Rotor Module for the Structural Dynamics Toolbox

For Use with MATLAB

THESE CAPABILITIES ARE DISTRIBUTED EXPERIMEN-TALLY. NOT ALL FUNCTIONALITY IS DOCUMENTED

User's Guide

Version 1.0

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Installation

1

The SDT/Rotor toolbox is intalled as a patch on an SDT installation. You can download the toolbox from

http://www.sdtools.com/distrib/beta/rotor_patch_dis.p

Then within MATLAB,

```
% 1/ cd to location where you saved the patch
% MUST differ from SDT path
cd tempdir
% 2/ check that you will overwrite the expected SDT installation
path
which('feplot');
% 3/ install the patch (you must have write permission on the directory)
rotor_patch_dis
```

END OF CHAPTER

Theoretical reminders

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The following sections give a number of theoretical reminders on things used for the toolbox. THIS IS VERY INCOMPLETE AND NOT VERY ORDERED.

2.1 Rotating bodies

2.1.1 Problem definition in a rotating frame

The developments of this section are derived from internal work on the SDT Rotor module which is currently only distributed to SNECMA. The results shown here can be seen as a summary of those found in Ref. [1] which treats the problem with a strong emphasis on the theoretical formalism. Other classical references that treat of the problem of rotating bodies are [2],[3], [4].

Particles located in point p of the body fixed frame are at location x at time t. One defines the displacement u by

$$\{x(p,t)\} = \{p\} + \{u(p,t)\}$$
(2.1)

At time t, a reference point of the rotating body is assumed to have a rigid rotation speed $\{\omega\}$ with respect to the reference frame (in the present study, this speed is related to a global rotation around a fixed axis characterized by angle θ). The velocity is thus given by

$$\{v_x\}(p,t) = \frac{\partial\{x\}}{\partial t} + \{\omega(t)\} \land \{x(p,t)\}$$

$$(2.2)$$

This expression can easily be derived by decomposing the position in body fixed coordinates $\{x\} = x_i \{e_{bi}\}$ and noting that the derivatives the base vectors $\partial \{e_{bi}\} / \partial t = \{\Omega\} (t) \land \{e_{bi}\}$. In implementations, one replaces the vector product $\omega(t) \land$ by the product by the corresponding skew-symmetric matrix

$$\Omega(t) = \begin{bmatrix} 0 & -\omega_z(t) & \omega_y(t) \\ \omega_z(t) & 0 & -\omega_x(t) \\ -\omega_y(t) & \omega_x(t) & 0 \end{bmatrix}.$$
(2.3)

The acceleration, derived from the velocity expression, is given by

$$\{a\}(p,t) = \{\ddot{x}\} + \left[\dot{\Omega}(t)\right]\{x\} + 2\left[\Omega\right]\{\dot{x}\} + \left[\Omega^{2}\right]\{x\} = \{\ddot{u}\} + \left[\dot{\Omega}(t)\right](p+u) + 2\left[\Omega\right]\{\dot{u}\} + \left[\Omega^{2}\right]\{p+u\}$$

$$(2.4)$$

where three contributions (rows of the equation) are typically considered : the acceleration in the rotating frame, the Coriolis acceleration and the centrifugal acceleration.

The virtual work of acceleration quantities is thus typically expressed as

$$\{\hat{q}\}\left([M]\{\ddot{q}\} + [D_g]\{\dot{q}\} + [K_a + K_c]\{q\} + f_c + f_g\right)$$
(2.5)

with the following element level expressions. The displacement within an element is given by the position and the element shape functions [N] in the three directions xyz

$$\{x\} = \{p\} + [N_{xyz}]\{q\}$$

The matrices and loads are integrated over the volume S_0 in the reference configuration and are given by

- $[M] = \int_{\mathcal{S}_0} \rho_0 [N_{xyz}]^\top [I] [N_{xyz}] d\mathcal{S}_0$ the mass in the rotating frame
- $[D_g] = \int_{\mathcal{S}_0} 2\rho_0 \left[N_{xyz} \right]^\top \left[\Omega \right] \left[N_{xyz} \right] \, d\mathcal{S}_0$ the gyroscopic coupling
- $[K_c] = \int_{\mathcal{S}_0} \rho_0 [N_{xyz}]^\top [\Omega^2] [N_{xyz}] d\mathcal{S}_0$ the centrifugal softening/stiffening
- $[K_a] = \int_{\mathcal{S}_0} \rho_0 [N_{xyz}]^\top \frac{\partial [\Omega]}{\partial t} [N_{xyz}] d\mathcal{S}_0$ the centrifugal acceleration
- $f_c = \int_{\mathcal{S}_0} \rho_0 \left[N_{xyz} \right]^\top \left[\Omega^2 \right] \{ p \} \ d\mathcal{S}_0$ the centrifugal load
- $f_g = \int_{\mathcal{S}_0} \rho_0 \left[N_{xyz} \right]^\top \frac{\partial [\Omega]}{\partial t} \{p\} \ d\mathcal{S}_0$ the Coriolis load

It is acknowledged that the notations used can be somewhat confusing. Indeed, in a discretized vectors DOFs are placed either sequentially x,y,z at all nodes of the element, or separated x at all nodes, ... while the operations $[N_{xyz}]^{\top}[I][N_{xyz}]$ imply the use of vectors. This 2 dimensional product notation however directly reflects the numerical implementation as is thus deemed preferable.

In the applications considered in this study, one will use a fixed axis of rotation $\Omega = \omega(t) \{e_z\} \wedge$. The matrices and loads are thus proportional to the scalars ω , ω^2 and $\dot{\omega}$. One will thus simply use

$$[D_g(\omega)] = \omega [D_g(1)]$$

which results in significant computational cost savings since the matrix only needs to be computed for a single velocity. One proceeds similarly for the other matrices and loads.

2.2 Problem definition in a fixed frame

Fully axisymmetric rotors can be modeled in a fixed frame using an Eulerian representation, where particles are moving under a deformed mesh. Particles located at point $\{p(p_0, t)\} = \{p_0\} + \{u(p_0, t)\}$ in the deformed Eulerian frame have a velocity given by xxx

$$\{v(x,t)\} = \frac{\partial x(p+u(p,t),t)}{\partial t} + \frac{\partial x(p+u(p,t),t)}{\partial x}$$
(2.6)

The body fixed frame verifies $\theta = \phi + \Omega t$, which can be written as $p_G = \{r, \theta, z\} = R_{\Omega t} \{p\}$. The matching of displacements in both frames is given by

$$\{u(p,t)\} = \{u_G(p_G,t)\}$$
(2.7)

The velocity of a particle in the disk is given by

$$\{v(p_G, t)\} = [\Omega] \{p_G\} + [\Omega] \left\{\frac{\partial u}{\partial p_G}\right\} + \left\{\frac{\partial u_G}{\partial t}\right\}$$
(2.8)

Validation example : One first considers a disk that has a steady state deformation in the global frame. That is the Eulerian frame, one has $\frac{\partial u}{\partial t} = 0$ and $\{u(r, \theta, z)\} = \{u(p)\} = (2 - \cos^2\theta) \{e_{\theta}\}$.

One now considers a disk that has a steady state deformation in the rotating frame: $\frac{\partial u}{\partial t} = 0$ and $\{u(r,\phi,z)\} = (1 - \cos^2 \phi) \{e_{\phi}\}$. The displacement of a particle located at at time t is given by

$$\left\{u_G(p_G, t)\right\} = \left(1 - \cos^2(\theta - \Omega t)\right) \left\{e_{(\theta - \Omega t)}\right\} = \left[R_{\Omega t}\right] \left\{u(p)\right\}$$

its velocity is

2.3 Fourier analysis of structures

For more details, you can refer to[5] that is available on-line.

2.3.1 cyclic structure basics

For cyclic system with N sectors (angle $\alpha = 2\pi/N$), a point load placed at angle $n\alpha$ is associated with the harmonic load

$$Y(k) = \frac{1}{N} \sum_{n=0}^{N-1} y(n) e^{-j\alpha nk}$$
(2.9)

this can be used to simply compute the k diameter response. Using the symmetry of the spectrum, one can recover the full spatial response by inverse fourier transform. For N odd

$$y(n) = Y(0) + \sum_{k=1}^{(N-1)/2} 2\operatorname{Re}(Y(k)e^{j\alpha nk})$$
(2.10)

for N even

$$y(n) = Y(0) + (-1)^{n} Y(N/2) + \sum_{k=1}^{N/2-1} 2\operatorname{Re}(Y(k)e^{j\alpha nk})$$
(2.11)

The displacements y(n) of each sector are however expressed in local coordinates, when applying conditions a transformation $\theta^{\alpha n}$ to global coordinates is thus needed

$$\left\{ \begin{array}{c} q(n)_x \\ q(n)_y \\ q(n)_z \end{array} \right\}_{Glob} = \left[\begin{array}{c} \cos\alpha & -\sin\alpha & 0 \\ \sin\alpha & \cos\alpha & 0 \\ 0 & 0 & 1 \end{array} \right]^n \left\{ \begin{array}{c} q(n)_x \\ q(n)_y \\ q(n)_z \end{array} \right\}_{Loc}$$
(2.12)

For a point on the axis, the in plane response is given by two components u(0), v(0), the response at other points of the series is given by (2.10) but the point being coincident on has $\{u(0), v(0)\} = (\theta^{\alpha})^n \{u(n), v(n)\}$. It follows that Y(k) = 0 for all $k \neq 1$.

The intersector continuity condition says that the nodes that are common to two sectors have the same motion. By convention the left nodes in SDT rotor are those with the lowest clockwise polar angle, the left nodes of sector 0 have thus equal motion than the right nodes or sector N-1. Thus $c_L - c_R(Y(k)e^{-j\alpha k}) = 0$. Which leads to the constraint

$$\begin{bmatrix} [c_l] - \cos(k\,\alpha) [c_r] & -\sin(k\,\alpha) [c_r] \\ \sin(\delta\,\alpha) [c_r] & [c_l] - \cos(\delta\,\alpha) [c_r] \end{bmatrix}_{2N_r \times 2N} \begin{cases} Re(q) \\ Im(q) \end{cases} = 0$$
(2.13)

The Fourier transform being a linear relation, one can actually rewrite the relation as a $\{y\} = [E] \{Y\}$. For N even

$$\begin{cases} y_{0} \\ \vdots \\ y_{n} \\ \vdots \\ y_{N-1} \end{cases} = \begin{bmatrix} 1 & \cdots & 2 & 0 & \cdots & 1 \\ \vdots & \vdots & \vdots & \vdots & \vdots \\ 1 & \cdots & 2cos(\alpha kn) & -2sin(\alpha kn) & \cdots & (-1)^{n} \\ \vdots & \vdots & \vdots & \vdots & \vdots \end{bmatrix} \begin{cases} Y_{0} \\ \vdots \\ Re(Y_{k}) \\ Im(Y_{k}) \\ \vdots \\ Y_{N/2} \end{cases}$$
(2.14)

Note that $E^T E$ is a diagonal matrix with $[N \ 2N \ \cdots \ 2N \ N]$ on the diagonal.

Posing $[I_{NQ}]$ is the identity matrix whose size is the number of physical DOF of the sector and \otimes is the Kronecker product, the fourier DOFs Q_k (sorted as in (2.14)) and physical DOFs on the whole disk are related by

$$\{q\} = [E \otimes I_{NQ}] \{Q_k\}$$

$$(2.15)$$

When performing harmonic computations, one typically uses the fact that the model matrices are identical for each sector. As a result, the transformed matrices are block-diagonal, which is the basis for the ability to compute full responses based on independent computation of each Fourier solution Q_k .

When adding a matrix to sector n, the relation to to the Fourier DOFs is given by a row E_n , the product $E_n^T E_n$ is not block diagonal as a result one has coupling between the Fourier harmonics.

2.3.2 Fourier transform for shaft computations

From mono-harmonic modes stored in SDT, you can obtain the physical response using for complex value shape pairs

$$\Re\left(\sqrt{2/N}\left\{q\right\}e^{shift}\right) \tag{2.16}$$

and for real valued vectors associated with 0 and N/2 diameters.

$$\left(\sqrt{1/N}\right)\left\{q\right\}e^{shift}\tag{2.17}$$

2.3.3 Solutions in periodic media

This really does not apply to rotors but is implemented in fe_cyclic, and thus documented here. One considers a model whose properties are spatially periodic. For a physical response, known at regularly spaced positions $n\Delta x$, one can compute the its fourier transform

$$U(\kappa) = \sum_{n=-\infty}^{+\infty} u(n\Delta x)e^{-jn(\kappa\Delta x)}$$
(2.18)

 $U(\kappa)$ is a complex shape defined on the mesh of the repeated cell. One actually uses two cells to represent the real and imaginary parts of U. The wave number κ varies in the $[0, \frac{2\pi}{\Delta x}]$ interval (or any interval of the same length, since $U(\kappa)$ is periodic in the wavelength domain).

Given the fourier transform $U(\kappa)$, on can recover the physical motion by computing the inverse Fourier transform

$$u(n\Delta x) = \frac{\Delta x}{2\pi} \int_0^{\frac{2\pi}{\delta x}} U(\kappa) e^{j\kappa n\Delta x} d\kappa$$
(2.19)

For a mono-harmonic response (fixed wave number κ), the spatial transform is given by $u(n\Delta x) = (U(\kappa)e^{jn(\kappa\Delta x)})$, using the continuity condition linked to the fact that $u_{left}(n\Delta x) = u_{right}((n-1)\Delta x)$, one thus has $[c_L] \{U(\kappa)\} = [c_R] \{U(\kappa)\} e^{-j(\kappa\Delta x)}$. Hence in the real/imaginary format, one has the constraint equation

$$\begin{bmatrix} [c_l] - \cos(\kappa \Delta x) [c_r] & -\sin(\kappa \Delta x) [c_r] \\ \sin(\kappa \Delta x) [c_r] & [c_l] - \cos(\kappa \Delta x) [c_r] \end{bmatrix}_{2N_r \times 2N} \begin{cases} Re(U) \\ Im(U) \end{cases} = 0$$
(2.20)

and one solves for $U(\kappa)$ knowning the transform of the applied loads. For a discrete load on the first slice n = 0, the fourier coefficients of the load is $F(\kappa) = f(0)$.

One can usefully note that the wave length $L = \pi/\kappa$ covers the full interval of positive lengths, that $U(\frac{2\pi}{\delta x} - \kappa) = \bar{U}(\kappa)$, and that the half spectrum corresponds to a wavelength of Δx . Using the property of symmetry allows the use of computations for wavelengths larger than Δx only. In the fe_cyclic eig calls, one specifies the period as a number of steps $p\Delta x$. The phase angle is thus $\kappa\Delta x = \frac{2\pi}{n}$ and the spectrum is symmetric for p' = 2/p.

```
mo1=femesh('testhexa8b divide 2 1 1');
mo1=fe_cyclic('build -1 1 0 0',mo1);
```

```
% symmetry along right edge
r2=fe_cyclic('build -1 0 1 0',mo1);
r2=fe_case(r2,'getdata','Symmetry');
mo1=fe_case(mo1,'FixDof','y_symmetry',r2.IntNodes(:,1)+.02);
```

```
mo1=stack_set(mo1,'info','EigOpt',[2 2 1e3]);
```

```
range=1./linspace(.01,.99,21);
def=fe_cyclic(['teig -all' sprintf(' %g',range)],mo1);
figure(1);plot(1./def.data(:,2),def.data(:,1),'x')
feplot(mo1,def)
[r1,i1]=unique(def.data(:,2),'first');
Ek=fe_caseg('enerm -bygroup',mo1,fe_def('subdef',def,i1));
if norm(Ek.Y*4-1)>sqrt(eps); error('Inconsistent energy');end
if 1==2 % manual verification
  [m,k,mdof]=fe_mkn1(mo1,'assemble NoT');
  def=feutilb('placeindof',mdof,def);
  feutilb('dtkt',real(def.def),m)+feutilb('dtkt',imag(def.def),m)
end
```

Toolbox tutorial

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3.1. ROTOR MESHING

ferrotor module can use 2 different frames to describe rotating effects (rotating frame and fixed frame). Rotors can also be describes at 3 levels of modelization : 1d, 2d or 3d.

The next section illustrates meshing capabilities, supported computations are described next

- Frequency domain analyses (section 3.4) : Campbell diagram building, Direct computation of critical speeds, response to unbalanced mass, asynchronous load, and harmonic loads on bearings.
- Time domain simulation accounting for non-linear bearings is under development (these are performed with the shaft in a rotating frame and the stator fixed).

3.1 Rotor meshing

3.1.1 Meshing utilities

1D

The SDT/Rotor toolbox supports analysis of 1D models of symmetric rotors composed of

- shafts represented by **beam1** element (see sdtweb('beam1') and sdtweb('p_beam')). The rotation axis is taken to be that of the beam.
- disks represented by mass1 (see sdtweb('mass1')). The rotation axis is the one whose moment of inertia is different from the 2 others that are equal.
- bearings supports by celas.

You can generate a beam model of your rotor by providing a skyline (points not on the axis defining the radius at various locations). Use NaN to define 2 segments. See rotor1d Skyline for more details.

```
xy=[0 0; 0 .1;1 .1;1 1;1.1 1;1.1 .1;2 .1];
xy(:,3)=0; figure(1);plot(xy(:,1),xy(:,2));
% Mesh as beams
mo1d=rotor1d('skylineToBeam',xy);
% Add bearings as spring elements:
```

```
mold=rotor1d('AddBearing DOF -123 k 1e4 -keep',mold,[0.1 0 0]);
mold=rotor1d('AddBearing DOF -123 k 1e4 -keep',mold,[1.9 0 0]);
```

```
% now view as 3D model
mo2d=rotor1d('1To2d lc 5e-2',mo1d);
cf=feplot;cf.model=rotor2d('buildFrom2D nsec16',mo2d);
```

You can also add beam through rotor1d Addbeam command.

```
mold=rotor1d('AddBeam x1 0 x2 0.7 r2 0.77 MatID 1',[]); % add rod
mold=rotor1d('AddBeam x1 0.5 x2 0.8 r1 0.77 r2 0.90 MatID 1',mold); %add tube
mo2d=rotor1d('1To2d lc 5e-2',mold);
cf=feplot; cf.model=mo2d; fecom colordatapro;
```

2D

For 2D rotor representations, the SkyLineTo2d command eases the generation of simple rotors. Note how in the following example Nan separators are used to generate a rotor in multiple parts : center shaft first, then after the separator various disks.

```
xy=[ 0 0;0 0.00945;0.0088 0.00945; 0.0088 0.0057;
0.042 0.0057:0.042 0.00938: 0.057 0.00938:
0.057 0.008; 0.0637 0.008; 0.0637 0.00595;
0.0919 0.00595;0.0919 0.00925; 0.0979 0.00925;
0.0979 0.006;0.121 0.006; 0.121 0.008;
0.127 0.008;0.127 0.0095; 0.142 0.0095;
0.142 0.006; 0.175 0.00565; 0.226 0.00565; 0.226 0.0088;
0.232 0.0088;
NaN 0.0057; % Disk
0.00952 0.0057;0.00952 0.008;
 0.0209 0.008; 0.0209 0.0369;
 0.0242 0.0369; 0.0242 0.0572; 0.0266 0.0572; 0.0266 0.0369;
 0.0299 0.0369; 0.0299 0.008; 0.0413 0.008
NaN .00565; % Separator giving non zero internal diameter .00565
0.177+.01 0.0057; 0.177+.01 0.02; 0.191 0.02; 0.191 0.0057;
NaN .00565; % Separator giving non zero internal diameter .00565
0.197+.005 0.0374; 0.215-.005 0.0374; % Impeller
0.215-.005 0.006+.003;0.222-.002 0.006+.003; % between sub disks
0.222-.002 0.0775; 0.2259 0.0775;
];
xy(:,3)=0;
```

```
mo2d=rotor1d('skyline To2d -lc .005',xy); % x axis
mo2d.pl=[ ...
    1 fe_mat('m_elastic',1,1) 20000000000 0.29 7800
    2 fe_mat('m_elastic',1,1) 7.17e10 0.33 2830 %2830
    3 fe_mat('m_elastic',1,1) 3.07e12 0.3 .78 % 7800
];
mo2d.Elt(feutil('findelt matid 2 3 4 5',mo2d),5:6)=2;
mo2d.Elt(feutil('findelt innode {x>=.15 & y<=.0057}',mo2d),5:6)=3;
mo2d.Elt(feutil('findelt innode {x>=.22 & y<=.01}',mo2d),5:6)=3;
feplot(mo2d); fecom colordatamat
cf=feplot;cf.model=rotor2d('buildFrom2D nsec16',mo2d);
cf.sel={'-disk1', 'ColorDataMat'};fecom(';view1')
```

Figure 3.1: Example rotor generated with meshing utilities

3.1.2 Basic 1D rotor example

Following example builds by hand a simple 1D rotor, with one shaft, one disk and 2 bearing stifnesses. It is almost the same as one accessible through d_rotor TestShaftDiskMdl.

```
model=fe_case(model,'fixdof','Ends',...
    [1.01;1.02;1.03;4.01;4.02;4.03]);
% Assemble nominal matrices:
model=fe_caseg('assemble -reset -secdof -matdes 2 1 70',model);
cf=feplot(model);
% For solution see sdtweb('freqstud')
```

Note that at this time only fixed frame representation is available for such 1d rotors (beam1 and mass1 elements). Gyroscopic coupling is then computed under MatType 70 (The formula for gyroscopic coupling can be found in [6]) The nominal representation for these models is then the Eulerian point of view where the displacement of the rotor is expressed in a non-rotating frame. For the 1D representation, the model nodes are always placed on the nominal rotation axis. Thermal and pre-stress effects are not accounted for.

3.1.3 Meshing 3D rotor from 1D and 2D models

The rotor module supports all 3D elements of SDT. 2D models are only considered through an extrusion and 3D cyclic symmetry. One can import a volume model, or mesh it using feutil meshing commands. This section describes procedures to mesh volumes from 1D and 2D models.

From 1D rotor model meshed using beam1 elements, one can create a 2D model using rotor1d 1To3D command. Note that mass1 elements can not be converted for the moment. For example, with section ?? rotor model

```
xy=[0 0; 0 .1;1 .1;1 1;1.1 1;1.1 .1;2 .1];
mold=rotor1d('skyline ToBeam',xy); % Mesh as beams
% Add bearings as spring elements:
mold=rotor1d('AddBearing DOF -123 k 1e4 -keep',mold,[0.1 0 0]);
mold=rotor1d('AddBearing DOF -123 k 1e4 -keep',mold,[1.9 0 0]);
mdl3d=rotor1d('1To3d-quad-lc0.02-div24',mold); % build 3d rotor
cf=feplot(mdl3d)
```

Command option -quad force the use of quadratic elements (hexa20) instead of linear elements (hexa8).

Shaft is meshed using hexa8 degenerated elements. The edges on the axis of the shaft are using the same nodes as the beam nodes so that bearings described by celas in 1D initial rotor can remain the same. RBE3 rings are created at each bearing celas. Concentrated masses (no inertia) on the axis are also left.

THIS IS NO LONGER TRUE BUT SHOULD BE REACTIVATED XXX: mass1 elements with inertia that represents disks, are meshed as volume disks of arbitrary thickness (dt) 0.005, with radius R_2 computed so that inertia along rotation axis is the same $(Ir = 0.5m(R_2^2 + R_1^2))$ and then density

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is computed to match the mass m $(m = \pi * (R_2^2 - R_1^2) * dt)$. (R_1 is shaft radius). Young modulus of disk is taken at 100*steel modulus.

Gyroscopic coupling and centrifugal stifness for 3D elements are only described in the rotating frame, that is to say under MatType 7 and MatType 8.

```
mdl3d=fe_mat('defaultil',mdl3d); % default element pro
mdl3d=fe_mat('defaultpl',mdl3d); % default material pro
% Assemble nominal matrices:
mdl3d=fe_caseg('assemble -reset -secdof -matdes 2 1 7 8',mdl3d);
```

One can also find an example of a 3D rotor in d_rotor('TestVolShaftDiskMdl').

```
data={'Rs' ,0.01 , 'shaft radius';
    'Rd' ,0.15 , 'disk radius';
    'Ls' ,0.4 , 'shaft length';
    'Ld' ,0.03, 'disk thickness';
    'yd' ,0.4/3, 'disk position on the shaft'
    'yb' ,0.4*2/3,'bearing stifness position on the shaft'
    'Tol' ,0.05, 'elt length Tol'
    };
model=d_rotor('TestVolShaftDiskMdl',data);
```

rotor2d BuildFrom2D convert 2d model to 3d model using cyclic symetry. For example, to convert previous 2D model:

```
mdl2d=rotor1d('1To2d-quad-lc0.02',mo1d);
mdl3d=rotor2d('buildFrom2D -close nsec3 div8',mdl2d);
```

3.1.4 From sector to shaft in the case of cyclic symmetry

Closing a disk

The SDT fe_cyclic function only handles single sector models using symmetry conditions. SDT/Rotor extends the capabilities by dealing with a full (possibly multi-stage, then called shaft) rotor model.

Building a disk/shaft model is done in two steps. For each sector, one defines matching edges using a fe_cyclic Build command, the generates a disk/shaft model using fe_cyclicb DiskFromSector.

```
demosdt(['download-back donnees_secteur.dat'])% xxx missing patchfile
% -removeface : removes skin elements
samcef('read-removeface',fullfile(sdtdef('tempdir'),'donnees_secteur.dat'));
```

```
cf=feplot;
% Fix sector edges
fe_cyclic('buildepsl .1 fix',cf.mdl);
fecom('curtabCases','Symmetry')
fecom(';view3;proViewOn');
```



Figure 3.2: Left and right sides of the sector

The left and right edges of sectors should be conform ! Practically you need to mesh the two surfaces first (possibly generate the second by rotation of the first). Then mesh the interior. You will note that this typically requires at least two layers of element for tetra10 meshing.

3.1.5 Utilities for handling slanted blades

Typically blades have an angle with respect to the e_r, e_z plane. In a number of cases, removing this angle makes node and element selection easier. With two nodes the FixTheta command modifies node positions by removing the angle. The transformation is saved in info,FixTheta stack entry and the back transformation is obtained using a FixTheta call with no argument.

```
cf=feplot(2);cf=d_rotor('TestSector');%sdtweb('d_rotor.m#TestSector')
n1=cf.mdl.Node; % save nodes
cg=feplot(5); % place rectified model in figure(5)
cg.mdl=fe_cyclicb('MeshFixTheta 10061 10086 -offset 80',cf.mdl);
fecom(cg,'view2')
% verify that the back step works
cf.mdl=fe_cyclicb('MeshFixTheta',cg.mdl);fecom(cf,'view1')
if norm(cf.mdl.Node-n1)>1e-10; error('Back transform failed');end
```

3.1.6 Disk connections in multi-stage cyclic symmetry

In multi-stage cyclic symmetry, each stage is modeled using a superelement called disk*i* (see fe_cyclicb DiskFromSector). Coupling between stages is done using elements. The most consistent approach is to use a physical area that is properly meshed for the full 360 degrees, but this may be difficult in particular when the mesh refinement is notably different between the two stages, so that a node to surface penalized connection is also implemented and an example given at the end of this section.

Two steps are required:

- 1. the first step is a manual declaration of the nodes that belong to the two regarding surfaces. The declaration of the nodes in the 2D cut provided by fe_cyclicb DisplayAllEdges command is sufficient (and therefore recommended).
- 2. the second step is the automatic reconstruction of the rims as volumes (using automated 3D tessellation) or **penalty** based node to surface bilateral contact in the MeshRim command.

Note that **fe_shapeoptim** can be used to local deform the disk in order to allow rim meshing.

```
% load two disk example with space between disks
cf=demo_cyclic('TestForMeshRimVol')
% RimStep1: select nodes at the matching interfaces
fe_cyclicb('DisplayAllEdges',cf);
% start cursor to pick values : fe_fmesh('3dlineinit')
n1=[12 18 24 1127 1133 1139];
% RimStep2: build rims and tessellate
fe_cyclicb('MeshRimStep2 epsl.1',cf,n1);
```

```
cf.sel='EltName~=SE';fecom('showpatch');
```



Figure 3.3: Selected disk edge nodes for rim tesselation

If you choose the **penalty** approach here is a working example where the edges of sector edges are assumed coincident thus allowing an automated search of intersection with the -FindIntersect option.

```
% load two disk example with space between disks
cf=demo_cyclic('testrotor 7 10 3 -NoRim -RimH .1 -blade -cf 2 reset');
% RimStep1: find nodes at the matching interfaces
n1=fe_cyclicb('DisplayAllEdges -FindIntersect epsl .2',cf);
% RimStep2: build rims as springs
fe_cyclicb('MeshRimStep2 epsl.1 -kp 1e12 -slavedisk 1 3 -masterdisk 2',cf,n1);
cf.sel={'groupall','colordatagroup -edgealpha .05 -alpha.1'}
def=fe_cyclicb('shaft eig 0',cf.mdl);
sel={'disk1','groupall';'disk2','groupall';'disk3','groupall';'rim',''};
cf.def=fe_cyclicb('DisplaySel',cf,def,sel);
fecom('ColorDataEvalTanz')
```

3.1.7 View meshes for cyclic symmetry

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Figure 3.4: Sample viewing mesh for post-processing multi-stage computations

Meshing tools also include procedures to build viewing meshes from the finite element mesh. fe_cyclicb(' aims to build viewing meshes made of surface elements connecting selected nodes of the true mesh of the rotor. sel can be:

- an array of lines connecting nodes of the inner disk parts in the 2D cut of the rotor (returned by a previous fe_fmesh('3dLineInit')). Note that this function automatically removes the segments that connect nodes of two different disks.
- an array of elements connecting nodes of the blades of each disk (returned by a previous fe_fmesh('3dLineInitAddInfo Quad4')).

The selection is stored in the cf.mdl.Stack{'info', 'ViewMesh'}. If the -reset option is not specified, the current selection is appended to the existing one. For each project, you should typically edit a script similar to the following

```
cf=demo_cyclic('BuildStepO');
% Viewing mesh step 1: disk elements
fe_cyclicb('DisplayAllEdges',cf);
%fe_fmesh('3dLineInit',cf); % right click 'Type' or 'Done'
```

```
L=[1 3 5 15 26 0 1121 1123 1125 1135 1146 0 13 15 18 1133 1135 1138];
fe_cyclicb('MeshRimLine2Patch -reset', cf,L);
% Viewing mesh step 2: blade elements
% disk1
fe_cyclicb('DisplayFirst', cf, {'disk1'});
%fe_fmesh('3dLineInitAddInfo Quad4',cf); % pick four nodes to form a quad
% use right click 'Type' or 'Done' to display
Elt=[Inf abs('quad4');
154 156 152 148 1 1;148 146 150 154 1 1;
146 142 144 150 1 1;142 111 83 144 1 1];
fe_cyclicb('MeshRimLine2Patch', cf, Elt);
% disk2
fe_cyclicb('DisplayFirst', cf, {'disk2'});
fe_fmesh('3dLineInitAddInfo Quad4',cf); % pick four nodes to form a quad
model.Elt=[Inf abs('quad4');
1274 1276 1272 1270 1 1;1270 1272 1268 1266 1 1
1266 1268 1264 1262 1 1;1262 1231 1203 1264 1 1];
fe_cyclicb('MeshRimLine2Patch', cf, model.Elt);
fecom('ShowPatch');
% save cf.Stack{'info','ViewMesh'}
% Once the geometry generated typical calls are
fesuper('Sebuildsel -initrot', cf, cf.Stack{'info', 'ViewMesh'})
fe_cyclicb('Displaysel 2',cf,def,cf.Stack{'info','ViewMesh'})
```

3.2 Bearing and support representations

3.2.1 Linear bearing

One can simply represent a support or a bearing by a linear spring that is to say a celas element (for more details see sdtweb('celas')). A celas element is announced in the model element matrix by the header [Inf abs('celas')] and format is as follow: [NodeId1 NodeId2 -DofID1 DofID2 ProID EltID Kv Mv Cv] NodeId1 and NodeId2 define between which nodes celas is connected (if NodeId2=0 celas is grounded), DofID1 and DofID2 which dof are connected, and Kv Mv and Cv give respectively stiffness mass and damping. A typical bearing for a rotor turning around x will then

be defined by



Figure 3.5: Simple disk on long rod model.

The following generates a spring bearing connected to node n1, stiffness is 1e9 and damping 1e2 in the xy plane. The use of a spring property is necessary for non linear time domain applications

```
model=d_rotor('TestDiskLongBeam');
n1=feutil('findnode z==.71 & x==0 & y==0',model);
```

```
model=feutil('AddElt',model,'celas',[n1 0 -12 0 100]);
model.il=p_spring(model.il,'dbval 100 1e10 0 1e2');
fe_simul('mode',model)
r1=struct('Origin',[20 10 0],'axis',[0 0 1],'radius',3, ...
'planes',[1.5 1 111 1 3.1;
5.0 1 112 1 4;], ...
'MasterCelas',[0 0 -123456 123456 10 0 1e14], ...
'NewNode',0);
links={'connection','bearing',...
struct('C1',1,'C2',r1,...
'link',{{'celas',[0 0 23 0 103 0 1e5 0 0.1]}},...
'NLdata',struct('type','RotCenter','lab','Bearing',...
'sel','ProId 1','JCoef',1,'drot',n1+.06))}
[model,RunOpt.idof]=nl_spring('ConnectionBuild',model,links)
```

[cf.mdl,RunOpt.idof]=nl_spring('ConnectionBuild',cf.mdl,links);

For 3d rotors, if there is no node on the axis of the shaft where one want to model the bearing, one can add a node whose displacement depends on displacements of nodes on a ring of the shaft using (see sdtweb('fe_case#connectionscrew')). Then one can connect a grounded celas to this node. xxx a full example is needed xxx

3.2.2 Non-linear bearings in the time domain

Documentation of nl_spring capabilities should go here.

One has a particular example, with a simple rotor (beam1+mass), non linear bearings (not the same in y and z direction), using the rotcenter non linearity, spurious mass for rotation DOF (no basis transformation). The strategy to have different NL according to Y and Z is to add a mass connected to the rotcenter extremity usually connected to stator, and to use this mass motion in global basis to apply NL bearing in the 2 different directions Y and Z. See sdtweb d_rotor rotcenter.

3.3 Gyroscopic effects

SDT supports gyroscopic matrices in both rotating (body-fixed) and fixed frames (Eulerian representation of an axisymmetric structure). When performing assembly, the matrices in the local rotating frame are obtained with MatType 7 gyroscopic and MatType 8 centrifugal softening. MatType 70 corresponds to the gyroscopic matrix in the global fixed frame. There is no centrifugal softening in this frame.

3.3.1 Fixed frame models

The mass1 and beam1 elements gyroscopic matrices are only available in the global fixed frame (MatType 70). The rotation axis is assumed to be the axis of the beam for beam1 elements. Moments of inertia must be equal (axisymmetry). For mass1 elements the rotation axis is assumed to be the one whose rotation inertia is different from the 2 others that must be equal.

mass1 .Elt format: [n mxx myy mzz ixx iyy izz EltId]

One can build simple models of 1d rotor using mass1 elements to represent rigid disk and beam1 to represent the shaft. One can find an example of such a rotor in d_rotor('TestShaftDiskMdl'). See section ?? for more details on how to mesh such a rotor.

For volume and shell elements, the formulation of gyroscopic matrices in global fixed frame is unclear and thus not currently implemented.

3.3.2 Rotating frame models

For all volume elements, one can compute gyroscopic (MatType 7) and centrifugal softening (MatType 8) matrices in the local rotating frame.

Elements under development are

- mass1 : Only point masses with same mass along the 3 translation DOF and no rotation inertia are considered. Other are ignored.
- beam1 : Not supported.
- **shell** : Not supported.

In that case rotation axis must be given as a vector in info,Omega stack entry in the model. For example model=stack_set(model,'info','Omega',[0 1 0]) will define a rotation axis along Y. Note that the norm of this vector is assumed to be the rotation speed in rad/s. The norm of the rotation vector should be 1 so that matrices are assembled for a rotation speed of 1 rad/s. Indeed

it is assumed that gyroscopic and centrifugal softening matrices have been assembled for a rotation speed of 1 rad/s in many functions (such as fe_rotor etc. ...).

Finer definition of the rotation speed is possible using a **struct** input with fields

- data the rotation axis defined as described above.
- unit optional, definiton of the rotation speed unit as a string, either rad/s or RPM.
- group optional, to apply gyroscopic coupling only to specified element groups.

The definition of the group field can be handled by fe_cyclic('OmegaGroup', model,findElt,[0 1 0]'. The findElt argument can be either a FindElt string or a vector of EltId values. In both cases, gyroscopic coupling will be applied to the groups containing the element founds, with the rotation axis [0 1 0] in this example. Warning: the whole group will be impacted even if the findElt input only selects parts of it.

3.4 Frequency domain analysis, full model

Supported computations in fe_rotor, are

- Campbell diagram building
- Direct computation of critical speeds
- Frequency domain response to unbalance and harmonic loads on bearings
- Time domain simulation accounting for non-linear bearings is under development (these are performed with the shaft in a rotating frame and the stator fixed).

3.4.1 Campbell diagrams, full model

Campbell diagrams are implemented in the fe_rotor Campbell command.

```
model=d_rotor('TestShaftDiskMdl');
fe_rotor('Campbell -crit -cf100',model,linspace(0,8000,100));
figure(100);set(gca,'ylim',[0 200])
```

Command options are -cfi to define figure where to plot, -crit to compute critical speeds, -stability to display a stability diagram (damping-rotation speed).

The following is an example of a simple disk rotating at the end of a long rod. For other details see rotor2d.

3.4.2 Blade with centrifugal stiffening

One considers stiffness matrices that are dependent on the rotation speed. Assuming that a second order polynomial representation is sufficient, one can build a vector of weighing coefficients

$$\left\{ \begin{array}{c} \alpha_1\\ \alpha_2\\ \alpha_3 \end{array} \right\} = \left[\begin{array}{ccc} 1 & 1 & 1\\ \Omega_1 & \Omega_2 & \Omega_3\\ \Omega_1^2 & \Omega_2^2 & \Omega_3^2 \end{array} \right]^{-1} \left\{ \begin{array}{c} 1\\ \alpha_2\\ \alpha_3 \end{array} \right\}$$
(3.1)

Such that the stiffness at rotation speed Ω is approximated by

$$[K(\Omega)] = \sum_{\alpha_i} [K\Omega_i] \tag{3.2}$$

The zCoef uses velocity Omega in rad/s.

This example now treats computation at variable rotation speeds

```
% Model Initialization
model=demo_cyclic('testblade');cf=feplot(model);
% Compute matrix coefficients for a multi-stage rotor
range=struct('data',[0 0 1;0 0 8e3;0 0 16e3],'unit','RPM');
% Assembling in the feplot figure, allows memory offload
fe_cyclicb('polyassemble -noT',cf,range);
X=struct('data',linspace(0,16e3,10),'unit','RPM');
fe_rotor('Campbell -cf1',cf.mdl,X);set(gca,'ylim',[0 3000])
```

Now a more standard non-linear static computation for a range or rotation speeds.

```
model=demo_cyclic('testblade');cf=feplot(model);
r2=linspace(1e3,15e3,20)';
range=struct('data',r2*[0 0 1],'unit','RPM');
model=stack_set(model,'info','Omega',range);
model=fe_cyclic('LoadCentrifugal',model)
opt=fe_simul('NLStaticOpt');
opt.MaxIter=100;opt.JacobianUpdate=1;
def=fe_time(opt,model);def.data(:,1)=range.data(:,3);
def.LabFcn='sprintf(''%.1f RPM'',def.data(ch,1))';
d_rotor('viewMises',cf,def);
```

3.4.3 Complex modes

To compute complex modes at a given rotation speed, one can use **fe_rotor** Campbell command with option -cmode. Complexe modes are the computed using a real mode basis projection. Mode options should be stored in the 'info' 'EigOpt' model stack entry. The complexe modes are return for the first rotation speed given as input argument.

Once modes are computed, you can display them in feplot. The **fecom** ShowTraj command displays trajectories. For example:

```
model=d_rotor('TestShaftDiskMdl')
model=fe_caseg('assemble -SE -secdof -matdes 2 1 70',model); % assemble model matrices
def=fe_rotor('Campbell -cmode -cf100',model,100); % C modes at 100 RPM.
cf=feplot(model,def); fecom('view3');iimouse('resetvie');
fecom(cf,'ShowTraj')
```



Figure 3.6: Complex mode and trajectory.

3.4.4 Forced frequency response to unbalanced load

One can compute the frequency response to an unbalanced load, or to an asynchronous load using **fe_rotor** RotatingLoad and ForcedResponse command (see sdtweb('fe_rotor') for more details). Definition of the load is different in local rotating frame or global fixed frame. One can see the validation part of this document for various examples. Following example computes the response to an unbalanced mass on the 1D rotor model:

```
model=d_rotor('TestShaftDiskMdl')
model=fe_caseg('assemble -SE -secdof -matdes 2 1 70',model); % assemble model matrices
mb=1e-4; db=0.15; % mass, distance to axis
om=linspace(0,6000,201); % RPM
model=fe_rotor(sprintf('RotatingLoad 2 %.15g -90 2',mb*db),model); % unbalanced mass
r1=struct('Omega',om/60*2*pi,'w',om/60*2*pi); % Range
model=stack_set(model,'info','Range',r1);
R1=fe_rotor('ForcedResponse',model); % compute forced response
iiplot(R1) % plot response
```

3.5 Solvers for models with cyclic symmetry

The SDT/Rotor module contains

- classical cyclic symmetry solvers, where one assumes the solution to be associated with a specific number of diameters (spatial harmonic associated to the Fourier transform of a periodic geometry, see [5] for more details.)
- full rotor reduced models where cyclic symmetry solutions are used to build a reduced model for various stages. The associated solvers are discussed in section 3.6 .

3.5.1 Static response

Resolution of static responses is performed using fe_cyclicb ShaftSolve. You should be aware, that non-linear static iterations may fail to converge if you have rigid body modes in your system. In the example below, this is the reason for fixing the axial motion of point 136 in disk1 and using the -FixTan which adds a tangential translation constraint on the first disk.

```
if 1==1 % One disk case
    if ishandle(2);delete(2);end
```

```
cf=demo_cyclic('testrotor 7 -blade -cf 2');
 Sel={'disk1','groupall'};
else % Two disk case
 cf=demo_cyclic('testrotor 7 10 -blade -cf 2 reset');
 Sel={'','EltName SE';
      'disk1(1:2)', 'groupall'; 'disk2(1:3)', 'groupall'};
end
cf.Stack{'disk1'}=fe_case(cf.Stack{'disk1'}, 'FixDof', 'Axial', 136.03);
% Linear static response
cf.Stack{'info','Omega'}=struct('data',250,'unit','RPM');
d0=fe_cyclicb('shaftsolvestatic 0 -FixTan',cf.mdl)
d_rotor('viewMises');
% Non-linear static response
cf.Stack{'info', 'Omega'}=struct('data', 250, 'unit', 'RPM');
d0=fe_cyclicb('shaftsolvestatic 0 -FixTan -nlstep 1e-10',cf.mdl)
fe_cyclicb('displaySel',cf,d0,Sel)
fecom('ColorDataEvalRadZ')
```

XXX example with thermal loading xxx

Example with temperature dependent properties.

3.5.2 Single stage mode computations

As a first example one will consider computations of a single disk using mono harmonic solutions

Call shaftTeig in fe_cyclicballows to compute the specified mono-harmonic normal modes.

Mono-harmonic normal modes are recovered to the rotor with the help from command shaftdispay. If command option sel is specified and a selection of elements is provided, the shapes are recovered to that selection only. Selections are cell arrays with the typical entry Sel={'disk1', sel1} where sel1 is either a string (to select a subset of elements in the true mesh) or a list of elements (to build a reduced viewing mesh).

```
% Model Initialization
cf=demo_cyclic('testrotor 7 -blade -cf 2 reset');
% Mono-harmonic Solutions
```
```
model=stack_set(cf.mdl,'info','EigOpt',[5 5 -1e3 11 1e-8]);
def=fe_cyclicb('shaft Teig 0 5',model);
```

```
% partial display and computation of strain energy
Sel={'disk1','withnode {r<130}'};
fe_cyclicb('Displaysel',cf,def,Sel,'enerkdens');
if 1==2 % total display if needed
cf.sel='groupall';cf.def=fe_cyclicb('Display',cf,def);
end
```

```
cf.Stack{'info','StressCritFcn'}='fe_cyclicb(''StressRR'');'
[dfull,STRESS]=fe_cyclicb('Displaysel',cf,def,Sel,'stress-gstate');
z=STRESS.GroupInfo{1,5};figure(1);plot(squeeze(z.Y(1,1,:,7)))
```

This will be extended to full disk computations in section ?? .

3.5.3 Multi-stage harmonic mode computations

Call shaftTeig in fe_cyclicb allow to compute the specified mono-harmonic solutions (multi-stage solutions for which disks share the same Fourier harmonic coefficient δ) in a single job.

Mono-harmonic eigensolutions are recovered using Dispay. If command option sel is specified and a selection of elements is provided, the shapes are recovered to that selection only. Selections are cell arrays with the typical entries:

- Sel={'disk1', sel1} where sel1 is either a string (to select a subset of elements in the true mesh) or a list of elements (to build a reduced viewing mesh),
- Sel={'', selg} where selg is a string that selects elements of the global mesh (selg is often 'eltname==SE' so that only disks are displayed).

In the same fashion, mono-harmonic static responses are returned by ShaftSolveStatic. This is of particular interest to compute the static deformation under the centrifugal loading (known to have a Fourier harmonic coefficient of $\delta = 0$) and built with command LoadCentrifugal within fe_cyclic.

```
% Model Initialization
cf=demo_cyclic('testrotor 7 10 -blade -cf 2');
```

```
model=stack_set(cf.mdl.GetData,'info','EigOpt',[5 5 -1e3 11 1e-8]);
def=fe_cyclicb('shaft Teig 0 2',model);
demo_cyclic('RefcheckDisk710',def) % non regression check
cf.Stack{'disk1'}=fe_case(cf.Stack{'disk1'}, 'park', 'blade', 'innode{r>=200}');
Curve=fe_cyclicb('fourier 1:13 -mono -egyfrac',cf,def); % check energies
iiplot(Curve);colormap(flipud(hot));
cf.def=fe_cyclicb('Display',cf,def);
% static responses : sdtweb('freqcyclic#cyclic_static')
model=stack_set(model,'info','Omega',struct('data',1000,'unit','RPM'));
model=stack_set(model,'curve','StaticState', ...
  fe_cyclicb('shaftsolvestatic 0',model));
fe_cyclicb('DisplayFirst',model, ...
  stack_get(model,'curve','StaticState','get'));
d_rotor('viewradz')
% Pre-stressed modes
dp=fe_cyclicb('shaft Teig 0 2',model);
[def.data dp.data]
```

After full rotor assembly restitution is performed using _SeRestit.

3.5.4 Campbell diagrams

First example of the beam with single disk

```
% Model Initialization
model2D=rotor2d('test simpledisk -back');
cf=feplot(rotor2d('buildFrom2D',model2D));
SE=cf.Stack{'disk1'}; % enforce boundary cond. on sector and assemble
SE=fe_case(SE,'FixDof','Base','z==1.01'); cf.Stack{'disk1'}=SE;
```

```
% Compute matrix coefficients for a multi-stage rotor
range=struct('data',[0 0 1; 0 0 1000/2/pi*60],'unit','RPM');
fe_cyclicb('polyassemble -noT',cf,range);
```

```
% Now run a mono-harmonic computation returning reduced model
cf.Stack{'info','Omega'}=struct('data',range.data(1,:),'unit','RPM');
```

```
cf.Stack{'info','EigOpt'}=[5 20 0]; % define eigenvalue options
MVR=fe_cyclicb('shafteig 1 -ReAssemble 2 -NoN -buildMVR',cf);
MVR.Stack{end}(end+[-1:0],4)={'1';'0'}; % skip problem with geometric softening
rc=struct('data',linspace(0,8000*2*pi,50),'unit','RPM');
hist=fe_rotor('Campbell',MVR,rc);
fe_rotor('plotCampbell',hist,struct('fig',100,'axProp',{{'ylim',[0 250]}}))
```

Another example will be needed to treating the multi-stage case. This example needs further validation and rewrite.

```
% Model Initialization
model=demo_cyclic('testrotor 7 10 -blade');
cf=feplot(model);
% Compute matrix coefficients for a multi-stage rotor
range=struct('data',[0 0 1;0 0 800;0 0 1600],'unit','RPM');
% Assembling in the feplot figure, allows memory offload
model=fe_cyclicb('shaftRimAsSe',model); % Needed for PolyAssemble
fe_cyclicb('polyassemble -noT',cf,range);
% Now run a mono-harmonic multi-speed computation
cf.Stack{'info', 'Omega'}=struct('data',range.data(1,:),'unit','RPM');
cf.Stack{'info','EigOpt'}=[5 20 -1e3]; % define eigenvalue options
MVR=fe_cyclicb('shafteig 1 -ReAssemble 2 -NoN -buildMVR',cf);
%MVR.Stack{end}(end+[-2:0],4)={'1';'0';'0'}; % skip problem with geometric softening
rc=struct('data',linspace(1,8000,50),'unit','RPM');
hist=fe_rotor('Campbell',MVR,rc);
fe_rotor('plotCampbell',hist,struct('fig',100,'axProp',{{'ylim',[0 3e3]}}))
%Sel={'disk1','groupall';'disk2','groupall'};
```

```
%fe_cyclicb('Displaysel',cf,def,Sel)
```

3.5.5 Complex modes

Need documentation here.

3.5.6 Forced frequency response to unbalanced load

Need documentation here.

3.6 Full rotor model from cyclic computation

3.6.1 Single stage full rotor example

Starting from the mono-harmonic computation in section 3.5.2. One builds a full shaft model that will allow prediction of all the modes.

```
%H5.close;fclose all; % may be needed for overwrite
cf=demo_cyclic('testrotor 7 -blade -cf 2');
cf.Stack{'info', 'Omega'}=struct('data', 250, 'unit', 'RPM');
d0=fe_cyclicb('shaftsolvestatic 0',cf.mdl) % auto-display with no arg
cf.Stack{'curve','StaticState'}=d0;
RunOpt.Root=fullfile(sdtdef('tempdir'), 'Disk_7');
RunOpt.FileSe=fullfile(sdtdef('tempdir'), 'Disk_7_SE');
cf.mdl=fe_cyclicb('shaftSeAssemble -reset',cf.mdl,RunOpt.FileSe);
RunOpt.FileTeig=fullfile(sdtdef('tempdir'), 'Disk_7_TEIG.mat');
% See sdtweb('fe_cyclicb#ShaftEig') for -batch option
def=fe_cyclicb('shaft Teig 0 1 5',cf.mdl);
% Now build a multi-harmonic model
fe_cyclicb('ShaftPrep -handle',cf,def);
fesuper('fassemble',cf);
% Mode Computations
defr=fe_eig(cf.Stack{'MVR'}, [5 50 1e3 11 1e-8]);
cf.sel='reset';cf.def=fesuper('sedef',cf,defr);
def_ext=fe_cyclicb('DefList',RunOpt.FileTeig(1:end-4))
\% the same results should be achived by assembling the prestress inside
```

% shaftTeig xxx update matrices only once, not for each diameter

```
cf=demo_cyclic('testrotor 7 -blade -cf 2');
cf.Stack{'info','Omega'}=struct('data',250,'unit','RPM');
cf.mdl.il=[1001 fe_mat('p_super','SI',1) 1 0 0 1];
fe_cyclicb('shaft Teig 0 1 5 -batch -reassemble',cf.mdl,RunOpt.FileTeig);
def_int=fe_cyclicb('DefList',RunOpt.FileTeig(1:end-4))
```

%if norm(def_int.data(def_int.data(:,1)>1,1)-def_ext.data(def_ext.data(:,1)>1,1),inf)>
%error('ShaftTEig internal and external prestress produce different results'); end;

Note that to build a disk model from a sector, you should use model=fe_cyclicb('DiskFromSector 1', model, sector). xxx Arnaud : need to check adaptation for multi-disk models.

3.7 Time domain analysis

Once rotor model is created, one can perform time computation. **nl_spring** function is very useful in that case. Gyroscopic effects can be treated as non linear load depending on instant value of a model DOF (observation of the rotation speed for example). Links between fixed and rotating parts can also be modelized as non linear loads.

3.7.1 Simple example

Following example simply computes the response to an impact on the disk, assuming that rotation speed is 1000 RPM:

```
model=d_rotor('TestShaftDiskMdl'); % build simple 1D rotor
% Define Time option and other parameters:
opt=fe_time('TimeOpt Newmark .25 .5 0 1e-4 1e4'); % TimeOpt
Range=struct('Omega',1000/60*2*pi);
model=stack_set(model,'info','Range',Range);
opt.matdes=[2 1 70];
model=stack_set(model,'info','TimeOpt',opt);
model=fe_rotor('TimeOptAssemble',model);
% initial impact:
model=fe_case(model,'DofLoad','In',...
struct('def',1,'DOF',2.02,...
'curve',sprintf('TestStep%.15g',opt.Opt(4)*5)));
```

```
% Compute response:
def=fe_time(model); iiplot(def);
```

3.7.2 Gyroscopic effects

Gyroscopic coupling is represented by a load $-\Omega(t).D(\Omega = 1) * V$ where Ω is the rotation speed and D the gyroscopic coupling matrix. That can be applied at each step of time using the DofKuva nl_spring non linearity (see sdtweb('nl_spring')).

An unbalanced mass is a load proportional to rotation speed. In the local rotating frame it can be described using nl_spring DofV non linearity.

Following example deals with the simple 1D rotor and performs a time integration in fixed global frame taking in account the gyroscopic effect, for an initial impact on the disk. Note that the non linear holding of the gyroscopic effect is not necessary here since the global frame is considered, and rotation speed is assumed to be constant (1000 RPM). The gyroscopic effects (in green in figure below) are coupling the y and z motion.



Figure 3.7: y (top) and z (bottom) deflection, with (green) and without (blue) gyroscopic effect.

```
model=d_rotor('TestShaftDiskMdl');
model=fe_case(model,'FixDof','Ends',...
                               [1.01 1.02 1.03 4.01 4.02 4.03 0.01]);
% TimeOpt:
opt=d_fetime('TimeOpt -gamma .501');
opt.NeedUVA=[1 1 1];
opt.IterEnd='if ite>90;evalin(''caller'',''assignin(''''base'''',''''def'''',out)'');e
```

```
opt.Follow=1; opt.RelTol=-1e-5;
opt.Opt(7)=-1; % factor type sparse
opt.Opt(4)=1e-4; opt.Opt(5)=0.6e4; % NSteps
% Initial impact:
model=fe_case(model, 'DofLoad', 'In',...
    struct('def',1e3,'DOF',2.02,...
           'curve',sprintf('TestStep%.15g',opt.Opt(4)*5)));
% Without gyroscopic effect:
def0=fe_time(opt,model) % compute
% With NL gyroscopic effects:
model=stack_set(model,'pro','DofKuva1005', ... % gyroscopic effects
    struct('il',[1005 fe_mat('p_spring','SI',1) 0 0 0 0 0],...
    'type', 'p_spring', 'NLdata', struct(...
        'type', 'DofKuva', 'lab', 'gyroscopic effect', ...
        'Dof', [], 'Dofuva', [0 1 0], 'MatTyp', 70, ...
        'factor',-1*1000/60*2*pi,'exponent',1,'uva',[0 1 0])));
def=fe_time(opt,model) % compute
% Display results as curves
ci=iiplot;
iicom('sub 2 1');
iicom(ci,'IIxOnly',{'disp(1)';'disp(2)'});
i1=fe_c(ci.Stack{'disp(1)'}.DOF,2+[.02;.03],'ind');
iicom(ci,sprintf(';cax1;chc%i;cax2;chc%i',i1(1),i1(2)));
% comgui('ImWrite',comgui('cd','o:\sdt.cur\rotor\plots\shaftdisk_withandwithoutgyro.pn
% display deformation
cf=feplot(model); % display model
cf.def=def0; % without gyrocopic effects
cf.def=def; % with gyroscopic effects
```

3.7.3 Other representations of bearings

In the fixed basis the simplest bearing model is a linear spring (celas element. In rotating basis, using a spring to model bearing is not possible using a celas element. The nl_spring RotCenter non linearty is useful to modelize link between rotating and non rotating parts.

In the fixed basis one can use a non linear spring to model the bearing. The **nl_maxwell** non linearity describes a set of generalized maxwell rheological models that can be used as non linear bearing in

the global frame. As an illustration, following example replaces the linear bearing of the 1D rotor by a zener model of the link.

```
model=d_rotor('TestShaftDiskMdl'); % Build model
model=fe_case(model,'FixDof','Ends',...
                    [1.01 1.02 1.03 4.01 4.02 4.03 0.01]); % Clamp ends
% TimeOpt:
opt=d_fetime('TimeOpt');
opt.Follow=1; opt.RelTol=-1e-5;
opt.Opt(7)=-1; % factor type sparse
opt.Opt(4)=1e-4; opt.Opt(5)=0.6e4; % NSteps
opt.IterInit='model.FNL0=model.FNL+0;';
% Initial impact:
model=fe_case(model, 'DofLoad', 'In',...
    struct('def',1e3,'DOF',2.02,...
           'curve',sprintf('TestStep%.15g',opt.Opt(4)*5)));
% NL bearings
% define nl_maxwell data
model.il(ismember(model.il(:,1),[100;101]),:)=[]; % remove properties
k0=500000; k1=k0/2; c1=600; % 20.41% for f=54.15 Hz
data=nl_maxwell(sprintf('db Fu"zener k0 %.15g k1 %.15g c1 %.15g"',k0,k1,c1));
data.Sens{2}=3.02; % translation sensor defining nl_maxwell inputs
r1=p_spring('default'); r1=feutil('rmfield',r1,'name');
r1.NLdata=data; r1.il(3)=k0; % translation sensor defining nl_maxwell inputs
r1.il(1)=100; model=stack_set(model,'pro','bearingy',r1);
r1.NLdata.Sens={'trans',3.03}; % translation sensor defining nl_maxwell inputs
r1.il(1)=101; model=stack_set(model,'pro','bearingz',r1);
ci=iiplot(3); % results will be store there
```

```
def0=fe_time(opt,model); % compute
```

Validation

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4.1 Rigid disk example



Figure 4.1: Simple rigid disk.

We consider a simple rigid disk of axis Y, thickness h, radius R, mass m. This simple example is very useful because we can easily compute matrices in both of global and local frame for a simple description of motion with 4 DOF. Besides we can compute in SDT equivalent matrices for a mass1 rigid disk, for a disk described by a beam1 element and for a volume disk in hexa8 elements. Then we can compare gyroscopic matrices and make sure that their implementation for each element is correct.

4.1.1 Matrices in rotating frame

The motion of the disk is described by 2 translations (u_c and w_c) and 2 rotations DOF (θ_u and θ_w). Disk is assumed to be rigid so displacements at each point of the disk are given by the shape functions

$$\left\{ \begin{array}{c} u \\ v \\ w \end{array} \right\} = \left[\begin{array}{ccc} 1 & 0 & 0 & -v \\ 0 & 0 & -w & u \\ 0 & 1 & v & 0 \end{array} \right] \left\{ \begin{array}{c} u_c \\ w_c \\ \theta_u \\ \theta_w \end{array} \right\} = \left[N_{xyz} \right] \{q\}$$
(4.1)

Rotation matrix (2.3) along Y axis is given by

$$[\Omega] = \Omega \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ -1 & 0 & 0 \end{bmatrix}$$
(4.2)

Using matrix expression given in section 2.1.1 in body fixed local rotating frame, and projecting the integrals by assuming rigid disk motion

$$[M] = \begin{bmatrix} m & 0 & 0 & 0 \\ 0 & m & 0 & 0 \\ 0 & 0 & I_u & 0 \\ 0 & 0 & 0 & I_w \end{bmatrix}$$
(4.3)

with $I_u=I_w=\frac{1}{4}mR^2+m\frac{h^2}{12}$

$$[D] = 2m\Omega \begin{bmatrix} 0 & 1 & 0 & 0 \\ -1 & 0 & 0 & 0 \\ 0 & 0 & 0 & \frac{h^2}{12} \\ 0 & 0 & -\frac{h^2}{12} & 0 \end{bmatrix}$$
(4.4)
$$[Kc] = -m\Omega^2 \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & \frac{h^2}{12} & 0 \\ 0 & 0 & 0 & \frac{h^2}{12} \end{bmatrix}$$
(4.5)

$$[M] \{ \ddot{q}_L \} + D \{ \dot{q}_L \} + Kc \{ q_L \} = F_L$$
(4.6)

An unbalanced mass is represented by a static load in the local rotating frame

$$\{F_L\} = \left\{ \begin{array}{c} 1\\0\\0\\0 \end{array} \right\} \tag{4.7}$$

4.1.2 Matrices in global fixed frame

Local frame (indexed with $_L$) is rotating in global fixed frame (indexed with $_G$) according to rotation matrix

$$[R_t] = \begin{bmatrix} \cos(\Omega t) & -\sin(\Omega t) \\ \sin(\Omega t) & \cos(\Omega t) \end{bmatrix}$$
(4.8)

so that we have $\{q_L\} = [R_t] \{q_G\}$ $\{\dot{q}_L\} = [R_t] \{\dot{q}_G\} + \left[\dot{R}_t\right] \{q_G\}$ $\{\ddot{q}_L\} = [R_t] \{\ddot{q}_G\} + 2\left[\dot{R}_t\right] \{\dot{q}_G\} + \left[\ddot{R}_t\right] \{q_G\}$

and equation (4.6) can thus be rewritten as

$$\begin{bmatrix} R^{T}MR \end{bmatrix} \{ \ddot{q}_{G} \} + \begin{bmatrix} 2R^{T}M\dot{R} + R^{T}DR \end{bmatrix} \{ \dot{q}_{G} \} + \begin{bmatrix} R^{T}M\ddot{R} + R^{T}D\dot{R} + R^{T}K_{c}R \end{bmatrix} \{ q_{G} \} = F_{G}$$

$$\begin{bmatrix} M_{G} \end{bmatrix} \{ \ddot{q}_{G} \} + \begin{bmatrix} D_{G} \end{bmatrix} \{ \dot{q}_{G} \} + \begin{bmatrix} Kc_{G} \end{bmatrix} \{ q_{G} \} = F_{G}$$

$$(4.9)$$

One has $R^T \dot{R} = \Omega \begin{bmatrix} 0 & -1 \\ 1 & 0 \end{bmatrix}$ and $R^T \ddot{R} = \Omega^2 \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}$ thus in the gyroscopic effect the translation terms disappear and one has

with $I_v = \frac{1}{4}mR^2 = I_u - 2mh^2/12$ and the xxx UNEXPECTED xxx centrifugal softening in the global frame

An unbalanced mass is represented by a rotating load in the global fixed frame

$$\{F_G\} = \begin{cases} \sin(\Omega t) \\ \cos(\Omega t) \\ 0 \\ 0 \end{cases}$$

$$(4.10)$$

4.1.3 Validation with 3D model disk

This validation example considers a disk meshed with hexa8 volume elements. Local frame matrices are computed and projected on the geometrical rigid body modes (x and z translations, and corresponding rotation). One checks matching of

- matching of theoretical and numerical matrices
- response amplitudes of the disk with unbalanced mass in global and local frame
- mass1 and beam1 elements gives the same gyroscopic matrices in the global frame (mattype 70)

```
% model of disk:
R1=.01;R2=.15;h=0.03;rho=7800;
md=d_rotor(sprintf('testvoldisk -dim %.15g %.15g %.15g 5 24',R1,R2,h))
md.DOF=[];md=fe_caseg('assemble -se -matdes 2 1 7 8 not',md);
% disk assumed to be rigid:
rb=feutil('geomrb',md); cf=feplot(md); cf.def=rb;
md.K=feutil('tkt',rb.def(:,[1 3 4 6]),md.K);
md.DOF=[1.01 1.03 1.04 1.06]';
md.K{2}=0*md.K{2}; % rigid disk
md.K{2}(1,1)=5e5;md.K{2}(2,2)=5e5; % bearing
md.K{2}(3,3)=1e5;md.K{2}(4,4)=1e5; % bearing rot /XXX
% Campbell in local rotating frame:
fe_rotor('campbell-critical',md,linspace(0,9000,31));
% Unbalanced mass:
w=1; t=12; wrange=logspace(0,3,100); Q=[];
for j1=1:length(wrange);w=wrange(j1);
 % rotation matrix
 R=[\cos(w*t) - \sin(w*t); \sin(w*t) \cos(w*t)]; R=[R \text{ zeros}(2); \text{zeros}(2) R];
 Rp=w*[-sin(w*t) - cos(w*t); cos(w*t) - sin(w*t)]; Rp=[Rp zeros(2); zeros(2) Rp];
 Rpp=-w^2*R;
 % Global matrices
 MG=R'*md.K{1}*R;
 DG=2*R'*md.K{1}*Rp + R'*w*md.K{3}*R;
 KG=R'*md.K{1}*Rpp + R'*w*md.K{3}*Rp + R'*w^2*md.K{4}*R + R'*md.K{2}*R;
 % compare to theoretical values
 m=pi*(R2^2-R1^2)*3e-2*7800;
```

```
Iv=0.5*m*(R2<sup>2</sup>+R1<sup>2</sup>); Iu=0.5*Iv+m*h<sup>2</sup>/12;
 DGth=zeros(4); DGth(4,3)=Iv; DGth(3,4)=-Iv; DGth=w*DGth;
 KGth=diag([0 0 1 1]); KGth=-w^2*Iv/2*KGth;
 if norm(KG-R'*md.K{2}*R-KGth)/norm(KGth)>5e-2;error('Cent. soft mismatch');end
 if norm(DG-DGth)/norm(DGth)>5e-2; error('gyro matrix mismatch'); end
 % unbalanced mass:
 XOL=inv(md.K{2}+w^2*md.K{4})*w^2*[1 0 0 0]'; % local
 XOL=abs(XOL(1));
 XOG=inv(-w^2*MG+1i*w*DG+KG) *w^2*[-1i 1 0 0]'; % global
 XOG=abs(XOG(1));
 Q(1,j1)=XOL; Q(2,j1)=XOG;
end
figure(4);semilogy(wrange*60/2/pi,Q(1,:),wrange*60/2/pi,Q(2,:));
legend('local', 'global'); title('Unbalanced mass');setlines
xlabel('Rotation velocity [RPM]'); ylabel('Amplitude [m]')
if norm(Q(1,:)-Q(2,:))/norm(Q(1,:))>1e-8; error('Global/Local error'); end
% check mass1 gyroscopic matrix 70:
mdmass=struct('Node', [1 0 0 0 0 0 0], ...
              'Elt', [Inf abs('mass1') 0; 1 m m m Iu Iv Iu]);
mdmass=fe_case(mdmass,'fixdof','Base',[1.02 ;1.05])
mdmass=fe_caseg('assemble -se -matdes 2 1 70',mdmass)
if norm(w*mdmass.K{3}-DGth)/norm(DGth)>1e-8; sdtw('_err', 'mass1 gyro 70 unmatch'); end
% check beam1 gyroscopic matrix 70:
Omega=[0 1 0]; % axis of the disk
mdbeam1=struct('Node', [1 0 0 0 -h/2*0mega; 2 0 0 0 h/2*0mega],...
               'Elt', [Inf abs('beam1') 0; 1 2 1 1 0 0 0]);
mdbeam1.il=p_beam('convertto1',p_beam(sprintf('dbval 1 TUBE %.15g %.15g',R2,R1)));
mdbeam1.pl=m_elastic('dbval 1 steel');
mdbeam1=fe_case(mdbeam1,'fixdof','fix1',...
 [1+(find(Omega)+[0;3])/100;2+(find(Omega)+[0;3])/100]);
rb=feutil('geomrb',mdbeam1); mdbeam1=fe_case(mdbeam1, 'DofSet', 'fix2',rb);
rb=fe_simul('static',mdbeam1); mdbeam1=fe_case(mdbeam1,'remove','fix2');
mdbeam1.DOF=[]; mdbeam1=fe_caseg('assemble -se -matdes 2 1 70', mdbeam1);
mdbeam1.DOF=fe_case('gettdof',mdbeam1);
rb=feutil('placeindof',mdbeam1.DOF,rb)
mdbeam1.K=feutil('TKT',rb.def(:,setdiff(1:6,find(Omega)+[0 3])),mdbeam1.K)
if norm(w*mdbeam1.K{3}-DGth)/norm(DGth)>1e-8;
 sdtw('_err', 'beam1 gyro 70 unmatch');
```

4.1. RIGID DISK EXAMPLE

end

The response to the unbalanced mass is the same in both of the 2 frames. The maximum of response matches rotation speed found as critical speed for forward whirl in local and global Campbell diagram.

Note that we can compute Campbell in local frame from Campbell in global frame with:

 $f_L^{(F)} = \|f_G^{(F)} - \Omega\| \text{ for forward modes and}$ $f_L^{(B)} = \|f_G^{(B)} + \Omega\| \text{ for backward modes.}$

The 2 Campbell diagrams have been computed using matrices in corresponding frame and we check that we can pass from one to other using these formulas.

XXX In this example:

- Centrifugal softening in global fixed frame? Not 0

- In GLOBAL frame gyroscopic matrix DG do not modify unbalanced response because only translation dof are affected... - In LOCAL frame gyroscopic D do not modify unbalanced response: $XOL=inv(md.K2+w^2*md.K4)*w^2*[1 \ 0 \ 0 \ 0]$; % local



Figure 4.2: Top: Unbalanced mass response amplitude computed in local and global frame. Bottom left: Campbell diagram in local rotating frame, right: in global frame.

4.2 Simple 2DOF model of shaft with disk

This section is about the simple 2 DOF rotor model described in chapter 2 of Lalanne and Ferraris [7].



Figure 4.3: Simple model of rotor.

The disk is rigid, the shaft is described by 2 sin shape functions $f(y) = sin(\frac{\pi y}{L})$ where L is the length of the shaft.

Bearing at $y = \frac{L}{3}$ is represented by 2 additional stiffness and damping. $K_{bearing} = sin(\frac{2\pi}{3}) \begin{bmatrix} kxx & kxz \\ kzx & kzz \end{bmatrix}$ $C_{bearing} = sin(\frac{2\pi}{3}) \begin{bmatrix} cxx & cxz \\ czx & czz \end{bmatrix}$

The 2 DOF are considered in the global fixed frame.

$$\begin{bmatrix} m & 0 \\ 0 & m \end{bmatrix} \{ \ddot{q} \} + \Omega \begin{bmatrix} 0 & -a \\ a & 0 \end{bmatrix} \{ \dot{q} \} + \begin{bmatrix} k & 0 \\ 0 & k \end{bmatrix} \{ q \} = F$$
(4.11)

For an unbalanced mass mb at a position of L/3 on the shaft and at a distance of db of the rotation axis $F = m^* db \Omega^2 \left\{ \begin{array}{c} \sin(\Omega t) \\ \cos(\Omega t) \end{array} \right\}$ where $m^* = \sin(\frac{\pi}{3})mb$.

For a simple asynchronous load of amplitude f0 rotating at $s\Omega F = f0 \begin{cases} sin(s\Omega t) \\ cos(s\Omega t) \end{cases}$.

Frequency analysis is performed.

Harmonic solutions are computed under the form $\Re(Q_0 exp(\imath s\Omega t))$ (s = 1 for unbalanced mass and s <> 1 for asynchronous load).

Implementation of this case can be found in d_rotor Lalanne2DOF. One can define values of pa-

rameters (kxx, czx, m, a, mb, f0, s etc. ...) as fields of a data structure **r1** given as argument : **d_rotor('Lalanne2DOF',r1)**. A campbell and frequency forced response are then computed at the rotation speeds defined (in RPM) in the field .om of the parameter data structure. One can ask only for the model using model=d_rotor('Lalanne2DOFMdl',r1);.

If this first case, no bearing is considered $(K_{bearing} = 0 \text{ and } C_{bearing} = 0)$.



Figure 4.4: Campbell diagram. Symmetric rotor without bearing.



Figure 4.5: Response to an unbalanced load (left) and asynchronous load (right). Symmetric rotor without bearing.

Maximum response for unbalanced mass is obtained for the exact critical rotation speed of forward mode computed in the Campbell (3089 RPM). This speed is the same as in Lalanne theoretical expression.

Bearing stiffness along z is now taken in account so that rotor is not symmetric. kzz = 5e5 and kxx = kxz = kzx = 0. There is no damping in the bearing: $C_{bearing} = 0$).



Figure 4.6: Campbell diagram. Asymmetric rotor with z stiffness bearing.

Now the 2 modes are not at the same frequency for $\Omega = 0$.



Figure 4.7: Response to an unbalanced load (left) and asynchronous load (right). Asymmetric rotor with z bearing stiffness.

Now there are 2 maximums of response that match critical rotation speeds for both forward and backward modes. This is the same for the asynchronous load.

Bearing stiffness along z is taken in account so that rotor is not symmetric. kzz = 5e5, kxx = 1e5 and kxz = kzx = 0. Damping in the bearing is also considered with $cxx = 100\beta$, $czz = 500\beta$ and cxz = czx = 0.

```
beta=15;
r1=struct('om',linspace(0,20e3,201),'s',1,...
'kxx',1e5,'kzz',5e5,'kxz',0,'kzx',0,...
'cxx',1e2*beta,'czz',5e2*beta,'cxz',0,'czx',0);
d_rotor('TestLalanne2D0F',r1)
```



Figure 4.8: Campbell (left) and Responses to unbalanced load (right) for different damping (top: $\beta = 1$, middle: $\beta = 15$ and bottom: $\beta = 26$). Asymmetric rotor with z bearing stiffness and damping.

Backward and forward mode can cross each other in the Campbell diagram. Asymmetry leads to the excitation of the backward mode. Damping leads to a more spread resonance response.

4.3 1D models

4.3.1 1D example in a fixed frame

This first example treats the simple case, taken from [6], of a shaft with a rotating disk at one third the length.

In order to compare this model to the simple 2 DOF model of Lalanne (see section 4.2) we project the matrices on the 2 sin shape function of the shaft. Relative error for mass matrix is 0.01%, 4.59% for stiffness and 1.64% for gyroscopic matrix. Campbell for the projected model and the Lalanne 2 DOF model are almost the same. For the full 1d model (not projected) the increasing of the frequency of 1rst forward whirl mode tends to an asymptote.



Figure 4.9: Campbell diagrams. Left: 1D rotor (up: projected on 2 DOF, down: full model), right: Lalanne 2 DOF rotor.

Following example performs preceding comparison :

```
% - LALANNE 2DOF
model2DOF=[]; m=14.29; k=1.195e6; a=2.871;
kxx=0; kzz=0; kxz=0; kzx=0;% bearing
% define model matrices :
model2DOF.K={m*eye(2), k*eye(2)+sin(pi*2/3)^2*[kxx kxz;kzx kzz],a*[0 -1;1 0]};
model2DOF.DDF=1+[.01;.03];
model2DOF.Klab={'m','k','Dg'};model2DOF.Opt=[1 0 0;2 1 70];
```

```
% rotate model so that x=rotating axis:
model2DOF.K=feutil('tkt',[1 0 0; 0 0 1],model2DOF.K);
model2D0F.K=feutil('tkt',[0 -1 0;1 0 0; 0 0 1],model2D0F.K);
for j1=1:3; model2D0F.K{j1}=model2D0F.K{j1}([2 3],[2 3]); end
model2D0F.D0F=[1.02;1.03];
model2D0F=stack_set(model2D0F, 'info', 'Omega', [1 0 0]);
% - SHAFTDISK beam1+mass1 gyro 70 - - - - -
 [model,TR]=d_rotor('TestshaftdiskMdl -TRLalanne')
 model.Elt=feutil('removeelt eltname celas',model);
 % project matrices on 2 sin shape functions equivalent to lalanne 2DOF model
 model=fe_caseg('assemble -secdof -matdes 2 1 70 3',model);
 TR=feutilb('placeindof',model.DOF,TR);
 model.K=feutil('TKT',TR.def,model.K); model.DOF=[2.02;2.03];
 % - compare with reference matrices from Lalanne
 for j1=1:3 % Compare matrices [model.K{j1} model2DOF.K{j1}]
  if normest(model.K{j1}-model2DOF.K{j1})/normest(model.K{j1})>0.05;
  sdtw('_err', '2DOF and shaftdisk unmatch for mat %s', model.Klab{j1});
  end
 end
% Campbell with SDT matrices reduced on Lalanne shapes
r1=struct('data',linspace(0,9e3),'unit','RPM');
fe_rotor('campbell -nodir -crit -cf1',model,r1);
% - Now compare full beam model and 2D shapes
model.DOF=[]; model.K={};if ishandle(1);close(1);end
fe_rotor('campbell -nodir -crit -cf1',model2DOF,r1);
set(findall(1,'type','line'),'color','k', ...
 'linewidth',2,'linestyle','--');
hold on;
fe_rotor('campbell -nodir -crit -cf1',model,r1);
hold off;set(gca,'ylim',[0 160]);
h=findall(1,'type','line');legend(h(end+[0 -3]),'2 DOF','beam')
```

Following example computes frequency response to unbalanced or asynchronous load:

```
model=d_rotor('TestShaftDiskMdl'); % Model Initialization
```

```
% Assemble nominal matrices:
model=fe_caseg('assemble -reset -secdof -matdes 2 1 70',model);
```

4.3. 1D MODELS

```
% Campbell diagram and critical speeds:
fe_rotor('campbell -full -critical',model,linspace(0,20000,30));
set(gca,'ylim',[0 350]);
% Unbalanced mass or asynchronous load :
mb=1e-4; db=0.15; % mass, distance to axis
s=1; f0=1; % s=1, unbalanced load. s<>1, asynchronous load
om=sort([3296 3200:10:3500 linspace(0,20000,101)]); % RPM
% RotatingLoad NodeId f0 theta0 exponent : define complex rotating load
if s==1 % unbalanced mass
model=fe_rotor(sprintf('rotatingload 2 %.15g -90 2',mb*db),model);
else % asynchronousload
model=fe_rotor(sprintf('rotatingload 2 %.15g -90 0',f0),model);
end
r1=struct('Omega',om/60*2*pi,'w',s*om/60*2*pi); % Range
model=stack_set(model,'info','Range',r1);
R1=fe_rotor('forcedresponse',model); % compute forced response
iiplot(R1) % plot response
% Post (radial deformation):
Q=max(abs(R1.Y),[],2); figure;semilogy(om,Q);
xlabel('Rotation speed [RPM]'); ylabel('radial def amplitude [m]')
if s==1; title('Unbalanced mass')
        title(sprintf('Asynchronous load %.15g\\Omega',s))
else:
end
```



Figure 4.10: Top: Campbell diagrams. Bottom: Responses to unbalanced mass and asynchronous load.

Fig 4.10 shows the radial deformation response for an unbalanced load and for an asynchronous load rotating at $-\Omega$ speed. The unbalanced load excites the forward modes (3296 RPM) whereas the asynchronous load excites the backward modes (2785 RPM and 4697 RPM). Frequencies match those computed as critical frequencies in the Campbell diagram.

4.3.2 1D models in a rotating (body-fixed) frame

While this representation is not very classical, it corresponds to the nominal choice when doing time integration of a rotor that is not axisymmetric.

4.4 3D rotor

The same rotor as described in lalanne (see fig 4.3) is meshed using hexa8 elements. Use model=d_rotor()



Figure 4.11: 3d model of Lalanne rotor.

Matrices are defined in the local rotating frame. We described the unbalanced load by a static load, and we use in following example the same procedure as for local frame 1d rotor, at w = 0.

```
model=d rotor('testvolshaftdiskmdl')
% Assemble nominal matrices:
model.DOF=[];model=fe_caseg('assemble -se -matdes 2 1 7 8',model);
model.DOF=fe_case('gettdof',model);
% Campbell diagram:
model=stack_set(model,'info','eigopt',[5 20 1e3]);
fe_rotor('campbell -critical',model,linspace(0,20000,30));
% Unbalanced mass or asynchronous load :
mb=1e-4; db=0.15; % mass, distance to axis
s=0; f0=1; % s=1, unbalanced load. s<>1, asynchronous load
om=sort([2789 2750:10:2820 11760:10:11840 linspace(0,20000,101)]); % RPM
model=fe_rotor(sprintf('rotatingload 180 %.15g 0 2',f0),model);
 r1=struct('Omega',om/60*2*pi,'w',s*om/60*2*pi); % Range
 model=stack_set(model,'info','Range',r1);
 R1=fe_rotor('forcedresponse',model); % compute forced response
 iiplot(R1) % plot response
```

```
% Post (radial deformation):
Q=abs(R1.Y(:,1)); % unbalanced along x
figure;semilogy(om,Q);
xlabel('Rotation speed [RPM]'); ylabel('radial def amplitude [m]')
if s==0; title('Unbalanced mass')
else; title(sprintf('Asynchronous load %.15g\\Omega',s))
end
```



Figure 4.12: Left: Campbell diagram. Right: Response to unbalanced mass.

Unbalanced mass excites the forward whirl modes. Maximum response is found at critical speeds (rotation speeds that induce a complex mode of 0 Hz frequency in the rotating frame). Campbell critical speed (2789 RPM) matches computed frequency response.

4.5 Data structure reference

xxx

Function reference

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fe_cyclic _

Purpose

Support for cyclic symmetry computations.

Syntax

```
model=fe_cyclic('build NSEC',model,LeftNodeSelect)
def=fe_cyclic('eig NDIAM',model,EigOpt)
```

Description

fe_cyclic groups all commands needed to compute responses assuming cyclic symmetry. For more details on the associated theory you can refer to [8].

Assemble [,-struct]

This command supports the computations linked to the assembly of gyroscopic coupling, gyroscopic stiffness and tangent stiffness in geometrically non-linear elasticity. The input arguments are the model and the rotation vector (in rad/s)

```
model=demosdt('demo sector all');
[K,model,Case]=fe_case('assemble -matdes 2 1 NoT -cell',model);
SE=fe_cyclic('assemble -struct',model,[0 0 1000]); %
def=fe_eig({K{1:2},Case.T,model.DOF},[6 20 0]);% Non rotating modes
def2=fe_eig({K{1},SE.K{4},Case.T,model.DOF},[6 20 0]); % Rotating mode shapes
[def.data def2.data]
```

Note that the rotation speed can also be specified using a stack entry model=stack_set(model, 'info', 'Omega',[0 0 1000]).

Build ...

 $model=fe_cyclic('build nsec epsl len', model, 'LeftNodeSelect')$ adds a cyclic symmetry entry in the model case. It automatically rotates the nodes selected with LeftNodeSelect by $2\pi/nsec$ and finds the corresponding nodes on the other sector face. The default for LeftNodeSelect is 'GroupAll' which selects all nodes.

The alternate command

model=fe_cyclic('build nsec epsl len -intersect',model,'LeftNodeSelect') is much faster
but does not implement strict node tolerancing and may thus need an adjustement of epsl to higher
values.

Command options are

- *nsec* is the optional number of sectors. An automatic determination of the number of angular sectors is implemented from the angle between the left and right interface nodes with the minimum radius. This guess may fail in some situations so that the argument may be necessary.
- nsec=-1 is used for periodic structures and you should then provide the translation step. For periodic solutions,

model=fe_cyclic('build -1 tx ty tz epsl len -intersect',model,'LeftNodeSelect')
specifies 3 components for the spatial periodicity.

- Fix will adjust node positions to make the left and right nodes sets match exactly.
- epsl*len* gives the tolerance for edge node matching.
- -equal can be used to build a simple periodicity condition for use outside of fe_cyclic. This option is not relevant for cyclic symmetry.
- -ByMat is used to allow matching by MatId which allows for proper matching of coincident nodes.

```
model=demosdt('demo sector 5');
cf.model=fe_cyclic('build epsl 1e-6',model);
```

LoadCentrifugal

The command is used to build centrifugal loads based on an info, Omega stack entry in the form

```
data=struct('data',[0 0 1000],'unit','RPM');
model=stack_set(model,'info','Omega',data);
model=fe_cyclic('LoadCentrifugal',model);
```

Eig

def=fe_cyclic('eig ndiam',model,EigOpt) computes ndiam diameter modes using the cyclic symmetry assumption. For ndiam;0 these modes are complex to account for the inter-sector phase shifts. EigOpt are standard options passed to fe_eig.

This example computes the two diameter modes of a three bladed disk also used in the d_cms2 demo.

```
model=demosdt('demo sector');
model=fe_cyclic('build 3',model,'groupall');
fe_case(model,'info')
def=fe_cyclic('eig 2',model,[6 20 0 11]);
fe_cyclic('display 3',model,def)
```

The basic functionality of this command is significantly extended in fe_cyclicb ShaftEig that is part of the SDT/Rotor toolbox.

Omega[,Group,GroupSet]

Handling of dynamic rotating bodies. **Warning** At the moment only one rotation vector can be defined. It can either be applied to the whole model or to specified groups. At low level, information is located in the info,Omega entry of an SDT model. This entry is a structure with fields

- .data provides the angular rotation vector whose norm is the angular velocity, defining the rotation axis.
- .unit provies the unit system associated to the amplitude, eitherrad/s or RPM.
- .group (optional) defines the model groups affected by the rotation, if omitted or left empty the whole model is affected.
- .orig (optional) defines the origin rotation (a point of the axis).

Command Omega provides the current data associated to a model.

[omega,rot,data]=fe_cyclic('Omega',model);

model is a standard SDT model. The outputs are omega the rotation vector, rot the rotation matrix, and data a reconstructed info,Omega stack entry based on the current state.

Commands OmegaGroup provides tools for definition of models with specific rotor areas.

• OmegaGroupSet provides an integrated definition forcing groups to be reset to conform with any FindElt selection. The specific group assignment is required due to low level assembly implementations.

```
model=fe_cyclic('OmegaGroupSet',model,list);
```

Input model is a standard SDT model, list is a three column cell-array with as many lines as declarations following the format {FindEltStr, Amplitude, Axis, Orig;...} respectively providing an element selection string, the angular velocity amplitude (scalar), the rotation axis (only the direction is used here), nx,ny,nz, and an origin point of the rotation axis ox,oy,oz. data can be directly placed as a stack entry named info,OmegaData in the model. The last

column can be omitted, in which case the origin considered is the global frame one. At the moment all rotation axes and amplitudes must be the same for all lines. The output model is then a model with separated groups for (one for each element type) affected to the rotation and with a new stack entry info,Omega. Command option First will force the new groups to be the first ones in the model.

• OmegaGroup is a lower level command without group modification. model=fe_cyclic('OmegaGroup',model,sel,data); Input model is a standard SDT model, sel is an element selection string, data is the omega structure with fields .data as defined at this command header.

See also

 ${\tt fe_cyclicb}$

fe_rotor

Purpose

The fe_rotor function implements classical solutions for rotor dynamics applications.

AddMass

mdl=fe_rotor('AddMass',mdl,RO);

This command can be used to add a local mass on a 3d rotor mesh. The mass is added, and linked to existing node using a MPC connection.

 ${\tt RO}$ is a data structure with fields

- .mxyz matrix whose first 3 columns define x y and z coordinates of added mass, and the 4th defines the mass.
- .ProId the ProId of the rotor where mass is added.

Command options are

- -DofLoad defines a 6 direction load on added masses (usefull for reduction purpose).
- -mpcmaster modifies MPC connection so that mass DOF are master rather than slave.

Campbell

fe_rotor('Campbell',model,RunOpt) computes the Campbell diagram and displays it with fe_rotor('p
if no output argument is requested.

RunOpt can be a vector of rotation speeds (RPM) or a data structure with at least field .Omega containing rotation speeds and other fields giving other variables used in zCoef (for example .par field).

Accepted command options are

- -cf i can be used to force display of the diagram in a specific figure.
- -nodir avoids the call to determine the rotation direction.

- -full forces the use of full matrices. Without the argument, full matrix eigenvalue calls are only performed with less than 100 DOFs.
- -crit overlays the critical speed computation.
- -stability displays stability diagram (damping/Omega).

Examples can be found in section 3.4.1 .

Critical

Computation of critical speeds. It can be called when computed Campbell using command option -crit (see Campbell).

Critical speed are computed assuming deformation is a complex mode at the same frequency as the rotation speed $(w = \Omega)$ in the global fixed frame or equal to 0 (w = 0) in the local rotating frame. In the global fixed frame:

$$\begin{split} M\{\ddot{X}\} &+ (C + \Omega D(\Omega = 1))\{\dot{X}\} + (K + \Omega^2 Kc(\Omega = 1))\{X\} = 0\\ \text{With } X &= X_0 exp(i\Omega t)\\ (\Omega^2(-M + iD(\Omega = 1)) + i\Omega C + K)\{X_0\} = 0\\ \text{In the local rotating frame:}\\ M\{\ddot{X}\} + (C + \Omega D(\Omega = 1))\{\dot{X}\} + (K + \Omega^2 Kc(\Omega = 1))\{X\} = 0\\ \text{With } X &= X_0 \text{ constant}\\ (K + \Omega^2 Kc(\Omega = 1))\{X_0\} = 0 \end{split}$$

Examples can be found in section 3.4 xxx.

Whirldir

Internal command to display the direction of the modes on the Campbell diagrams. xxx details

Assemble

model=fe_rotor('Assemble',model,zCoef,r1); This command is used to assemble mass stiffness
and damping matrices taking in account all the rotor matrices (gyroscopic coupling, centrifugal
softening, etc. ...).

Default zCoef can be obtain using model=fe_def('zCoef-default', model); r1 is a data structure whose fields are parameters used in zCoef. In particular one should specify the rotation speed r1.Omega (rad/s). Command option -cell can be used to return only matrices in a cell array.

RotatingLoad

This command builds a rotating DofLoad on a model node for frequency analysis.

model=fe_rotor('RotatingLoad NodeId f0 theta0 wexponent',model);

NodeId is the id of the node where load will be applied, f0 is the amplitude of the load, theta0 is the angle formed by the load initial direction and the first global direction in the plane orthogonal to rotation axis (x for y and z rotation and y for x rotation). The amplitude of the load can depend on a power of pulsation defined with wexponent.

Resulting complex load is of the form

$$f0w^{\text{wexponent}} \left\{ \begin{array}{c} 1\\ \pm i \end{array} \right\}$$

in the plane orthogonal to the rotation axis (sign of +/-i depends on rotation axis).

Forced response to this load can be computed using ForcedResponse.

ForcedResponse

This command can be used to compute forced response to a frequency load (for example a rotating load built using RotatingLoad).

R1=fe_rotor('ForcedResponse',model);

Complex load must be prior defined as a 'In' entry in the case of the model.

Observation can be defined as 'output' SensDof entry. If not, only dof corresponding to the 'In' load will be returned.

Range of computation must be defined in the 'info' 'Range' Stack entry of the model as a structure of data (with fields .Omega defining rotation speeds and .w defining pulsations of the load at corresponding rotation speeds. These fields must be of the same length. If necessary one can add other fields for variables used in a defined zCoef). If .w is equal to zero, frequency dependence of load is assumed to concern the .Omega vector (it is useful to described unbalanced mass in the local rotating frame since load amplitude depends on Ω^2 but load is static in this frame so that w = 0).

Following example defines an unbalanced load on the simple 1d ShaftDisk model defined in d_rotor and computes the forced response (local frame):

```
model=d_rotor('TestShaftDiskMdl') % build simple model
model=fe_caseg('assemble -se -matdes 2 1 70',model); % assemble model
mb=1e-4; db=0.15; % mass, distance to axis
om=linspace(0,6000,201)'; % RPM
model=fe_rotor(sprintf('RotatingLoad 2 %.15g -90 2',mb*db),model); % unbalanced mass
```

```
r1=struct('Omega',om/60*2*pi,'w',om/60*2*pi); % Range
model=stack_set(model,'info','Range',r1);
R1=fe_rotor('forcedresponse',model); % compute forced response
iiplot(R1) % plot response
```

In global rotating frame, load is static (w = 0). Following example deals with a rigid disk in global rotating frame:

```
R1=.01;R2=.15;h=0.03;
md=d_rotor(sprintf('TestVolDisk -dim%.15g %.15g %.15g 2 16',R1,R2,h));
md.DOF=[]; md=fe_caseg('assemble -se -matdes 2 1 7 8',md);
% disk assumed to be rigid :
rb=feutil('geomrb',md); cf=feplot(md); cf.def=rb;
md.K=feutil('tkt',rb.def(:,[1 3 4 6]),md.K);
md.DOF=[1.01 1.03 1.04 1.06]';
md.K{2}=0*md.K{2}; % rigid disk
md.K{2}(1,1)=5e5;md.K{2}(2,2)=5e5; % bearing
md.K{2}(3,3)=1e5;md.K{2}(4,4)=1e5; % bearing rot
% Unbalanced mass or asynchrone load :
mb=1e-4; db=0.15; % mass, distance to axis
s=0;
om=sort([2789 2750:10:2820 11760:10:11840 linspace(0,20000,101)]); % RPM
md=fe_rotor(sprintf('RotatingLoad 1 %.15g 0 2',mb*db),md);
r1=struct('Omega',om/60*2*pi,'w',s*om/60*2*pi); % Range
md=stack_set(md, 'info', 'Range', r1);
R1=fe_rotor('forcedresponse',md); % compute forced response
iiplot(R1) % plot response
```

SEBuild

This command can be used to build a superelement of rotor from a 3d rotor model. It includes Craig Bampton reduction, bearing rings adding, ...

SE=fe_rotor('SEBuild',model,RO);
model is a 3d mesh of rotor.
RO is a data structure with following fields

.xbea vector defining position of bearings according to rotor axis.

.bea data structure of bearing infos. bea.length is the length of the bearings, bea.Npt the number of points on one bearing, bea.ProId is the ProId of the journal.

.mxyz is optional. Defines the mass position (reduction is performed so that DOF are created at these positions).

.EigOpt eig option for mode computation for Craig-Bampton reduction.

One can use following options:

-noreduce no reduction is performed.

-rigid use rigid ring for bearings instead of rbe3 rings.

TimeOpt

model=fe_rotor('TimeOpt',model);

Defines a default TimeOpt in the 'info' 'TimeOpt' model stack entry for time integration with rotor models. Following command options are available:

• Assemble Defines AssembleCall field in order to take in account gyroscopic effects in time computation. TimeOpt field .matdes should be filled by user in order to define what matrices will be assembled. Default is [2 1 7] for mass stiffness and gyroscopic coupling matrix, see sdtweb('MatType') for available matrices (for example 8 for centrifugal softening...). Before time integration desired matrices will be assembled taking in account the 'info' 'Omega' model stack entry that defines rotation vector (norm should be equal to one so that matrices can be assembled once for different computation with different rotation speed). Rotation speed is taken as first parameter on the 'info' 'Range' model stack entry (in that case rotation
vector norm must be one). If there is no Range entry, rotation is taken equal to the norm of rotation vector. Corresponding M C K matrices are then built (fe_rotor Assemble call) and will be used for Newmark time integration.

rotor1d

Purpose This function gives some commands to help the meshing of 1D rotor. See section ?? for general details about 1d rotors and section 3.1.2 for a full example of 1d rotor meshing.

1To2D

From a 1d model with **beam1** elements (masses are not converted for the moment), builds a 2d rotor. LC l defines the max length of elements.

1To3D

From a 1d model with **beam1** elements (masses are not converted for the moment), builds a 3d rotor. one can see section 3.1.3 for a tutorial.

- LC l defines the max length of elements.
- div n defines the number of sector of 3d mesh.

The grounded celas are assumed to be bearings. RBE3 rings are defined at each bearing.

AddBearing

This command add a rotor bearing as a celas element. mold=rotorld('AddBearing ...',mold,pos); mold is a ld rotor model pos is the x y z position of the bearing.

Command options are

- -keep if not present remove existing celas at given position.
- DOF *DOF* that defines the DOF concerned by bearing link.
- k k the spring stiffness.
- c c the spring damping.

• **ProID** *ProID* the spring ProID.

```
For example:
mold=rotor1d('AddBearing DOF -123 k 1e4 -keep',mold,pos);
```

AddBeam

mold=rotor1d('AddBeam ...',mold);

This command adds a beam on the axis of the rotor, accordin to following argument specified as command option:

- x1 x1 beginning of the beam in the axis direction
- x2 x2 end of the beam in the axis direction
- r1 r1 inner radius of the beam. By default 0.
- r2 r2 outer radius of the beam.
- MatID MatID MatID. If not given new MatID is used.
- **ProID** ProID ProID. Note that the associated element property contains the section information according to r1 and r2, so that existing ProID will be lost. If not given new ProID is used.
- -refine Refine existing beams to ensure that beams have coincident nodes.

For example mold=rotor1d('AddBeam x1 0.5 x2 0.7 r1 0.11 r2 0.77 MatID 1',mold);

AddNodeRefine

[mold,ind]=rotor1d('AddNodeRefine ...',mold,xyz);

This low level call command adds nodes at given positions xyz (one row per node, 3 columns x y and z, or one column to define distance along the rotor axis) refining beams of the model mold if needed.

Command options are:

• -epsl epsl Tolerance for new node adding.

```
For example
[mold,ind]=rotor1d('AddNodeRefine',mold,[0.2 0 0]);
```

Skyline

mdl=rotor1d('Skyline...',xy);

This command can be used to generate rotor model from skyline description. xy is a 2 column matrix whose 1st column defines position according to rotation axis, and 2nd column defines radius at corresponding positions. One can define multiple parts rotor using separator NaN in the 1st column. For 2D rotor meshing, use [NaN r] to define internal radius r of the corresponding part.

- SkylineToBeam builds 1d beam rotor model. See example in 1D subsection of section ?? .
- SkylineTo2D a 2D rotor model (that can be used to generate 3D model by revolution, or using cyclic symetry, see rotor2d BuildFrom2D)), see example in 2D subsection of section ?? .

Following command options are accepted

• -Lc l specify maximum length l of elements.

rotor2d

Purpose

The **rotor2d** function lets you build a superelement representation for rotor applications starting from a 2D model. A tutorial is given in section **??**

teig

Needs description.

test

Simpledisk

BuildFrom2D

model=rotor2d('buildFrom2D', model2D); builds a 3D sector model by revolution of a 2D section. In the rotor module, the symmetry axis is always z so that if the 2D mesh is given in xy coordinates a permutation is performed.

- -nsec N number of sectors. By default, one considers 10 sectors, for a different value use buildFrom2D -nsec N.
- -div N number of divisions in each sector.

The sector is modeled as a superelement called disk1.

```
% Model Initialization
model2D=rotor2d('test simpledisk -back');
cf=feplot(rotor2d('buildFrom2D',model2D));
SE=cf.Stack{'disk1'}; % enforce boundary cond. on sector and assemble
SE=fe_case(SE,'FixDof','Base','z==1.01');
SE=stack_set(SE,'info','Omega',[0 0 0;0 0 1000]); % define speed range
SE=fe_cyclic('assemble -se',SE);
cf.Stack{'disk1'}=SE; fecom('view1');
cf.Stack{'info','EigOpt'}=[5 20 0]; % define eigenvalue options
RunOpt=struct('targ',1, ... % define target diameter
'Range',linspace(0,1,30)); % define speed points relative to range
```

```
[cf.def,hist]=rotor2d('teig',cf,RunOpt);
figure(1);rotor2d('plot',hist);set(gca,'ylim',[0 250])
```

demo_cyclic

Purpose

Combines examples of the use of fe_cyclicb commands.

\mathbf{Syntax}

```
demo_cyclic('testdisk nsec')
demo_cyclic('testrotor nsec1 nsec2 ...')
```

Disk

Moved to section 3.5.2 .

${\tt ShaftMono}$

Moved to section 3.5.3.

Variable speed

One considers stiffness matrices that are dependent on the rotation speep. Assuming that a second order polynomial representation is sufficient, one can build a vector of weighing coefficients

$$\left\{ \begin{array}{c} \alpha_1\\ \alpha_2\\ \alpha_3 \end{array} \right\} = \left[\begin{array}{ccc} 1 & 1 & 1\\ \Omega_1 & \Omega_2 & \Omega_3\\ \Omega_1^2 & \Omega_2^2 & \Omega_3^2 \end{array} \right]^{-1} \left\{ \begin{array}{c} 1\\ \alpha_2\\ \alpha_3 \end{array} \right\}$$
(5.1)

Such that the stiffness at rotation speed Ω is approximated by

$$[K(\Omega)] = \sum_{\alpha_i} [K\Omega_i]$$
(5.2)

The zCoef uses velocity Omega in rad/s.

This example now treats computation at variable rotation speeds

```
% Model Initialization
model=demo_cyclic('testblade');cf=feplot(model);
% Compute matrix coefficients for a multi-stage rotor
```

demo_cyclic _

```
range=struct('data',[0 0 1;0 0 800;0 0 1600],'unit','RPM');
% Assembling in the feplot figure, allows memory offload
fe_cyclicb('polyassemble -noT',cf,range);
X=struct('data',linspace(0,1600,10),'unit','RPM');
fe_rotor('Campbell -cf1',model,X)
```

Another example will be needed to treat the multi-stage case

```
% Model Initialization
model=demo_cyclic('testrotor 7 10 -blade');
model=fe_cyclicb('shaftRimAsSe',model);
cf=feplot(model);
% Compute matrix coefficients for a multi-stage rotor
range=struct('data',[0 0 1;0 0 800;0 0 1600],'unit','RPM');
% Assembling in the feplot figure, allows memory offload
fe_cyclicb('polyassemble -noT',cf,range);
% Now run a mono-harmonic multi-speed computation
cf.Stack{'info', 'Omega'}=struct('data', range.data(1,:), 'unit', 'RPM');
def=fe_cyclicb('shafteig 0 -ReAssemble 2 -NoN',cf);
Sel={'disk1','groupall';'disk2','groupall'};
fe_cyclicb('DisplaySel',cf,def,Sel)
% Reduce the full model
fe_cyclicb('ShaftPrep -svdtruncate -mseq1 -handle -norestit',cf,def);
fesuper('fassemble',cf);
% Force single harmonic
```

% xxx

Rewrite needed here.

ShaftMulti

The second example is a non-monoharmonic shaft computation.

The following example builds a reduced order model from a set of mono-harmonic modeshapes whose Fourier harmonics are 0, 1 and 2 and sector modes with fixed interfaces. The latter are computed within the framework of mono-harmonic computations, they are called with -1 in shaftTeig. No confusion is possible since the true mono-harmonic solutions with $\delta = -1$ are solutions with $\delta = 1$. However, the restitution of fixed interface solutions with Display[Sel] has no sense.

Call shaftprep aims to build the reduced kinematic subspaces of the sector super-elements from the specified target solutions. Prior to that, it separates the sector mesh into two: the slice with the left-interface nodes (the inter-sector super-element) and the remaining elements (the sector superelement). It projects the matrices of the sector super-elements onto their individual subspaces and the matrices of the inter-sector slices onto the subspaces of its two neighbouring sector superelements.

Command fassemble in fesuper first projects the finite elements matrices of the inter-disk volumic interface onto the subspaces of its neighbouring disks. It then assembles the reduced matrices of the sector super-elements and inter-sector slices to form the reduced matrices of the disks. Finally, it assembles these reduced matrices and that of the volumic interfaces to form the reduced matrices of the whole rotor.

Here too, a selection can be specified so that the generalized modeshapes can be recovered to a subset of physical DOF (relying on the true mesh or a viewing mesh). The selections are defined like in DisplaySel, however both the sector and inter-sector super-elements have to be considered so that the recovery concerns the whole underlying bladed sector.

The Fourier harmonic contents of the generalized modeshapes can be obtained without recovery with the help from command fourier of fe_cyclicb. When specified, -egyfrac returns the fraction of strain energy per harmonic per disk so that energy localization within a disk can be achieved with the supplementary information of which harmonics are involved in the response. The graph displays the disks (from top to bottom) and for each disk the possible harmonics between 0 and N/2 (if applicable). Another way to display the same information is to group the harmonics first and then the disks. This is done with the -sortbyd option. It adds the proper amount of zeros for harmonics that are not present on a given disk.

_mono.mat to name a model that contains the mono-harmonic description.

```
if any(r1(1:6,1)>r1(7,1)/1e5);error('Missing rigid modes');end
 % Model Reduction
 fe_cyclicb('ShaftPrep -svdtruncate -mseq1 -handle -norestit',cf,def);
 fesuper('fassemble',cf);
 % Mode Computations
 defr=fe_eig(cf.Stack{'MVR'}, [5 50 1e3 11 1e-8]);
 Sel={'disk2','innode {r>150}';
      'disk21','innode {r>150}';
      'disk1','groupall';
      'disk1l','groupall';
      '', 'withnode {x>0}'};
fesuper('sebuildsel -initrot',cf,Sel);
cf.def=feutilb('placeindof',cf.sel.cna{1}.adof,defr);
fecom('colordataevala');
% Post-processing: spatial spectra
cf.sel='reset';cf.def=[];
fe_cyclicb('fourier 7:25 -red -egyfrac -cf 4',cf,defr);
fe_cyclicb('fourier 7:25 -red -egyfrac -sortbyd -cf 5',cf,defr);
% focus on mode 7
fe_cyclicb('fourier 7 -red -egyfrac -cf 6',cf,defr);
```

Now one wants to treat the case of a forced response for loads defined on the disks only. The command ShaftLoadMulti allows to build one mono-harmonic excitation per disk. For each disk i, the shape def_i and harmonic coefficient delta_i are specified in a cell array whose a typical line is {'diski',def_i,delta_i}.

Call **fourier** provides a means to checkout the spatial harmonic content of the generalized load with respect to the set of generalized coordinates (option **-red** has to be specified). Energy-based computations are not available for generalized loads.

fe_cyclicb Mesh

Purpose

MeshAddRim

MeshAddRim deals with already existing inter-disk volumic rings. It renumbers their model to be integrated with the already defined disk super-elements. Note that adjusting the geometric tolerance with option epsl val can be important.

```
cf=demo_cyclic('buildstep0');% See sdtweb('demo_cyclic.m#Step0')
% Extract rim model into mo1 (would come from other reading)
[cf.mdl.Elt,mo1.Elt]=feutil('RemoveElt EltName~=SE',cf.mdl);
mo1.Node=cf.mdl.Node;mo1.Node=feutil('GetnodeGroupAll',mo1);
cf.mdl.Node=[];
% Combine mo1 (rim) and cf.mdl (disks)
fe_cyclicb('MeshAddRim epsl 1e-3',cf,mo1);
cf.sel='EltName~=SE';fecom('showpatch');
```

Other commands documented in the tutorial are

- MeshRim described below and illustrated in section 3.1.6 , is an automated procedure to mesh rims form a selection of nodes on two disks.
- MeshRimLine2Patch, described in section 3.1.6 is used to build view meshes of large models.

MeshRim

This command is used to mesh rims as volumes or penalty springs.

- -kp val is used to specify the penalty stiffness.
- epsl val specifies length tolerance for node matching.
- -masterdisk val specifies the disk(s) on whose surface the slave nodes will be matched.
- -masterdisk val specifies the slave disk(s) whose edge nodes will be connected to the master disk surface.

MeshSecAddNode

MeshSecAddNode [epsl espl] is used to place sensors relatively to a disk/rotor model. It uses a cell array to define sensors (see sdtweb('sensor#scell')). It first matches each sensor to a surface in a sector of a disk with an optional tolerance epsl that allows to tune the research. It then adds these surface elements to the selection. The latter serves as the input of the subsequent SensMatch call that

- 1. automatically deals with the specified direction of measure (column 'DirSpec') by detecting the normals, computing vectors from the latter, *etc.*,
- 2. builds the observation matrix at each sensor (cta entries) that relates the DOF of the matched surface to the observation at this sensor along the measurement direction.

```
% Load 360 reduced shaft model, sdtweb('demo_cyclic.m#Step2')
cf=demo_cyclic('buildstep2');
sensors=cf.Stack('Test').tdof;
disp(sensors); % display sensor properties
% match sensors
[sens,sel]=fe_cyclicb('MeshSecAddNode epsl 20',cf,sensors);
cf.mdl=fe_case(cf.mdl,'SensDof','Test',sens);
% display sensors
fecom('showpatch');
fecom(cf,'curtab Cases','Test');
```

The model to which the sensors are matched is made of super-elements already reduced, call to MeshSecAddNode with the supplementary input of a generalized quantity (with a list of generalized coordinates stored in .DOF) automatically builds the observation matrix that expands the generalized modeshapes to the sensors along the measurement direction.

A first application of this is to animate generalized modeshapes on the experimental wire frame as proposed in the example below.

```
cf=demo_cyclic('buildstep4');
def=cf.Stack('def_mvr');
% match sensors
sensors=stack_get(cf.mdl,'info','Test','GetData');
```

A second application is to display and animate generalized response or transfer functions.

```
cf=demo_cyclic('buildstep6');
xF=cf.Stack('xF_mvr');
```

```
% match sensors
sensors=stack_get(cf.mdl,'info','Test','GetData');
[r1,sel,sens]=fe_cyclicb('MeshSecAddNode epsl 20',cf,sensors.tdof,xF);
% add wireframe
r1.Elt=[r1.Elt zeros(size(r1.Elt,1),1);cf.Stack('WireFrame')];
r1.Elt=feutil('setgroupall egid -1',r1);
cf.mdl=fe_case(cf.mdl,'SensDof','Test',r1);
% display responses
xF=cf.Stack('xF_mvr');
ci=iiplot(3);iicom(ci,'curvereset');
r1=struct('X',{{xF.data(:) sens.lab}},'Xlab',{{'Freq','DOF'}}, ...
  'Y',(sens.cta*xF.def).')
iicom('curveInit','Test',r1)
iicom('SubMagPha');
% animate response at sensors
cf.sel(1)='groupall';
cf.sel(2)='-Test';
cf.def=struct('def',sens.cta*xF.def,...
              'DOF', cf.CStack('Test').tdof(:,1),...
              'data', xF.data);
```

cf.o(1)={'sel 2 def 1 ch 1 ty8 scc10', 'edgecolor', 'r', 'linewidth',2};

${\tt MeshSensMatch}$

supports topology correlation for a SensDof entry defined on a multi-stage shaft model.

MeshSurfSet

generates standard surface sets for a sector.

MeshCylSurf

model=fe_cyclicb('MeshCylSurf epsl *val*', model, NodeList) makes a surface perfectly cylindrical. The node list of nodes to be used on the left edge may be omitted (it will be found automatically).

MeshFixTheta

provides handling utilities for slanted sectors, see section 3.1.5.

MeshFixRadial

Generates MPC constraints for using DOFs given in radial coordinates: .01 is radial, .02 tangential, .03 axial.

The following example illustrates a case where tangential motion of nodes 5 and 6 is set.

```
cf=demo_cyclic(sprintf('testrotor %i -cf 5',5));
cf.Stack{'disk1'}=fe_cyclicb('MeshFixRadial', ...
cf.Stack{'disk1'},'radial',[5.02;6.02]);
def=fe_cyclicb('shaftteig 0 1', ...
stack_set(cf.mdl.GetData,'info','EigOpt',[5 3 0]))
fe_cyclicb('Display',cf,def);
fecom('ColorDataEvalTanZ -ColorBarTitle"TanZ"');
```

fe_cyclicb ____

Purpose

Support for advanced cyclic symmetry computations.

Description

fe_cyclicb groups advanced commands used to build and manipulate reduced order models of single symmetric structures and their assemblies. For more details on the associated theory you can refer to [8].

Rotor Construction

DiskFromSector

This command builds a disk/rotor model from (a) physical sector model(s). Shafts can be generated by combining multiple calls to disk from sector. Once, disks are combine, their connection trough rim models is described in section 3.1.6.

Command DiskFromSector also handles the construction of the cyclic sector models. Cyclic symmetry information can be already given in the sector model (calls to fe_cyclic('build') done beforehand) or done in the command. In the later case, an optional epsl tol can be declared so that it is propagated to the subsequent call to fe_cyclic('build epsl tol',...), where tol is the desired tolerance for left-right interface node matching.

The example below demonstrates the capability of the function for two disks with 7 and 10 blades respectively.

```
cf=demo_cyclic('buildstep0');
sector1=cf.Stack{'disk1'};
sector2=cf.Stack{'disk2'};
% build disk1 from sector1
model=fe_cyclicb('DiskFromSector epsl 1e-6',[],sector1);
% build disk2 from sector2 and append to disk1
model=fe_cyclicb('DiskFromSector epsl 1e-6',model,sector2);
fe_cyclicb('DisplayFirst',model) % Avoids full display for large models
```

In cases when **cf** already contains one sector per disk, the shaft model can be created in a single operation with the command **fe_cyclicb('diskfromsector', [], cf, {'sel_disk1', 'sel_disk2',...})**; where **sel_disk***i* selects the sector model of disk *i*. The example below illustrate this by putting the two sector models into a single one prior to the rotor assembly.

```
fe_cyclicb .
```

```
cf=demo_cyclic('buildstep0'); % See sdtweb demo_cyclic('Step0')
sectors=cf.Stack{'disk1'}; % Build a model with two sectors
sectors=feutil('addtest',sectors,cf.Stack{'disk2'});
sectors.Stack={};
cf.model=sectors;
```

```
% build rotor from sectors and auto display
fe_cyclicb('DiskFromSector epsl 1e-3',cf,{'group1:2','group3:4'});
```

During the build process, sectors are automatically renumbered so that node numbers are left interface, interior, right interface (in order matching that of the left interface). The renumbering can be forced with the **-renumber** option. This allows to have nodal overlap between the superelements of two adjacent sectors. The command then adds a mpc,diskiend multiple point constraint to account for the fact that the disk is closed circumferentially.

Mesh

Meshing utilities See fe_cyclicb Mesh.

ConnectionRing

ConnectionRing builds a "ring connection" where the structure is fixed axially and radially on a set of nodes and first point only in tangential direction

Display

Display commands group tools to build mesh views specific to disk assemblies.

• [def,ENER] = fe_cyclicb('Displaysel', cf,def,Sel, 'enerkdens') is used to recover monoharmonic solutions on a partial selection For details on monoharmonic solutions see section 3.5 . Examples can be found in section 3.5.2, section 3.5.3, ...

 $\tt Sel$ is a cell array specifying how each stage is displayed. In the example from section 3.5.1 , one uses

```
Sel={'', 'EltName SE'; % Keep only SE for display (no interstage rim)
    'disk1(1:2)', 'groupall'; % all elements from sectors 1:2
    'disk2(1:3)', 'groupall'};% all elements from sectors 1:3
```

See sdtweb fesuper#SeBuildSel for details on partial superelement display and more examples on the way to define Sel). The last command can be any valid fe_stress command. Without output argument the result is displayed.

```
[cf,def]=demo_cyclic('buildstep1');
def=fe_def('subdef',def,def.data(:,2)>=0); % remove fixed edge solutions
fe_cyclicb('Display',cf,def);fecom('ColorDataEvalA');
```

```
fe_cyclicb('DisplaySel',cf,def,cf.Stack('ViewMesh'));
fecom('ColorDataEvalA');
```

During the command one defines SE.cGLO corresponding to a rotation by one sector. And the SE.Alpha for the harmonic shift. fesuper('sedefinit -rot',cf); is then used to define a restitution by disk. The SeRestit then contains the def so that SeDefinit is performed for every restitution.

- fe_cyclicb('Display', cf, def) defines a full disk selection within feplot. def=fe_cyclicb('Disp is used to recover full motion from mono-harmonic solutions. For large models, restitution on the full shaft model may be very costly (remember that one vector for 1e6 DOF requires 7.6 MB) and DisplaySel is typically preferred.
- $fe_cyclicb('DisplayAllEdges', cf[,sel])$ displays 2D cuts of the disks specified in the cell array sel, with a typical entry {'diski'} to display only disk *i* (default displays all disks). This cut is basically the projection of the right interface (or equivalently the left interface when meshes are compatible) to the plane $\theta = 0$. Such a view is particularly well suited to the definition of the inter-stage rim nodes in MeshAddRim as well as to the construction of viewing meshes in MeshRimLine2Patch. Note finally that this view keeps the elements in their original groups.

```
cf=demo_cyclic('buildstep0');
fe_cyclicb('DisplayAllEdges',cf);
fecom('colordatag');
```

• fe_cyclicb('DisplayFirst', cf[,sel]) provides a simple command to display the first sector of each stage (cf can be replaced by a model resulting from fe_cyclicb DiskFromSector). A selection can also be specified to restrict the view to a subset of stages.

```
cf=demo_cyclic('buildstep0');
fe_cyclicb('DisplayFirst',cf,{'disk2'});
```

• fe_cyclicb('DisplaySkin', cf[,sel]) displays the outer enveloppe of the selected disks in sel with the inter-sector common surfaces removed.

```
cf=demo_cyclic('buildstep0');
fe_cyclicb('DisplaySkin',cf,{'disk1'});
fecom('showpatch');set(cf.o(1),'FaceAlpha',.33);
```

fe_cyclicb _

• fe_cyclicb('DisplayInterDisk', cf, nodes) displays the two ring surfaces used to define the inter-stage volumic interface.

```
cf=demo_cyclic('buildstep0');
% rim nodes
n1=feutil('getnode NodeId',cf.mdl,[12 18 24 1127 1133 1139]');
fe_cyclicb('DisplayInterDisk',cf,n1);
```

• fe_cyclicb('DisplaySymmetrySurface', cf) displays the nodes in a cyclic symmetric condition as colored surfaces.

Mono-harmonic solutions

These commands apply to sector models used to compute mono-harmonic eigensolutions.

ShaftEig, ShaftTEig, ShaftSolve

These commands compute mono-harmonic solutions with specified Fourier harmonics (classical cyclic solution for single stage models). For a *tutorial on generating the proper models*, see section 3.1.6. For the associated theory, please refer to [9].

The calling format is def=fe_cyclicb('shaft Teig delta_list', model);. ShatTeig accepts multiple diameters in the delta_list and packages individual calls that to ShatTeig. For a disk example section 3.5.2.

The main command options are

- Diameter -1 is used to ask for the computation of fixed edge modes.
- The eigenvalue options 'info', 'EigOpt' should be set in the model stack.
- -ReAssemble forces reassembly rather than reuse of disk matrices that may have been precomputed and saved in the sector superelements.
- -thermal takes thermal loading into account. Thermal state should be stored as a case entry called DofSet, ThermalState. See example in section 3.5.1.
- for computations at prestressed state curve, StaticState should be defined.
- -all xxx

- -FixTan is used to enforce no tangential motion of one interface node. This is used for static analysis of freely rotating rotors (example in section 3.5.1).
- -NoN stores the imaginary component of the eigenvector using a DOF shift by 50 (thus .51 is the imaginary x translation). This is necessary if further computations require complex fields (by considering different components there is no difficulty storing the spatial Fourier transform of a complex field).
- model.Dbfile when the field is defined to a proper file name, intermediate matrices above the preference getpref('SDT', 'OutOfCoreBufferSize', 100) (in MB) are stored in the database file which uses the standard HDF5 based .mat format (MATLAB i = 7.3).
- -nlstep tol is used to compute the large non-linear large transformation problem with a tolerance $\delta q/q < \text{tol}$. See an example in section 3.5.1.
- -soft if present include centrifugal softening effects
- For static computations, the centrifugal load is rebuilt at each step using model=fe_cyclic('loadcer for the rim and sectors.
- -batch option is used to compute eigenvectors with multiple diameter in a single run. For large models, this can take many hours so that intermediate file saves are used to allow restarts.

One then typically expects to have set a cf.mdl.Dbfile='file_DB.mat' to allow memory off-loading during the computation.

```
cf=demo_cyclic('testrotor 7 -blade -cf 2');
root=fullfile(sdtdef('tempdir'),'Disk_7_Batch')
%setpref('SdtRuntime','ExecLocal',1) % May be needed
% cf.mdl.Dbfile=[root '_DB.mat']; % out of core matrices
fe_cyclicb('shaft Teig 0 1 5 -batch',[root '.mat']);
% multiple Disk_7_batch_diam.mat files are generated
% Now reload pointers to selected solutions
R0=struct('Fmax',8000,'diams',[0 1 5]);
d1=fe_cyclicb('DefList',root,R0);
```

ShaftSeAssemble

fe_cyclicb('ShaftSeAssemble -force', cf.mdl, fname) is used to assemble superelement matrices of each of the disk* superelements. If a curve, StaticState is defined in the model stack, fe_cyclic assemble is used, otherwise SE=fe_mknl(SE, 'NoT') is called. If the -force option is omitted an attempt to reload the variable Stack_SE_disk*i* from the file is first made and assembly is only performed if that variable does not contain the matrices.

If a mdl.Dbfile field is defined, the argument fname may be omitted.

-reset xxx.

Deflist,Def ...

Cyclic symmetry results can be stored in three main forms

- the basic form, valid for real vectors only, stores real and imaginary components for the spatial fourier transform as real and imaginary components at a single DOF
- the long vector form uses additionnal DOFs shifted by 50. Starting from the basic form one would have d1.DOF=[d1.DOF;d1.DOF+.5]; d1.def=[real(d1.def);imag(d1.def)]

The test for usage of this format is that the last dof is above .05 rem(d2.DOF(end),1)>.5.

• the double vector form uses the nominal DOFs of the first sector but store the real and imaginary parts as consecutive vectors.

d1.def=[real(d1.def(:,1)); imag(d1.def(:,1)]; d1.data=[d1.data(1,:); NaN NaN]

DefDouble, DefLong, DefBasic commands allow transformations between formats while handling out-of-core files properly.

When reading results, def=fe_cyclicb('DefList', 'root'); reads all root*.mat files and combines the vectors into a single deformation set in the double vector format. Selection of diameters and frequency range during the read process is peformed using

```
RO=struct('Fmax',8000,'diams',[0 1 5 18]);
d1=fe_cyclicb('DefList','root',RO);
```

Specific cases require to sort the output vectors according to the list of diameters specified in the .diams field, especially when one wants to put fixed interfaces solutions first for reduction purposes. The command to use becomes DefListSortDiam.

Full rotor SE model

ShaftPrep

fe_cyclicb('ShaftPrep', cf, def); generates reduced sector super-elements.

Each bladed sector is divided into two regions. A first super-element is attached to the elements with the left interface nodes, it is called the inter-sector super-element. A second one is attached to the remaining elements to form the sector super-element.

A reduced kinematic subspace of the sector super-element (with the definition above) is built from def, disk by disk.

Vectors are first sorted with respect to their contribution to the considered disk if the -svdtruncate option is used.

Then, they are sorted according to their contribution to subsets of physical DOF of the initial sector. If one specifies <code>-mseq 0</code> (default call), these subsets are

- 1. DOFs within either the inter-sector interface (right interface) or the inter-stage interface(s),
- 2. remaining DOFs (left interface and interior DOF).

If -mseq 1 is enforced, these subsets are

- 1. DOFs common to the inter-stage and inter-sector interface(s),
- 2. DOFs within the inter-sector interface (right interface),
- 3. DOFs within the inter-stage interfaces,
- 4. remaining DOFs (left interface and interior DOFs).

Both these sortings make the subsets of vectors linearly independent from each other. They require that fixed edge solutions are stored at the beginning in def.

The following step is to make the vectors linearly independent within each set. Vectors in sets (1.), (2.) and (4.), when applicable, are processed with an Iterative Maximum Sequence Algorithm ([10]). Vectors in set (3.), when applicable, are processed with a Singular Value Decomposition.

See Ref. [5] for details.

-handle option controls whether the resulting bases are stored in memory or on the disk.

-norestit suppresses the explicit construction of the *Restit* variable, normally stored in cf.mdl.Stack.

Once the sector superelements have been generated, the disk model is assembled using the subsequent fesuper('fassemble',cf) call which generates the reduced disk model in cf.Stack{'SE','MVR'}.

A compact example is provided below. A fully developed example can be found in demo_cyclic ShaftMulti.

```
[cf,def]=demo_cyclic('buildstep1');
fe_cyclicb('ShaftPrep -svdtruncate -mseq1 -norestit',cf,def);
fesuper('fassemble',cf);
```

ShaftLoad, ShaftSELoad

fe_cyclicb('ShaftLoadMulti', cf,data); generates a reduced mono-harmonic load for each disk specified in data. It is a cell array whose a typical line is {'diski',def_i,delta_i} where for each disk i, the shape def_i and harmonic coefficient delta_i are specified.

fe_cyclicb('ShaftSELoad',cf,def); generates a reduced load from its physical counterpart. In
practice, it is used for very specific loading cases, e.g. single DOF load or random load.

Developed examples are presented in demo_cyclic ShaftMulti.

```
% define model
cf=demo_cyclic('buildstep4');
% build the excitation on first all sectors, with specified
% diameters 3 on disk1 and 4 on disk 2
data={'disk1',cf.Stack{'disk1'}.Stack{'Enforce_mode_7'},3;
      'disk2', cf.Stack{'disk2'}.Stack{'Enforce_mode_7'},4
      'disk1l',struct('def',1,'DOF',cf.Stack{'disk1l'}.Elt(2,1)+.01),3;
      'disk2l', struct('def',1,'DOF', cf.Stack{'disk2l'}.Elt(2,1)+.01),4};
def34=fe_cyclicb('ShaftLoadMulti',cf,data);
def34.def=sum(def34.def,2); % one column in def34.def per row in data
def34.data=[0 0];
fe_cyclicb('fourier 1 -red -cf 1',cf,def34); % checkout shape
% now keep the same shapes but force delta=0 on both disks
data(:,3)=\{0\};
def00=fe_cyclicb('ShaftLoadMulti',cf,data);
def00.def=sum(def00.def,2); % one column in def00.def per row in data
def00.data=[0 0];
fe_cyclicb('fourier 1 -red -cf 1',cf,def00); % checkout shape
% now try a random load
r1=fesuper('fnode',cf.mdl);
r1=[r1(:,1)'+.01;r1(:,1)'+.02;r1(:,1)'+.03];
defrnd=struct('DOF',r1(:),'def',[]);
defrnd.def=randn(length(defrnd.DOF),1);
```

```
defrnd=fe_cyclicb('ShaftSELoad',cf,defrnd);
defrnd.data=[0 0];
fe_cyclicb('fourier 1 -red -cf 1',cf,defrnd); % checkout shape
% now consider a physical load on first sector of disk 1
r1=fesuper('fnode',cf.mdl);
r1=[r1(:,1)'+.01;r1(:,1)'+.02;r1(:,1)'+.03];
defsp=struct('DOF',r1(:),'def',[]);
defsp.def=fe_c(defsp.DOF,1+[.01;.02;.03])'*randn(3,1);
defsp=fe_cyclicb('ShaftSELoad',cf,defsp);
defsp.data=[0 0];
fe_cyclicb('fourier 1 -red -cf 1',cf,defsp); % retrieve a Dirac's comb
```

ShaftFRF [, D, MS, -rest]

This command allows to build the Frequency Response Functions of a rotor model, either full or reduced. A load and a set of observation DOF have to be defined and added to the model with fe_case. The frequency range is stored in the stack as a 'info', 'Freq' entry.

The general call is

```
xF=fe_cyclicb('ShaftFRFD',disk,lossfac)
xF=fe_cyclicb('ShaftFRFD -rest',disk,lossfac,cf,sel)
xF=fe_cyclicb('ShaftFRFMS',disk,def,damp)
xF=fe_cyclicb('ShaftFRFMS -rest',disk,def,damp,cf,sel)
```

The command FRFD assembles the matrices of the model then uses them to compute the response. An optional loss factor can be specified.

The command FRFMS synthetizes the response from a set of modeshapes. A damping ratio for all modes can be specified.

The option **-rest** restores the response computed on the reduced model to a given selection of physical DOF. Without selection, the response is restored to the whole physical DOF set. This option must be disabled when dealing with a full rotor model.

The example developed in demo_cyclic ShaftMulti builds the sythetized response of a reduced rotor model to a random excitation.

```
% define model
[cf,load,def]=demo_cyclic('buildstep5');
```

% Compute the response to the random excitation

```
mdl=fe_case(cf.Stack('mvr'), 'dofload', 'Load', load);
mdl=stack_set(mdl,'info','Freq',linspace(900,2000,2201));
xF=fe_cyclicb('shaft frfms',mdl,def,.001);
% select where to restore (upper blade corner)
Sel1={'','eltname SE';'disk1','selface & withnode{NodeId 154}'};
Sel1=fesuper('SeBuildSel -initrot', cf, Sel1);
Sel2={'','eltname SE';'disk2','selface & withnode{NodeId 154}'};
Sel2=fesuper('SeBuildSel -initrot', cf, Sel2);
% ... and do restore
xF1=fesuper('SeDef',Sel1.cna{1},xF);
xF2=fesuper('SeDef',Sel2.cna{1},xF);
% plot responses
ci=iiplot(3);
XF=iicom(ci,'curveXF');
XF('Disk1')=struct('w',xF.data,'xf',xF1.def','dof',xF1.DOF);
XF('Disk2')=struct('w',xF.data,'xf',xF2.def','dof',xF2.DOF);
iicom('subMagPha')
iicom(ci,'IIxOnly',{'Disk1','Disk2'});
ii_plp(def.data);
```

Fourier [,ind], ... [-phys, -mono, -red, -egy, -egyfrac, -sortbyd]

This command allows to perform a 3D Fourier analysis of given modeshapes. The maximum norm of each harmonic is plotted against the harmonic coefficient. The plot is different when dealing with a single modeshape or a set of modeshapes.

Accepted options are

- ind is an optional selection of deformations. See also the alternate fe_def SubDef.
- -phys, -mono and -red are used to distinguish between an analysis of physical modeshapes (full 3D), mono-harmonic modeshapes (the DFT step is omitted) and generalized modeshapes (reduced multi-harmonic model). In all cases, the user has to check that the physical or reduced models are geometrically periodic, *i.e.* that DOF come in repetitive groups, except for -mono where the concept of mono-harmonic modeshapes assumes that structures are periodic.
- -egy and -egyfrac provides means for energy-based computations. Option -egy displays the fraction of energy in each existing harmonic within each disk so that the total amount of

energy within each disk is 1. Option -egyfrac displays the fraction of energy in each existing harmonic within each disk so that that total amount of energy within the rotor is 1. The default displays these quantities disk by disk (from top to bottom) and for each disk, all the possible harmonics are displayed (from bottom to top), as depicted in the figure below.

• -sortbyd groups these quantities first by harmonics (from bottom to top) and then by disk (from top to bottom), with the appropriate number of zeros for non present harmonics (typically when $\delta > N/2$ for a given disk), as displayed in the figure below.

```
[cf,def]=demo_cyclic('buildstep1'); % sdtweb demo_cyclic('step1')
Curve=fe_cyclicb('fourier 1:50 -mono -egyfrac',cf,...
fe_def('subdef',def,def.data(:,2)~=-1));
'xxx'%fe_cyclicb('fourier -mono -egyfrac -cf 3',Curve);
% sdtweb demo_cyclic('step4') multi-harmonic analysis
[cf,def]=demo_cyclic('buildstep4'); %
fe_cyclicb('fourier 7:25 -red -egyfrac -cf 11',cf,def);
fe_cyclicb('fourier 7:25 -red -egyfrac -sortbyd -cf 13',cf,def);
```

See also

fe_cyclic

obsolete

Purpose

Obsolete functionality

Syntax

```
model=fe_cyclicb('Basis [, -norm, -all, -int, -rb]',model,orders,omegas,opt)
def=fe_cyclicb('DiskEig',DISK)
def=fe_cyclicb('DiskEngineLoad EO',model)
def=fe_cyclicb('DiskFRFD [, -rest]',DISK,lossfac,cf,sel)
def=fe_cyclicb('DiskFRFMS [, -rest]',DISK,def,damp,cf,sel)
K=fe_cyclicb('DiskMatrices [ mk]',DISK,Eltselection)
fe_cyclicb('Fourier MODENUM [, -phys, rotor, disk, -red, -test TOL]',fignr,model,def)
model=fe_cyclicb('PolyAssemble [, NoT]',model,params)
DISK=fe_cyclicb('Reduce NODEIDO ELTIDO [, -int]',sector,def)
model=fe_cyclicb('Renumber',model)
def=fe_cyclicb('Display',cf,def)
def=fe_cyclicb('ShaftEngineLoad EO',model)
def=fe_cyclicb('ShaftFRFD [-rest]',model,lossfac,cf,sel)
def=fe_cyclicb('ShaftFRFMS [-rest]',model,def,damp,cf,sel)
fe_cyclic('ShaftPrep',model,def)
mdl=fe_cyclic('ShaftRing [-round N1 N2 N3 -autoclose N4]',rim1,rim2)
fe_cyclicb('ShaftLoad',model)
[def,mdl]=fe_cyclicb('ShaftTEig ORDERS',model)
```

OBSOLETE Basis [, -all, -norm, -int, -rb]

This command allows to build a set of modes :

- with the harmonics specified in an array $[\delta_1, \delta_2, ...]$,
- for the rotation vectors specified in a cell $\{ [\omega_{1x}, \omega_{1y}, \omega_{1z}], [\omega_{2x}, \omega_{2y}, \omega_{2z}], ... \}.$

The general call is

[model,def]=fe_cyclicb('basis -all -int -norm -rb',sector,orders,omegas);

The output is a struct array containing the modeshapes. If only one output is required, the basis is added to the model as a TR field. The field data refers to the harmonic in column 2 and the rotation speed in column 3. The number of computed modes is controlled by the field 'info', 'EigOpt' in the stack.

```
cf=demo_cyclic('testload disk 5 -nor reset') % reset file (rather than load)
% Set of Cyclic Modes / Fixed Interface Modes
sector=stack_set(cf.Stack{'disk1'},'info','EigOpt',[5 4 0 11 1e-8]);
sector=stack_set(sector,'info','EigOptFixInt',[5 4 0 11 1e-8]);
sector=fe_cyclicb('basis -all -int -norm',sector,[0:3],...
{[0 0 0],[0 0 250]});
cf.model=sector;cf.def=sector.TR;
```

The -all option is used to get both modeshapes associated with a double eigenvalue in the case where harmonics are not 0 or half the number of sectors, when applicable. For more information on cyclic symmetry, please refer to [...].

The -norm option ensures that modes are orthonormalized in mass and stiffness because of convergence problems caused by the presence of double eigenvalues. This option is not required when the eigenvalue problem is solved with Nastran (fe_eig method 50). xxx discuss with EB

When the -int option is added, modes of the initial sector with its left and right interfaces fixed (clamped) are also computed and added at the beginning of the output. These modes have a -1 in the field data. The computation parameters are specified in the 'info', 'EigOptFixInt' stack entry. When this option but no harmonic are given, it computes only fixed interface modes.

The six rigid body modes of a cyclic symmetric structure are mono-harmonic with harmonic 0 (1T along and 1R around the axis of symmetry) and harmonic 1 (2T along and 2R around the other axes). Thus, the -rb option is used to compute two more flexible modes with harmonic 0 and four more flexible modes with harmonic 1.

OBSOLETE DiskEig [, -ord]

This command allows to compute the approximate modes of a reduced disk model built with the command Reduce of fe_cyclicb.

Command fails and is no longer maintained zzz see with arnaud

cf.def=fe_cyclicb('Display',cf,def);

```
cf=demo_cyclic('testload disk 5 -nor') % reload model
fe_cyclicb('reduce 1 1 -int',cf);disk=cf.Stack{'diskmodel'};
% Mode computations
disk=stack_set(disk,'info','EigOpt',[5 30 1e3 11 1e-8]);
[def,disk]=fe_cyclicb('diskeig -ord',disk);
disp(def.data);
```

The -ord option is used to identify the Fourier harmonic coefficient associated with each mode when dealing with the reduced model of a tuned disk. When dealing with a mistuned disk (whose modes are multi-harmonic), this returns the coefficient whose harmonic is the greatest.

DiskEngineLoad EO [, sel]

This commands builds a physical load, spatially mono-harmonic, on a specified set of nodes. If no selection is present, all nodes are used.

```
% Model Initialization
cf=demo_cyclic('testdisk 7 -blade -cf 2');
% External Load
Load=fe_cyclicb('DiskEngineLoad 3 r > 201',cf);
cf.def=Load;fecom showdefarrow;
```

OBSOLETE DiskFRF [, D, MS, -rest]

This command allows to build the Frequency Response Functions of a disk model, either full or reduced. A load and a set of observation DOF have to be defined and added to the model with fe_case. The frequency range is stored in the stack as a 'info', 'Freq' entry.

The general call is

```
xF=fe_cyclicb('DiskFRFD',disk,lossfac)
xF=fe_cyclicb('DiskFRFD -rest',disk,lossfac,cf,sel)
xF=fe_cyclicb('DiskFRFMS',disk,def,damp)
xF=fe_cyclicb('DiskFRFMS -rest',disk,def,damp,cf,sel)
```

The command FRFD assembles the matrices of the model then uses them to compute the response. An optional loss factor can be specified.

The command FRFMS synthetizes the response from a set of modeshapes. A damping ratio for all modes can be specified.

The option **-rest** recovers (xxxEB recovers, expands, interpolates, ...) the response computed on the reduced model to a given selection of physical DOF. Without selection, the response is expanded to the whole physical DOF set. This option must be disabled when dealing with a full disk model.

The following example builds both direct and sythetized responses of a reduced disk model to a 2EO excitation.

```
cf=demo_cyclic('testload disk 5 -nor') % reload model
```

```
% the call to fe_cyclicb basis is already done
fe_cyclicb('reduce 1 1 -int',cf);disk=cf.Stack{'diskmodel'};
% External Load
Load=fe_cyclicb('DiskEngineLoad 2',cf);
fe_cyclicb('DiskSeDefInit',cf);
Rload=fe_cyclicb('DiskSeLoad', cf, Load);
disk=fe_case(disk,'dofload','Blade_load',Rload);
freq=[1500:5:3000]';
disk=stack_set(disk,'info','Freq',freq);
% Restitution to Blade Dofs
tips=feutil('FindNode r>201',cf.mdl);
disk=fe_case(disk,'SensDof','Sensors',disk.DOF);
% Transfert Functions / Direct
xFd=fe_cyclicb('disk frfd -rest',disk,.002,cf,tips);
% Transfert Functions / Modal Synthesis
disk=stack_set(disk,'info','EigOpt',[5 50 -1e3 11 1e-8]);
[def,disk]=fe_cyclicb('diskeig',disk);
xFms=fe_cyclicb('disk frfms -rest',disk,def,.001,cf,tips);
% Response Plots
ci=iiplot;
XF=iicom('curveXF');
XF('Blade resp.D')=struct('w',xFd.data,'xf',xFd.def.','dof',xFd.DOF);
XF('Blade resp.MS')=struct('w',xFms.data,'xf',xFms.def.','dof',xFms.DOF);
iicom('subMagPha')
iicom(ci,'IIxOnly',{'Blade resp.D','Blade resp.MS'})
```

OBSOLETE DiskPlot

This commands provides a graphical representation of the generalied quantities (modeshapes or load) on a patch whose nodes represent the generalized DOFs.

The following example deals with such representation for both the modes and a 3EO excitation.

```
% Model Initialization
```

obsolete .

```
cf=demo_cyclic('testload disk 5 -nor reset')
% Model Reduction
fe_cyclicb('reduce 1 1 -int',cf);disk=cf.Stack{'diskmodel'};
% Mode Computations
disk=stack_set(disk,'info','EigOpt',[5 50 -1e3 11 1e-8]);
[def,disk]=fe_cyclicb('diskeig',disk);
fe_cyclicb('diskplot',3,cf,def);
% External Load
Load=fe_cyclicb('DiskEngineLoad 3',cf);
fe_cyclicb('DiskSeDefInit',cf);
Rload=fe_cyclicb('DiskSeLoad',cf,Load);
fe_cyclicb('diskplot',4,cf,Rload);
```

```
OBSOLETE Disk ... (internal commands)
```

DiskRest computes the Fourier Recovery Matrix associated with the specified harmonic. This function is used internally by fe_cyclicb.

DiskMatrices returns the reduced matrices assembled from the sector superelement matrices. When the option mk is specified, it returns the mass and stiffnes matrices only.

```
cf=demo_cyclic('testload disk 5 -nor reset') % reload model
fe_cyclicb('reduce 1 1 -int',cf);disk=cf.Stack{'diskmodel'};
```

```
Kr0=fe_cyclicb('diskmatrices',disk);
disk.il=[1001 1 1 250^2 250^4];
Kr=fe_cyclicb('diskmatrices mk',disk);
```

PolyAssemble [, noT] supports the computations of the coefficients of the matrix polynomial from the assembly of the stiffness matrices (including gyroscopic and nonlinear tangent stiffnesses) at the given values of the parameters. Three values are required.

The following example assembles the mass and full stiffness matrices at 0, 500 and 1000 rps, then computes the modes of the free sector at 250 rps.

```
% Model Initialization
cf=demo_cyclic('testdisk 5 -blade noK -nor -cf 2');
sector=fe_cyclicb('polyassemble noT',cf.Stack{'disk1'},[0 500 1000]);
```

```
[Case,sector.DOF]=fe_mknl('init',sector);
K={sector.K{1},feutilb('sumkcoef',{sector.K{2:4}},[1 250^2 250^4])};
% Mode Computations
def=fe_eig({K{1},K{2},Case.T,sector.DOF},[5 10 0 11]);
cf.model=sector;cf.def=def;
```

OBSOLETE DiskSeDefInit

This command initializes the restitution of the generalized modes computed with the DiskEig command on the physical DOF set. The restitution bases are stored in the Stack as a 'info', 'SeRestit' entry which contains the following fields:

- a field .DOF that contains the physical DOF set,
- a field .adof that contains the generalized DOF set,
- a cell .Restit where the first column gives the indices in the physical DOF vector associated with each sector, the second column gives the indices in the generalized DOF vector associated with each sector and the third column contains the transformation matrix from physical to generalized DOF coordinates.
- a cell . cGL that contains one matrix per sector which is a local to global frame transformation matrix.

The restitution is performed with the command SeDef of fesuper. In this command, a patch can be defined as a selection of elements so that the modeshapes are expanded only to the physical DOF of their nodes.

The following example builds a reduced model, computes a set of generalized modes, then the modeshapes are recovered on the blades only.

```
% Model Initialization
cf=demo_cyclic('testload disk 5 reset') % reset file (rather than load)
% Mode Computations
def=fe_eig(cf.mdl,[5 20 -1e3 11 1e-8]);
% Restitution
cf.def=fesuper('SeDef',cf,def);
```

OBSOLETE DiskSeLoad

This commands transforms an external load, expressed on the physical DOF set, into a generalized load, expressed on the generalized DOF set. It is necessary to initialize the restitution of the underlying disk model, since this transformation is the inverse operation.

```
% Model Initialization
cf=demo_cyclic('testload disk 5 -nor')
%xxx obsolete
%Model Reduction
fe_cyclicb('reduce 1 1 -int',cf);
% External Load
Load=fe_cyclicb('DiskEngineLoad 1',cf);
fe_cyclicb('DiskSeDefInit',cf);
Rload=fe_cyclicb('DiskSeLoad',cf,Load);
```

Reduce [, -int]

The **Reduce** command is used to generate a disk model from a set of cyclic modes associated with multiple harmonics. The general call is

```
[disk,SEsector]=fe_cyclicb('reduce NodeId0 EltId0 [-int]',model);
fe_cyclicb('reduce NodeId0 EltId0 [-int]',cf);
```

The second call uses directly a global model stored in the variable cf and stores the reduced model as an entry 'info', 'diskmodel' in cf.Stack.

If required, new starting points for numbering the generalized DOF and the associated elements are defined in the command string through the two integers NodeId0 and EltId0.

The initial set of modes has to be given as a field TR of the model. This operation can be performed directly by using the command Basis of fe_cyclicb, as shown in the following script.

```
% Model Initialization
cf=demo_cyclic('testload disk 5 -nor -cf reset');
fe_cyclicb('reduce 1 1 -int',cf);
disk=cf.Stack{'diskmodel'};
```

The reduction basis is built by separating right, left and interior motion from the cyclic modeshapes. When the -int option is invoked, fixed interface modes of the sector are added to the set of interior modes. For more information about this procedure, refer to [11].

nl_spring

Purpose

Non linear links/force modeling for time simulation

Syntax

```
model=nl_spring('tab',model);
...= nl_spring('command', ...)
```

Description

nl_spring supports non-linear connections and loads for transient analysis. Non linear springs between 2 DOF (see nlspring). loads which depend on DOF values (see DofKuva, DofV), springs between 2 nodes in different bases (see RotCenter), etc. ...). A full list of non-linearities is given in nllist

Standard non-linear simulations are handled by nl_solve. Below is a description of the inner mechanisms of a non-linear simulation with the non-linear toolbox.

After the non linearity definition, a proper TimeOpt is required to set the good fe_time calls to perform a non linear Newmark time integration. A default TimeOpt can be set using nl_spring TimeOpt. It is possible to save transient results on the fly using a properFinalCleanup call, see nl_spring fe_timeCleanupCall, and to reload the same results using fe_simul fe_timeLoad.

The following steps are required for a time simulation

• Definition of non-linear properties. These are stored as pro entries of the model stack. The associated property function must handle non-linearities which is currently only the case for p_spring and p_contact.

A non-linearity is always associated with elements or superelements (typically a celas element. A given group of elements can only be associated with a single non-linearity type.

The information needed to describe the non linearity is stored in a .NLdata field.

- Model initialization using the an fe_case('assemble') call in fe_time, is followed by the building of a model.NL stack that describes all non-linearities of the model in a format that is suitable for efficient time domain integration. This translation is performed by the nl_spring NL command.
- Jacobian computation, see nl_spring NLJacobianUpdate.
- Residual computations are performed through mkl_utils. The nominal residual call is r=-fc; mkl_utils('residual', r,model,u,v,a,opt,Case);.

Supported non linearities

See nllist for supported non linearities, and nl_fun to add your own non-linearities.

ConnectionBuild

One can define a set of non linear links between 2 parts of a model using a call of the form [model,idof]=nl_spring('ConnectionBuild',model,data);

idof is a second optional out argument. It returns the list of DOF concerned by links (it can be useful in order to reduce super elements keeping idof as interfaces DOF for instance). data contains all the information needed to define links. It is a 3 column stack like cell array. First column contains the string 'connection', the second the name of the non linear link described in the third column that contains a data structure with following fields:

- .Ci define nodes to connect in first (.C1) and second component (.C2). It can be a vector of NodeId or a screw data structure (slave nodes of the model nodes via RBE3 links, see see sdtweb('fe_case#connectionscrew').
- .link defines how to link component 1 to component 2. It is a 1x2 cell array. First cell defines the type of link ('EqualDof' or 'Celas') and the second gives information about the link. For celas link it is a standard element matrix row with 0 replacing NodeId :[0 0 DofId1 DofId2 ProId EltId Kv Mv Cv Bv].
- .NLdata (optional) defines non linearity associated to celas link. See the list in list of supported NL. If this field is not present or empty, only linear link is considered.
- .PID (optional) is a 1x2 line vector that defines PID (second column of .Node matrix, see sdtweb('node') of connected node (1rst column for 1rst component).
- .DID (optional) is the same as above, defining DID (third column of .Node matrix, see sdtweb('node') of connected nodes.

Following example defines a model with a cylinder and a hole in a block. The cylinder is linked to the block by 3 celas preserving the pivot link.

```
mo1=demosdt('demoConnection-vol'); % meshes models
mo1=fe_case(mo1,'fixdof','base','z==-1'); % clamps the cylinder base
r1=struct('Origin',[0.5 0.5 0.5],'axis',[0 0 1],...
'radius',.1,'rtol',.01,'length',1,'Npt',-3,...
'ProId',111,'planes',[]); % Cylinder-side
r1=nl_spring('ConnectionCyl',r1); % defines planes
```

See also t_nlspring('2beam') example.

ConnectionCyl

Utility to fill the .planes field of a cylinder connection in the standard connection screw data structure format (see fe_caseg ConnectionScrew).

dataOut=nl_spring('ConnectionCyl',dataIn);

The dataIn uses fields:

- .Origin origin of the cylinder axis, .axis orientation of the cylinder
- .rtol radius tolerance for cylinder selection.
- .length length of the cylinder.
- .Npt number of planes (equally distributed on the whole length). If Npt<0, ends of the cylinder are included in the connection points.
- .ProId ProId of the elements containing nodes to connect.



Figure 5.1: ConnectionCyl

InitV

q0=nl_spring('InitV',model,d0,R0);

InitV computes the initial static displacement and velocity associated to a DOF initial position and velocity. d0 is a data structure with field .DOF containing the DofId where initial value is applied and .def containing initial displacement and velocity at this DOF. RO is a optional input argument data structure with following fields that define:

- .dt time step for time integration.
- .dq increment for initial vel computation.
- .Nv] number of time steps to reach d0.def(1) (displacement is imposed as a 0.5(1 cos) time function on these time steps).
- .Np number of steps to stabilize at d0.def(1) and d0.def(1)+dq.

If input argument RO omitted, options are get from 'info' 'initvopt' Stack entry. If there is no such entry, InitV parameters are computed using -optim process (see below).

Displacement at q0 and q0+dq is obtained meaning the last Np/10 steps of each stabilization period, and initial velocity is computed from those 2 displacements to match d0.def(2) at d0.DOF.
[q0,R0]=nl_spring('InitV-optim',model,d0); can be used to find input parameters R0. Optimization of dt and Np is performed from given or default values. Parameters dq and Nv are kept at given or default value. First dt is optimized. dt is increased (multiplied by 4) until time integration of the InitV process diverge and last dt that leads to convergence is kept. Then Np is increased by 100 steps until the deformation is converged on the stabilization periods, that is to say that a criteria taking in account standard deviation/mean of the deformation and the ratio of the last Np/10 steps upon previous Np/10 steps on each Np period is less than a tolerance (2.0).

See also t_nlspring('2beam') example.

NL

model=nl_spring('NL',model)

This command is used to build .NL field data for time integration from NLdata field in NL p_spring property entries in the input model Stack. The command option -storefnl can be used to specify the way of computing and storing a non linear effort associated to NL (for those which support it).

NLJacobianUpdate

opt.Jacobian=nl_spring('JacobianCall') returns the callback used to update or initialize the Jacobian ki used in iterative methods. This is the low level implementation of calls documented in nl_solve TgtMdl. The said Jacobian must take non-linearities into account and is thus of the form

$$k_j = [b] \left[\frac{\partial s_{nl}}{\partial u_{nl}} \right] [c] \tag{5.3}$$

the output is controlled by the value of NL.Jacobian.

- O gives no Jacobian.
- 1 use finite differences to evaluate Jacobian.
- 2 fixed Jacobian.

For the case of a non-linear spring, the most important gradient of the tabulated law Fu is added as stiffness between the 2 DOF to the stiffness matrix and the most important gradient of Fv to the damping matrix.

For non-linear iterations in a Newmark scheme, the Jacobian is given by

ki=(model.K{3}+kj)+ (opt(2)/opt(1))/dt*(model.K{2}+cj) + 1/opt(1)/dt^2*model.K{1};

nl_spring

Accepted command options, associated to variants of the call are

- There are three outputs accessible, being [ki,mo1,C1]=nl_spring('NLJacobian'...).
- -noFact not to factorize the output Jacobian. This is useful if further actions are performed on the Jacobian after the standard call.
- -TangentMdl to return tangent model. It is assumed that model.K(1:3) correspond to M, C, and K (in this order). u and v variables of caller workspace can be needed.
- -TangentMdl-back to return a superelement containing the tangent matrices.
- -TangentMdl-back-sepKj to return a superelement containing the tangent matrices split by non linearities.
- -ener to compute for each def stored in model.d1 def structure (that is typically computed modes), some associated energies:
 - freq frequency in Hz.
 - damping damping ratio: $(\phi_j^T[C]\phi_j)/(2\omega_j)$.
 - enerK total strain energy: $\phi_j^T[K]\phi_j$.
 - enerC $\phi_j^T[K]\phi_j$.
 - *NLlink*-enerK strain energy for each NL link: $\phi_i^T[K_{NLlink}]\phi_j$.
 - *NLlink*-enerK for each NL link: $\phi_j^T[C_{NLlink}]\phi_j$.

${\tt SetPro}$

model=nl_spring('SetPro ProId i ParamName1 Value1 ...',model)

This command is used to change some nl_spring properties parameters. *i* is the ProId of corresponding p_spring property, *ParamName* the name of parameter to change (k for il(3), c for il(5) or the field name in NLdata) and *Value* the value to assign.

It is possible to define a new property by specifying an NLdata structure in third argument: model=nl_sprinered i', model, NLdata). If the property already exists, the NLdata is interpreted as a string of parameters and parsed to define the fields specified in the given NLdata to the existing one. Command option Edit allows directly merging the existing NLdata to the provided NLdata with priority given to the new fields.

```
model=nl_spring('Demo1DOF');
\% define a non linearity with partial definition of parameters and other by default
NLdata=nl_fun('db data 4') % standard NLdata defintion
% NLdata has fields data, Jacobian (by default) and type
% set in model
model=nl_spring('setpro proid201',model,NLdata);
% edit the nl_fun nl by string keyword
model=nl_spring('setpro proid201 data2',model);
% edit the nl_fun with struct input
% property will be parsed using nl_fun('paramedit')
model=nl_spring('setpro proid201',model,struct('Jacobian',2));
% field Jacobian has been edited, other fields are kept unchanged
model.Stack{end,3}.NLdata
model=nl_spring('setpro proid201',model,struct('NewField','test'));
% you can see that in this case NewField was not set
% as it is not referenced in the nl_fun parameters
model.Stack{end,3}.NLdata
% Force the with struct input with no check
model=nl_spring('setproedit proid201',model,struct('data',10,'NewField','test'));
% in this mode the NewField is propagated regardless of the
% standard nl_fun input
model.Stack{end,3}.NLdata
```

Standard NLdata structures depend on the non-linear function, see nllist for more details. They can be obtained through the nl_function command db, see nl_fun for more details.

In the case where

${\tt GetPro}$

pro=nl_spring('GetPro',model)

This command is used to get non linear properties in the model stack.

- Command option ID allows getting a specific non linear property by specifying its ProId.
- Command option type '`nl_fun'' allows getting the non linear properties of a specific type. See nllist for more details on types of non-linearities.

nl_spring

Follow

The Follow mechanism can be used to observe some variable evolution during the time integration. opt=fe_simul('Followi', opt);

1st Follow consists in monitoring the number of iteration, the residual norm and displacement increment norm at each time step.

```
model=nl_spring('Demo1DOF')
opt=stack_get(model,'info','TimeOpt','GetData');
opt=fe_simul('Follow1',opt); % niter norm(r) norm(dq)
def=fe_time(opt,model);
```

2nd Follow consists in monitoring the def.FNL in iiplot. For the moment the mechanism is different (so note that you can't both tracker niter and FNL), and you only have to specify the field .Fnlliplot equal to 1 in the 'info', 'OutputOptions' stack entry of the input model, as in following example :

```
model=nl_spring('Demo1DOF');
r1=stack_get(model,'info','OutputOptions','GetData');
r1.FnlIiplot=1; % define FNL tracker
model=stack_set(model,'info','OutputOptions',r1);
opt=stack_get(model,'info','TimeOpt','GetData');
def=fe_time(opt,model);
```

TimeOpt

This command returns usual default TimeOpt for non-linear simulations. By default the output is the same as the TimeOptNLNewmark presented below. See also fe_time for TimeOpt definition details.

Supported TimeOpt commands are

- TimeOptNLNewmark, or TimeOpt to obtain the TimeOpt for NLNewmark simulations. Use TimeOpt-gam .51 to introduce numerical damping by directly giving gamma.
- TimeOptStat to perform static simulations (see also fe_time nl_solve).
- TimeOptTheta to perform time simulations with the θ -method (see fe_time). Numerical damping can be introduced using TimeOptTheta-alpha .05, the specified α value will be added to θ , so that the coefficient used in the simulations will be $\theta_1 = \theta + \alpha$.
- TimeOptExplicit to perform time simulations with the explicit Newmark scheme.

The following command options allows setting other TimeOpt fields to their desired value.

- dtval time step.
- tsN number of time steps.
- tendval optional end time
- tInitval initial time.
- AlphaRval a global Rayleigh damping mass coefficient (applied to the model total mass).
- BetaRval a global Rayleigh damping stiffness coefficient (applied to the model total stiffness).
- maxNoutN requests an output subsampling strategy such that only N times equally spread over the simulation time span are output.
- RelTolval requests a specific relative tolerance for the convergence of iterative schemes.
- -gammaval requests a specific γ coefficient (default to .5) of the Newmark scheme. For the non explicit versions, β is adapted to ensure unconditional stability of the scheme.
- -thetaval requests a specific θ coefficient (default to .5) of the Theta method.
- -acallstr provides a series of command options applied to the AssembleCall generation.
- -fcleanstr provides a series of command options applied to the FinalCleanupFcn generation.
- -jcallmodel edits the jacobian call to allow late model modification.

Alternatively to providing all these command options in the command string, one can provide a MATLAB struct with equivalent fields as an additional argument.

By adding an SDT model as third argument, the generated **TimeOpt** will be directly integrated in the model, that will be output.

Sample calls :

```
% basic call
opt = nl_solve('TimeOpt dt1e-6 ts3e5 maxNout1e4 -acall"lumpedMass"');
% call with struct input
RO=struct('dt',1e-6,'ts',3e5,'maxNout',1e4,...
'acall','lumpedMass');
opt = nl_solve('TimeOptExplicit',RO);
% basic call with model input
```

```
model = nl_solve('TimeOpt dt1e-6 ts3e5 maxNout1e4 -acall"lumpedMass"',[],model);
% call with struct and model input
model = nl_solve('TimeOptExplicit',RO,model);
```

Convergence tests depend on the iteration algorithm and several behaviors can be obtained by modifying RelTol. In any case the absolute value of RelTol is used for the convergence test application; its sign is used to determine the convergence test to be used as described in the following.

- For algorithms using iterNewton as IterFcn, as is the case for methods newmark (explicit or not), NLNewmark, and staticNewton.
 - using RelTol > 0 tests the convergence of the mechanical residue, relative to value opt.nf. If opt.nf is not provided, the scheme takes in input the norm of the external forces fc at the first time step, or if zero the norm of the first residue of the first time step. If still zero, opt.nf is set to 1. This convergence test is the most widespread as it ensures mechanical stabilization. It is strongly recommended for static computations, or when using large time steps.
 - using RelTol < 0 tests the convergence of the displacement correction, relative to the current displacement norm. The idea of this mode is to stop iterating if the correction becomes negligible, this is very useful to limit iterations with little impact on the results in transient simulations with small enough time steps. This must be used with care as this criterion does not imply that the mechanical residue is converged at the end of the time step, it is thus strongly advised to check results convergence.</p>

iterNewton does not support the use of opt.cvg yet.

- For algorithms using itertheta_nl as IterFcn, as is the case for method theta,
 - using RelTol > 0 tests the convergence of the velocity field, its correction relative to the previous iteration velocity norm.
 - using RelTol < 0 tests the convergence of the velocity field, and the model.FNL vector, their correction relative to the previous iteration norm. Stabilization of the model.FNL field may be difficult to attain and very sensitive as this vector can contain heterogeneous data, this mode is then not recommended by default, and use of opt.cvg should be preferred.

iterthetal_nl supports the use of opt.cvg, that forces iteration if set to 1. It is reset to zero at the start of each iteration, but any non-linearity can alter its value by using sp_util('setinput',opt.cvg,ones(1),zeros(1));. Each non-linearity can thus internally test the convergence of its fields of interest and apply a convergence veto if its convergence is not satisfied.

From standard fe_time simulations, the following TimeOpt fields are added or modified

- Jacobian field is modified to take into account non linearities, see NLJacobianUpdate.
- Residual field is modified to take into account non linearities, and to use mkl_utils to improve computation times, see sdtweb mkl_utils. This should be initialized by nl_spring('ResidualCall'
- AssembleCall field is modified, to perform non-linearities initialization after assembly. AssembleCal is the string passed to fe_case, generated by nl_spring('AssembleCall').
- OutputInit field is modified to also check non linearities and initialize non-linearities related outputs, this is a callback generated by nl_spring('OutputInitCall').
- FinalCleanUpFcn field is modified to perform cleanup on non linearities as well, this is realized through the ExitFcn command option of fe_simulfe_timeCleanUp (see fe_timeTimeOpt), using '-ExitFcn"nl_spring(''fe_timeCleanUp'')"'. This should be initialized by nl_spring('fe_t
- OutputFcn The output function should be generated by the OutputInit command, since it handles proper interpolation of output as function of the time step, and requires fine tuning in the case of non linear simulations. If nl_spring handles the OutputInit call, OutputFcn is thus reset during initializations. Handling of output time steps using a time vector in OutputFcn is supported.

AssembleCall

The TimeOptAssembleCall must use the -InitFcn callback of fe_caseg Assemble to perform initialization of the non linearities.

Command options are available to tweak the assemble call with minimal user input

- MVR To adapt the assemble call for preassembled reduced models. This typically removes the -load command option of the call as this has to be recovered in the MVR itself.
- skipMKL No to transform the model matrices into mkls objects.
- lumpedMass To adapt the mass matrix mattype to 20 and get a lumped mass matrix.
- compose For more complex calls one can redefine from scratch the assemble call line to which the ad hoc initFcn will be added.

nl_spring

ResidualCall

The **TimeOptResidual** callback should be a call to **mkl_utils**, that performs optimized matrix vector products, and the computation of non linear forces handled by **nl_functions**. Command options allows choosing a call adapted to the type of simulations

- by default a call adapted to the nlnewmark scheme.
- ResidualCallStatic provides a residual adapted to the newton-Raphson schem.
- ResidualCallExplicit provides a residual adapted to the newmark explicit scheme.

fe_timeCleanupCall

The TimeOptFinalCleanupFcn callback must use the -ExitFcn of fe_simulto perform post treatments of non linearities. Custom options classical to the fe_simulFinalCleanup call can be added either in the command string or as a string in second argument.

```
opt.FinalCleanupFcn=nl_spring('fe_timeCleanupCall -cf-1-fullDOF');
% equivalent call with second argument
opt.FinalCleanupFcn=nl_spring('fe_timeCleanupCall','-cf-1-fullDOF');
```

In addition to the standard fe_simulFinalCleanup, the following command options are available (to be specified outside the ExitFcn callback.

- -HDFSave To save the output in a temporary file, and output a v_handle pointer to the saved data. This is useful for RAM optimization matters.
- -HDFfname fname In combination to -HDFSave, to specify the file in which the output will be saved.
- -Save To save the output in a temporary file, but keep the results.
- -fname fname In combination to -Save, to specify the file in which the output will be saved.

OutputInitCall

The OutputInit callback is locked for internal nl_spring use. Several command options are available that will be forwarded to the OutputInit procedure

• -BlockSaveN To initialize a bufferization of the output of size N. Results will be saved as blocks containing each N saved time steps.

- -exit To force exit after initialization. This can be used to check the output format without performing the simulation.
- -postFcn To provide a callback that can tweak the output at the end of the OutputInit procedure. This can be used for example to initialize out.Post post treatments.

TimeOutputOptions

Fine tuning of fe_time output can be achieved by specifying an 'info', 'OutputOptions' case entry.

Accepted fields for the OutputOptions structure are

- .FnlAllT if defined and equal to 1, non-linear loads are saved at all time steps.
- .Fnlliplot if defined and equal to 1, non linear loads are displayed in an iiplot figure as curve FNL. If the display timer associated with this figure does not stop automatically, you can stop it with cingui('TimerStop').

mkl_utils

Non linearities are treated by mkl_utils mex file. Details are provided in mkl_utils.

rheo2NL

OBSOLETE. Use now nl_spring NL. NL=nl_spring('rheo2NL',model,DOF,offset);

This command is used to convert rheological data into a structure of data understandable for NLforce command. DOF is the list of the DOF coherent with u and v arguments of NLforce command. Offset is optional. It is a structure of data with fields .DOF and .def that defined 0 reference for Fu and Fv tab laws.

tab

model=nl_spring('tab',model);

This command is used to convert formal rheological description data stored in model.Stack to a tabulated law description. The format is likely to change due to optimization of the compiled functionality in mkl_utils (see mkl_utils).

BlockSave,BlockLoad

Undocumented intermediate save of a time block for long simulations that do not fit in memory.

mkl_utils

Purpose

For detailed callback information see sdtweb('nlspring_timeopt').

Residual

Residual command is used to compute standard residue.

mkl_utils('residual',r,model,u,v,a,opt,Case); call modifies variable r in memory according to following standard residue computation (implicit Newmark).

```
r = model.K{1}*a + model.K{2}*v + model.K{3}*u + b*(snl_fun) + fnl_int -fc;
```

In this formula, **b***(**snl_fun**) refers to the legacy m file implementations (non **chandle**) which expect the **nl_** function to fill a **unl** vector multiplying **b** (.**snl** field support was not activated). This was incorrectly noted -fnl in earlier documentation the minus sign coming from the convention of considering non-linear forces as external rather than internal (the convention used from now on). fnl_int refers to chandle implementations that match classical conventions : traction leading to positive stress in continuous mechanics, positive over-closure (negative gap) leading to positive pressures in contact mechanics.

Typically in fe_time computations one has

```
opt.Residual='r=-full(fc);mkl_utils(''residual'',r,model,u,v,a,opt,Case);';
```

with fc the time load (resulting from DofLoad entries in model Case) and fnl is the sum of the non linear efforts (if any) computed directly by mkl_utils (rotcenter, mocirc2), in the non linear functions (see sdtweb nl_fun) or in nl_spring. mkl_utils then calls the adequate nl_fun function (nl_spring by default) automatically.

Such call stored in opt.Residual is filled by nl_spring('TimeOpt') for default simulations.

Model information specifically supported by the residual command are

- opt.Rayleigh if the field exists defines a global Rayleigh damping and opt.Rayleigh(1)*model.K{1 is added to the residual.
- $model.K{2}$ can be a data structure describing modal damping with following fields:
 - .def : $M\Phi$ vectors as columns.
 - .data : c_j modal damping coefficients as a vector. $c_j = 2\omega_j\zeta_j$. A second column has to be set to zero for transient applications.

- .type : Onl_modaldmp handling function for callbacks. The following callbacks must be handled
 - * matrix projection $tkt = T^T KT$: tkt=feval(K.type, 'getTKT', K,T,Tt,typ) with K the implicit matrix, T the right projection matrix, Tt the left projection matrix (can be empty or skipped if $Tt = T^T$, typ the output type, either *imp* to keep the implicit format (by default), or *full* to recover a full numeric matrix (to be reserved for small output sizes).
 - * vector application f = Kq: f=feval(K.type,'getForce',K,q) with K the implicit matrix, q a deformation vector.
- .UseDiag : to be set to one if one wants the output of getTKT to be diagonal (as for a standard dtkt call).
- .K : optional additional damping matrix. This matrix must be in a mkl transposed v_handle format (use v_handle('mklst',K) to convert a matlab matrix to this format). Note that model.K{2}.K is taken in account for the Jacobian computation whereas modal damping is not.
- .defT : the resitution matrix (left side $M\Phi$), that can occur mainly in the case where a non-symmetric projection has been carried out. *E.g.*, the implicit representation of $T_l^T M\Phi \left[{}^2\zeta_j \omega_{j_k} \right] \Phi^T MT_R$ will use field .def to store $T_R^T M\Phi$ and .defT to store $T_l M\Phi$.

Corresponding additional residue term is $\sum_{j} [M] \phi_{j} * c_{j} * \phi_{j}^{T} [M]^{T} * v.$

• model.NL can be a stack of non linearities. Column 3 provides a structure with the following standard fields, see nldata.

Typically, fnl is computed by non linearity functions, see nl_fun for details on these functions. The non linear functions are called by mkl_utils to provide the value of fnl at a given state. Two implementations are supported

- An optimized *input-output* formulation, using observation and command matrices c and b documented in nldata. The computation of the observation is possible either on the displacement, the velocity or both, and the command is added to the residual using r = r + b*unl. With unl a vector depending on the observation (c*u, c_v v).
- A used defined addition (older format, that should be only used when the generic *b,c* format fails to be relevant. In this mode the non linear function must add fnl by itself, choosing the sign convention, using a call of type of_time(-1,fc,fc-fnl);. One will note that the residue vector is named fc in the non linear functions.

chandle

chandle objects are used to streamline communication between mex and MATLAB in iterative processes. They are used in various nl_solve calls and in particular for ModalNemwark and ExpNewmark.

chandle

Purpose

chandle objects are used to streamline communication between mex and MATLAB in iterative processes.

Creation generates a C copy of the matlab array and returns a **vhandle.chandle** object containing the ID. Register the chandle object for mexAtExit.

• chandle.numType lists currently implemented chandle subtypes.

DiagNewmark

DiagNewmark is an implementation of the Newmark scheme when assuming a fixed diagonal full Jacobian as occurs in modal domain transients (explicit or implicit).

ExpNewmark

ExpNewmark is an implementation of the Newmark scheme when assuming a fixed diagonal mass matrix for large explicit dynamic problems.

nl_inout

Support for observation performed in C. .iopt for standard integer options.

```
.N field : Nunl, (c,1),(c,2),cTrans, (b,2),(b,1),bTrans, Nopt[8],Niopt[9],size(unl,3)[
.opt field ? tc[1] dt0[2] K[3] Fmax[4] Fu functions currently implemented in C are listed
under nl_inout fun
```

The header of the associated class is

```
// nl_inout non linearity 1003
class chandleNl_inout: public chandle {
  public:
    int *irc, *jcc,*irb, *jcb,*iopt;
    double *prc,*pic,*prb,*pib,*unl,*vnl,*snl,*opt;
    int N[11]; // Nunl, (c,1),(c,2),cTrans, (b,2),(b,1),bTrans, Nopt[7],Niopt[8],size(ur
    __Fu Fu;// (*Fu)(chandleNl_inout*,struct _ROr);
    mxArray* MexData[2];
    chandleNl_inout();
```

```
~chandleNl_inout();
void Residual(struct _ROr ROr, double* fc);
void initCpt(); // Initialize pointers
void EndStep(); // propagate internal states using StoreType strategy
};
// Residual structure ------
struct _ROr {
    int _Nk,Nnl;
    double RayleighM,RayleighK,tc;
    double *u,*v,*a,*FNL;
    };
// Default function handle
typedef void(*__Fu)(chandle* ph, struct _ROr ROr);
```

Non linearities list

Purpose

List of supported non linearities. It is possible to create new ones (sdtweb nl_fun)

nl_inout

nl_inout is the more general non linearity, using observation and command matrix associated with elements supporting the kinematics (cbush for point connections, see section ?? , zero thickness volumes for surface connections (where two layers of coincident nodes are considered for a hexa8 or penta6 element, see section ??), volume elements for 3D applications (see section ??) or the deprecated observation/sensor command/loads as detailed in section ?? .

The general form of the non-linearity $f_{NL} = b \times f(C.u, C.v)$ is detailed in section ?? . For a list of implemented non-linear constitutive laws xxx

The pro.NLdata structure has fields described in section ?? (with the need to distinguish the form for model declaration and during time integration).

By default, no Jacobian is computed for this non-linearity. Experimental Jacobian are computed according 3 methods according to the NL.Jacobian value:

- 0 : no Jacobian. (default).
- 1 : tangent matrices.
- 2 : fixed Jacobian (can be max stiffness / damping or mean, ...).

Then computed matrices are then multiplied by NL.alphaJK factor for Jacobian stiffness, and NL.alphaJD factor for Jacobian damping.

${\tt nl_contact}$

Supports non conform fixed matching contact (squeal applications for example) and (surface contact large displacement, as in rail/wheel interaction for example). For conform meshes, zero thickness elements associated with p_zt can be used.

See p_contact, ctc_utils.

nl_modaldmp

Implementation of modal damping. Although modal damping is not a non-linear feature in itself, its implementation requires it to be declared as a non-linearity.

The concept is to provide shapes defined on a part of a model with associated damping ratios. nl_modaldmp handles the kinematic projection on the model which can contain superelements. In the case where superelements are used and concerned with modal damping, the shapes provided must be written on the physical DOF of the superelements.

The set of shapes must be stacked in model with a valid ID field. It is a common deformation SDT data structure (see sdtweb def), with an additional .ID field. The .data field is equivalent to the ones of complex modes (see fe_ceig). It is a matrix of two columns respectively giving the frequency and the target damping ratio for each mode.

Since modal damping implies a modal sensor, the features performs both by default. It is however possible to simplify it as a pure modal sensor. The theory around modal sensing/damping can be found in [12].

The pro.NLdata structure has fields

- type: string 'nl_modaldmp'.
- CurveId: the curve ID stacked in model which provides the shapes and their damping ratios.
- SensorOnly: to use the feature only as a modal sensor in a def data structure.

The NLdata structure generation can be integrated using an nl_modaldmp('db') call. See sdtweb nl_spring#setpro for this integration. This is used in transient simulations, and in complex mode computations, see nl_solve.

nl_inout _

Purpose

The generic form of NLdata specification is discussed in section ?? .

DofSet

Implementation of linear or large rotation setting of DOF values from time curves of a fe_case DofSet entry. sdtweb('_eval;','d_fetime.m#NLNewmark_LrDofSet').

Supported variants are

- large rotation trajectories (using MBBryan) uses .unl(:,:,1) corresponding to large displacement and updates .unl(:,:,j1) when changing time step to allow velocity computations (currently only valid for fixed time step). Requires setting .KeepDof=1 and enforcing 6 DOF associated with the master node.
- translation trajectories (xxx).

Power

```
uses NLdata.opt=[comstr('Power',-32) k n] and s_{nl} = k u_{nl}^{n}.
```

FuTable

uses tabular definitions in either NLdata.Fu or NLdata.Fv. If you want a case with both simultaneously to be re-implemented please provide a test case. Implements a Jacobian using tangent stiffness. Run example with sdtweb('_eval;','d_fetime.m#ModalNew_FuTable').

$K_{-} \texttt{t}$

Implement time varying matrices and DofSet xxx.

MexIOa

Implements callback to user defined .m file implementations of non-linearities as detailed in section ??.

SCLd

Large displacement surface contact supported as part of the **contact** module (see **sdtweb(xxx)**). Possible application : rail/wheel contact. This supports low pass filtering of contact forces, using an evolution equation of the form $\dot{F}_c/\omega_c + F_c = k_c g$ (for a positive constant) or $\dot{F}_c/\omega_c + F_c = k_c \sqrt{(g)}$ for a negative constant. This avoids incorrect bouncing of stiff contacts in implicit computations and ω_c should be a fraction of the sampling frequency (inverse of time step) or the maximum mesh frequency.

LRFu

Large rotation tabular spring for multi-body applications. cbush kinematics are expected and rotations are assumed using Bryan angles in radians. Requires one non-linearity for each spring (not currently vectorized). The NL.Node field gives slave and master node for each body (4 nodes). The observed motion is given at the master node and large rotation is used to determine the current position of the slave node. With single axis springs (large rotation rod) the position of the two slave nodes should be disjoint.

The stress vector used by this non-linearity contains 24 elements

- sx1,sy1,sz1,srx1,sry1,srz1 forces and moments on node 1
- sx,sy,sz,srx,sry,srz forces and moments on node 2
- PX1, PY1, PZ1, rx, ry, rz absolute position of node 1 and rotations
- ux, uy, uz relative displacements in x, y, z directions, sex, sey, sez force in element frame.

slab sensor non-linearity

This non-linearity supports observation of various quantities during FEM computations. The base definition is associated with the sdtweb('sensor# scell')

• For direct resultant sensors in time integration schemes using enforced displacement -fieldOut4, leading to NL.iopt(6+isens)==4. See more details in xxx

Needs documentation, see sdtweb d_fetime('slab').

nl_inout _

Note that in predictor/corrector schemes, it may be necessary to recompute the residual to obtain the correct residual.

temp still undocumented

- FuExpon $s_{nl} = a \ b^{u_{nl}}$.
- uMaxw generic C implementation of nl_maxwell.m file
- FuDahlC Dahl model with constant force
- $\bullet~{\tt STS}$ PSA scalar STS
- CLIMA2 connector non-linearity developed with Marco Rosatello
- FuFric basic friction model with $F_k = Ku$ bounded by $\pm F_{max}$.
- nl_bset xxx
- DofSet xxx

Non linearities list (deprecated)

Purpose

nl_maxwell (deprecated)

nl_maxwell describes rheological models using stiffness and damping. Deprecated implementation that should now be called with an nl_inout.

.type 'nl_maxwell'

- .lab Label of the non linearity.
- .Sens Observation definition. Cell array of the form {SensType,SensData} where SensType is a string defining the sensor type and SensData a matrix with the sensor data (see sdtweb sensor).
- .Load data structure defining the command as a load (with .DOF and .def fields).
- .SE superelement that defines the rheological model. Only matrices are used (.K field). Mass matrix is ignored. The .DOF field is unused and first DOF are assumed to be the observations defined, and following correspond to internal states.
- .NLsteps Number of sub steps for the integration.
- .StoreFNL strategy to store FNL output, should be initialized from StoreType.
- Ncell number of cells.

Jacobian is computed using a Guyan condensation keeping only the observation (internal states are condensed) to obtain tangent damping and stiffness.

Internal states are integrated using an independent finite differences explicit scheme, with the same step of time as the main scheme, or a subsampling NL.NLsteps times.

At the first residue computation, the initial internal states are computed according to initial condition in terms of displacements and velocities through a time integration until variation of speed between the 2 last computed steps is lower than opt.RelTol.

Force on the observation DOF (F), displacement (Qc) and velocity (dQc) of the internal DOF, displacement and velocity observations are stored in the NL output.

The command nl_spring db Fu" type" is a database of generalized Maxwell rheological models. type can be:

• zener standard viscoelastic model. Parameter k0, k1 and c1 can be given as a string of the form db Fu"zener k0 k0 k1 k1 c1 c1" in the command.

The example of the standard viscoelastic model is detailed here as an illustration. The standard viscoelastic model, also known as Zener model, is composed by a spring (K_0) in parallel with another spring (K_1) and a serial dashpot (C_1) as displayed figure 5.2.



Figure 5.2: Standard viscoelastic model.

In the Laplace domain, the relation between the relative load and the relative displacement is given by

$$F(s) = K(s)X(s) = \frac{K_0K_1 + (K_0 + K_1)C_1s}{K_1 + C_1s} = K_0\frac{1 + s/z}{1 + s/p}$$
(5.4)

where p and z are respectively the pole and the zero of the model

$$p = \frac{K_1}{C_1} \tag{5.5}$$

$$z = \frac{K_0 K_1}{\left(K_0 + K_1\right) C_1} \tag{5.6}$$

The maximum loss factor is

$$\eta_m = \frac{p-z}{2\sqrt{pz}} = \frac{1}{2} \frac{K_1}{\sqrt{K_0 \left(K_0 + K_1\right)}}$$
(5.7)

and obtained for pulsation

$$\omega_m = \sqrt{pz} = \frac{K_1}{C_1} \sqrt{\frac{K_0}{K_0 + K_1}}$$
(5.8)

 K_0 is the static stiffness of the model. Typically $K_1 = \frac{K_0}{2}$ and C_1 is defined so that the damping is maximal for the frequency of interest.

Following example considers $K_0 = 1000N/m$, $K_1 = 500N/m$ and $C_1 = 1.4Ns/m$. These parameters lead to a maximum loss factor of 20.14% for a frequency of 46.41Hz. The module and the loss factor are represented in figure 5.3.



Figure 5.3: Module and loss factor.

Following example consists in a mass of 1e-2kg linked to the ground by the Zener model. Initial displacement corresponding to a 1N load on the mass is imposed and then a time simulation is performed.

Non linearities list (deprecated).

```
r1=p_spring('default'); r1=feutil('rmfield',r1,'name');
r1.NLdata=data; r1.il(3)=param.k0;
r1.il(1)=100; model=stack_set(model,'pro','zener',r1);
% define option for time integration
opt=d_fetime('TimeOpt');
opt.NeedUVA=[1 1 1];
opt.Follow=1; opt.RelTol=-1e-5;
opt.Opt(7)=-1; % factor type sparse
opt.Opt(4)=param.dt; opt.Opt(5)=param.N; % NSteps
%opt.IterEnd='eval(opt.Residual)'; % to compute real FNL for current state
% Initial state
r1=data.SE.K{3}\[1;0]; r1=r1(1); % initial displacement for 1N load
model=stack_set(model,'curve','q0',struct('def',r1,'DOF',1.03));
% Time computation
def0=fe_time(opt,model); ci=iiplot; % compute
\% The same but NL as a model
SE2=data.SE;
SE2.Elt(end+1:end+2,1:6)=[Inf abs('mass1'); 1 0 0 param.m 0 0];
SE2=fe_caseg('assemble -secdof -matdes 2 3 1 -reset',SE2);
r1=SE2.K{3}\[1;0]; %r1=r1(1);
SE2=stack_set(SE2,'curve','q0',struct('def',r1,'DOF',SE2.DOF));
def20=fe_time(opt,SE2); % compute
F20=SE2.K{2}*def20.v+SE2.K{3}*def20.def; F20=F20(1,:);
% zener labs: {'zener-F1', 'zener-q1', 'zener-q1-1', 'zener-dq1', 'zener-dq1-1'}
NL20=struct('X',{{def20.data {'LIN-F1';'LIN-Qc1';'LIN-dQc1';'LIN-unl1';'LIN-vnl1';'ft'}
 'Xlab', {fe_curve('datatypecell', 'time')},...
 'Y', [F20' (fe_c(def20.DOF,1.03)*def20.def)'...
     (fe_c(def20.DOF,3.03)*def20.def)'...
     (fe_c(def20.DOF,1.03)*def20.v)'...
     (fe_c(def20.DOF,3.03)*def20.v)' zeros(size(def20.def,2),1) ]);
NL20.name='NLfromLIN';
iicom('curveinit',{'curve','NL(1)',ci.Stack{'NL(1)'};
        'curve',NL20.name,NL20});
A=ci.Stack{'NL(1)'}.Y(2:end,:);B=NL20.Y(2:end,:);t=NL20.X{1}(2:end);i2=any(A);
if norm(A(:,i2)-B(:,i2),'inf')/norm(B,'inf')>0.01
 figure(1);plot(t,A,'--o',t,B,'-')
 sdtw('_err', 'something has changed')
end
```

DofKuva

DofKuva defines a non linear load of the form

 $av_{Dof}^{e}[K]\{V\}$ with a scalar coefficient *a*, a scalar v_{Dof} extracted from displacement, velocity or acceleration, and *V* a field specified as follows

.type	'DofKuva'
.lab	Label of the non linearity.
.Dof	Dof of Case.DOF.
.Dofuva	[1 0 0] for displacement Dof, [0 1 0] for velocity and [0 0 1] for acceleration.
.MatTyp	Type of the matrix K (see MatType). Desired matrix is automatically assembled
	before time computation.
.factor	Scalar factor a.
.exponent	Exponent of the DOF.
.uva	Type of vector V: [1 0 0] for displacement, [0 1 0] for velocity and [0 0 1] for
	acceleration.

For example one can take in account gyroscopic effect in a time computation with a NL of the form

```
model=stack_set(model,'pro','DofKuva1005', ... % gyroscopic effects
    struct('il',[1005 fe_mat('p_spring','SI',1) 0 0 0 0 0],...
    'type','p_spring','NLdata',struct(...
        'type','DofKuva','lab','gyroscopic effect', ...
        'Dof',1.06,'Dofuva',[0 1 0],'MatTyp',7,...
        'factor',-1,'exponent',1,'uva',[0 1 0])));
```

DofV

DofV defines a non linear effort of the following form (product of a fixed vector and a dof) $(u)^{exponent}$. V NDdata fields for this non-linearity are

.type	'DofV'
.lab	Label of the non linearity.
.Dof	Dof of Case.DOF.
.Dofuva	[1 0 0] for displacement Dof, [0 1 0] for velocity and [0 0 1] for acceleration.
.exponent	Exponent of the DOF.
.def	data structure with fields .def which defines vector $\tt V$ and .DOF which defines corresponding DOF.

nl_spring defines a non linear load from rheological information (stop, tabulated damping or stiffness laws etc.) between 2 DOF.

To define a non linear spring, one has to add a classic **celas** element, linear spring between only 2 DOF. The non linear aspect is described by associated properties as a 'pro' entry in the model **Stack**.

One can describe non linearity by a formal rheological description using one or more of following fields in the pro Stack entry:

- .But : [dumax k0 c0 dumin k1 c1] bumpstop. For du from dumin to dumax, f=0. For du>dumax, k0 stiffness is applied to du-dumax, and for du<dumin, k1 stiffness is applied to du-dumin. Damping is not taken in account at this time (due to tabulated law strategy).
- .Fsec : [fsec,cpenal]. For dv<-fsec/cpenal or for dv>fsec/cpenal, f=fsec is applied. For -fsec/cpenal<dv<fsec/cpenal,f=cpenal*dv is applied. If omitted, cpenal=1e5.
- .K
- .C

This information will be converted in tabulated laws Fu and Fv using nl_spring tab (low level call that should be automatically called at the beginning of time computation).

One can also describe non linearity with a tabulated effort / relative displacement and effort / relative velocity law between the DOF (dof2-dof1), respectively in the Fu and Fv fields of the pro Stack entry. First column of Fu (resp. Fv) gives the relative displacements (resp. velocities) and second column gives the efforts. One can give a coefficient av factor of Fv depending on relative displacement as a third column of Fu. It can be useful to describe a non linearity depending on relative displacement and relative velocity. Force applied is F=av(du) .Fv(dv). It is used in particular to describe damping in a stop (.But NL).

Following example performs a non linear time computation on a simple 2-node model:

```
li{3}='SimuCfg{Exp{1m,10,uva110}}'; RT.nmap('CurExp')=li;% Same using explicit
sdtm.range(RT);mo2=RT.nmap('CurModel');d2=RT.nmap('CurTime');
```

```
figure(10);clf;% plot some comparison between results
subplot(211);plot(def.data,[def.def' d2.def']);xlabel('Time [s]');ylabel('displacement'
subplot(212);plot(def.data,[def.v' d2.v']);xlabel('Time [s]');ylabel('velocity')
legend('Implicit','Explicit');setlines;
sdth.os(10,'@OsDic',{'ImGrid','ImSw80','ImTight'})
```

Following example deals with a clamped-free beam, with a bilateral bump stop at the free end.

```
% define model:
L=1; b=1e-2; h=2e-2; e=1e-3; % dimensions
model=[];
model.Node=[1 0 0 0 0 0 0; 2 0 0 0 L 0 0];
model.Elt=[Inf abs('celas') 0 0;
           20 20
                        100 1 110 0; % linear celas
           Inf abs('beam1') 0 0;
           1 2 1 1
                       0 1 0 0
           ];
model=feutil(sprintf('RefineBeam %.15g',L/20),model);
model=fe_case(model,'FixDof','base',1); % clamps 1st end
model=fe_case(model,'FixDof','2D',[0.03;0.04;0.05]); % 2D motion
% model properties:
model.pl=m_elastic('dbval 1 steel');
model.il=p_beam(sprintf('dbval 1 BOX %.15g %.15g %.15g %.15g',b,h,e,e));
% Bump stop NL:
model=stack_set(model,'pro','celas1',...
        struct('il',[100 fe_mat('p_spring','SI',1) 1e-9 0 0 0 0],...
               'type', 'p_spring',...
               'NLdata', struct('type', 'nl_inout',...
                               'but', [0.02 5e2 0 -0.02 5e2 0], ... % gap knl cnl...
                               'umin',3)));
if 1==1
 model=fe_case(model,'DofLoad','in',struct('DOF',2.02,'def',50));
 model=fe_curve(model,'set','input','TestStep t1=0.02');
else
 f=linspace(12,18,3);
model=fe_case(model,'DofLoad','in',struct('DOF',2.02,'def',1));
 model=fe_curve(model,'set','input',sprintf('Testeval cos(%.15g*t)',f(1)*2*pi));
end
```

```
model=fe_case(model,'setcurve','in','input');
% Time computation:
opt=d_fetime('TimeOpt dt=1e-3 tend=10'); opt.NeedUVA=[1 1 0];
def=fe_time(opt,model);
```

RotCenter

The **Rotcenter** joint is used to introduce a penalized translation link between two nodes A and B (rotation DOFs of NL entry are ignored), where the motion of A is defined in a rotating frame associated with angle θ_A and large angle rotation $R_{LG}(\theta_A)$. The indices G and L are used to indicate vectors in global and local coordinates respectively.

The positions of nodes are given by

$$\{x_A\}_G = [R_{GL}] (\{p_A\} + \{u_A\}_L) \{x_B\}_G = (\{p_B\} + \{u_B\}_G)$$
(5.9)

which leads to expressions of the loads as

$$\{F_A\}_L = [R_{LG}] \left(K \left(\{x_B\}_G - \{x_A\}_G \right) \right) \{F_B\}_G = K \left(\{x_A\}_G - \{x_B\}_G \right)$$
 (5.10)

To account for viscous damping loads in the joints, one must also compute velocities. Using (2.2), one obtains

$$\{\dot{x}_A\}_G = [R_{GL}] (\{\dot{u}_A\}_L + \{\omega(t)\} \land \{p_A + u_A\}_L) \{\dot{x}_B\}_G = \{\dot{u}_B\}_G$$

$$(5.11)$$

Velocity computations are currently incorrect with u_A ignored in the rotation effect. So that viscous damping loads can be added

$$\{FC_A\}_L = [R_{LG}] \left(K \left(\{\dot{x}_B\}_G - \{\dot{x}_A\}_G \right) \right) \{FC_B\}_G = K \left(\{\dot{x}_A\}_G - \{\dot{x}_B\}_G \right)$$
 (5.12)

For a linearization around a given state (needed for frequency domain computations or building a sensor observation matrix),

$$\left\{ \begin{array}{c} q_{AG} \\ q_{BG} \end{array} \right\} = \left[\begin{array}{c} R_{GL} & 0 \\ 0 & I \end{array} \right] \left\{ \begin{array}{c} q_{AL} \\ q_{BG} \end{array} \right\}$$
(5.13)

In global basis, stiffness matrix of a celas link is given by

$$k \begin{bmatrix} I & -I \\ -I & I \end{bmatrix}$$
(5.14)

which leads to the following stiffness matrix

$$\begin{bmatrix} R_{GL}^T & 0\\ 0 & I \end{bmatrix} k \begin{bmatrix} I & -I\\ -I & I \end{bmatrix} \begin{bmatrix} R_{GL} & 0\\ 0 & I \end{bmatrix} = k \begin{bmatrix} I & -R_{GL}^T\\ -R_{GL} & I \end{bmatrix}$$
(5.15)

where q_A DOFs are in the local basis (motion relative to the shaft in its initial position) and q_B are in the global frame.

data describing this link is stored in model stack as a p_spring pro entry. Stiffness and damping are stored respectively as 3rd and 5th column of the data.il field (standard linear spring, see sdtweb('p_spring')).

NDdata fields:

- .type string 'RotCenter'.
- .sel a FindElt command to find celas of RotCenter type.
- .k this field should not be used. .JCoef field should be used instead and has priority. Stiffness used for Jacobian computation. Damping is not taken in account in Jacobian in this case.
- .JCoef coefficient of celas stiffness and damping for Jacobian computation. Default is 1.
- .drot the rotation DOF.
- .lab label.

nl_rotCenter

This non linearity can be used to connect 2 points A and B, where the motion of A is defined in a rotating frame associated with angle θ_A and large angle rotation $R_{LG}(\theta_A)$. More generally A and B are no real nodes but defined implicitly as observation matrices. **nl_rotcenter** is an extension of **RotCenter** documented above, using observation matrices which is more general.

- .type string 'nl_rotcenter'.
- .sel a FindElt command to find elements associated to the NL link
- .JCoef coefficient of celas stiffness and damping for Jacobian computation. Default is 1.
- .drot the rotation DOF.
- .lab label.

Non linearities list (deprecated) _

- .Weights (optional) Weight of the stiffness in a pivot link (in fact computed force is multiplied by the weight factors before being applied so that the sum of weight coef divided by number of points by pivot should be equal to 1).
- .Stack Stack of cta coupling. Of the form {'cta', 'name', {r1,r2}}, where 'cta' is a constant string defining the type of the link, 'name' a string containing the name of corresponding links. r1 is the observation in the first (rotating) part. It is a data structure with fields .Node defining the nodes involved, .cta defining the observation matrix, .DOF defining corresponding DOF (as many columns as in .cta) and .SeName defining as a string the name of the superelement where cta is defined (if omitted, it is assumed that DOF and cta are defined on the model.DOF - no superelement -). r2 is the same for the non rotating part.

An example can be found in t_nlspring 2beam.

Default uses the damping and stiffness defined in the il field of the p_spring pro entry to model a linear spring/damper between the 2 parts (stiffness il(3) and damping il(5)).

Defining a xb parameter, the Excite NONL law will be applied instead of the spring/damper. Parameter that are to be defined are

- .xb Radial clearance.
- .kb Stiffness at radial clearance.
- .cb Damping at radial clearance.

Stiffness and damping at initial position are given in corresponding p_spring properties il(3) and il(5). For example:

```
cf.mdl=nl_spring('setpro ProId 103 k 371 c 2000e-3 xb 0.03 kb 37100 cb 5',cf.mdl);
```

rod1

The rod1 non-linear connection is a simple penalized rigid link. One considers two nodes A and B (see figure 5.4).



Figure 5.4: Large rotation rod functional representation.

Currently, one can introduce masses at points A and B. mass2 elements should be used to account for the actual position of the center of gravity.

The global non linear load associated with the rod is thus

$$F_{rod} = k_r \left(\| \{ x_B - x_A \} \| - L_0 \right) \frac{\{ x_B - x_A \}}{\| \{ x_B - x_A \} \|}$$
(5.16)

which accounts for a load proportional to the length fluctuation around L_0 (penalized rod model).

When linearizing, one considers a strain energy given by $k_r ||q_B - q_{A'}||^2$ with the motion at node A' being related to the 6 DOFs at node A by

$$\{q_{A'}\} = \begin{bmatrix} I & \left[\vec{AB}\land\right]\\ 0 & I \end{bmatrix} \{q_A\}$$
(5.17)

Node A node is free to rotate. The linearized stiffness thus corresponds to an axial stiffness in the direction of the rod. The computation of the stiffness is however based on the current position of the extremity nodes, a difficulty in model manipulations is thus to translate these nodes.

data describing this link is stored in model stack as a p_spring pro entry. Stiffness and damping are stored respectively as 3rd and 5th column of the data.il field (standard linear spring, see sdtweb('p_spring')). NL information is stored in the data.NLdata field which has itself following fields :

- type : string 'rod1'.
- sel: a FindElt command to find associated celas of rod1 type (('proid100').
- ulim : build tabulated law from -ulim to ulim. Default is 1e3.
- lab : label.

nl_gapcyl



This non-linearity implements non-linear contact between two cylinders of radius R_O for the outer cylinder and R_I for the inner cylinder with motion defined on the cylinder center line. Assuming the mesh to be defined in an updated Lagrangian configuration where the center lines X_I and X_O at not coincident, the positions in a deformed states are given by $x_I = X_I + u_I$ and $x_O = X_O + u_O$.

Contact may only occur when the cylinders are not centered. When the two cylinders are not centered the non-linear observation is given by

$$u_{NL} = x_O - x_I = u_O - u_I + (X_O - X_I) = [c] \{q\} + u_{NL0}$$
(5.18)

From this distance between the center lines, a cylindrical basis is defined with $e_r(q)$ along the direction from x_I to x_O and e_θ forming a direct basis with the cylinder axis (kept constant from the initial value of the updated lagrangian position).

The functional definition of the contact force uses the gap in the e_r direction defined by

$$g = \{e_r\}^T \{u_{NL}\} - (R_O - R_I)$$
(5.19)

where $R_O - R_I = d$ is stored as parameter NLdata.d and the contact leads to two opposite forces on the cylinder center lines

$$\{f_I\} = \{-f_O\} = f(g)\{e_r\}$$
(5.20)

The current implementation assumes rotations to be small enough to ignore the difference between e_r and the corresponding vector E_r in the upated Lagrangian configuration.

If update of mesh leads to lateral slip, then u_I may account for longitudinal position of the contact point along the beam using shapes functions.

When linearizing the contact around a given point, the stiffness $\partial f \partial g$ only occurs in the e_r direction.

Creating a new non linearity: nl_fun.m ____

Purpose

The structure of nl_spring allows creating any new non-linearity through the use of a dedicated function, named nl_fun.m. This function which non-linearity name will be fun, will be automatically called by nl_spring for classical operations.

The function structure has been designed to comply with specific needs. Standard calls have been defined, which are detailed below:

• Residue computation, called by mkl_utils (sdtweb mkl_utils), must output the entry force minus the non linear force computed. The call performed is

```
nl_fun(r2,fc,model,u,v,a,opt,Case)
```

This call is low level and must modify fc using sp_util('setinput') as fc-fnl where fnl is the non linear force computed. Note that this is the only possible call for nargin==8. Note that mkl_utils allows a formalism with precomputed observations, using fields unl.

• Jacobian computation, must output the tangent stiffness and tangent damping matrices associated to the non linearity. The call performed is

```
[kj2,cj2]=nl_fun(NL,[],model,u,v,[],opt,Case,RunOpt);
```

This call must output either empty matrices if no tangent nor Jacobian matrix is associated to the non linearity, or matrices expressed on the DOF vector of Case.DOF. The first matrix is the tangent stiffness matrix, the second one is the tangent damping matrix. Typically there are 3 normalized methods to be defined (but not all of them must be defined, and more can be defined) according to the NL.Jacobian value:

- -0: no Jacobian. (default).
- -1: tangent matrices.
- 2 : fixed Jacobian (can be max stiffness / damping or mean, ...).

Then computed matrices are then multiplied by NL.alphaJK factor for Jacobian stiffness, and NL.alphaJD factor for Jacobian damping.

• Initializations for fe_time, must initialize the model non-linearity for non linear forces computation

The call must generate the non linearity stored in model.NL, it can optionally generate non linear DOF and labels. The call performed is of the type.

NL=nl_fun('init',data,mo1);

NL is a struct containing at least the field type with the nl_fun handle (*e.g.* NL.type=@nl_fun). data contains the Stack, pro entry, and mo1 is the model, named mo1 where the call is performed.

• **ParamEdit** returns the **ParamEdit** string allowing integrated parameters interpretation (for internal SDT use).

The call performed is of the type.

st=nl_fun('ParamEdit');

• db returns default NLdata fields for a non linearity. This allows integrated building of nonlinearities in a model. This function can call ParamEdit to allow interactive setup.

This call must return a NLdata field and is of the type

NLdata=nl_fun('db data 0');

• Energy post treatments capability, should return the elastic energy stored in the nonlinearity as a vector with as many lines as time steps in the output.

The call performed by nl_solvePost is of the form

r2 = nl_fun('PostEnerNL');

The non linearity function can access in caller fields RO, out, model, NL, i1 with

- RO a structure with fields EnerP and EnerK respectively containing the potential and kinetic energy.
- out the fe_timeoutput.
- model the model used in the simulation.
- NL the NL containing the data of the non-linearity called.
- i1 the row index of out.Post that is currently generated.
- **Renumbering** capability, must return the non-linearity written for the new renumbered nodes, elements, dof, . . .

The call performed (by feutilbfor example) is of the type

NL=nl_fun('renumber',NL,nind);

nind is the renumbering vector.

The designed nl_fun template is given in the non-linear toolbox, sdtweb nl_fun.m#1. It is a functional non linear function, computing a zero non linear force. The definition of a non linearity using nl_fun in a standard SDT model is given in the following.

```
% A standard SDT model
model=struct('Node', [1 0 0 0 0 0 0; 2 0 0 0 0 1],...
             'Elt', [Inf abs('celas') 0 0;
             1 2 3 -3 0 1 0 10; % linear celas
            ]);
% Define a non linearity of type nl_fun
model=nl_spring('SetPro ProId 100',model,nl_fun('db data0'));
%Equivalent to
% model=stack_set(model,'pro','nl_fun',...
% struct('il',[100 fe_mat('p_spring','SI',1)],...
%
         'type', 'p_spring',...
         'NLdata', struct('type', 'nl_fun', 'data', [])));
%
% Define the case
model=fe_case(model, 'FixDof', 'base', 1);
model=fe_case(model,'DofLoad','in',struct('DOF',2.03,'def',1));
model=fe_curve(model,'set','input','TestStep t1=0.02');
model=fe_case(model,'setcurve','in','input');
% Define the TimeOpt and compute the solution
opt=nl_solve('TimeOpt'); opt.Opt([4 5])=[1e-3 1e4];opt.NeedUVA=[1 1 0];
def=fe_time(opt,model);
```

nl_solve

Purpose

Integrated non linear simulations

Description

The simulation of non linearities require special handling in SDT, which is packaged in the non linear toolbox. This function aims at performing classical studies, such as done by fe_simulfor classical SDT models with this special handling.

See **nllist** for the list of supported non linearities.

TimeOpt

nl_solve('TimeOptMethod',RO) used to initialize fe_time options for later simulation. Currently implemented methods

- Explicit Newmark scheme
- Stat non-linear static Newton
- Theta method time integration.
- ModalNewmark uses an optimized fully C based integration for the case where DOF correspond to modal degree of freedom. The stepped sine strategy is discussed in section ?? .
- NLNewmark default implicit Newmark scheme.

Associated options provided in ${\tt RO}$ or in the command are

• .tend end time of simulation. Used to initialize .ts=ceil(tend/dt).

Static

To compute the static state of a model with non-linearities.

q0=nl_solve('static',model);

It is possible to use custom fe_time simulation properties using the model stack entry info,TimeOptStat. See nl_spring TimeOpt for fields and defaults.
It is possible to use as command option any field from the usual static simulation option, see sdtweb nl_spring#TimeOpt to have more details. *E.g.* To redefine on the fly the maximum number of iteration, one can enter [q0,opt]=nl_solve('static maxiter 100',model);.

By default, the staticNewton algorithm implemented in fe_time is called.

An Uzawa algorithm is also implemented in nl_solve, under the method static nl_solve uzawa. This algorithm is very different from the staticNewton one since here the solution is not incremented but fully re-computed at each iteration. This is useful when some non-linear forces do not derive from potentials. Command StaticUzawa can be used in nl_solve - to access it: q0=nl_solve('static Uzawa', model);.

Mode

The definition of modes for non-linear models is not straight forward. This command aims at computing tangent modes as function of a non-linear model current state. The resolution thus concerns a linear model with tangent stiffness, damping matrices corresponding to the model current displacement, velocity, acceleration state. The eigenvalue solvers used are then fe_eigfor real modes and fe_ceigfor complex modes.

By default, modes tangent to a static state are computed. A static simulation is performed to produce a model state from which tangent matrices are computed. It is also possible to compute tangent modes at specific instants during a transient simulation, at SaveTimes instant, and to store frequency/damping data and deformations.

A set of command options allows detailing the mode computation wanted and the output.

Accepted command options to control the model computation itself are

- -allmatdes to ask for an assembly with all matrix types assembled, the default assembly command used is -matdes 2 3 1. This command can be used to keep specific matrix types defined in pre-assembled superelements.
- cpx for complex mode computation (default is real mode computation).
- -evalFNL (in combination with command traj) asks to recompute the FNL field on the fly based on displacements prior to mode computation. This command is useful when solutions used for the tangent state have been imported from an external solver.
- skip skips fe_timesimulations and performs the complex mode computation based on the zero deformation and with initialized values of non linearities. The behavior will thus depend on the non linearity initialization strategy. *E.g.* for contact see (p_contact), the -skip option will consider a full contact state.

- stat for mode computation based on a static state (typically after a fe_time staticNewton simulation). Uses model stack entry info,TimeOptStat.
- time for mode computations during a transient simulation (exclusive with the default -stat option). Uses model stack entry info,TimeOpt.
- traj for mode computations based on states provided as an additional argument.

The -stat and -time options are mutually exclusive and define the base solver options to be used by fe_timefor the preliminary state computation. With -stat option (default) the stack entry info, TimeOptStat will be sought and used if found. With -time option, the stack entry info,TimeOpt will be used if found.

The -traj option is complementary and is used to force the complex mode computation on provided states. On can either provide the state in deformation curve format, see sdtweb def as a last argument, or use predefined stack entries. In -stat mode (default), the model stack entry curve,q0 will be sought and used if found, if not the result will use the -skip mode. In -time mode, the model stack entry curve,TSIM will be sought and used. If not found an error will occur.

Accepted command options to control the output format are

- -addedOnly (in combination with backTgtMdl) only outputs the tangent matrices as a superelement that would have been added to the base matrices for the mode computation.
- -alpha (requires -cpx) to also output the real mode participation to the complex modes. This is in fact the projection of the complex modes on the real mode basis.
- -backTgtMdl outputs the tangent model that would have been used for mode computation.
- -dataOnly to save only the frequency, damping data (does not store the deformation field). The output is then under a frequency tracking curve in the iiplotformat.
- -fullDOF to output the deformation fields restituted on the unconstrained DOF.
- -keepTval (requires -cpx) to allow keeping the underlying real mode basis when computing complex modes. With val set to 1, the initial real mode basis will be kept under field def.T, as an additional independent output, coherent with the -alpha command option. With val set to 2, the complex modes will not be restituted but expressed on the subspace used for their computation, the subspace basis will be output in def.Mode.TR, allowing a complete compatibility with feploton-the-fly restitution strategy for display. This latter option is the most complete and efficient strategy. Complete subspace information is kept and can be used for further exploitation, complex mode projection on real mode (-alpha) is naturally obtained, and memory footprint is optimized as the storage size of the subspace is commonly lower by a factor 1.5 to 2 than the complex mode basis;

- -noPost is used to skip any solution post treatment, and outputs the raw mode structure straight from the solver.
- -PostFcn'' cam'' is used to perform specific post-treatments on the mode output after computation.
- -real "ModeBas" (requires -cpx) to specify a particular real mode basis on which the complex modes will be computed. The real mode basis is supposed to be stored in the model stack entry curve, ModeBas.

Internally, the solver defines and uses the model stack entry info,SolveOpt structure to handle the options documented above. One can define it as a structure with the fields documented (case sensitive) and provide it instead of the EigOpt input. Additional advanced field are then accessible

- EigOpt a vector providing eigenvalue computation options following the fe_eigformat.
- cpx'' command'' to externalize the mode computation. This command is by default a Boolean telling the solver whether to perform a complex mode computation (set to 1) or a real mode computation (set to 0). If a string is provided, the solver will evaluate it as an external command instead of performing mode computation. One then gets access to the nl_solve mode computation framework for ones' own solver.
- ind provides a vector of indices that will be used to restrict the output to the indexed modes.
- SubDef provides a command that will be evaluated to perform a dynamic user defined restriction to the output modes, it is thus more general than the ind option. The result of the command has to be a vector of indices.
- AssembleCall to force a specific AssembleCall strategy.

The various input and output strategies allow for the support of several input syntax. The following calls are thus accepted, with model a standard SDT model, Case a standard SDT case structure, eigopt either a vector providing options for fe_eigor a structure with optional fields defined above, def a standard SDT deformation field structure used by -traj when necessary.

```
nl_solve('mode',model);
nl_solve('mode',model,eigopt);
nl_solve('mode',model,Case,eigopt);
nl_solve('mode',model,def);
nl_solve('mode',model,Case,def);
nl_solve('mode',model,eigopt,def);
nl_solve('mode',model,Case,eigopt,def);
```

Sample calls using command options to extract tangent modes are given below.

```
def0=nl_solve('Mode',model)
def0=nl_solve('Mode',model,[5 20 1e3]) % with eigopt
def0=nl_solve('Mode-stat-fullDOF',model);
defT=nl_solve('Mode-time',model);
hist=nl_solve('Mode-time-dataOnly',model);
histC=nl_solve('Mode-cpx-time-dataOnly',model);
defC=nl_solve('Mode-cpx-time-alpha-real''MyBas''-fullDOF',model);
def1=nl_solve('Mode-skip-fullDOF',model);
```

Post

The Post command allows performing energy and potential further post treatments of a non-linear simulation. The output is integrated in the standard fe_timesimulation outputs in field out.Post that is a three columns cell array directly compatible with the iiplot format.

To obtain the post treatments, one must define them prior to starting the simulation. Direct computation of the post-treatments *a posteriori* is also possible.

• Command PostDefine adapts the TimeOpt structure to initialize fields in the output and trigger post treatments in the final cleanup phase. The PostDefine call must thus be performed after the TimeOpt call. Using this command itself prior to a time simulation is enough to obtain the post treatments.

```
opt=nl_solve('PostDefine keys',opt); % adapts the opt structure.
model=nl_solve('PostDefine keys',model); % adapts the opt structure contained in mediate
```

- Command PostLab provides the list of available post treatment keywords. The input is a structure with fields the post treatment keywords and a logical.
- Command PostHist provides an iiplot curve structure adapted to the post treatments. On can provide a PostLab structure with fields assigned to 1 for desired posts to obtain the corresponding curve.
- Command PostCompute computes the post treatments and store them in out.Post. This command is internally called if the PostDefine command was used prior to the time simulation. For *a posteriori* computations, the user must provide the out as a standard fe_time format initialized with Post field and the assembled model. The model must feature a stack entry info, OutputOptions with field Post containing the PostLab structure.

```
% Generate a TimeOpt
opt=nl_solve('TimeOpt');
Perform the time simulation
def=fe_time(opt,model);
% Initialize for post treatments
[def,RO]=nl_solve('PostInit EnerM',def);
model=stack_set(model,'info','OutputOptions',...
struct('Post',RO));
% Assemble model with non linearities
model=fe_case(opt.AssembleCall,model);
% Compute post treatments
def=nl_solve('PostCompute',def,model);
% display in iiplot
iiplot(def.Post);
```

• Command PostInit is an internal function that initializes the output Post field at the start of the simulation. Early initialization is useful if the post treatments are performed on the fly by the OutputFcn.

The following post treatments are available

- EnerP The linear potential, or strain energy.
- EnerK The kinetic energy.
- EnerNL The elastic or strain energy stored in the non linearities.
- EnerM The mechanical energy, defined as EnerP + EnerK + EnerNL.
- PDiss The instant dissipated power.
- EnerDiss The cumulated dissipated energy over time.

Command PostEstimate allows analyzing the energy curves to compute

- Fest an estimation of the vibration frequency (based on quasi-sinusoidal oscillations)
- DmpR an estimation of the damping ratio based on the estimated frequency by computing the dissipated mechanical energy. $\zeta = \frac{1}{4\pi} \log \frac{E_m(t_0)}{E_m(t_1)}$
- Emax the maximum mechanical energy identified on the cycle analyzed.

• EDiss the dissipated mechanical energy over the cycle analyzed.

The following command options allow altering the estimation

- -cfi to specify the iiplot figure with handle i.
- -bandpass fmax to perform a bandpass from 0 to fmax Hz filtering prior to the analysis.
- -curveName'' name'' to provide the iiplot stacked curve name to exploit.
- -baseOn''name'' to specify on which post treated curve the frequency estimation is made.
- -globalMaxTolval to provide a relative tolerance over which a point is detected as close to the global maximum. This is exploited to detect the peaks over the energy signal analyzed.
- -localMax to estimate the frequency by detecting the zeros of the signal derivative (less robust).
- -unit'', *II*'' to provide an output unit system.
- XFcn''str'' to provide a function call to be evaluated that can perform further post treatments(e.g. model specific posts). The called function can access out, outLab, st, j1 with out a matrix containing the output with as many lines as provided curves and as many columns as outputs data, outLab a cell array containing the labels of each column, st the curve list (either names or the curves themselves), j1 the curve currently treated.

```
r1 = nl_solve('PostEstimate',def);
r1 = nl_solve('PostEstimate',def.Post{1,3});
r1 = nl_solve('PostEstimate',{'disp(1)'});
r1 = nl_solve('PostEstimate',{'Post_NLsolve(1)'});
```

TgtMdlBuild, Assemble

Integrated command to generate linearized models around a specific working point. This command packages the tangent model generation procedures of nl_solve Mode-backTgtMdl. The low level implementations are documented in nl_spring NLJacobianUpdate (for example keepLin interaction are documented there).

• TgtMdlAssemble command outputs a fully linearized assembled model, based on the static state provided.

- TgtMdlBuild command generates a linearized model with superelement coupling containing the tangent stiffness and damping contributions of all non-linearities. The following command options are supported
 - -keepName allows naming the superelements with the non-linearity name.
 - -evalFNL forces recomputation of non-linearities states before generation.
 - SEPro to initialize p_superentries for the generated superelements.
 - setPar to setup parameters associated to the generated superelements, conforming to the content of the nlpro.
 - keepNLProId not to generate tangent superelements for the non-linear properties specified by their ProId.
 - -staticInterp generates a tangent model allowing tangent matrix interpolation between different static states. The procedure requires the definition of parameters and a method to compute static states. Static states for MinMax configurations of each parameters is then performed. Matrices showing differences as function of parameters are kept and an interpolation rule is defined using the linear finite element functions of a 2^{npar} vertices hypercube. The output model has stack fields curve,q0 the series of static states with q0.data providing the parameter points, and info,sCoef providing interpolation rules for each matrix.

```
RA=struct('par',Ra,'q0cbk',{{@my_fun,'ComputeStatic'}});
mo1=nl_solve('TgtMdlBuild-staticInterp',model,RA);
```

Ra is either a Range structure or the content of Range.param (see sdtweb fe_range), q0cbk is a callback in cell-array format.

staticInterp2 implements a more generic method using vhandle.matrix NLJac types for richer interpolation. In such case one can provide a field .Range in the input options to force a richer DoE for the static basis. By default griddedInterpolant is used so that the given Range must be compatible with the interpolation strategy.

```
% Linearized model generation
% sample model with cubes in contact
model=d_contact('cubes cbuild');
% resolve static state
q0=nl_solve('static',model);
% linearized model
mo1=nl_solve('TgtMdlBuild',stack_set(model,'curve','q0',q0));
% check the result
feutil('info',mo1)
SE=stack_get(mo1,'SE'); SE{1,3}
```

nl_mesh

Purpose

Integrated mesh modifications and case handling for non-linear applications

Description

Integrated case handling for constraint penalization and coupling component splitting hare implemented in this function.

Some non-linearities require surface/volume remeshing (*e.g.* definition of conforming interfaces for contact) or adaptations (*generation of thin interface layers*). This function regroups such functionalities. Mesh generation are performed by fe_gmsh(interface to gmsh) and fe_tetgen (interface to tetgen, see help fe_tetgen).

Conform

The Conform call is an integrated call to generate conforming meshes between two facing interfaces. The command generates a conforming surface mesh of the face to replace, merges it with the conform mesh of the second interface, replaces the model face mesh and remeshes the model volume to yield a new equivalent volume with a conform face mesh.

```
mo1=nl_mesh('conform eltsel"FindElt"',model,sel);
% sel={eltSelToReplace eltSelForReplacement;...}
```

model is a standard SDT model. sel is a cell array containing in each line two FindElt commands specifying the element selection face to remesh and the element selection face to use for the conforming interface for replacement.

- Command option eltsel allows specifying in a string a FindElt command restraining the working area in the original model.
- Command option **smartSize** allows generating a conforming mesh with a coherent mesh characteristic length.
- Command option gmsh allows using gmsh to mesh the final volume.
- Command option tetgen allows using tetgen to mesh the final volume (by default).
- Command option output asks to output the generated mesh in a .mat file.
- Command option **OrigContour** asks to keep original positions of mid-nodes of the quadratic faces delimiting the volume to remesh. This may however yield mesh wrapping problems when the face to remesh is much coarser than the mesh trace to place for conformity.

- Command option mergeTo allows specifying a FindElt selection command in a string to replace the mesh on another model selection than the one used to generate the conforming interface (which uses eltsel.
- It is also possible to provide additional arguments, which will be passed the the nl_meshcover call performed in the procedure.

Limitations: The Conform call only supports generation of conforming interfaces when one interface contour fully contains the other interface contour. Handling of more complex contour configurations has not been implemented. Besides, this function has been designed to handle planar surfaces. Additional operation to work on non planar surfaces are left to the user (*e.g.* pre/post projections of the surfaces on a plane).

Contour

Call ContourFrom generates SDT beam1or beam3 contour models for CAD definitions. All formats readable by gmsh can theoretically be used. Only the .geo, .stp and .igs are tested.

Since .geo files can contain geometric yet non discretized objects, a 1D meshing pass is performed with gmsh to provide an SDT contour model. This is not supported for other file types.

```
model=nl_mesh('contourFrom','file.stp'); % not specifying the type
```

Call Contour generates an SDT face mesh from an SDT beam1or beam3 contour.

model=nl_mesh('contour',model);

model is an SDT beam model defining a closed contour.

- Command option lcval allows specifying a characteristic length for Gmsh.
- Command option lcminval allows specifying a minimal characteristic length for Gmsh.
- Command option quad allows generating quadratic meshes.
- Command option keepNode asks to keep the original contour NodeId for the contour command.
- Command option diag asks to output the Gmsh log file for diagnostic problems.
- Command option single tells nl_mesh that a single contour is defined. This is useful when several closed contours are defined since it is impossible to automatically decide whether each contour is independent or if they define a single complex contour.

nl_mesh_

- Command option groupval is used in combination to the single command option. This allows specifying which contour group will be meshed, while other possible contours will define holes.
- Command option algo ''val'' allows specifying which algorithm gmsh must use (this depends on the gmsh version, report to the gmsh documentation for more details).
- Command option AllowContourMod allows gmsh adding nodes on the contour provided. By default gmsh is forced not to add nodes to the lines defining the contour to mesh.

Cover

The Cover call is designed to mesh the interstice between two closed planar contours, when one fully contains the other. The call is performed as

[newModel,opt,largeContour]=nl_mesh('cover',model,{eltsel_large,eltsel_small});

model is a standard SDT model. Variables eltsel_large and eltsel_small are FindElt calls defining the element selection of the respectively large surface and small surface (the small being contained in the large).

The output **newModel** is the mesh generated from the surface contours.

opt outputs additional information about the mesh generation, it is a struct containing fields .NodeAdd specifying the potential nodes added in the interstice space meshed,.nodeEdgeSel1 specifying the NodeId of the nodes located on the eltsel_large contour, .nodeEdgeSel2 specifying the NodeId of the nodes located on the eltsel_small contour, and .tname the name of the temporary file containing the generated mesh.

largeContour provides the original contour in beam elements of the eltsel_large selection.

The following command options are available

- merge allows merging the interstice mesh with the inner mesh of the eltsel_small selection.
- quad allows generating proper quadratic meshes.
- **smartSize** allows generating an interstice mesh with a characteristic length in coherence with the contour mesh length.
- lcval allows setting the characteristic length to Val to the interstice mesher.
- algo ''name'' allows specifying the meshing algorithm name to the gmsh mesher. See the gmsh documentation for more information.

Hole[,Groups,Diff,Drill,Gen]

The Hole command series aims at handling hole detection on surfaces and bore drilling generation. The following functionalities are avaiable

Command HoleGroups detects holes on a closed surface and outputs a contour model with element groups relative to each isolated contour. A second output provides the GroupId corresponding to detected holes.

Command HoleDiff provides surface elements that are inside the holes of a given contour. You should better exploit lsutilto get a robust result.

Command HoleGen generates a planar surface with a ruled mesh featuring a hole and controlled radial positions.

- .len length of plate
- .wid width of plate
- .rAnulus radii for base positions
- .ND angular refinement
- .NRext external to bolt radius refinement
- .MatId assign mat/pro id to meshed part
- .noExt remove exterior side
- .Center
- .normal

Command HoleDrill generates cylindrical drills in a model with the possibility to integrate a ruled bolt mesh.

Replace

The call **Replace** is designed to replace parts of a model mesh with new given meshes, mesh parts conformity is assumed. It is performed as

model=nl_mesh('replace',model,nodesToReplace,NewModel,nodeIDtoKeep)

model is a standard SDT model. nodesToReplace is a cell array containing vectors of NodeId specifying the areas to be replaced. NewModel is a cell array containing the new models which will be merged to the mesh in coherence with the removed elements (specified by nodesToReplace). nodeIDtoKeep is an optional argument specifying NodeId of the original model for nodes whose NodeId must not change in the transformation.

Control of nodeIDtoKeep per NewModel part is possible by providing a cell array of NodeId list of the same size than NewModel.

The following command options are available

- setMat allows defining a specific MatId to the output mesh.
- setPro allows defining a specific ProId to the output mesh.
- eltset*FindEltString* can be provided to provide an element selection for MatId and ProId assignment.
- keepNoCheck in combination with the use of a third argument nodeIDtoKeep assumes the nodes numbering is correct and forces the nodes original numbering without check.
- -jAll asks to join all elements per type, then separated by MatId
- -inSet asks to maintain coherence with EltId sets. EltId sets for which the totality of a given removed part belonged to will be updated to contain the EltId of the replacement mesh.

Rivet

This command generates rivet drills in a specified contour. A model containing a beam contour can be provided, or an EltSel string generating a surface selection (see section and the selface option) on a bigger model. A data structure providing the origins, and rivet radius and washer (or rivet head radius). The mesh generated between both radius is structured.

The data structure must contain fields

- Orig providing the rivet centers in an [x y z;...] matrix.
- **radHole** providing the rivet hole radius, either a scalar if all rivets have the same radius, or a line vector providing each rivet radius separately.
- radWash providing the rivet washer (or head) radius, either a scalar if all rivets have the same washer radius, or a line vector providing each rivet washer radius separately.

and can optionally contain fields

- plane To directly provide the contour plane normal to define the drilling, in an [nx ny nz; ...] matrix.
- Ns To define the number of mesh segments in the rivet to washer radius area (default 10), either a scalar if all rivet heads have the same properties, or a line vector defining the property for each rivet separately.
- Nr To define the number of mesh radial nodes in the rivet to washer radius area (default 2), either a scalar if all rivet heads have the same properties, or a line vector defining the property for each rivet separately.
- Command option MatIdval allows setting the modified mesh MatId to val.
- Command option ProId val allows setting the modified mesh ProId to val.
- Command option –fill outputs in second argument a compatible mesh of the rivet bores.
- Command option –allQuad outputs the remeshed model with elements only.

Following example meshes a rectangular contour with a few rivet drilling inside.

```
% Generate a global contour
model=struct('Node',[...
1 0 0 0 0 0 0;
2 0 0 0 10 0 0;
3 0 0 0 10 2 0
4 0 0 0 0 2 0], 'Elt',[]);
model.Elt=feutil('ObjectBeamLine 1 2 0 2 3 0 3 4 0 4 1',model);
model=feutil('refinebeam .2',model);
%feplot(model)
% define rivet positions, eventually planes
R0=struct('Orig',[ 3 1 0;6 1 0;9 1 0],...
'radHole',[.2;.2;.2],...
'radHole',[.8;.8;.8]);
model=nl_mesh('Rivet',model,R0);
cf=feplot(model);
```

ShellSkin

Generation of a skin mesh from shell elements.

Syntax: mo1=nl_mesh('ShellSkin,model,RO. model is a model with shell elements, RO is a facultative option structure input. Output mo1 is a model with added volume elements.

By default, all shell elements are selected, and the average thickness is computed using the associated integration rules. A symmetric extrusion following the shell normals is performed.

Command option Get outputs an exteral model rahter than an addition to the existing model. The following options are available:

- .sel to provide a Findelt command to select target shell elements.
- .twoSided to generate a volume extrusion from both sides.
- .vol to define the volume mesh format: 0 will generate shell skin meshes linked with rigid elements, 1 will generate volume elements, 2 will generate volume elements linked with rigid elements to the shell.
- .MatId to assign a new material identifier to the new elements. Set to NaN to leave initial MatId.
- .ProId to assign a specific ProId to the new elements, one can set to 0 if only the topology is used.
- .New to add the new elements to the existing model, with new EltId.

```
% Generate a shell mesh
model=femesh('testquad4 -divide 4 4');
% Generate a wolume mesh with rigid connections to the shell nodes.
RA=struct('sel','groupall','vol',2,'twoSided',1);
mo1=nl_mesh('ShellSkinGet',model,RA);
cf=feplot(mo1); fecom(cf,'colordagagroup')
```

GmshVol

This call integrates the generation of a volume mesh from a face mesh with gmsh.

```
model=nl_mesh('GMSHvol',model);
```

model is a standard SDT face mesh model.

- Command option setmat allows specifying a specific MatId to the output mesh.
- Command option setpro allows specifying a specific Prold to the output mesh.
- Command option keepFaces asks to keep original NodeId of the nodes located on the face mesh.
- Command option lc specifies a characteristic length for gmsh.
- Command option clmin specifies a minimal mesh length for gmsh.
- Command option clmax specifies a maximal mesh length for gmsh.

ExtrudeLayer

This command generates a non trivial extrusion of a face mesh following the face normal at each node, to generate a volume layer.

model=nl_mesh('ExtrudeLayer thick Val',model);

model is an SDT model with shell elements (a surface definition).

Command option thick specifies the extrusion thickness. Command option setmat allows specifying a specific MatId to the output. Command option setpro allows specifying a specific ProId to the output.

StackClean

This call cleans up a model stack when mesh modifications have been performed. It cleans up stack entries definition that became incoherent with some mesh modifications.

model=nl_mesh('StackClean',model);

Command option **rmuns** removes stack entries that could not be sorted out. Command option **rmmod** removes stack entries affected by the model modifications.

See also celas, p_spring, fe_gmsh

spfmex_utils

Purpose

OfactOptim

This command can be used to set spfmex parameters in order to optimize computation speed for factorization and / or solving.

spfmex_utils('OfactOptim',ki,RO,ofact(1,'lu'));

 $\tt ki$ is the matrix that is used for the optimization. $\tt RO$ is a data structure defining options with following fields:

- .nCompt Number of computation for result averaging.
- .maxDomain Max size of blocks of the elimination tree (fraction of matrix size).
- .maxZeros Max number of zeros in the blocks of the resolution tree (fraction of matrix size).
- .refineStep Number of step to refine the optimal parameter pair found in the first step. Command option -refine must be added to perform the refine step.

The last argument ofact(1,'lu') is needed in order to call directly spfmex_utils.

Available command options are

- -setopt use default RO.
- -refine performs refine step for optimal search.
- fact to benchmark factorization step.
- solve to benchmark resolution step.
- -plot to plot history in iiplot

Following example optimize only solving:

nl_bset

Purpose

Non linearity to support handling of enforced displacement

This is implemented in nl_spring('nl_bset'). This currently assumes the existence of a stiffness and xxx viscous damping xxx.

extrotor

Purpose

External links

- https://www.sdtools.com/help/mpc.html
- https://www.sdtools.com/help/p_spring.html
- https://www.sdtools.com/help/beam1.html
- https://www.sdtools.com/help/case.html
- https://www.sdtools.com/help/celas.html
- https://www.sdtools.com/help/findnode.html
- https://www.sdtools.com/help/findelt.html
- https://www.sdtools.com/help/fe_gmsh.html
- https://www.sdtools.com/help/fe_gmsh.html
- https://www.sdtools.com/help/fe_mknl.html#MatType
- https://www.sdtools.com/help/fe_time.html#newmark
- https://www.sdtools.com/help/fe_time.html#NLNewmark
- https://www.sdtools.com/help/fe_time.html#staticNewton
- https://www.sdtools.com/help/fe_case.html#ConnectionScrew
- https://www.sdtools.com/help/fesuper.html
- https://www.sdtools.com/help/feplot.html
- https://www.sdtools.com/help/iiplot.html
- https://www.sdtools.com/help/ctc_utils.html
- https://www.sdtools.com/help/fe_eig.html
- https://www.sdtools.com/help/fe_ceig.html
- https://www.sdtools.com/help/fe_time.html
- https://www.sdtools.com/help/fe_simul.html
- https://www.sdtools.com/help/feutilb.html

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extrotor _____