, and control the type of solver used for the block inverse operation by the **Solver** list. If you choose GMRES, then you can control the convergence tolerance and the number of iterations before restart by the parameters in the **Tolerance** and **Number of iterations before restart** edit fields, respectively. Control the Vanka update relaxation parameter ω by the parameter in the **Relaxation factor** edit field. For the SSOR update, control the number of SSOR sweeps by the parameter in the **Number of secondary iterations** edit field and control the SSOR relaxation factor, used in these sweeps, with the parameter in the **Relaxation factor** edit field.

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The COMSOL Multiphysics Files

This chapter describes the COMSOL Multiphysics files in binary format and text format.

Overview

A COMSOL Multiphysics file is used to store COMSOL Multiphysics data. The format is suitable for exchange of mesh or CAD data between COMSOL Multiphysics and other software systems. It is possible to save a COMSOL Multiphysics file in a text file format, using the extension .mphtxt, or a binary file format, using the extension .mphbin. The file formats contain the same data in the same order.

File Structure

The COMSOL Multiphysics file format has a global version number, so that it is possible to revise the whole structure. The first entry in each file is the file format, indicated by two integers. The first integer is the major file version and the second is referred to as the minor file version. For the current version, the first two entries in a file is 0 1.

The following sections describe the file structure of the supported version.

FILE VERSION 0.1

After the file version, the file contains three groups of data:

- A number of *tags* stored as strings, which gives an identification for each *record* stored in the file.
- A number of *types*, which are strings that can be used in serializing the object. The tag should be used as an indication of the contents of the file, but can also be an empty string.
- The *records* containing the serialized data in the file.

Example When using flsave from COMSOL Script to save a COMSOL

Multiphysics mesh, the tag equals the variable name (m1) in COMSOL script, the type is set to obj (but this is not used), and the record contains the serialization of the mesh object, including point coordinates and element data of the mesh. See "Examples" on page 571 for more examples of COMSOL Multiphysics text files.

```
# Created by COMSOL Multiphysics Fri Aug 26 14:19:54 2005
```

```
# Major & minor version
0 1
######### Tags
1 # number of tags
```

```
2 m1
######## Types
1 # number of types
3 obj
######## Records
# A planar face object
0 0 1
4 Mesh # class
...
```

Records

The record contains the serialization data in the file and additional information on how to process the serialized data. It also has a version number.

The record is a wrapper for *serializable types* stored in the file. The reason for having this wrapper is to be able to use a version number, so that the serialization can be revised in the future while maintaining backward compatibility.

The following sections describe the format of the supported version:

RECORD VERSION 0

This record is a wrapper for *serializable types* stored in the file. The following table contains the attributes of the records:

ENTITY/OBJECT	VARIABLE	DESCRIPTION
Integer		Version
Integer		Not used
Integer	type	Serialization type, I for Serializable
Serializable	obj	If type equals 1, this field follows

Serialization type 1 indicates that the following field is a subtype to Serializable. COMSOL Multiphysics uses type equal to 0 internally, but such files are only used for temporary purposes.

Terminology

The following data types are used in the serialization:

- Boolean refers to an 8-bit signed character which must be 0 or 1.
- Character refers to an 8-bit signed character.
- Integer refers to a 32-bit signed integer.
- *Double* refers to a 64-bit double.

Matrices are stored in row-major order. In this documentation we use brackets to indicate a matrix. Hence, integer[3][4] means that 12 integers representing a matrix are store in the file. The first three entries corresponds to the integers in the first row of the matrix, and so on.

Text File Format

COMSOL Multiphysics text file, using the file extension .mphtxt, are text files where values are stored as text separated by whitespace characters.

Lexical conventions:

- Strings are serialized as the length of the string followed by a space and then the characters of the string, for example, "6 COMSOL". This is the only place where whitespace matters.
- The software ignores everything following a # on a line except when reading a string. This makes it possible to store comments in the file.

Binary File Format

COMSOL Multiphysics binary file, using the extension .mphbin, are binary files with the following data representation:

- · Integers and doubles are stored in little-endian byte order.
- Strings are stored as the length of the string (integer) followed by the characters of the string (integers).

SAVING AND LOADING

You can import COMSOL Multiphysics files into the COMSOL Multiphysics GUI or load them as variables into COMSOL Script (or MATLAB).

To load and save COMSOL Multiphysics files in COMSOL Script, use the functions flsave and flload, respectively.

To export to a COMSOL Multiphysics file from the GUI, select **File>Export>Geometry to File**. There is also a corresponding import menu. Note that the multiphysics files do not describe a complete model, so it is not possible to open them from the standard **Open File** dialog box.

Serializable Types

Attribute on page 541 BezierCurve on page 542 BezierMfd on page 543 BezierSurf on page 544 BezierTri on page 545 BSplineCurve on page 546 BSplineMfd on page 547 BSplineSurf on page 549 Ellipse on page 550 GeomO on page 552 Geom1 on page 553 Geom2 on page 554 Geom3 on page 556 GeomFile on page 557 Manifold on page 558 Mesh on page 559 MeshCurve on page 561 MeshSurf on page 562 Plane on page 563 PolChain on page 564 Serializable on page 565 Straight on page 566 Transform on page 567 VectorDouble on page 568 VectorInt on page 569 VectorString on page 570

Supported Versions	0				
Subtype of	Serializable				
Fields	The class is defined by the following fields:				
	ENTITY/OBJECT VARIABLE DESCRIPTION integer Version				
	integer n Number of attribute fields				
	Serializable[n]		Each entity field is stored as a serializable		
Description	An Attribute is a general purpose class from which different subtypes can be derived. Each such subtype should then be serialized using the serialization of the Attribute class, which means that all that it should add to the serialization is the version number. Attributes are used in COMSOL Multiphysics for internal purposes, and these attributes are not documented. However, because Attribute has a strict serialization structure, the serialization of these attributes is well documented.				
Example	This is a serialization of an attribute used internally in COMSOL Multiphysics. It is serialized as a vector of integers. 11 AssocAttrib # class 0 0 # version 1 # nof attribute fields 9 VectorInt # class 18 3 2 2 2 1 0 1 1 1 1 1 0 1 1 1 1 0				

Supported Versions	0		
Subtype of	BezierMfd		
Fields	The class is defined by the following fields:		
	ENTITY/OBJECT	DESCRIPTION	
	integer	Version	
	BezierMfd	Parent class containing common data	
Description	A rational Bézier	curve is a parameterized curve of the form	

$$\mathbf{b}(t) = \frac{\displaystyle\sum_{i=0}^{p} \mathbf{b}_{i} w_{i} B_{i}^{p}(t)}{\displaystyle\sum_{i=0}^{p} w_{i} B_{i}^{p}(t)} \quad , 0 \leq t \leq 1$$

where the functions

$$B_i^p(t) = {p \choose i} t^i (1-t)^{p-i}$$

are the *Bernstein basis* functions of *degree* p, $\mathbf{b}_i = (x_1, ..., x_n)$ are the control vertices of the *n*-dimensional space, and w_i are the weights, which should always be positive real numbers to get a properly defined rational Bézier curve. A rational Bézier curve has a direction defined by the parameter t.

Example

The following illustrates a linear Bézier curve.

11 BezierCurve # class
0 0 # version
2 # sdim
0 2 1 # transformation
1 0 # degrees
2 # number of control points
control point coords and weights
0 0 1
1 1 1

See also

BSplineCurve

Subtype of

Fields

Manifold

0

The class is defined by the following fields:

ENTITY/OBJECT	VARIABLE	DESCRIPTION		
integer		Version		
integer	d	Space dimension		
Transform		Transformation class		
integer	m	Degree in first parameter		
integer	n	Degree in second parameter		
integer	k	Number of control points		
double[k][d+1]	Р	Matrix of control points with the weights in the last column		

Description

The BezierMfd type is the abstract base class for the different type of Bézier surfaces and curves that are supported. These can all be represented in using the generalized equation

$$\mathbf{S}(s,t) = \frac{\displaystyle\sum_{i=0}^{m} \sum_{j=0}^{n} \mathbf{b}_{i,j} w_{i,j} B(s,t)}{\displaystyle\sum_{i=0}^{m} \sum_{j=0}^{n} w_{i,j} B(s,t)}$$

where B are functions as described in the respective entry, and **b** are the control point coordinates in P and w are the weights stored in the last column of P.

See also

BSplineMfd

Supported Versions	0		
Subtype of	BezierMfd		
Fields	The class is defined by the following fields:		
	ENTITY/ DESCRIPTION OBJECT		
	integer Version BezierMfd Parent class containing common data		
Description	A rectangula	r rational Bézier patch of degree <i>p</i> -by- <i>q</i> is described by	

$$\mathbf{S}(s,t) = \frac{\sum_{i=0}^{p} \sum_{j=0}^{q} \mathbf{b}_{i,j} w_{i,j} B_{i}^{p}(s) B_{j}^{q}(t)}{\sum_{i=0}^{p} \sum_{j=0}^{q} w_{i,j} B_{i}^{p}(s) B_{j}^{q}(t)}, \ 0 \le s, t \le 1 \ ,$$

where B_i^p and B_j^q are the Bernstein basis functions of degree p and q, respectively, as described in the entry of BezierCurve. This surface description is called rectangular because the parameter domain is rectangular, that is, the two parameters s and t can vary freely in given intervals.

See also

BSplineSurf, BezierTri

Subtype of

BezierMfd

0

Fields

The class is defined by the following fields:

ENTITY/ OBJECT	DESCRIPTION
integer	Version
BezierMfd	Parent class containing common data

Description

Another form of surface description is the triangular patch, also called a *Bézier* triangle. A triangular rational Bézier patch is defined as

$$\begin{split} \mathbf{S}(s,t) &= \frac{\displaystyle\sum_{i+j \leq p} \mathbf{b}_{i,j} w_{i,j} B_{i,j}^p(s,t)}{\displaystyle\sum_{i+j \leq p} w_{i,j} B_{i,j}^p(s,t)} \quad, 0 \leq s,t \leq 1 \end{split}$$

which differs from the Bézier curve description only by the use of *bivariate* Bernstein polynomials instead of *univariate*, for the curve case. The bivariate Bernstein polynomials of degree *p* are defined as

$$B_{i,j}^{p}(s,t) = \frac{p!}{i!j!(p-i-j)!}s^{i}t^{j}(1-s-t)^{p-i-j}, \quad i+j \le p$$

where the parameters s and t must fulfill the conditions

$$\begin{cases} 0 \le s, t \\ s+t \le 1 \end{cases}$$

which form a triangular domain in the parameter space, therefore the name of this surface description.

See also BezierSurf

Supported Versions	0			
Subtype of	BSplineMfd			
Fields	The class is defined by the following fields:			
	ENTITY/ DESCRIPTION OBJECT			
	integer	Version		
	BSplineMfd Parent class containing common data			
Description	The BSpline	Curve, describes a general spline curve, using B-spline basis functions,		

The BSplineCurve, describes a general spline curve, using B-spline basis functions, as defined in BSplineMfd. Splines on this form are often referred to as B-splines.

A *p*th-degree spline curve is defined by

$$\mathbf{C}(u) = \frac{\sum_{i=0}^{n} N_i^p(u) w_i \mathbf{P}_i}{\sum_{i=0}^{p} N_i^p(u) w_i} , a \le u \le b$$

where \mathbf{P}_i are the control points., the wi are the weights and the N_i^p are the *p*th degree B-spline basis functions defined in the nonperiodic and nonuniform knot vector

$$U = \{a, ..., a, u_{p+1}, ..., u_{m-p-1}, b, ..., b\}$$

For non-rational B-splines, all weights are equal to 1 and the curve can be expressed as

$$\mathbf{C}(u) = \sum_{i=0}^{n} N_{i}^{p}(u) w_{i} \mathbf{P}_{i}, a \le u \le b$$

See also

BezierCurve

Subtype of

Fields

Manifold

0

The class is defined by the following fields:

ENTITY/ OBJECT	VARIABLE	DESCRIPTION	
integer		Version	
integer	d	Space dimension	
Transform		Transformation class	
integer		Dimension (1 if curve, 2 if surface)	
integer	p, q	Degree in each dimension (1 or 2 integers)	
boolean		If rational equal to I	
integer		Number of knot vectors (1 for curves, 2 for surfaces)	
integer	ml	Length of first knot vector	
double[m1]	U	First knot vector	
integer	m2	Length of second knot vector (not for curves)	
double[m2]	V	Second knot vector (not for curves)	
integer	nl	Number of control points in first parameter direction	
integer	n2	Number of control points in second parameter dimension	
integer	n3	Number of coordinates per control point	
double [n1][n2][n3]	Ρ	Matrix of coordinates where the last dimension i increased by 1 to store the weights if the manifold is rational	

Description

The BSplineMfd type is the abstract base type for BSplineCurve and BSplineSurf, that represents general spline curves and surfaces respectively.

They are represented using *B-spline basis functions*. Let $U = \{u_0, ..., u_m\}$ be a non decreasing sequence of real numbers. U is called the *knot vector* and the elements u_i of U are called *knots*. The *i*th B-spline basis function of p-degree, $N_i^p(u)$, is defined as

$$N_{i}^{0}(u) = \begin{cases} 1 \ u_{i} \le u < u_{i+1} \\ 0 \ \text{otherwise} \end{cases}$$
$$N_{i}^{p}(u) = \frac{u - u_{i}}{u_{i+p} - u_{i}} N_{i}^{p-1}(u) + \frac{u_{i+p+1} - u}{u_{i+p+1} - u_{i+1}} N_{i+1}^{p-1}(u)$$

A general B-spline can be described by

$$\mathbf{S}(u,v) = \frac{\sum_{i=0}^{n} \sum_{j=0}^{m} N_{i}^{p}(u) N_{j}^{q}(v) w_{i,j} \mathbf{b}_{i,j}}{\sum_{i=0}^{n} \sum_{j=0}^{m} N_{i}^{p}(u) N_{j}^{q}(v) w_{i,j}}, a \le u \le b, c \le v \le d$$

where

 $U = \{a, ..., a, u_{p+1}, ..., u_{m-p-1}, b, ..., b\}$

and

$$V = \{c, ..., c, v_{p+1}, ..., v_{m-p-1}, d, ..., d\}$$

are the two knot vectors stored in the entry, and \mathbf{b} and w are the control points coordinates and weights stored in P.

For periodic splines, the first and last parameter values in the knot vectors are not duplicated.

Subtype of

Fields

BSplineMfd

0

The class is defined by the following fields:

ENTITY/ OBJECT	DESCRIPTION
integer	Version
BSplineMfd	Parent class containing common data

Description

The generalization of B-spline curves to surfaces is a tensor product surfaces given by

$$\mathbf{S}(u,v) = \frac{\sum_{i=0}^{n} \sum_{j=0}^{m} N_{i}^{p}(u) N_{j}^{q}(v) w_{i,j} \mathbf{P}_{i,j}}{\sum_{i=0}^{n} \sum_{j=0}^{m} N_{i}^{p}(u) N_{j}^{q}(v) w_{i,j}}, a \le u, v \le b$$

See also

BezierSurf

Subtype of	Manifold	Manifold				
Fields	The class is d	The class is defined by the following fields:				
	ENTITY/ OBJECT	VARIABLE	DESCRIPTION			
	integer		Version			
	integer	d	Space dimension			
	Transform		Transformation class			
	double[d]	center	Center coordinate			
	boolean		Equal to 1 if clockwise rotation (only if $d==2$)			
	double[d]	normal	Normal vector coordinates,			
	double[d]	М	Major axis			
	double	rat	Ratio of minor axis length to major axis length			
	double	offset	Parameter at the end of major axis			
	boolean		Equal to 1 if ellipse is degenerated			
Description	This manifol	This manifold defines an ellipse in the two or three dimensional space.				
	axis M of the the minor ax	In 2D, an ellipse is defined by a center point center, a vector defining the major axis M of the ellipse (including the magnitude of the major axis), the radius ratio o the minor axis length to the major axis length rat, the direction of the ellipse, and the parameter offset at the major axis offset.				
	plane of the (including th	In 3D, an ellipse is defined by a center point center, a unit vector normal to the plane of the ellipse normal, a vector defining the major axis of the ellipse M (including the magnitude of the major axis), the radius ratio, and the parameter offset at the major axis offset. The direction of the ellipse is defined by the right hand rule using the normal vector.				
		ing the norma	l vector.			
	hand rule usi	-	l vector. that has a period of 2π . It is parameterized as:			
	hand rule usi An ellipse is :	a closed curve	that has a period of 2π . It is parameterized as:			
	hand rule usi An ellipse is a point =	a closed curve center + M				

0 # reverse? 2 0 # major axis 0.5 # minor axis length / major axis length 0 # parameter value at end of major axis 0 # degenerated? # Attributes 0 # nof attributes

Supported Versions	1			
Subtype of	Serializable			
Fields	The class is defined by the following fields:			
	ENTITY/OBJECT VARIABLE DESCRIPTION			
	integer		Version	
	integer	type	Geometry type	
	double		Relative geometry tolerance	
	integer	Р	Number of points (0 or 1)	
	integer	na	Number of attributes	
	Attribute[na]		Vector of attributes	
Description	geom2, geom3	on page 205.	netry class, as described in the entry geom0, geom1, solid or –1 for general object.	
Example	A solid OD geometry object. 5 GeomO # class 1 O 1e-O1O 1 O # nof attributes			

Supported Versions	1			
Subtype of	Serializable			
Fields	The class is defined by the following fields:			
	ENTITY/OBJECT	VARIABLE	DESCRIPTION	
	integer		Version	
	integer	type	Geometry type	
	double		Geometry tolerance	
	integer	nv	Number of vertices	
	double[nv]	vtx	Vector of vertex coordinates	
	integer[nv][2]	ud	Matrix of integers giving subdomains on up and down side of each vertex	
Description	The type represent a 1D geometry class, as described in the entry geom0, geom2, geom3 on page 205. The type can be either 0, 1, or -1 for point, solid or general objects.			
Example	A solid 1D object. 5 Geom1 # class 1 # version 1 # type 1e-010 # gtol 3 # number of vertices # Vertex coordinates 0 1 3 # Vertex up/down 1 0 2 1 0 2 # Attributes 0 # nof attributes			

Supported Versions	1			
Subtype of	Serializable			
Fields	The class is defined by the following fields:			
	ENTITY/OBJECT	VARIABLE	DESCRIPTION	
	integer		Version	
	integer		Туре	
	double		Relative geometry tolerance	
	integer	nv	Number of vertices	
	integers/doubles [nv][4]	vertex	Matrix of vertex data	
	integer	ne	Number of edges	
	integers/doubles [ne][8]	edge	Matrix of edge data	
	integer	nc	Number of curves	
	Manifold[nc]	curve	An array of Manifold objects	
	geom2, geom3 of The type can be e		or –1 for point, curve, solid or general objects.	
Example	<pre>5 Geom2 # class 1 # version 1 # type 1e-010 # gtol 0.0001 # resTol 2 # number of vertices # Vertices # Vertices # X Y sub tol 0 0 -1 NAN 1 2.29999999999999998 -1 NAN 1 # number of edges # Edges # vtx1 vtx2 s1 s2 up down mfd tol 1 2 0 1 0 0 1 NAN 1 # number of manifolds 11 BezierCurve # class 0 0 # version 2 # sdim 0 2 1 # transformation</pre>			

1 0 # degrees
2 # number of control points
0 0 1
1 2.299999999999998 1
0 # nof attributes

555

ubtype of	Serializable	Serializable			
Fields	The class is defined by the following fields:				
	ENTITY/OBJECT	VARIABLE	DESCRIPTION		
	integer		Version		
	integer	type	Туре		
	double		Relative geometry tolerance		
	double		Relative resolution tolerance		
	integer	nv	Number of vertices		
	integers/doubles [nv][5]	vertex	Matrix of vertex data		
	integer	npv	Number of parameter vertices		
	integers/doubles [npv][6]	pvertex	Matrix of parameter vertex data		
	integer	ne	Number of edges		
	integers/doubles [ne][7]	edge	Matrix of edge data		
	integer	npe	Number of parameter edges		
	integers/doubles [nep][10]	pedge	Matrix of parameter edge data		
	integer	nf	Number of faces		
	integers/doubles [nf][4]	face	Matrix of face data		
	integer	nm	Number of 3D manifolds, curves and surfaces		
	Manifold[nmfd]	mfd	Vector of manifolds		
	integer	npc	Number of parameter curves		
	Manifold[npc]	pcurve	Vector of parameter curves		

The type can be either 0, 1, 2, 3 or -1 for point, curve, shell, solid or general objects.

Supported Versions	1		
Subtype of	Manifold		
Fields	The class is defined by the following fields:		
	ENTITY/ OBJECT	VARIABLE	DESCRIPTION
	integer		Version
	string		M-file name
	integer		Boundary number in M-file
	double		Start parameter value
	double		End parameter value
Description	The bounda	ry index, and s	med part of a boundary described by a geometry M-file. start and end parameters of the trimming parts are store tails on Geometry M-files, see the entry geomfile on
Example	A curve representation using the cardg.m Geometry M-file. 8 GeomFile # class 0 0 # version 2 # sdim 0 2 1 # transformation 5 cardg # filename 1 # boundary number 1.5707963267948966 3.1415926535897931 # parameter range		

Manifold

Supported Versions	0
Subtype of	Serializable
Fields	This is an abstract class with no fields.
Description	A manifold is the common supertype for curve and surface types. It is used by the geometry types.
See also	Geom0, Geom1, Geom2, Geom3

Subtype of

Serializable

1

Fields

The class is defined by the following fields:

ENTITY/OBJECT	VARIABLE	DESCRIPTION Version		
integer				
integer	d	Space dimension (if equal to 0 no more fields)		
integer	np	Number of mesh points		
integer		Lowest mesh point index		
double[d][np]	Р	Mesh points		
integer	nt	Number of element types (fives the number of repeats of the following fields)		
string		Element type		
integer	nep	Number of nodes per element		
integer	ne	Number of elements		
integer[ne][nep]	elem	Matrix of point indices for each element.		
integer	ner	Number of parameter values per element		
integer	nr	Number of parameter sets		
double[nr][ner]	par	Matrix of parameter values		
integer	ndom	Number of domain values		
integer[ndom]	dom	Vector of domain labels for each element		
integer	nud	Number of up/down boundary relations		
integer[nud]	ud	Matrix of integers stating subdomain number or up and down side of the boundary		

Description

This type represent a mesh that can be used by COMSOL Multiphysics. The entries p, elem, par, dom, and ud are all described in the reference entry femmesh on page 134.

Example

The following displays a mesh with triangular elements on a unit square. Neither point or edge elements are present.

4 Mesh # class
2 # sdim
5 # number of mesh points
0 # lowest mesh point index
Mesh point coordinates on unit square
0 0

```
1 0
1 1
0 1
0.5 0.5
1 # number of element types
3 tri # type name
3 # number of nodes per element
4 # number of elements
# Elements, 4 triangular elements
0 1 4
304
234
124
6 # number of parameter values per element
0 # number of parameters
4 # number of domains
# Domains
1
1
2
2
0 # number of up/down pairs
```

Subtype of

Manifold

1

Fields

The class is defined by the following fields:

ENTITY/OBJECT	VARIABLE	DESCRIPTION
integer		Version
integer		Space dimension
Transform		Transformation
integer	np	Number of points
double[np][d]	Р	Matrix of point coordinates
double[np]	par	Vector of point parameters
Manifold	intcurve	Interpolating curve

Description

Mesh structures can also be used to define manifolds. Because meshes contain a number of nodes and, in the case of COMSOL Multiphysics, corresponding parameter values, a good geometric representation can be obtained using a suitable interpolation method for evaluating the values of the manifolds and its derivatives on parameter values that are inside the intervals between the given nodes. Mesh curves are handled by cubic spline interpolation.

The matrix **p** and the vector **par** corresponds to the structures corresponding structures in an edge mesh representation. For the MeshCurve, they serve as the interpolation data to obtain intcurve.

See also BSplineCurve

Supported Versions	1			
Subtype of	Manifold			
Fields	The class is defined by the following fields:			
	ENTITY/OBJECT	VARIABLE	DESCRIPTION	
	integer		Version	
	integer		Space dimension	
	Transform		Transformation	
	integer	nv	Number of vertices	
	double[nv][3]	Р	Matrix of mesh vertex coordinates	
	double[nv][2]		Matrix of mesh vertex parameters	
	integer	nt	Number of triangles	
	integers[nv][3]	elem	Matrix of vertex indices for each element	
Description	ion Mesh structures for surface meshes can be used to make a geon unstructured data. Since a mesh type in COMSOL Multiphysics well as parameter values for each element, interpolation can be smooth surfaces.		sh type in COMSOL Multiphysics have coordinates as	
	A quadratic interpolation is used to define a parametric surface from a surface mesh.			
	The matrix p of the interpolation		and the triangles elem with indices into p are used as	
See also	Mesh			

Supported Versions	1			
Subtype of	Manifold			
Fields	The class is defined by the following fields:			
	ENTITY/ OBJECT	VARIABLE	DESCRIPTION	
	integer		Version	
	integer	d	Space dimension	
	Transform		Transformation	
	double[d]	Р	The point in the plane with parameter value $(0,0)$	
	double[d]	n	Normal vector	
	double[d]	b	Direction of first parameter axis	
Description	This manifold defines a plane in the three dimensional space. It is represented by a point, a unit vector normal to the plane, and the vector of the u derivative. A plane is open in both parameter directions and neither periodic nor singular at any point. It is parameterized as: $pos = p + u*b + v*(n \times b)$			
Example	5 Plane # class 0 0 # version 3 # sdim 0 3 1 # transformation 1.3 0.8000000000000004 1.60000000000001 # root point -6.1257422745431001e-017 0 1 # normal -1 0 -6.1257422745431001e-017 # derivatives 0 # degenerated?			

Supported Versions	1			
Subtype of	Manifold			
Fields	The class is defined by the following fields:			
	ENTITY/ OBJECT	VARIABLE	DESCRIPTION	
	integer		Version	
	integer	d	Space dimension	
	Transform		Transformation	
	integer	nl	First dimension of matrix of point coordinates (equal to d)	
	integer	n2	Second dimension of matrix point coordinates, number of points	
	doubles	pol	n1-by-n2 matrix of point coordinates	
Description	Polygon chains are piece wise linear curves, that are used as approximations of curves in the decomposition algorithm. They have an implicit parameter representation, that is $[(i-1) \& (p-1), i \& (p-1)]$ on the <i>i</i> th interval in a polygon chain with <i>p</i> points. This is not a suitable representation because the derivatives may vary substantially along the curve.			
See also	MeshCurve			

Supported Versions	0
Subtype of	
Fields	This is the abstract base type of all other types. It has no fields.
Description	Serializable is the abstract base type. It is used to indicate that a field can contain all supported types, as is the case for the Attribute type.
See also	Attribute

Supported Versions	1		
Subtype of	Manifold		
Fields	The class is defined by the following fields:		
	ENTITY/ OBJECT	VARIABLE	DESCRIPTION
	integer		Version
	integer	d	Space dimension
	Transform		Transformation
	double[d]	root	The point from which the ray starts
	double[d]	dir	The direction in which the ray points
	double	pscale	Parameter scale
	It is represented by a point and a unit vector specifying the direction. A straigh has a scale factor for the parameterization, so the parameter values can be madinvariant under transformation. If not specified the value of this parameter is s 1.0. A straight line is an open curve that is not periodic. It is parameterized as: pos = root + u*pscale*dir		
	where u is the	e parameter.	
Example	8 Straight # class 0 0 # version 3 # sdim 0 3 1 # transformation 1.3 0.8 0.0 # root point -1 0 0 # direction 1 # parameter scale		
See also	Plane		

Supported Versions

Subtype of Serializable

Fields

The class is

1

ENTITY/ OBJECT	VARIABLE	DESCRIPTION
integer	d	Space dimension
boolean		l if transformation is a unit transformation, 0 otherwise. If the value is 1, no more fields are present
double [d+1][d+1]	М	Values in transformation matrix
boolean		I if determinant is positive, 0 otherwise
boolean		I if matrix is isotropic, 0 otherwise

Description

The transformation class is defined by the transformation matrix, that operates as a pre-multiplier on column vectors containing homogeneous coordinates thus

$$\begin{bmatrix} x' \ y' \ z' \ s' \end{bmatrix} = M \cdot \begin{bmatrix} x \ y \ z \ s \end{bmatrix}'$$

where the conventional 3D coordinates are

$$\frac{x}{s} \frac{y}{s} \frac{z}{s}$$

The matrix thus consists of

$$\begin{bmatrix} & T_x \\ \mathbf{R} & T_y \\ & T_z \\ 0 & 0 & 0 \end{bmatrix}$$

where \mathbf{R} is a non singular transformation matrix, containing the rotation, reflection, non-uniform scaling and shearing components, \mathbf{T} is a translation vector and S is a global scaling factor greater than zero.

Supported Versions			
Subtype of	Serializable		
Fields	The class is defined by the following fields:		
	ENTITY/ OBJECT	VARIABLE	DESCRIPTION
	integer	n	Number of elements
	double[n]	d	Elements
Description	This is just a wrapper for a vector of doubles, that can be used to store fields in the Attribute class.		
See also	Attribute		

Supported Versions

Subtype of	Serializable		
Fields	The class is defined by the following fields:		
	ENTITY/ OBJECT	VARIABLE	DESCRIPTION
	integer	n	Number of elements
	integer[n]	d	Elements
Description	This is just a wrapper for a vector of integers, that can be used to store fields in the Attribute class.		
See also	Attribute		

VectorString

Supported Versions			
Subtype of	Serializable		
Fields	The class is defined by the following fields:		
	ENTITY/ OBJECT	VARIABLE	DESCRIPTION
	integer	n	Number of elements
	string[n]	d	Elements
Description	This is just a wrapper for a vector of strings, that can be used to store fields in the Attribute class.		
See also	Attribute		

Examples

To illustrate the use of the serialization format, some text files are listed in this session.

A Mesh with Mixed Element Types

A file containing a 3D mesh with 2nd order tetrahedral, prism and block elements. Some rows in the file are removed and replaced by an ellipsis(...).

```
# Created by COMSOL Multiphysics Fri Aug 26 12:43:42 2005
# Major & minor version
01
1 # number of tags
# Tags
7 fem35.0
1 # number of types
# Types
3 obj
# A mesh object
001
4 Mesh # class
3 # sdim
1503 # number of mesh points
0 # lowest mesh point index
# Mesh point coordinates
000
2.5 0 0
500
. . .
12.5 28.33333000000004 15
12.5 30 13.125
12.5 28.33333000000004 13.125
7 # number of element types
# Type #0
4 tet2 # type name
10 # number of nodes per element
318 # number of elements
# Elements
926 18 13 17 971 958 61 967 66 60
11 345 918 342 950 951 1137 949 373 1129
924 164 345 5 1026 1138 384 938 385 378
```

```
20 339 15 16 352 68 356 69 960 64
. . .
287 919 930 285 1100 1102 1152 317 1096 1098
3 227 4 8 936 28 243 35 945 36
30 # number of parameter values per element
0 # number of parameters
# Parameters
318 # number of domains
# Domains
1
. . .
1
1
0 # number of up/down pairs
# Up/down
# Type #1
6 prism2 # type name
18 # number of nodes per element
96 # number of elements
# Elements
85 174 90 476 474 557 221 118 237 499 1171 494 1170 1172 566 496 563 597
174 225 90 474 588 557 238 237 244 494 1173 579 1172 1174 566 598 597 604
174 175 225 474 472 588 222 238 239 494 1175 489 1173 1176 579 491 598 599
. . .
654 528 530 404 409 408 693 692 541 703 1333 538 1332 1334 543 460 459 465
654 504 528 404 405 409 687 693 694 703 1324 520 1333 1335 538 453 460 461
504 506 528 405 410 409 523 694 536 520 1336 525 1335 1337 538 462 461 466
54 # number of parameter values per element
0 # number of parameters
# Parameters
96 # number of domains
# Domains
2
2
2
. . .
2
2
0 # number of up/down pairs
# Up/down
# Type #2
4 hex2 # type name
27 # number of nodes per element
36 # number of elements
# Elements
```

```
410 506 409 528 762 760 831 859 525 466 1337 536 538 780 1339 775 1338 1343
1340 840 1341 853 777 837 1342 865 867
506 507 528 529 760 758 859 860 526 536 1281 537 539 775 1344 770 1340 1348
1345 853 1346 854 772 865 1347 866 868
. . .
893 785 809 787 718 719 722 723 908 910 1446 804 815 916 1487 801 1495 1502
1499 817 1500 806 747 750 1501 751 754
81 # number of parameter values per element
0 # number of parameters
# Parameters
36 # number of domains
# Domains
3
3
. . .
3
3
0 # number of up/down pairs
# Up/down
# Type #3
3 vtx # type name
1 # number of nodes per element
16 # number of elements
# Elements
0
4
20
. . .
723
0 # number of parameter values per element
0 # number of parameters
# Parameters
16 # number of domains
# Domains
0
1
. . .
14
15
0 # number of up/down pairs
# Up/down
# Type #4
4 edg2 # type name
```

```
3 # number of nodes per element
102 # number of elements
# Elements
4 9 37
9 14 50
14 19 63
. . .
175 174 222
174 85 221
3 # number of parameter values per element
102 # number of parameters
# Parameters
0 5 2.5
5 10 7.5
10 15 12.5
. . .
7.5 11.25 9.375
11.25 15 13.125
102 # number of domains
# Domains
0
0
. . .
27
27
27
0 # number of up/down pairs
# Up/down
# Type #5
4 tri2 # type name
6 # number of nodes per element
224 # number of elements
# Elements
18 17 13 66 61 60
164 5 345 385 384 378
20 339 15 352 68 356
. . .
404 409 405 460 453 461
405 409 410 461 462 466
12 # number of parameter values per element
0 # number of parameters
# Parameters
224 # number of domains
# Domains
12
7
```

```
7
. . .
5
5
5
224 # number of up/down pairs
# Up/down
10
10
10
1 0
. . .
2 0
2 0
# Type #6
5 quad2 # type name
9 # number of nodes per element
102 # number of elements
# Elements
85 90 476 557 118 499 1170 566 563
85 476 174 474 499 221 1171 496 494
. . .
809 787 722 723 815 817 1500 806 754
718 722 719 723 750 747 1501 754 751
18 # number of parameter values per element
0 # number of parameters
# Parameters
102 # number of domains
# Domains
1
4
4
. . .
8
2
102 # number of up/down pairs
# Up/down
2 0
2 0
. . .
3 0
3 0
```

A Planar Face

The following listing is a complete representation of a planar 3D face.

```
# Created by COMSOL Multiphysics Fri Aug 26 14:19:54 2005
# Major & minor version
0 1
######## Tags
1 # number of tags
# Tags
2 b1
####### Types
1 # number of types
# Types
3 obj
####### Records
# A planar face object
001
5 Geom3 # class
1 # version
2 # type
1e-010 # gtol
0.0001 # resTol
4 # number of vertices
# Vertices
# X Y Z sub tol
0 0 0 -1 1e-010
1 0 6.1257422745431001e-017 -1 1e-010
0 1 0 -1 1e-010
1 1 6.1257422745431001e-017 -1 1e-010
4 # number of parameter vertices
# Parameter vertices
# vtx s t fac mfd tol
1 0 0 -1 1 NAN
2 1 0 -1 1 NAN
3 0 1 -1 1 NAN
4 1 1 -1 1 NAN
4 # number of edges
# Edges
# vtx1 vtx2 s1 s2 sub mfd to1
1 2 0 1 -1 2 NAN
2 4 0 1 -1 3 NAN
4 3 0 1 -1 4 NAN
3 1 0 1 -1 5 NAN
4 # number of parameter edges
# Parameter edges
# edg v1 v2 s1 s2 up down mfdfac mfd tol
1 1 2 0 1 1 0 1 1 NAN
2 2 4 0 1 1 0 2 1 NAN
343011031 NAN
4 3 1 0 1 1 0 4 1 NAN
1 # number of faces
```

```
# Faces
# up down mfd tol
0 0 1 NAN
5 # number of 3D manifolds
# Manifold #0
5 Plane # class
0 0 # version
3 # sdim
0 3 1 # transformation
0 0 0 # root point
-6.1257422745431001e-017 0 1 # normal
1 0 6.1257422745431001e-017 # derivatives
0 # degenerated?
# Manifold #1
8 Straight # class
0 0 # version
3 # sdim
0 3 1 # transformation
0 0 0 # root point
1 0 6.1257422745431001e-017 # direction
1 # parameter scale
# Manifold #2
8 Straight # class
0 0 # version
3 # sdim
0 3 1 # transformation
1 0 6.1257422745431001e-017 # root point
0 1 0 # direction
1 # parameter scale
# Manifold #3
8 Straight # class
0 0 # version
3 # sdim
0 3 1 # transformation
1 1 6.1257422745431001e-017 # root point
-1 0 -6.1257422745431001e-017 # direction
1 # parameter scale
# Manifold #4
8 Straight # class
0 0 # version
3 # sdim
0 3 1 # transformation
0 1 0 # root point
0 -1 0 # direction
1 # parameter scale
4 # number of parameter curves
# Paramerer curve #0
8 PolChain # class
0 0 # version
```

```
2 # sdim
0 2 1 # transformation
2 2 0 0 1 0 # chain
```

```
# Paramerer curve #1
8 PolChain # class
0 0 # version
2 # sdim
0 2 1 # transformation
2 2 1 0 1 1 # chain
```

```
# Paramerer curve #2
8 PolChain # class
0 0 # version
2 # sdim
0 2 1 # transformation
2 2 1 1 0 1 # chain
```

Paramerer curve #3

8 PolChain # class 0 0 # version 2 # sdim 0 2 1 # transformation 2 2 0 1 0 0 # chain

Attributes
0 # nof attributes

COMSOL Engine API

Introduction

The COMSOL Engine API (Application Programming Interface) makes it possible to set up, solve, and postprocess a PDE problem using COMSOL as a "black box." The API uses a Java VM to communicate with the COMSOL shared library. To create an application with the API requires the Java SDK 1.4.2, which you must download and install separately.

We recommend you run the COMSOL Engine API from the Java VM (Java Virtual Machine) that ships with COMSOL. The Java VM needs access to the COMSOL class library and the COMSOL shared libraries. The COMSOL API lets you control COMSOL through commands similar to those in COMSOL Script. In fact, to run a COMSOL model through the API, you can use the exact same command sequence that you get by saving a Model M-file from the COMSOL user interface. COMSOL commands sent to the Java VM describe the geometry, mesh, PDE, and boundary conditions; you send additional commands to solve the model and perform postprocessing. Finally, other COMSOL Engine API methods fetch the data into Java.

The next section, "Using the COMSOL Engine API," explains how to get started using the API. Additionally, the *COMSOL Multiphysics Scripting Guide* is useful reading for working with the API, and the "Command Reference" chapter on page 1 in this manual describes the COMSOL Multiphysics commands in detail.

Note that it is possible to start the COMSOL API from a C program, as well. You must use the JNI (Java Native Interface) to start a Java VM from the C program, and then send commands to the Java VM using the JNI. Also use the JNI to fetch numerical results from COMSOL. It is clearly easier to use the COMSOL API from a program that is already integrated with Java.

Using the COMSOL Engine API

An external program can communicate with COMSOL through a Java API that is based on executing COMSOL Script commands. The interface classes are declared in the Java package com.femlab.script.api.

When it receives commands through the API, COMSOL evaluates them sequentially. The subset of MATLAB from which these commands come consists of those that can appear in Model M-files; for additional details in this regard as well as a description of which COMSOL functions that you can call and in what order, see "The Structure of a Model M-file" on page 65 in the *COMSOL Multiphysics Scripting Guide* that is part of the COMSOL Multiphysics documentation set.

Classes

THE APIWORKSPACE CLASS

ApiWorkspace is the main class for communicating with COMSOL. It corresponds to the root workspace in MATLAB. It provides the following interface:

- ApiWorkspace() creates a new COMSOL workspace.
- void destroy() deactivates the workspace and frees all memory it has allocated.
- void eval(String command) executes a COMSOL command written in the MATLAB language. The command can contain several MATLAB statements separated by semicolons, commas, or newline characters.
- boolean isNumeric(String var) returns True if a variable called var exists and its value is or can be converted to a real or complex matrix.
- boolean isComplex(String var) returns True if a variable called var exists and its value has a non-zero imaginary part.
- double[][] getMatrixReal(String var) returns a matrix containing the real parts of the matrix var. If no such variable exists an exception is thrown.
- double[][] getMatrixImag(String var) returns a matrix containing the imaginary parts of the matrix var. If no such variable exists, an exception is thrown.
- String getVar(String var) returns the value of a the variable var as a string.
- void setVar(String var, String value) assigns the variable var the value of the expression value.

THE APIEXCEPTION CLASS

ApiException is an exception thrown by all member functions of ApiWorkspace. It provides the following interface:

• String getMessage() returns a descriptive error message.

Some common situations where this exception is thrown include:

- The command sent to ApiWorkspace.eval() could not be parsed.
- The command sent to ApiWorkspace.eval() contains MATLAB commands, expressions, or functions not supported by COMSOL.
- The command sent to ApiWorkspace.eval() contains invalid calls to COMSOL functions.
- The variable name passed to ApiWorkspace.getMatrixReal() or ApiWorkspace.getVar() does not correspond to an existing matrix variable.

Example

The following code snippet illustrates how it is possible to use the API.

```
import com.femlab.script.api.*;
import java.io.*;
public class Example {
  public static void main(String[] args) {
    try {
      ApiWorkspace ws = new ApiWorkspace();
      // Create geometry: A square
      ws.eval("g1 = rect2(1, 1, 'base', 'corner', 'pos',[0, 0]);");
      ws.eval("s.objs = {g1};");
      ws.eval("s.name = { 'R1 ' }; ");
      ws.eval("s.tags = {'g1'};");
      ws.eval("fem.draw = struct('s', s);");
      ws.eval("fem.geom = g1;");
      // Create mesh
      ws.eval("fem.mesh = meshinit(fem);");
      // Use the Electrostatics application mode
      ws.eval("appl.mode.class = 'Electrostatics';");
      ws.eval("appl.assignsuffix = ' es';");
```

```
// Set boundary constraints: Potentials 0 and 10V on
      // the left and right sides, respectively.
      ws.eval("bnd.V0 = {0, 0, 10};");
      ws.eval("bnd.type = {'V0', 'nD0', 'V'};");
      ws.eval("bnd.ind = [2, 3, 2, 1];");
      ws.eval("appl.bnd = bnd;");
      ws.eval("fem.appl{1} = appl;");
      // Solve the problem using the linear solver
      ws.eval("fem = multiphysics(fem);");
      ws.eval("fem.xmesh = meshextend(fem);");
      ws.eval("fem.sol = femlin(fem);");
      // Retrieve the potential in the midpoint of the square
      ws.setVar("coord", "[0.5; 0.5]");
      ws.eval("sol = postinterp(fem, 'V', coord);");
      double[][] sol = ws.getMatrixReal("sol");
      ws.destroy();
      // Print the results
      for (int i=0; i<sol.length; i++) {</pre>
        for (int j=0; j<sol[i].length; j++)</pre>
          System.out.print(sol[i][j]);
        System.out.println();
      }
    } catch (ApiException e) {
      System.out.println(e.getMessage());
    }
  }
}
```

Running the Example

WINDOWS

To run this example under Windows, you must first have COMSOL 3.4 installed. Next download and install Java SDK 1.4.2 for Windows. You also need the batch files comsolapic.bat and comsolapi.bat (available in the COMSOL bin directory) and the example file Example.java (available in the COMSOL api/engine directory). Put all three of these files in a new directory. Edit the variable SDKROOT in the batch file comsolapic.bat to reflect the installation directory of Java SDK 1.4.2.

To compile the function enter

comsolapic.bat Example.java and to run the model enter

comsolapi.bat Example

You can determine exactly which parameters Java needs in order to run a model using the COMSOL API by looking at the file comsolapi.bat. You must also perform these tasks: provide the correct class path to Java, update the PATH variable, and set the variable LMCOMSOL_LICENSE_FILE.

LINUX/SUN/MAC

To run the example just presented under Linux, Sun, or the Mac, you must first have COMSOL 3.4 installed. Next download and install Java SDK 1.4.2 for your platform. You also need the shell scripts comsolapic and comsolapi plus the example file Example.java. Edit the variables FLROOT, SDKROOT, and ARCH in the shell scripts to reflect the installation directories of COMSOL 3.4, Java SDK 1.4.2, as well as the computer architecture.

To compile the function enter

comsolapic Example.java

and to run the model enter

comsolapi Example

You can determine exactly which parameters Java needs in order to run a model using the COMSOL API by looking at the file comsolapi. You must also perform these tasks: provide the correct class path to Java, update the LD_LIBRARY_PATH variable (DYLD_LIBRARY_PATH on Mac), and set the variable LMCOMSOL_LICENSE_FILE.

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